Amazon SageMaker: Developer Guide

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# Table of Contents

**What is Amazon SageMaker?** ........................................................................................................... 1

Pricing for Amazon SageMaker ........................................................................................................... 1
Are you a first-time user of Amazon SageMaker? .............................................................................. 1
Overview of machine learning with Amazon SageMaker ................................................................. 2

**SageMaker Features** ......................................................................................................................... 5

New features ........................................................................................................................................ 5
Machine learning environments ............................................................................................................. 7
Major features ....................................................................................................................................... 8

**Get started** ...................................................................................................................................... 12

Set Up Amazon SageMaker Prerequisites ............................................................................................ 13
Create an AWS Account ....................................................................................................................... 13
Create an Administrative User and Group ....................................................................................... 14
AWS CLI Prerequisites ....................................................................................................................... 15

Domain overview ................................................................................................................................ 16
SageMaker domain ............................................................................................................................... 18
Quick onboarding ................................................................................................................................. 50
Custom onboarding using IAM Identity Center .................................................................................. 53
Custom onboarding using IAM ........................................................................................................... 61
Choose an Amazon VPC ....................................................................................................................... 67

Supported Regions and Quotas ............................................................................................................ 68
Quotas .................................................................................................................................................. 69

**Use automated ML, no-code, or low-code** ..................................................................................... 70

SageMaker Autopilot ............................................................................................................................. 70

Create a Regression or Classification Job Using the AutoML API ..................................................... 74
Create an Image Classification job using the AutoML API .................................................................. 159
Create a Text Classification job using the AutoML API .................................................................... 170
Create a Time-series Forecasting job using the AutoML API ............................................................ 181
Create an LLM fine-tuning job using the AutoML API ....................................................................... 221
Create a Regression or Classification Job Using the Studio Classic UI ............................................ 246

Example Notebooks ............................................................................................................................. 257
Quotas .................................................................................................................................................. 260
API reference ...................................................................................................................................... 262

SageMaker JumpStart ........................................................................................................................... 264
Open and use JumpStart in Studio ..................................................................................................... 265
Open and use JumpStart in Studio Classic ........................................................... 267
Foundation Models ................................................................................................. 271
Task-Specific Models ............................................................................................... 318
Shared Models and Notebooks .............................................................................. 338
Solution Templates .................................................................................................. 344
SageMaker JumpStart Industry: Financial ............................................................ 356

**Use machine learning environments offered by SageMaker** .................................. 362

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Studio</td>
<td>363</td>
</tr>
<tr>
<td>Migrating from Amazon SageMaker Studio Classic</td>
<td>365</td>
</tr>
<tr>
<td>Launch Amazon SageMaker Studio</td>
<td>382</td>
</tr>
<tr>
<td>Amazon SageMaker Studio UI overview</td>
<td>384</td>
</tr>
<tr>
<td>Applications supported in Amazon SageMaker Studio</td>
<td>388</td>
</tr>
<tr>
<td>Amazon SageMaker Studio spaces</td>
<td>389</td>
</tr>
<tr>
<td>Perform common tasks</td>
<td>392</td>
</tr>
<tr>
<td>Use NVMe stores with Amazon SageMaker Studio</td>
<td>393</td>
</tr>
<tr>
<td>Local mode support in Amazon SageMaker Studio</td>
<td>395</td>
</tr>
<tr>
<td>View, stop, or delete your Studio running instances, applications, and spaces</td>
<td>403</td>
</tr>
<tr>
<td>Amazon SageMaker Studio pricing</td>
<td>413</td>
</tr>
<tr>
<td>Troubleshooting</td>
<td>413</td>
</tr>
<tr>
<td>Studio Classic</td>
<td>414</td>
</tr>
<tr>
<td>Studio Classic Features</td>
<td>415</td>
</tr>
<tr>
<td>UI Overview</td>
<td>416</td>
</tr>
<tr>
<td>Launch Amazon SageMaker Studio Classic</td>
<td>423</td>
</tr>
<tr>
<td>JupyterLab Versioning</td>
<td>426</td>
</tr>
<tr>
<td>Use the Studio Classic Launcher</td>
<td>436</td>
</tr>
<tr>
<td>Collaborate with shared spaces</td>
<td>441</td>
</tr>
<tr>
<td>Use Studio Classic Notebooks</td>
<td>451</td>
</tr>
<tr>
<td>Customize Studio Classic</td>
<td>517</td>
</tr>
<tr>
<td>Perform Common Tasks</td>
<td>567</td>
</tr>
<tr>
<td>Studio Classic Pricing</td>
<td>581</td>
</tr>
<tr>
<td>Troubleshooting</td>
<td>581</td>
</tr>
<tr>
<td>SageMaker JupyterLab</td>
<td>587</td>
</tr>
<tr>
<td>JupyterLab user guide</td>
<td>588</td>
</tr>
<tr>
<td>JupyterLab administrator guide</td>
<td>597</td>
</tr>
<tr>
<td>Migrating from SageMaker Studio Classic to SageMaker Studio</td>
<td>624</td>
</tr>
<tr>
<td>SageMaker Notebook Instances</td>
<td>626</td>
</tr>
</tbody>
</table>
Amazon SageMaker

SageMaker Studio Lab ................................................................. 692
  Studio Lab components overview ........................................... 693
  Onboard to Studio Lab ............................................................ 698
  Manage your account ............................................................. 699
  Launch Studio Lab ................................................................. 701
  Use Studio Lab starter assets .................................................. 703
  Studio Lab pre-installed environments ...................................... 705
  Use the Studio Lab project runtime .......................................... 707
  Troubleshooting ................................................................. 731

SageMaker Canvas ...................................................................... 734
  Are you a first-time SageMaker Canvas user? ............................ 736
  Getting started ........................................................................ 737
  Setting Up and Managing Amazon SageMaker Canvas (for IT Administrators) ........................................ 744
  Import data into Canvas ........................................................... 800
  Prepare data .......................................................................... 842
  Use generative AI with foundation models ................................ 933
  Use Ready-to-use models ......................................................... 958
  Use custom models .............................................................. 969
  Logging out ........................................................................... 1112
  Limitations and troubleshooting .............................................. 1113
  Manage billing and cost ......................................................... 1124
  SageMaker geospatial capabilities ............................................ 1126
  How can I use SageMaker geospatial capabilities? ..................... 1127
First-time user? ................................................................................................................................. 1128
Getting started ................................................................................................................................. 1129
Geospatial processing job ................................................................................................................ 1145
Earth Observation Jobs ..................................................................................................................... 1161
Vector Enrichment Jobs .................................................................................................................... 1169
Visualization Using SageMaker geospatial capabilities ................................................................. 1170
Amazon SageMaker geospatial Map SDK ....................................................................................... 1174
SageMaker geospatial capabilities FAQ ........................................................................................... 1182
Security and Permissions .................................................................................................................. 1183
Types of compute instances ............................................................................................................ 1196
Data collections ................................................................................................................................ 1199
RStudio on Amazon SageMaker ........................................................................................................ 1204
  Region availability ........................................................................................................................... 1205
  RStudio components ....................................................................................................................... 1206
  Differences from Posit Workbench .................................................................................................. 1206
  Manage RStudio on SageMaker ..................................................................................................... 1207
  Use RStudio on Amazon SageMaker .............................................................................................. 1256
SageMaker Code Editor ...................................................................................................................... 1261
  Code Editor user guide .................................................................................................................... 1262
  Code Editor administrator guide ..................................................................................................... 1274
SageMaker HyperPod ........................................................................................................................ 1284
  Prerequisites .................................................................................................................................. 1286
  Getting started with SageMaker HyperPod .................................................................................... 1294
  Operate SageMaker HyperPod ........................................................................................................ 1302
  SageMaker HyperPod lifecycle configuration best practices ......................................................... 1312
  Run jobs on HyperPod clusters ...................................................................................................... 1325
  Cluster resiliency ............................................................................................................................ 1331
  Cluster management ....................................................................................................................... 1337
  References ..................................................................................................................................... 1339
  SageMaker HyperPod FAQ .............................................................................................................. 1344
  HyperPod release notes .................................................................................................................. 1346
Use generative AI in SageMaker notebook environments .............................................................. 1349
  Installation ...................................................................................................................................... 1350
  Features .......................................................................................................................................... 1351
  Model configuration ....................................................................................................................... 1353
  Use Jupyter AI .................................................................................................................................. 1360
<table>
<thead>
<tr>
<th>Label data with a human-in-the-loop</th>
<th>1365</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ground Truth</td>
<td>1365</td>
</tr>
<tr>
<td>Are You a First-time User of Ground Truth?</td>
<td>1366</td>
</tr>
<tr>
<td>Getting started</td>
<td>1367</td>
</tr>
<tr>
<td>Label Images</td>
<td>1374</td>
</tr>
<tr>
<td>Label Text</td>
<td>1399</td>
</tr>
<tr>
<td>Label Videos and Video Frames</td>
<td>1413</td>
</tr>
<tr>
<td>Label 3D Point Clouds</td>
<td>1461</td>
</tr>
<tr>
<td>Verify and Adjust Labels</td>
<td>1525</td>
</tr>
<tr>
<td>Creating Custom Labeling Workflows</td>
<td>1537</td>
</tr>
<tr>
<td>Create a Labeling Job</td>
<td>1584</td>
</tr>
<tr>
<td>Use Input and Output Data</td>
<td>1634</td>
</tr>
<tr>
<td>Enhanced Data Labeling</td>
<td>1743</td>
</tr>
<tr>
<td>Security and Permissions</td>
<td>1760</td>
</tr>
<tr>
<td>Monitor Labeling Job Status</td>
<td>1799</td>
</tr>
<tr>
<td>Ground Truth Plus</td>
<td>1803</td>
</tr>
<tr>
<td>Getting Started with Amazon SageMaker Ground Truth Plus.</td>
<td>1805</td>
</tr>
<tr>
<td>Request a Project</td>
<td>1807</td>
</tr>
<tr>
<td>Create a Project Team</td>
<td>1809</td>
</tr>
<tr>
<td>Open the Project Portal</td>
<td>1812</td>
</tr>
<tr>
<td>Create a Batch</td>
<td>1813</td>
</tr>
<tr>
<td>Review Metrics</td>
<td>1815</td>
</tr>
<tr>
<td>Review Batches</td>
<td>1817</td>
</tr>
<tr>
<td>Accept or Reject Batches</td>
<td>1820</td>
</tr>
<tr>
<td>Create and Manage Workforces</td>
<td>1820</td>
</tr>
<tr>
<td>Using the Amazon Mechanical Turk Workforce</td>
<td>1821</td>
</tr>
<tr>
<td>Managing Vendor Workforces</td>
<td>1826</td>
</tr>
<tr>
<td>Use a Private Workforce</td>
<td>1828</td>
</tr>
<tr>
<td>Crowd HTML Elements Reference</td>
<td>1860</td>
</tr>
<tr>
<td>SageMaker Crowd HTML Elements</td>
<td>1861</td>
</tr>
<tr>
<td>Augmented AI Crowd HTML Elements</td>
<td>1964</td>
</tr>
<tr>
<td>Augmented AI</td>
<td>1974</td>
</tr>
<tr>
<td>Get Started with Amazon Augmented AI</td>
<td>1976</td>
</tr>
<tr>
<td>Use Cases and Examples</td>
<td>2006</td>
</tr>
<tr>
<td>Create a Human Review Workflow</td>
<td>2017</td>
</tr>
<tr>
<td>Delete a Human Review Workflow</td>
<td>2044</td>
</tr>
<tr>
<td>Topic</td>
<td>Page</td>
</tr>
<tr>
<td>----------------------------------------------------------------------</td>
<td>------</td>
</tr>
<tr>
<td>Create and Start a Human Loop</td>
<td>2047</td>
</tr>
<tr>
<td>Delete a Human Loop</td>
<td>2054</td>
</tr>
<tr>
<td>Create and Manage Worker Task Templates</td>
<td>2058</td>
</tr>
<tr>
<td>Monitor and Manage Your Human Loop</td>
<td>2073</td>
</tr>
<tr>
<td>Output Data</td>
<td>2074</td>
</tr>
<tr>
<td>Permissions and Security</td>
<td>2089</td>
</tr>
<tr>
<td>CloudWatch Events</td>
<td>2097</td>
</tr>
<tr>
<td>API References</td>
<td>2100</td>
</tr>
<tr>
<td><strong>Prepare data</strong></td>
<td>2102</td>
</tr>
<tr>
<td>Prepare Data with Data Wrangler</td>
<td>2103</td>
</tr>
<tr>
<td>Get Started with Data Wrangler</td>
<td>2107</td>
</tr>
<tr>
<td>Import</td>
<td>2120</td>
</tr>
<tr>
<td>Create and Use a Data Wrangler Flow</td>
<td>2195</td>
</tr>
<tr>
<td>Get Insights On Data and Data Quality</td>
<td>2204</td>
</tr>
<tr>
<td>Automatically Train Models on Your Data Flow</td>
<td>2217</td>
</tr>
<tr>
<td>Transform Data</td>
<td>2218</td>
</tr>
<tr>
<td>Analyze and Visualize</td>
<td>2279</td>
</tr>
<tr>
<td>Reusing Data Flows for Different Datasets</td>
<td>2291</td>
</tr>
<tr>
<td>Export</td>
<td>2302</td>
</tr>
<tr>
<td>Use Data Preparation in a Studio Classic Notebook to Get Data Insights</td>
<td>2337</td>
</tr>
<tr>
<td>Security and Permissions</td>
<td>2343</td>
</tr>
<tr>
<td>Release Notes</td>
<td>2359</td>
</tr>
<tr>
<td>Troubleshoot</td>
<td>2365</td>
</tr>
<tr>
<td>Increase Amazon EC2 Instance Limit</td>
<td>2375</td>
</tr>
<tr>
<td>Update Data Wrangler</td>
<td>2376</td>
</tr>
<tr>
<td>Shut Down Data Wrangler</td>
<td>2378</td>
</tr>
<tr>
<td>Prepare Data at Scale with Studio Classic using Amazon EMR or AWS Glue</td>
<td>2379</td>
</tr>
<tr>
<td>Prepare data using Amazon EMR</td>
<td>2380</td>
</tr>
<tr>
<td>Prepare data using AWS Glue Interactive Sessions</td>
<td>2423</td>
</tr>
<tr>
<td><strong>Process data</strong></td>
<td>2433</td>
</tr>
<tr>
<td>Sample Notebooks</td>
<td>2434</td>
</tr>
<tr>
<td>CloudWatch Logs and Metrics</td>
<td>2435</td>
</tr>
<tr>
<td>Data Processing with Apache Spark</td>
<td>2435</td>
</tr>
<tr>
<td>Running a Spark Processing Job</td>
<td>2435</td>
</tr>
<tr>
<td>Data Processing with scikit-learn</td>
<td>2436</td>
</tr>
<tr>
<td>Data Processing with Framework Processors</td>
<td>2437</td>
</tr>
</tbody>
</table>
Time to live (TTL) duration for records .......................................................... 2539
Cross account feature group discoverability and access ................................ 2541
  Enabling cross account discoverability ......................................................... 2543
  Enabling cross account access ...................................................................... 2548
Feature Store storage configurations ............................................................. 2560
  Online store .................................................................................. 2560
  Offline store .................................................................................. 2562
  Throughput modes ............................................................................. 2563
Collection types ..................................................................................... 2566
Add features and records to a feature group ................................................ 2567
  API ............................................................................................ 2568
  Example code ................................................................................ 2568
Find features in your feature groups ............................................................. 2570
  How to search for your features ............................................................... 2571
Find feature groups in your Feature Store .................................................... 2576
  How to find feature groups .................................................................... 2578
Adding searchable metadata to your features ............................................. 2584
  How to add searchable metadata to your features .................................... 2585
Create a dataset from your feature groups .................................................. 2592
  Using the Amazon SageMaker Python SDK to get your data from your feature groups ... 2593
  Sample Amazon Athena queries .................................................................. 2598
Delete records from your feature groups ..................................................... 2600
  Delete records from the online store ......................................................... 2600
  Delete records from the offline store ......................................................... 2602
Logging Feature Store operations by using AWS CloudTrail ..................... 2605
  Management events ............................................................................ 2605
  Data events .................................................................................... 2605
Security and access control ....................................................................... 2607
  Using AWS KMS permissions for Amazon SageMaker Feature Store ....... 2607
  Authorizing use of a customer managed Key for your online store .......... 2608
  Using grants to authorize Feature Store ............................................... 2611
  Monitoring Feature Store interaction with AWS KMS ........................... 2611
  Accessing data in your online store ......................................................... 2611
  Authorizing use of a customer managed key for your offline store ......... 2611
Quotas, naming rules and data types .......................................................... 2612
Quota terminologies .................................................................................. 2612
Limits and quotas .............................................................................................................................. 2612
Naming rules .................................................................................................................................. 2613
Data types ......................................................................................................................................... 2613
Amazon SageMaker Feature Store offline store data format ...................................................... 2614
Amazon SageMaker Feature Store offline store URI structures ................................................. 2615
Amazon SageMaker Feature Store resources ................................................................................... 2616
Feature Store example notebooks and workshops ........................................................................ 2616
Feature Store Python SDK and API ................................................................................................ 2617

Train machine learning models ..................................................................................................... 2619
The simplest training workflow in SageMaker .............................................................................. 2619
Full view of the SageMaker Training workflow and features ...................................................... 2620
Before training ............................................................................................................................... 2622
During training ............................................................................................................................... 2624
After training .................................................................................................................................. 2626
Model Training .................................................................................................................................. 2628
Choose an Algorithm ..................................................................................................................... 2630
Choose an algorithm implementation ........................................................................................... 2631
Problem types for the basic machine learning paradigms .......................................................... 2634
Use Built-in Algorithms ................................................................................................................. 2637
Use Reinforcement Learning ......................................................................................................... 3076
Run local code as a remote job ....................................................................................................... 3084
Set up your environment .................................................................................................................. 3085
Invoking a function .......................................................................................................................... 3093
Configuration file ........................................................................................................................... 3104
Customize your runtime environment ............................................................................................ 3106
Container image compatibility ....................................................................................................... 3107
Logging parameters and metrics with Amazon SageMaker Experiments ................................... 3113
Using modular code with the @remote decorator ........................................................................ 3117
Private repository for runtime dependencies ............................................................................... 3120
Example notebooks ....................................................................................................................... 3122
Experiments ...................................................................................................................................... 3122
Supported AWS Regions ............................................................................................................... 3123
Create an experiment ..................................................................................................................... 3124
View, search, and compare experiment runs .................................................................................. 3132
SageMaker integrations .................................................................................................................. 3137
Tutorials ............................................................................................................................................ 3141
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>CloudTrail metrics</td>
<td>3142</td>
</tr>
<tr>
<td>Clean up experiment resources</td>
<td>3144</td>
</tr>
<tr>
<td>Additional supported SDK</td>
<td>3146</td>
</tr>
<tr>
<td>Experiments FAQs</td>
<td>3151</td>
</tr>
<tr>
<td>Search using the console and API</td>
<td>3154</td>
</tr>
<tr>
<td>Perform Automatic Model Tuning</td>
<td>3160</td>
</tr>
<tr>
<td>How Hyperparameter Tuning Works</td>
<td>3161</td>
</tr>
<tr>
<td>Define metrics and environment variables</td>
<td>3165</td>
</tr>
<tr>
<td>Define Hyperparameter Ranges</td>
<td>3168</td>
</tr>
<tr>
<td>Track and set completion criteria</td>
<td>3173</td>
</tr>
<tr>
<td>Tune Multiple Algorithms</td>
<td>3177</td>
</tr>
<tr>
<td>Example: Hyperparameter Tuning Job</td>
<td>3190</td>
</tr>
<tr>
<td>Stop Training Jobs Early</td>
<td>3205</td>
</tr>
<tr>
<td>Run a Warm Start Hyperparameter Tuning Job</td>
<td>3207</td>
</tr>
<tr>
<td>Resource Limits for Automatic Model Tuning</td>
<td>3213</td>
</tr>
<tr>
<td>Best Practices for Hyperparameter Tuning</td>
<td>3216</td>
</tr>
<tr>
<td>Refine data during training</td>
<td>3219</td>
</tr>
<tr>
<td>How SageMaker smart sifting works</td>
<td>3220</td>
</tr>
<tr>
<td>Supported frameworks and AWS Regions</td>
<td>3222</td>
</tr>
<tr>
<td>Apply SageMaker smart sifting to your training script</td>
<td>3223</td>
</tr>
<tr>
<td>Best practices, considerations, and troubleshooting</td>
<td>3233</td>
</tr>
<tr>
<td>Security in SageMaker smart sifting</td>
<td>3234</td>
</tr>
<tr>
<td>SageMaker smart sifting Python SDK reference</td>
<td>3235</td>
</tr>
<tr>
<td>Release notes</td>
<td>3238</td>
</tr>
<tr>
<td>Debug and improve model performance</td>
<td>3238</td>
</tr>
<tr>
<td>Use TensorFlow</td>
<td>3239</td>
</tr>
<tr>
<td>Use SageMaker Debugger</td>
<td>3258</td>
</tr>
<tr>
<td>Access a training container through SSM for remote debugging</td>
<td>3435</td>
</tr>
<tr>
<td>Release notes</td>
<td>3445</td>
</tr>
<tr>
<td>Profile and optimize computational performance</td>
<td>3447</td>
</tr>
<tr>
<td>Use SageMaker Profiler</td>
<td>3449</td>
</tr>
<tr>
<td>Monitor AWS compute resource utilization in SageMaker Studio Classic</td>
<td>3472</td>
</tr>
<tr>
<td>Release notes</td>
<td>3551</td>
</tr>
<tr>
<td>Distributed training</td>
<td>3552</td>
</tr>
<tr>
<td>Before you get started</td>
<td>3553</td>
</tr>
<tr>
<td>Get started with distributed training in Amazon SageMaker</td>
<td>3554</td>
</tr>
</tbody>
</table>
Asynchronous inference ........................................................................................................ 4286
How It Works ......................................................................................................................... 4286
How Do I Get Started? .......................................................................................................... 4287
Create, invoke, and update an Asynchronous Endpoint .................................................... 4288
Monitor asynchronous endpoint ......................................................................................... 4301
Check prediction results ...................................................................................................... 4305
Autoscale an asynchronous endpoint ................................................................................ 4309
Troubleshooting .................................................................................................................. 4313

Batch Transform .................................................................................................................. 4320
Use Batch Transform to Get Inferences from Large Datasets ............................................ 4321
Speed up a Batch Transform Job ......................................................................................... 4323
Use Batch Transform to Test Production Variants ............................................................. 4323
Sample Notebooks ............................................................................................................... 4324
Associate Prediction Results with Input ............................................................................. 4324
Storage in Batch Transform ............................................................................................... 4332
Troubleshooting .................................................................................................................. 4333

Model parallelism and large model inference ..................................................................... 4334
Deep learning containers for LMI ....................................................................................... 4335
SageMaker endpoint parameters for LMI ........................................................................... 4339
LMI tutorials ........................................................................................................................ 4340
Configurations and settings ............................................................................................... 4360
Choosing instance types for LMI ....................................................................................... 4385
Deploying uncompressed models ....................................................................................... 4391
LMI FAQs ............................................................................................................................ 4392
LMI troubleshooting .......................................................................................................... 4393
Release notes for LMI deep learning containers ............................................................... 4394

Update models in production ............................................................................................. 4404
How to get started ............................................................................................................... 4404
Auto-Rollback Configuration and Monitoring ................................................................... 4405
Blue/Green Deployments .................................................................................................... 4409
Rolling Deployments .......................................................................................................... 4424
Exclusions ............................................................................................................................ 4429

Shadow tests ....................................................................................................................... 4430
Create a shadow test ........................................................................................................... 4430
View, monitor, and edit shadow tests ................................................................................ 4435
Complete a shadow test ...................................................................................................... 4442
Best Practices .......................................................................................................................... 4445
Exclusions ............................................................................................................................... 4445
Access containers through SSM ......................................................................................... 4446
   Allowlist ............................................................................................................................. 4446
   Enable SSM access .......................................................................................................... 4447
   IAM configuration ........................................................................................................... 4447
SSM access with AWS PrivateLink .................................................................................... 4449
Logging with Amazon CloudWatch Logs ............................................................................. 4449
Accessing model containers .............................................................................................. 4449
Deploy models with model servers ................................................................................... 4450
   Deploy models with TorchServe ...................................................................................... 4450
   Deploy models with DJL Serving ...................................................................................... 4458
   Deploy models with Triton Inference Server ................................................................... 4463
Deploy models at the edge with SageMaker Edge Manager ............................................ 4473
   Why Use Edge Manager? ................................................................................................. 4473
   How Does it Work? ........................................................................................................... 4473
   How Do I Use SageMaker Edge Manager? ...................................................................... 4474
   Getting Started ............................................................................................................... 4474
Set Up Devices and Fleets .................................................................................................. 4497
Package Model ................................................................................................................... 4505
The Edge Manager Agent .................................................................................................... 4512
Manage Model ...................................................................................................................... 4533
SageMaker Edge Manager end of life .................................................................................. 4545
Optimize model performance using Neo .......................................................................... 4547
   What is SageMaker Neo? .................................................................................................. 4547
   How it Works ..................................................................................................................... 4548
Compile Models .................................................................................................................... 4548
Cloud Instances .................................................................................................................... 4570
Edge Devices ....................................................................................................................... 4611
Troubleshoot Errors ............................................................................................................ 4644
Elastic Inference .................................................................................................................... 4654
Migrate from Amazon Elastic Inference to other instances .............................................. 4656
How EI Works ....................................................................................................................... 4661
Choose an EI Accelerator Type .......................................................................................... 4662
Use EI in a SageMaker Notebook Instance .......................................................................... 4663
Use EI on a Hosted Endpoint ............................................................................................... 4663
Frameworks that Support EI ................................................................. 4663
Use EI with SageMaker Built-in Algorithms ........................................ 4664
EI Sample Notebooks ........................................................................ 4664
Set Up to Use EI .............................................................................. 4664
Attach EI to a Notebook Instance ..................................................... 4669
Endpoints with Elastic Inference ....................................................... 4672
Best practices .................................................................................. 4677
Best practices for deploying models on SageMaker Hosting Services .... 4677
Monitor Security Best Practices ......................................................... 4679
Low latency real-time inference with AWS PrivateLink ...................... 4679
Migrate inference workload from x86 to AWS Graviton ..................... 4681
Troubleshoot deployments ............................................................... 4684
Inference cost optimization best practices ........................................ 4687
Best practices to minimize interruptions during GPU driver upgrades .. 4689
Best practices for endpoint security ............................................... 4693
Supported features .......................................................................... 4695
Resources ....................................................................................... 4701
  Blogs, example notebooks, and additional resources ....................... 4702
  Troubleshooting and reference ...................................................... 4705
  Model Hosting FAQs ..................................................................... 4706
Implement MLOps ........................................................................... 4716
  Why MLOps? .................................................................................. 4716
  Challenges with MLOps ................................................................. 4717
  Benefits of MLOps ....................................................................... 4718
Experiments .................................................................................... 4719
Workflows ...................................................................................... 4719
  Amazon SageMaker Model Building Pipelines ................................. 4720
  Kubernetes Orchestration .............................................................. 4864
  Notebook Jobs ............................................................................. 4958
ML Lineage Tracking ......................................................................... 5021
  Tracking Entities .......................................................................... 5023
  SageMaker-Created Entities .......................................................... 5025
  Manually Create Entities .............................................................. 5027
  Querying Lineage Entities ............................................................. 5032
  Cross-Account Tracking .............................................................. 5041
Catalog models with Model Registry ............................................... 5045
Models, Model Versions, and Model Groups ................................................................. 5045
Collections .................................................................................. 5086
Model Registry FAQ ........................................................................ 5098
Model Deployment ............................................................................. 5100
Model Monitor .................................................................................. 5100
Projects ............................................................................................... 5101
  SageMaker Projects ........................................................................ 5102
  SageMaker Studio Permissions Required to Use Projects .................. 5105
  Create an MLOps Project ............................................................... 5107
Templates ............................................................................................. 5108
View Resources .................................................................................. 5123
Update an MLOps Project ................................................................. 5124
Delete an MLOps Project ................................................................. 5126
Project walkthrough ........................................................................ 5127
  Project Walkthrough Using Third-party Git Repos .......................... 5134
MLOps FAQ ....................................................................................... 5140

**Monitor data and model quality** ............................................................................... 5148
  Model Monitoring ............................................................................. 5149
  How It Works ................................................................................... 5149
    Sample Notebooks ....................................................................... 5152
  Capture data ..................................................................................... 5153
    Capture data from real-time endpoint ............................................ 5153
    Capture data from batch transform job ........................................ 5161
  Monitor data quality ........................................................................ 5165
    Create a Baseline .......................................................................... 5166
    Schedule data quality monitoring jobs ........................................... 5169
    Statistics ....................................................................................... 5170
    CloudWatch Metrics .................................................................. 5172
    Violations ..................................................................................... 5173
  Monitor model quality ..................................................................... 5175
    Create a Model Quality Baseline .................................................. 5176
    Schedule Model Quality Monitoring Jobs ..................................... 5179
    Ingest Ground Truth Labels and Merge Them With Predictions ...... 5181
    Model Quality Metrics ............................................................... 5183
    Model Quality CloudWatch Metrics ............................................. 5187
  Monitor bias drift .............................................................................. 5188
Evaluate, explain, and detect bias in models .................................................. 5269

Evaluate foundation models ................................................................. 5269

What are foundation model evaluations? .............................................. 5270

Get started ............................................................................................... 5272

Foundation model evaluation overview .................................................. 5274

Use a human evaluation ......................................................................... 5285

Use an automatic evaluation ................................................................. 5305
Customize your workflow using the fmeval library .......................................................... 5332
Notebook tutorials .................................................................................................................. 5339
Troubleshooting guide ......................................................................................................... 5356
Explain and detect bias ......................................................................................................... 5360
What is fairness and model explainability? ........................................................................... 5361
SageMaker Clarify Processing Jobs ....................................................................................... 5364
Configure a SageMaker Clarify Processing Job .................................................................... 5366
Run SageMaker Clarify Processing Jobs ............................................................................... 5438
Get Analysis Results ............................................................................................................ 5455
Troubleshoot Jobs ................................................................................................................ 5467
Sample notebooks ................................................................................................................ 5471
Detect Pre-training Data Bias ............................................................................................... 5473
Detect Post-training Data and Model Bias .............................................................................. 5494
Model Explainability ............................................................................................................. 5528
Use Explainability with Autopilot ......................................................................................... 5532

Use governance to document and track model performance .............................................. 5533
Amazon SageMaker Role Manager ......................................................................................... 5533
Amazon SageMaker Model Cards ......................................................................................... 5533
Amazon SageMaker Model Dashboard .................................................................................. 5533
Model Cards ......................................................................................................................... 5534
Prerequisites .......................................................................................................................... 5534
Intended uses of a model ....................................................................................................... 5535
Risk ratings ............................................................................................................................ 5535
Model card JSON schema ...................................................................................................... 5535
Create a model card ............................................................................................................... 5553
Manage model cards .............................................................................................................. 5561
Cross account support ............................................................................................................ 5563
SageMaker APIs ..................................................................................................................... 5568
Model card FAQs .................................................................................................................... 5569
Model Dashboard .................................................................................................................. 5571
Model Dashboard elements ................................................................................................. 5572
View Model Monitor schedules and alerts ............................................................................. 5574
View a model lineage graph .................................................................................................... 5577
View Endpoint Status ............................................................................................................ 5579
Model Dashboard FAQ ......................................................................................................... 5581

Use Docker containers to build models .............................................................................. 5585
<table>
<thead>
<tr>
<th>Scenario</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use cases for using pre-built Docker containers with SageMaker</td>
<td>5586</td>
</tr>
<tr>
<td>Use cases for extending a pre-built Docker container</td>
<td>5587</td>
</tr>
<tr>
<td>Use case for building your own container</td>
<td>5587</td>
</tr>
<tr>
<td>Docker Container Basics</td>
<td>5589</td>
</tr>
<tr>
<td>Use Pre-built SageMaker Docker images</td>
<td>5589</td>
</tr>
<tr>
<td>Prebuilt Deep Learning Images</td>
<td>5590</td>
</tr>
<tr>
<td>Prebuilt Scikit-learn and Spark ML Images</td>
<td>5591</td>
</tr>
<tr>
<td>Deep Graph Networks</td>
<td>5592</td>
</tr>
<tr>
<td>Extend a Pre-built Container</td>
<td>5596</td>
</tr>
<tr>
<td>Adapting your own Docker container to work with SageMaker</td>
<td>5609</td>
</tr>
<tr>
<td>Individual Framework Libraries</td>
<td>5609</td>
</tr>
<tr>
<td>SageMaker Training and Inference Toolkits</td>
<td>5610</td>
</tr>
<tr>
<td>Adapting your own training container</td>
<td>5612</td>
</tr>
<tr>
<td>Adapting Your Own Inference Container</td>
<td>5630</td>
</tr>
<tr>
<td>Create a container with your own algorithms and models</td>
<td>5647</td>
</tr>
<tr>
<td>Use Your Own Training Algorithms</td>
<td>5647</td>
</tr>
<tr>
<td>Use Your Own Inference Code</td>
<td>5665</td>
</tr>
<tr>
<td>Examples and more info</td>
<td>5681</td>
</tr>
<tr>
<td>Setup</td>
<td>5681</td>
</tr>
<tr>
<td>Host models trained in Scikit-learn</td>
<td>5682</td>
</tr>
<tr>
<td>Package TensorFlow and Scikit-learn models for use in SageMaker</td>
<td>5682</td>
</tr>
<tr>
<td>Train and deploy a neural network on SageMaker</td>
<td>5682</td>
</tr>
<tr>
<td>Training using pipe mode</td>
<td>5682</td>
</tr>
<tr>
<td>Bring your own R model</td>
<td>5683</td>
</tr>
<tr>
<td>Extend a pre-built PyTorch container Image</td>
<td>5683</td>
</tr>
<tr>
<td>Train and debug training jobs on a custom container</td>
<td>5683</td>
</tr>
<tr>
<td>Troubleshooting</td>
<td>5683</td>
</tr>
<tr>
<td><strong>Configure security in Amazon SageMaker</strong></td>
<td>5685</td>
</tr>
<tr>
<td>Data Privacy</td>
<td>5686</td>
</tr>
<tr>
<td>Types of information collected</td>
<td>5686</td>
</tr>
<tr>
<td>How to opt out of metadata collection</td>
<td>5686</td>
</tr>
<tr>
<td>Additional information</td>
<td>5688</td>
</tr>
<tr>
<td>Data Protection</td>
<td>5689</td>
</tr>
<tr>
<td>Protect Data at Rest Using Encryption</td>
<td>5690</td>
</tr>
<tr>
<td>Protecting Data in Transit with Encryption</td>
<td>5693</td>
</tr>
</tbody>
</table>
Sell algorithms and packages in the AWS Marketplace ................................................................. 6054

Topics ........................................................................................................................................... 6054
SageMaker Algorithms ...................................................................................................................... 6054
SageMaker Model Packages ............................................................................................................. 6055
Use your own algorithms and models with the AWS Marketplace .............................................. 6055
Create Algorithm and Model Package Resources ........................................................................... 6055
Use Algorithm and Model Package Resources ............................................................................. 6065
Sell Amazon SageMaker Algorithms and Model Packages ......................................................... 6076
Topics ........................................................................................................................................... 6076
Develop Algorithms and Models in Amazon SageMaker .............................................................. 6077
List Your Algorithm or Model Package on AWS Marketplace ..................................................... 6079
Monitor AWS resources provisioned while using Amazon SageMaker ........................................ 6082

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monitoring with CloudWatch</td>
<td>6083</td>
</tr>
<tr>
<td>Endpoint Invocation Metrics</td>
<td>6083</td>
</tr>
<tr>
<td>SageMaker Inference Component Metrics</td>
<td>6087</td>
</tr>
<tr>
<td>Multi-Model Endpoint Metrics</td>
<td>6088</td>
</tr>
<tr>
<td>Jobs and Endpoint Metrics</td>
<td>6091</td>
</tr>
<tr>
<td>Inference Recommender Metrics</td>
<td>6097</td>
</tr>
<tr>
<td>Ground Truth Metrics</td>
<td>6098</td>
</tr>
<tr>
<td>Feature Store Metrics</td>
<td>6101</td>
</tr>
<tr>
<td>Pipelines Metrics</td>
<td>6104</td>
</tr>
<tr>
<td>Logging with CloudWatch</td>
<td>6107</td>
</tr>
<tr>
<td>Log SageMaker API Calls with CloudTrail</td>
<td>6109</td>
</tr>
<tr>
<td>SageMaker Information in CloudTrail</td>
<td>6109</td>
</tr>
<tr>
<td>Operations Performed by Automatic Model Tuning</td>
<td>6110</td>
</tr>
<tr>
<td>Understanding SageMaker Log File Entries</td>
<td>6111</td>
</tr>
<tr>
<td>Monitoring user resource access from Amazon SageMaker Studio Classic</td>
<td>6113</td>
</tr>
<tr>
<td>Prerequisites</td>
<td>6113</td>
</tr>
<tr>
<td>Considerations when using sourceIdentity</td>
<td>6114</td>
</tr>
<tr>
<td>Turn on sourceIdentity</td>
<td>6115</td>
</tr>
<tr>
<td>Turn off sourceIdentity</td>
<td>6116</td>
</tr>
<tr>
<td>Automating with EventBridge</td>
<td>6117</td>
</tr>
<tr>
<td>Model state change</td>
<td>6118</td>
</tr>
<tr>
<td>Training job state change</td>
<td>6118</td>
</tr>
<tr>
<td>HyperParameter tuning job state change</td>
<td>6120</td>
</tr>
<tr>
<td>Transform job state change</td>
<td>6122</td>
</tr>
<tr>
<td>Endpoint state change</td>
<td>6123</td>
</tr>
<tr>
<td>Feature group state change</td>
<td>6124</td>
</tr>
<tr>
<td>Model package state change</td>
<td>6125</td>
</tr>
<tr>
<td>Pipeline execution state change</td>
<td>6127</td>
</tr>
<tr>
<td>Pipeline step state change</td>
<td>6128</td>
</tr>
<tr>
<td>Processing job state change</td>
<td>6129</td>
</tr>
<tr>
<td>SageMaker image state change</td>
<td>6131</td>
</tr>
<tr>
<td>SageMaker image version state change</td>
<td>6131</td>
</tr>
<tr>
<td>Endpoint deployment state change</td>
<td>6133</td>
</tr>
</tbody>
</table>
Model card state change ................................................................. 6136

Reference .................................................................................. 6137

ML Frameworks and Languages ......................................................... 6137
   Apache MXNet ................................................................. 6138
   Apache Spark ................................................................. 6139
   Chainer ................................................................. 6152
   Hugging Face ................................................................. 6153
   PyTorch ................................................................. 6157
   R ........................................................................ 6157
   Scikit-learn ........................................................................ 6161
   SparkML Serving ............................................................... 6163
   TensorFlow ................................................................. 6163
   Triton Inference Server ......................................................... 6164

API Reference ........................................................................ 6166
   Programming Model for Amazon SageMaker ......................................... 6166
   APIs, CLI, and SDKs ............................................................ 6168

SageMaker Distribution Images ...................................................... 6168
   Supported packages and versions .................................................. 6169

SageMaker Document History ....................................................... 6171

AWS Glossary ........................................................................ 6183
What is Amazon SageMaker?

Amazon SageMaker is a fully managed machine learning (ML) service. With SageMaker, data scientists and developers can quickly and confidently build, train, and deploy ML models into a production-ready hosted environment. It provides a UI experience for running ML workflows that makes SageMaker ML tools available across multiple integrated development environments (IDEs).

With SageMaker, you can store and share your data without having to build and manage your own servers. This gives you or your organizations more time to collaboratively build and develop your ML workflow, and do it sooner. SageMaker provides managed ML algorithms to run efficiently against extremely large data in a distributed environment. With built-in support for bring-your-own-algorithms and frameworks, SageMaker offers flexible distributed training options that adjust to your specific workflows. Within a few steps, you can deploy a model into a secure and scalable environment from the SageMaker console.

Topics

- Pricing for Amazon SageMaker
- Are you a first-time user of Amazon SageMaker?
- Overview of machine learning with Amazon SageMaker
- Amazon SageMaker Features

Pricing for Amazon SageMaker

For information about AWS Free Tier limits and the cost of using SageMaker, see Amazon SageMaker Pricing.

Are you a first-time user of Amazon SageMaker?

If you're a first-time user of SageMaker, we recommend that you complete the following:

1. Overview of machine learning with Amazon SageMaker – Get an overview of the machine learning (ML) lifecycle and learn about solutions that are offered. This page explains key concepts and describes the core components involved in building AI solutions with SageMaker.
2. Get started – Learn how to set up and use SageMaker based on your needs.
3. **Use automated ML, no-code, or low-code** – Learn about low-code and no-code ML options that simplify a ML workflow by automating machine learning tasks. These options are helpful ML learning tools because they provide visibility into the code by generating notebooks for each of the automated ML tasks.

4. **Use machine learning environments offered by SageMaker** – Familiarize yourself with the ML environments that you can use to develop your ML workflow, such as information and examples about ready-to-use and custom models.

5. **Explore other topics** – Use the SageMaker Developer Guide's table of contents to explore more topics. For example, you can find information about ML lifecycle stages, in *Overview of machine learning with Amazon SageMaker*, and various solutions that SageMaker offers.

6. **Amazon SageMaker resources** – Refer to the various developer resources that SageMaker offers.

### Overview of machine learning with Amazon SageMaker

This section describes a typical machine learning (ML) workflow and summarizes how to accomplish those tasks with Amazon SageMaker.

In machine learning, you *teach* a computer to make predictions or inferences. First, you use an algorithm and example data to train a model. Then you integrate your model into your application to generate inferences in real time and at scale.

The following diagram illustrates the typical workflow for creating a machine learning model. It includes three stages in a circular flow that we will cover in more detail below: generate example data, train a model, and deploy the model.
The diagram illustrates how to perform the following activities in most typical scenarios:

1. **Generate example data** – To train a model, you need example data. The type of data that you need depends on the business problem that you want the model to solve (the inferences that you want the model to generate). For example, suppose that you want to create a model to predict a number from an input image of a handwritten digit. To train such a model, you need example images of handwritten numbers.

   Data scientists often devote time exploring and preprocessing example data before using it for model training. To preprocess data, you typically do the following:

   a. **Fetch the data** – You might have in-house example data repositories, or you might use datasets that are publicly available. Typically, you pull the dataset or datasets into a single repository.

   b. **Clean the data** – To improve model training, inspect the data and clean it, as needed. For example, if your data has a country name attribute with values United States and US, you can edit the data to be consistent.

   c. **Prepare or transform the data** – To improve performance, you might perform additional data transformations. For example, you might choose to combine attributes. If your model predicts the conditions that require de-icing an aircraft, instead of using temperature and humidity
attributes separately, you can combine those attributes into a new attribute to get a better model.

In SageMaker, you can preprocess example data using SageMaker APIs with the SageMaker Python SDK in an integrated development environment (IDE). With SDK for Python (Boto3) you can fetch, explore, and prepare your data for model training. For information about data preparation, processing, and transforming your data, see Prepare data, Process data, and Create, store, and share features with Amazon SageMaker Feature Store.

2. **Train a model** – Model training includes both training and evaluating the model, as follows:

   • **Training the model** – To train a model, you need an algorithm or a pre-trained base model. The algorithm you choose depends on a number of factors. For a built-in solution, you can use one of the algorithms that SageMaker provides. For a list of algorithms provided by SageMaker and related considerations, see Use Amazon SageMaker Built-in Algorithms or Pre-trained Models. For a UI-based training solution that provides algorithms and models, see SageMaker JumpStart.

   You also need compute resources for training. Depending on the size of your training dataset and how quickly you need the results, you can use resources ranging from a single general-purpose instance to a distributed cluster of GPU instances. For more information, see Train a Model with Amazon SageMaker.

   • **Evaluating the model** – After you train your model, you evaluate it to determine whether the accuracy of the inferences is acceptable. To train and evaluate your model you can use the SageMaker Python SDK to send requests to the model for inferences through one of the available IDEs. For more information about evaluating your model, see Monitor data and model quality.

3. **Deploy the model** – You traditionally re-engineer a model before you integrate it with your application and deploy it. With SageMaker hosting services, you can deploy your model independently, which decouples it from your application code. For more information, see Deploy models for inference.

Machine learning is a continuous cycle. After deploying a model, you monitor the inferences, collect more high-quality data, and evaluate the model to identify drift. You then increase the accuracy of your inferences by updating your training data to include the newly collected high-quality data. As more example data becomes available, you continue retraining your model to increase accuracy.
Amazon SageMaker Features

Amazon SageMaker includes the following features.

Topics

• New features for re:Invent 2023
• Machine learning environments
• Major features

New features for re:Invent 2023

SageMaker includes the following new features for re:Invent 2023.

SageMaker Canvas chat for data prep

SageMaker Canvas chat for data prep helps you create data preparation flows using LLMs.

Code Editor

Code Editor extends Studio so that you can write, test, debug and run your analytics and machine learning code in an environment based on Visual Studio Code - Open Source ("Code-OSS").

Deep learning containers for large model inference

SageMaker has replace the default NCCL kernels with inference optimized kernels to improve GPU utilization and offer differentiating performance against OSS.

Deploy models for real-time inference

SageMaker Inference provides developer experience and user interface abstractions to help you get started more quickly with model deployment.

SageMaker customers can now improve the utilization of their accelerated compute instances by deploying up to thousands of models to a SageMaker endpoint with guaranteed throughput and auto-scaling on a per model basis.

SageMaker Distribution Images

SageMaker Distribution is a collection of Docker images designed for machine learning, data science, and data analytics. The images are available across Studio, Studio Lab, Studio notebooks and Github.
**domain onboarding simplification**

A simplified and guided Amazon SageMaker domain onboarding experience with new capabilities for single users and organization administrators. The capabilities includes direct IAM Identity Center integration, fine-grained access policy management, seamless SageMaker apps management and configurations, and VPC and storage configuration.

**Amazon S3 Express One Zone**

Amazon S3 Express One Zone is new storage class that provides single-digit millisecond access for the most latency-sensitive applications. Amazon S3 Express One Zone allows customers to collocate their object storage and compute resources in a single AWS Availability Zone, optimizing both compute performance and costs with increased data processing speed.

**Foundation model evaluations (FMEval)**

Foundation model evaluations (FMEval) helps you quantify the risk of providing inaccurate, toxic or biased content with your language model so that you can choose the best one for your use case. Bring your own custom dataset or use a built-in to evaluate any language model. FMEval is integrated with tens of text-based foundation models in SageMaker JumpStart or bring your own. You can also create customized evaluations using the FMEval library.

**SageMaker HyperPod**

SageMaker HyperPod is a capability of SageMaker that provides an always-on machine learning environment on resilient clusters that you can run any machine learning workloads for developing large machine learning models such as large language models (LLMs) and diffusion models.

**JupyterAI**

Jupyter AI and Code Whisperer have been included to SageMaker Distribution. With this update, users of Studio or Code Editor can easily use generative AI from their notebooks and take advantage of Code Whisperer's code completion feature.

**JupyterLab in Studio**

JupyterLab in Studio improves latency and reliability for Studio Notebooks

**SageMaker Notebook Jobs**

SageMaker Notebook Jobs provides SDK support for notebook jobs so you can schedule your notebook jobs programmatically.
SageMaker Pipelines

SageMaker Pipelines provides you the option to convert your local machine learning code to a SageMaker Pipeline step, from which you can create and run a pipeline.

SageMaker smart sifting

SageMaker smart sifting is a capability of SageMaker Training that improves the efficiency of your training datasets and reduces total training time and cost.

SageMaker Studio

Studio is the latest web-based experience for running ML workflows. Studio offers a suite of IDEs, including Code Editor, a new Jupyterlab application, RStudio, and Studio Classic.

Machine learning environments

SageMaker includes the following machine learning environments.

SageMaker geospatial capabilities

Build, train, and deploy ML models using geospatial data.

SageMaker Canvas

An auto ML service that gives people with no coding experience the ability to build models and make predictions with them.

SageMaker Studio

An integrated machine learning environment where you can build, train, deploy, and analyze your models all in the same application.

SageMaker Studio Lab

A free service that gives customers access to AWS compute resources in an environment based on open-source JupyterLab.

RStudio on Amazon SageMaker

An integrated development environment for R, with a console, syntax-highlighting editor that supports direct code execution, and tools for plotting, history, debugging and workspace management.
Major features

SageMaker includes the following major features in alphabetical order excluding any SageMaker prefix.

**Amazon Augmented AI**

Build the workflows required for human review of ML predictions. Amazon A2I brings human review to all developers, removing the undifferentiated heavy lifting associated with building human review systems or managing large numbers of human reviewers.

**AutoML step**

Create an AutoML job to automatically train a model in SageMaker Pipelines.

**SageMaker Autopilot**

Users without machine learning knowledge can quickly build classification and regression models.

**Batch Transform**

Preprocess datasets, run inference when you don't need a persistent endpoint, and associate input records with inferences to assist the interpretation of results.

**SageMaker Clarify**

Improve your machine learning models by detecting potential bias and help explain the predictions that models make.

**Collaboration with shared spaces**

A shared space consists of a shared JupyterServer application and a shared directory. All user profiles in a Amazon SageMaker domain have access to all shared spaces in the domain.

**SageMaker Data Wrangler**

Import, analyze, prepare, and featurize data in SageMaker Studio. You can integrate Data Wrangler into your machine learning workflows to simplify and streamline data pre-processing and feature engineering using little to no coding. You can also add your own Python scripts and transformations to customize your data prep workflow.

**Data Wrangler data preparation widget**

Interact with your data, get visualizations, explore actionable insights, and fix data quality issues.
**SageMaker Debugger**

Inspect training parameters and data throughout the training process. Automatically detect and alert users to commonly occurring errors such as parameter values getting too large or small.

**SageMaker Edge Manager**

Optimize custom models for edge devices, create and manage fleets and run models with an efficient runtime.

**SageMaker Elastic Inference**

Speed up the throughput and decrease the latency of getting real-time inferences.

**SageMaker Experiments**

Experiment management and tracking. You can use the tracked data to reconstruct an experiment, incrementally build on experiments conducted by peers, and trace model lineage for compliance and audit verifications.

**SageMaker Feature Store**

A centralized store for features and associated metadata so features can be easily discovered and reused. You can create two types of stores, an Online or Offline store. The Online Store can be used for low latency, real-time inference use cases and the Offline Store can be used for training and batch inference.

**SageMaker Ground Truth**

High-quality training datasets by using workers along with machine learning to create labeled datasets.

**SageMaker Ground Truth Plus**

A turnkey data labeling feature to create high-quality training datasets without having to build labeling applications and manage the labeling workforce on your own.

**SageMaker Inference Recommender**

Get recommendations on inference instance types and configurations (e.g. instance count, container parameters and model optimizations) to use your ML models and workloads.

**Inference shadow tests**

Evaluate any changes to your model-serving infrastructure by comparing its performance against the currently deployed infrastructure.
**SageMaker JumpStart**

Learn about SageMaker features and capabilities through curated 1-click solutions, example notebooks, and pretrained models that you can deploy. You can also fine-tune the models and deploy them.

**SageMaker ML Lineage Tracking**

Track the lineage of machine learning workflows.

**SageMaker Model Building Pipelines**

Create and manage machine learning pipelines integrated directly with SageMaker jobs.

**SageMaker Model Cards**

Document information about your ML models in a single place for streamlined governance and reporting throughout the ML lifecycle.

**SageMaker Model Dashboard**

A pre-built, visual overview of all the models in your account. Model Dashboard integrates information from SageMaker Model Monitor, transform jobs, endpoints, lineage tracking, and CloudWatch so you can access high-level model information and track model performance in one unified view.

**SageMaker Model Monitor**

Monitor and analyze models in production (endpoints) to detect data drift and deviations in model quality.

**SageMaker Model Registry**

Versioning, artifact and lineage tracking, approval workflow, and cross account support for deployment of your machine learning models.

**SageMaker Neo**

Train machine learning models once, then run anywhere in the cloud and at the edge.

**Notebook-based Workflows**

Run your SageMaker Studio notebook as a non-interactive, scheduled job.

**Preprocessing**

Analyze and preprocess data, tackle feature engineering, and evaluate models.
SageMaker Projects

Create end-to-end ML solutions with CI/CD by using SageMaker projects.

Reinforcement Learning

Maximize the long-term reward that an agent receives as a result of its actions.

SageMaker Role Manager

Administrators can define least-privilege permissions for common ML activities using custom and preconfigured persona-based IAM roles.

SageMaker Serverless Endpoints

A serverless endpoint option for hosting your ML model. Automatically scales in capacity to serve your endpoint traffic. Removes the need to select instance types or manage scaling policies on an endpoint.

Studio Classic Git extension

A Git extension to enter the URL of a Git repository, clone it into your environment, push changes, and view commit history.

SageMaker Studio Notebooks

The next generation of SageMaker notebooks that include AWS IAM Identity Center (IAM Identity Center) integration, fast start-up times, and single-click sharing.

SageMaker Studio Notebooks and Amazon EMR

Easily discover, connect to, create, terminate and manage Amazon EMR clusters in single account and cross account configurations directly from SageMaker Studio.

SageMaker Training Compiler

Train deep learning models faster on scalable GPU instances managed by SageMaker.
Get started

If you want to set up Amazon SageMaker for yourself as quickly as possible with no customization, follow the instructions in Quick setup for single users.

- Open the SageMaker console and access the SageMaker applications from the left navigation pane.
- Using the quick setup gets you set up with a Amazon SageMaker domain without the need to learn the details of a domain. For information about the domain default settings, see Default settings.
- If you do not intend to customize your domain any further, you can skip this Get started chapter and begin exploring SageMaker ML environment products at Use machine learning environments offered by SageMaker.

If you want to set up Amazon SageMaker for many users or an organization with customization, use one of the following step-by-step guides with contextual help for advanced and enterprise Machine Learning (ML) administrators. Administrators of organizations can onboard their users or groups through authentication, using AWS Identity and Access Management (IAM) or AWS IAM Identity Center, define scoped-down access policies, manage SageMaker applications and configurations, and set up a VPC configuration.

- To set up a custom domain for users using IAM, see Custom onboarding to Amazon SageMaker domain using IAM.
- To set up a custom domain for groups using authentication in IAM Identity Center, see Custom onboarding to Amazon SageMaker domain using IAM Identity Center.

Note

If you intend to use the Amazon SageMaker Studio Lab ML environment or if an AWS account and Amazon SageMaker domain has already been set up for you, you can skip this Get started chapter and begin exploring SageMaker ML environment products at Use machine learning environments offered by SageMaker.

If a sign-in URL is provided to you, you can set up by following the instructions in Access the domain after onboarding.
After the domain is set up, the administrative user can [View and edit domains](#) and the users can access the ML environments using the sign-in URL or opening the [SageMaker console](#) and choosing the environment from the left navigation pane.

**Topics**
- [Set Up Amazon SageMaker Prerequisites](#)
- [Amazon SageMaker domain overview](#)
- [Supported Regions and Quotas](#)

### Set Up Amazon SageMaker Prerequisites

To get set up with Amazon SageMaker:

- You will need to create an Amazon Web Services (AWS) account to get access to all of the AWS services and resources for the account.
- To adhere to the [Security best practices in IAM](#), create an administrative user to provision AWS resources, like creating and updating a Amazon SageMaker domain for users.
- Set up your AWS Command Line Interface (AWS CLI) to manage your AWS services and resources for the account.

**Topics**
- [Create an AWS Account](#)
- [Create an Administrative User and Group](#)
- [AWS CLI Prerequisites](#)

### Create an AWS Account

If you already have an AWS account, skip this step.

When you sign up for AWS, your AWS account is automatically signed up for all AWS services, including SageMaker. You are charged only for the services that you use. For details on pricing, see [AWS pricing](#) and [Amazon SageMaker pricing](#).

**To create an AWS account**

2. Follow the online instructions.

Part of the sign-up procedure involves receiving a phone call and entering a verification code on the phone keypad.

When you sign up for an AWS account, an *AWS account root user* is created. The root user has access to all AWS services and resources in the account. As a security best practice, assign administrative access to an administrative user, and use only the root user to perform tasks that require root user access.

Write down your AWS account ID because you need it for the next task.

### Create an Administrative User and Group

When you create an AWS account, you get a single sign-in identity that has complete access to all of the AWS services and resources in the account. This identity is called the AWS account *root user*. Signing in to the AWS console using the email address and password that you used to create the account gives you complete access to all of the AWS resources in your account.

We strongly recommend that you *do not* use the root user account for everyday tasks, even the administrative ones. Instead, adhere to the Security best practices in IAM, and create an administrative user to provision AWS resources, like creating and updating a Amazon SageMaker domain for users within the domain. Then securely lock away the root user credentials and use them to perform only a few account and service management tasks.

**To create an administrative user**

1. Create an administrative user in your AWS account. For instructions, see [Create an administrative user](#) in the *IAM User Guide*.

   **Note**

   We assume that you use administrator user credentials for the exercises and procedures in this guide. If you choose to create and use another user, grant that user minimum permissions. For more information, see Authenticating with Identities.

2. Ensure that your administrator user has the [AmazonSageMakerFullAccess](#) policy, as well as a policy with the following content needed to create a SageMaker domain. For more information about creating IAM policies, see Creating IAM policies.
AWS CLI Prerequisites

The following prerequisites are required to manage your domain and other AWS services and resources using the AWS CLI.

- Update the AWS CLI by following the steps in Installing the current AWS CLI Version.
- From your local machine, run aws configure and provide your AWS credentials.

If you do not already have AWS credentials, see Managing access keys for IAM users for information about creating and managing your access keys.

For information about AWS credentials, see AWS security credentials.
Amazon SageMaker domain overview

To have access to most Amazon SageMaker environments and resources, you must complete the Amazon SageMaker domain onboarding process using the SageMaker console or the AWS CLI. For a guide describing how to get started using SageMaker based on how you wish to access SageMaker, and if necessary how to set up a domain, see Get started.

An Amazon SageMaker domain consists of the following:

- An associated Amazon Elastic File System (Amazon EFS) volume
- A list of authorized users
- A variety of security, application, policy, and Amazon Virtual Private Cloud (Amazon VPC) configurations

The following diagram provides an overview of private apps and shared spaces within each domain.

When onboarding, you can choose to use either AWS Identity and Access Management (IAM) or AWS IAM Identity Center for authentication methods. When you use IAM authentication, you can choose either the Set up for single users or the Set up for organizations procedure. RStudio setup is only available when using the Set up for organizations procedure.
Note
If you onboard using IAM authentication and want to switch to authentication using IAM Identity Center later, you must delete the domain that you created. Then, you need to manually re-import all notebooks and other user data that you created. For more information, see Delete an Amazon SageMaker domain.

The simplest way to create an Amazon SageMaker domain is to follow the Set up for single users procedure from the SageMaker console. Set up for single users uses default settings. These settings include shareable notebooks and public internet access. For information on the default settings, see Default settings.

For more control, including the option of using authentication using IAM Identity Center and RStudio, use the Set up for organizations procedures.

Authentication using IAM Identity Center

To use authentication using IAM Identity Center with domain, you must onboard to an organization in AWS Organizations.

Note
The AWS Organizations account must be in the same AWS Region as the domain.

Authentication using IAM Identity Center provides the following benefits over IAM authentication:

• Members given access to domain have a unique sign-in URL that directly opens the domain, and they sign in with their IAM Identity Center credentials. When you use IAM authentication, you must sign in through the SageMaker console.

For more information on how to access your domain with IAM Identity Center authentication, see Access the domain after onboarding.

• Organizations manage their members in IAM Identity Center instead of the domain. You can assign multiple members access to the domain at the same time. When you use IAM authentication, you must add and manage members manually, one at a time, using the domain control panel.
Topics

- Amazon SageMaker domain
- Quick onboard to Amazon SageMaker domain
- Custom onboarding to Amazon SageMaker domain using IAM Identity Center
- Custom onboarding to Amazon SageMaker domain using IAM
- Choose an Amazon VPC

Amazon SageMaker domain

Amazon SageMaker domain supports SageMaker machine learning (ML) environments. A SageMaker domain is composed of the following entities. For onboarding steps to create a domain, see Amazon SageMaker domain overview.

- **domain**: An Amazon SageMaker domain consists of an associated Amazon Elastic File System (Amazon EFS) volume; a list of authorized users; and a variety of security, application, policy, and Amazon Virtual Private Cloud (Amazon VPC) configurations. Users within a domain can share notebook files and other artifacts with each other. An account can have multiple domains. For more information about multiple domains, see Multiple domains overview.

- **UserProfile**: A user profile represents a single user within a domain. It is the main way to reference a user for the purposes of sharing, reporting, and other user-oriented features. This entity is created when a user onboards to the Amazon SageMaker domain. For more information about user profiles, see Domain user profiles.

- **shared space**: A shared space consists of a shared JupyterServer application and shared directory. All users within the domain have access to the shared space. All user profiles in a domain have access to all shared spaces in the domain. For more information about shared spaces, see Collaborate with shared spaces.

- **App**: An app represents an application that supports the reading and execution experience of the user’s notebooks, terminals, and consoles. The type of app can be JupyterServer, KernelGateway, RStudioServerPro, or RSession. A user may have multiple apps active simultaneously.

The following tables describe the status values for the domain, UserProfile, shared space, and App entities. Where applicable, they also give troubleshooting steps.
### domain status values

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pending</td>
<td>Ongoing creation of domain.</td>
</tr>
<tr>
<td>InService</td>
<td>Successful creation of domain.</td>
</tr>
<tr>
<td>Updating</td>
<td>Ongoing update of domain.</td>
</tr>
<tr>
<td>Deleting</td>
<td>Ongoing deletion of domain.</td>
</tr>
<tr>
<td>Failed</td>
<td>Unsuccessful creation of domain. Call the DescribeDomain API to see the failure reason for domain creation. Delete the failed domain and recreate the domain after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Update_Failed</td>
<td>Unsuccessful update of domain. Call the DescribeDomain API to see the failure reason for domain update. Call the UpdateDomain API after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Delete_Failed</td>
<td>Unsuccessful deletion of domain. Call the DescribeDomain API to see the failure reason for domain deletion. Because deletion failed, you might have some resources that are still running, but you cannot use or update the domain. Call the DeleteDomain API again after fixing the error mentioned in FailureReason.</td>
</tr>
</tbody>
</table>

### UserProfile status values

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pending</td>
<td>Ongoing creation of UserProfile.</td>
</tr>
</tbody>
</table>
### Value Description

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>InService</td>
<td>Successful creation of UserProfile.</td>
</tr>
<tr>
<td>Updating</td>
<td>Ongoing update of UserProfile.</td>
</tr>
<tr>
<td>Deleting</td>
<td>Ongoing deletion of UserProfile.</td>
</tr>
<tr>
<td>Failed</td>
<td>Unsuccessful creation of UserProfile. Call the DescribeUserProfile API to see the failure reason for UserProfile creation. Delete the failed UserProfile and recreate it after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Update_Failed</td>
<td>Unsuccessful update of UserProfile. Call the DescribeUserProfile API to see the failure reason for UserProfile update. Call the UpdateUserProfile API again after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Delete_Failed</td>
<td>Unsuccessful deletion of UserProfile. Call the DescribeUserProfile API to see the failure reason for UserProfile deletion. Because deletion failed, you might have some resources that are still running, but you cannot use or update the UserProfile. Call the DeleteUserProfile API again after fixing the error mentioned in FailureReason.</td>
</tr>
</tbody>
</table>

**shared space status values**

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pending</td>
<td>Ongoing creation of shared space.</td>
</tr>
<tr>
<td>InService</td>
<td>Successful creation of shared space.</td>
</tr>
<tr>
<td>Value</td>
<td>Description</td>
</tr>
<tr>
<td>------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Deleting</td>
<td>Ongoing deletion of shared space.</td>
</tr>
<tr>
<td>Failed</td>
<td>Unsuccessful creation of shared space. Call the DescribeSpace API to see the failure reason for shared space creation. Delete the failed shared space and recreate it after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Update_Failed</td>
<td>Unsuccessful update of shared space. Call the DescribeSpace API to see the failure reason for shared space update. Call the UpdateSpace API again after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Delete_Failed</td>
<td>Unsuccessful deletion of shared space. Call the DescribeSpace API to see the failure reason for shared space deletion. Because deletion failed, you might have some resources that are still running, but you cannot use or update the shared space. Call the DeleteSpace API again after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Deleted</td>
<td>Successful deletion of shared space.</td>
</tr>
</tbody>
</table>

**App status values**

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pending</td>
<td>Ongoing creation of App.</td>
</tr>
<tr>
<td>InService</td>
<td>Successful creation of App.</td>
</tr>
<tr>
<td>Deleting</td>
<td>Ongoing deletion of App.</td>
</tr>
<tr>
<td>Value</td>
<td>Description</td>
</tr>
<tr>
<td>---------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Failed</td>
<td>Unsuccessful creation of App. Call the DescribeApp API to see the failure reason for App creation. Call the CreateApp API again after fixing the error mentioned in FailureReason.</td>
</tr>
<tr>
<td>Deleted</td>
<td>Successful deletion of App.</td>
</tr>
</tbody>
</table>

**Maintenance of applications**

At least once every 90 days, SageMaker performs security and performance updates to the underlying software for Amazon SageMaker Studio Classic JupyterServer and KernelGateway, SageMaker Canvas, and Amazon SageMaker Data Wrangler applications. Some maintenance items, such as operating system upgrades, require that SageMaker takes your application offline for a short time during the maintenance window. Because this maintenance takes the application offline, you cannot perform any operations while the underlying software is being updated. When the maintenance activity is in progress, the state of the application transitions from **InService** to **Pending**. When maintenance is complete, the status of the application transitions back to **InService**. If patching fails, then the status of the application becomes **Failed**. If an application is in the **Failed** state, we recommend creating a new application of the same type. For information about creating Studio Classic applications, see [Shut Down and Update SageMaker Studio Classic and Studio Classic Apps](#). For information about creating SageMaker Canvas applications, see [Manage applications](#).

For more information, contact [https://aws.amazon.com/premiumsupport/](https://aws.amazon.com/premiumsupport/).

**Topics**

- [Prerequisites](#)
- [Multiple domains overview](#)
- [Domain resource isolation](#)
- [Setting defaults for a domain](#)
- [Attaching a custom file system to a domain or user profile](#)
- [Environment](#)
- [View and edit domains](#)
• Delete an Amazon SageMaker domain

• Domain user profiles

• IAM Identity Center Groups in a domain

Prerequisites

To use the features available in an Amazon SageMaker domain, you must first onboard to a domain. For more information, see Onboard to Amazon SageMaker Domain.

If you are interacting with your domain using the AWS CLI, you must also complete the following prerequisites.

• Update the AWS CLI by following the steps in Installing the current AWS CLI Version.
• From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.

Multiple domains overview

Amazon SageMaker supports the creation of multiple Amazon SageMaker domains in a single AWS Region for each account. Additional domains in a Region have the same features and capabilities as the first domain in a Region. Each domain can have distinct domain settings. The same user profile cannot be added to multiple domains in a single Region within the same account. For more information about domain limits, see Amazon SageMaker endpoints and quotas.

Topics

• Automatic tag propagation
• Domain resource display filtering
• Backfilling domain tags

Automatic tag propagation

By default, any SageMaker resources that support tagging and are created from within the Studio Classic UI after 11/30/2022 are automatically tagged with a domain ARN tag. The domain ARN tag is based on the domain ID of the domain that the resource is created in. The following list describes the only SageMaker resources that do not support automatic tag propagation, as well as the impacted API calls where the tag is not returned because it was not automatically set.
You can also use these tags for cost allocation using AWS Billing and Cost Management. For more information, see Using AWS cost allocation tags.

**Note**

All SageMaker List APIs do not support tag-based resource isolation. The default app, which manages the Studio UI, is not automatically tagged.

<table>
<thead>
<tr>
<th>SageMaker resource</th>
<th>Affected API calls</th>
</tr>
</thead>
<tbody>
<tr>
<td>ImageVersionArn</td>
<td>• describe-image-version</td>
</tr>
<tr>
<td></td>
<td>• update-image-version</td>
</tr>
<tr>
<td></td>
<td>• delete-image-version</td>
</tr>
<tr>
<td>ModelCardExportJobArn</td>
<td>describe-model-card-export-job</td>
</tr>
<tr>
<td>ModelPackageArn</td>
<td>describe-model-package</td>
</tr>
</tbody>
</table>

### Domain resource display filtering

By default, SageMaker filters resources displayed within Studio Classic at the domain level. SageMaker implements resource filtration in Studio Classic using the `sagemaker:domain-arn` tag attached to SageMaker resources.

**Note**

This only applies to the Studio Classic UI. SageMaker does not support resource filtering using the AWS CLI by default.

Using this resource filtration, SageMaker only displays SageMaker resources created in the domain, as well as SageMaker resources that do not have a `sagemaker:domain-arn` tag associated to them. These untagged resources are either created outside the context of a domain or were created before 11/30/2022. You can add a tag to these untagged resources for better filtration by following the steps in Backfilling domain tags. Resources created in other domains are automatically filtered out.
All resources created in shared spaces are automatically filtered to that space.

**Backfilling domain tags**

If you have created resources in a domain before 11/30/2022, those resources are not automatically tagged with the domain Amazon Resource Name (ARN) tag.

To accurately attribute resources to their respective domain, you must add the domain tag to existing resources using the AWS CLI, as follows.

1. Map all existing SageMaker resources and their respective ARNs to the domains that exist in your account.
2. Run the following command from your local machine to tag the resource with the ARN of the resource's respective domain. This must be repeated for every SageMaker resource in your account.

   ```bash
   aws resourcegroupstaggingapi tag-resources \\
   --resource-arn-list arn:aws:sagemaker:region:account-id:space/domain-id/space-name \\
   ```

**Domain resource isolation**

You can isolate resources between each of the domains in your account and Region using an AWS Identity and Access Management policy. With resource isolation, SageMaker resources, such as models, experiments, training jobs, and pipelines created in one domain, cannot be accessed from other domains. The following topic shows how to create a new IAM policy that limits access to resources in the domain to user profiles with the domain tag, as well as how to attach this policy to the IAM execution role of the domain. You must repeat this process for each domain in your account. For more information about domain tags and backfilling these tags, see [Multiple domains overview](#).

**Console**

The following section shows how to create a new IAM policy that limits access to resources in the domain to user profiles with the domain tag, as well as how to attach this policy to the IAM execution role of the domain, from the Amazon SageMaker console.
This policy only works in domains that use Amazon SageMaker Studio Classic as the default experience.

1. Create an IAM policy named StudioDomainResourceIsolationPolicy-*domain-id* with the following JSON policy document by completing the steps in Creating IAM policies (console).

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "CreateAPIs",
            "Effect": "Allow",
            "Action": "sagemaker:Create*",
            "NotResource": [
                "arn:aws:sagemaker:*:*:domain/*",
                "arn:aws:sagemaker:*:*:user-profile/*",
                "arn:aws:sagemaker:*:*:space/*"
            ]
        },
        {
            "Sid": "ResourceAccessRequireDomainTag",
            "Effect": "Allow",
            "Action": [
                "sagemaker:Update*",
                "sagemaker:Delete*",
                "sagemaker:Describe*"
            ],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "aws:ResourceTag/sagemaker:domain-arn": "domain-arn"
                }
            }
        },
        {
            "Sid": "AllowActionsThatDontSupportTagging",
            "Effect": "Allow",
            "Action": [
```

2. Attach the StudioDomainResourceIsolationPolicy-\textit{domain-id} policy to the domain's execution role by completing the steps in Modifying a role (console).

\section*{AWS CLI}

The following section shows how to create a new IAM policy that limits access to resources in the domain to user profiles with the domain tag, as well as how to attach this policy to the execution role of the domain, from the AWS CLI.

\begin{note}

This policy only works in domains that use Amazon SageMaker Studio Classic as the default experience.

\end{note}

1. Create a file named StudioDomainResourceIsolationPolicy-\textit{domain-id} with the following content from your local machine.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "CreateAPIs",
            "Action": "sagemaker:CreateImageVersion",
            "Resource": "arn:aws:sagemaker:*:*:image-version",
            "Condition": {
                "StringEquals": {
                    "sagemaker:imageVersionName": "default"
                }
            }
        },
        {
            "Sid": "DeleteDefaultApp",
            "Effect": "Allow",
            "Action": "sagemaker:DeleteApp",
            "Resource": "arn:aws:sagemaker:*:*:app/\textit{domain-id}/*/jupyterserver/default"
        }
    ]
}
```
"Effect": "Allow",
"Action": "sagemaker:Create*",
"NotResource": [
    "arn:aws:sagemaker:*:*:domain/**",
    "arn:aws:sagemaker:*:*:user-profile/**",
    "arn:aws:sagemaker:*:*:space/**"
],
},
{
    "Sid": "ResourceAccessRequireDomainTag",
    "Effect": "Allow",
    "Action": [
        "sagemaker:Update*",
        "sagemaker:Delete*",
        "sagemaker:Describe*"
    ],
    "Resource": "*",
    "Condition": {
        "StringEquals": {
            "aws:ResourceTag/sagemaker:domain-arn": "domain-arn"
        }
    }
},
{
    "Sid": "AllowActionsThatDontSupportTagging",
    "Effect": "Allow",
    "Action": [
        "sagemaker:DescribeImageVersion",
        "sagemaker:UpdateImageVersion",
        "sagemaker:DeleteImageVersion",
        "sagemaker:DescribeModelCardExportJob",
        "sagemaker:DescribeAction"
    ],
    "Resource": "*"
},
{
    "Sid": "DeleteDefaultApp",
    "Effect": "Allow",
    "Action": "sagemaker:DeleteApp",
    "Resource": "arn:aws:sagemaker:*:*:app/domain-id/*/jupyterserver/default"
}

   ```bash
   ```

3. Attach the newly created policy to a new or existing role that is used as the domain's execution role.

   ```bash
   aws iam attach-role-policy --policy-arn arn:aws:iam::account-id:policy/StudioDomainResourceIsolationPolicy-\textit{domain-id} --role-name domain-execution-role
   ```

### Setting defaults for a domain

With SageMaker, you can set default settings for your resources at the Amazon SageMaker domain level. These default settings are used in the creation of resources within the domain. The following sections list default settings for domain and give information on using context keys when setting defaults.

#### Topics

- Domain default settings
- Context keys

#### Domain default settings

You can set the following defaults when creating or updating a domain. Values passed at the user profile and shared space level override defaults set at the domain level.

- DefaultUserSettings
- DefaultSpaceSettings
Note

DefaultSpaceSettings only supports the use of JupyterLab 3 image ARNs for SageMakerImageArn. For more information, see JupyterLab Versioning.

```json
"DefaultSpaceSettings": {
    "ExecutionRole": "string",
    "JupyterServerAppSettings": {
        "DefaultResourceSpec": {
            "InstanceType": "string",
            "LifecycleConfigArn": "string",
            "SageMakerImageArn": "string",
            "SageMakerImageVersionArn": "string"
        },
        "LifecycleConfigArns": [ "string" ]
    },
    "KernelGatewayAppSettings": {
        "CustomImages": [
            {
                "AppImageConfigName": "string",
                "ImageName": "string",
                "ImageVersionNumber": number
            }
        ],
        "DefaultResourceSpec": {
            "InstanceType": "string",
            "LifecycleConfigArn": "string",
            "SageMakerImageArn": "string",
            "SageMakerImageVersionArn": "string"
        },
        "LifecycleConfigArns": [ "string" ]
    },
    "SecurityGroups": [ "string" ]
}
```
Context keys

You can add context keys to the IAM policy that creates a domain. This restricts the values that users can pass for those fields. The following list shows the context keys that domain supports and where they’re implemented.

- **sagemaker:ImageArns**
  


- **sagemaker:VpcSecurityGroupIds**
  
  **Implemented as part of DefaultUserSettings:** `SecurityGroups` in `DefaultUserSettings`.

  **Implemented as part of DefaultSpaceSettings:** `SecurityGroups` in `DefaultSpaceSettings`.

- **sagemaker:DomainSharingOutputKmsKey**

  **Implemented as part of DefaultUserSettings:** `S3KmsKeyId` in `DefaultSpaceSettings.SharingSettings`.

You cannot restrict users to passing incompatible values when using context keys for the defaults. For example, the values for `SageMakerImageArn` set as part of `DefaultUserSettings` and `DefaultSpaceSettings` must be compatible. You cannot set the following incompatible default values. For more information about the available JupyterLab version ARNs, see [Setting a default JupyterLab version](#).

- Only a JupyterLab version 1 ARN can be used for the `SageMakerImageArn` value in `DefaultUserSettings`

- Only a JupyterLab version 3 ARN can be used for the `SageMakerImageArn` value in `DefaultSpaceSettings`
Attaching a custom file system to a domain or user profile

When you create a domain, Amazon SageMaker automatically associates it with an Amazon Elastic File System (Amazon EFS) volume that SageMaker creates for you. You also have the option to associate the domain with a custom Amazon EFS file system that you've created in your AWS account. This file system is available to any users who belong to the domain when they use Amazon SageMaker Studio. Users can attach the file system to any space that they create for the supported applications: JupyterLab and Code Editor. Then, after running the space and starting the application, they can access any data, code, or other artifacts that the file system contains.

If you don't want to permit all of the users for a domain to access the file system, you can attach it to a specific user profile instead. If you do that, the file system is available only in spaces that the associated user creates.

You can attach a custom file system by using the Amazon SageMaker API, the AWS SDKs, or the AWS CLI. You can't attach a custom file system by using the SageMaker console.

Prerequisites

Before you can attach a custom Amazon EFS file system to a domain, you must meet the following requirements:

- You have an Amazon EFS file system in your AWS account. For the steps to create one, see Create your Amazon EFS file system in the Amazon Elastic File System User Guide.

- Before Studio can access your file system, it must have a mount target in each of the subnets that you associate with the domain. For more information about assigning mount targets to subnets, see Creating and managing mount targets and security groups in the Amazon Elastic File System User Guide.

- For each mount target, you must add the security group that Amazon SageMaker created in your AWS account when you created the domain. The security group name has the format security-group-for-inbound-nfs- domain-id.

- Your IAM permissions must allow you to use the elasticfilesystem:DescribeMountTargets action. For more information about this action, see Actions, resources, and condition keys for Amazon Elastic File System in the Service Authorization Reference.
Attaching a custom file system with the AWS CLI

To attach a custom file system to a domain or user profile with the AWS CLI, you pass a `CustomFileSystemConfigs` definition when you use any of the following commands:

- `create-domain`
- `update-domain`
- `create-user-profile`
- `update-user-profile`

Example create-domain command with a custom file system

The following example attaches a file system to a new domain.

```
aws sagemaker create-domain --domain-name domain-name \ 
--vpc-id vpc-id --subnet-ids subnet-ids --auth-mode IAM \ 
--default-user-settings file://default-user-settings.json \ 
--default-space-settings "ExecutionRole=execution-role-arn"
```

In this example, the file `default-user-settings.json` has the following settings, which include the `CustomPosixUserConfig` and `CustomFileSystemConfigs` keys.

```
{
    "ExecutionRole": "execution-role-arn",
    "CustomPosixUserConfig": {
        "Uid": UID,
        "Gid": GID
    },
    "CustomFileSystemConfigs": [
        {
            "EFSFileSystemConfig": {
                "FileSystemId": "file-system-id",
                "FileSystemPath": "/
            }
        }
    ]
}
```
This example configuration has the following keys:

**ExecutionRole**

The default execution role for the users of the domain.

**CustomPosixUserConfig**

The default POSIX identities that are used for file system operations. You can use these settings to apply your existing POSIX permission structure to the user profiles that access the custom file system. At a POSIX permissions level, you can control which users can access the file system and which files or data they can access.

You can also apply CustomPosixUserConfig settings when you create a user profile by using the `create-user-profile` command. The settings that you apply to a user profile override those that you apply to the associated domain.

---

**Note**

You can apply CustomPosixUserConfig settings when you use the `create-domain` and `create-user-profile` commands. However, you can't apply these settings when you do the following:

- Use the `update-domain` command for a domain that is already associated with any user profiles. You can apply these settings only to domains that have no user profiles.

- Use the `update-user-profile` command. To apply these settings to a profile that you've already created, delete the profile, and create a new one that has the updated settings.

---

**Uid**

The POSIX user ID. The default is 200001.

**Gid**

The POSIX group ID. The default is 1001.

**CustomFileSystemConfigs**

Settings for custom file systems (only Amazon EFS file systems are supported).
You can also apply CustomFileSystemConfigs settings to a user profile when you use the `create-user-profile` or `update-user-profile` commands. The user profile will have access to those file systems as well as any that you attach to their domain.

**EFSFileSystemConfig**

Settings for custom Amazon EFS file systems.

**FileSystemId**

The ID of your Amazon EFS file system.

**FileSystemPath**

The path to the file system directory that is accessible to the domain users in their spaces in Studio. Permitted users can access only this directory and below. The default path is the file system root: `/`.

SageMaker creates a symbolic link at the following path: `/home/sagemaker-user/custom-file-systems/file-system-type/file-system-id`. With this, the domain users can navigate to the custom file system from within their home directory, `/home/sagemaker-user`.

After you attach a custom file system to a domain, the domain users can attach the file system to a space when they use the `create-space` command.

**Example create-space command with a custom file system**

The following example attaches a file system to a new space.

```
aws sagemaker create-space \
--space-name space-name \
--domain-id domain-id \
--ownership-settings "OwnerUserProfileName=user-profile-name" \
--space-sharing-settings "SharingType=Private" \
--space-settings file://space-settings.json
```

In this example, the file `space-settings.json` has the following settings, which include the CustomFileSystems configuration with the FileSystemId key.

```json
{
    "AppType": "JupyterLab",
    "JupyterLabAppSettings":
    {
```
Environment

This page gives information about modifications to the Amazon SageMaker domain environment. This includes custom images, lifecycle configurations, and git repositories attached to a domain environment. These can also be attached to a shared space using the AWS CLI by passing values to the create-space command using the space-settings parameter.

For more information about bringing a custom Amazon SageMaker Studio Classic image, see Bring your own SageMaker image.

For more information about bringing a custom RStudio image, see Bring your own image to RStudio on SageMaker.

For instructions on using a lifecycle configuration with Studio Classic, see Use Lifecycle Configurations with Amazon SageMaker Studio.

For information about attaching a git repository to a domain, see Attach Suggested Git Repos to SageMaker.

Complete the following procedure to view the custom images, lifecycle configurations, and git repositories attached to a domain environment.

Open the Environment page

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select a domain to open the Environment page.
5. On the domain details page, choose the Environment tab.

**View and edit domains**

This topic shows how to view a list of your Amazon SageMaker domains, view the details of a domain, and edit domain settings from the Amazon SageMaker console or AWS Command Line Interface (AWS CLI).

**Topics**
- View domains
- Edit domain settings

**View domains**

The following section shows how to view a list of your domains, and details of an individual domain from the SageMaker console or the AWS CLI.

**Console**

The console's domain overview page gives information about the structure of a domain, and it provides a list of your domains. The page's domain structure diagram describes domain components and how they interact with each other.

The following procedure shows how to view a list of your domains from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.

To view the details of the domain, complete the following procedure. This page gives information about the general settings for the domain, including the name, domain ID, execution role used to create the domain, and the authentication method of the domain.

1. From the list of domains, select the domain for which you want to open the domain settings page.
2. On the **domain details** page, choose the **domain settings** tab.

**AWS CLI**

Run the following command from the terminal of your local machine to view a list of domains from the AWS CLI.

```
aws sagemaker list-domains --region region
```

**Edit domain settings**

You can edit the settings of a domain from the SageMaker console or the AWS CLI. The following considerations apply when updating the settings of a domain.

- If `DefaultUserSettings` and `DefaultSpaceSettings` are set, they cannot be unset.
- `DefaultUserSettings.ExecutionRole` can only be updated if there are no applications running in any user profile within the domain. This value cannot be unset.
- `DefaultSpaceSettings.ExecutionRole` can only be updated if there are no applications running in any of shared spaces within the domain. This value cannot be unset.
- If the domain was created in **VPC only** mode, SageMaker automatically applies updates to the security group settings defined for the domain to all shared spaces created in the domain.
- `DomainId` cannot be updated.

The following section shows how to edit domain settings from the SageMaker console or the AWS CLI.

**Console**

You can edit the domain from the SageMaker console using the following procedure.

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the domain for which you want to open the **domain settings** page.
5. On the **domain details** page, you can configure and manage your domain details by choosing the appropriate tab.
6. To configure the general settings, on the **domain details** page choose the **domain settings** tab then choose **Edit**.

**AWS CLI**

Run the following command from the terminal of your local machine to update a domain from the AWS CLI. For more information about the structure of `default-user-settings`, see **CreateDomain**.

```
aws sagemaker update-domain \
--domain-id domain-id \
--default-user-settings default-user-settings \
--default-space-settings default-space-settings \
--domain-settings-for-update settings-for-update \
--region region
```

**Delete an Amazon SageMaker domain**

A domain consists of a list of authorized users, configuration settings, and an Amazon Elastic File System (Amazon EFS) volume. The Amazon EFS volume contains data for the users, including notebooks, resources, and artifacts. A user can have multiple applications (apps) which support the reading and execution experience of the user's notebooks, terminals, and consoles.

You can delete your domain using one of the following:

- AWS console
- AWS Command Line Interface (AWS CLI)
- SageMaker SDK

The following sections explain how to delete a domain and the requirements for doing so.

**Requirements**

You must satisfy the following requirements to delete a domain.

- You must have admin permission to delete a domain.
- You can only delete an app with the status `InService` displayed as `Ready` in the domain. To delete the containing domain, you don't need to delete an app whose status is `Failed`. In the domain, an attempt to delete an app in the failed state results in an error.
• To delete a domain, the domain cannot contain any user profiles or shared spaces. To delete a user profile or shared space, the user profile or space cannot contain any non-failed apps.

When you delete these resources, the following occurs:
• App – The data (files and notebooks) in a user's home directory is saved. Unsaved notebook data is lost.
• User profile – The user can no longer sign in to the domain. The user loses access to their home directory, but the data is not deleted. An admin can retrieve the data from the Amazon EFS volume where it is stored under the user's AWS account.
• To switch authentication modes from IAM to IAM Identity Center, you must delete the domain.

EFS files

Your files are kept in an Amazon EFS volume as a backup. This backup includes the files in the mounted directory, which is `/home/sagemaker-user` for Amazon SageMaker Studio Classic and `/root` for kernels.

When you delete files from these mounted directories, the kernel or app may move the deleted files into a hidden trash folder. If the trash folder is inside the mounted directory, those files are copied into the Amazon EFS volume and will incur charges. To avoid these Amazon EFS charges, you must identify and clean the trash folder location. The trash folder location for default apps and kernels is `~/.local/`. This may vary depending on the Linux distribution used for custom apps or kernels. For more information about the Amazon EFS volume, see [Manage Your Amazon EFS Storage Volume in SageMaker Studio Classic](#).

When you use the SageMaker console to delete the domain, the Amazon EFS volume is detached but not deleted. The same behavior occurs by default when you use the AWS CLI or the SageMaker Python SDK to delete the domain. However, when you use the AWS CLI or the SageMaker Python SDK, you can set the RetentionPolicy to `HomeEfsFileSystem=Delete` to delete the Amazon EFS volume along with the domain.

**Delete an Amazon SageMaker domain (console)**

**To delete a domain**

1. Open the [SageMaker console](#).
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Select the domain that you want to delete.

5. Repeat the following steps for each user in the User profiles list.
   a. Choose the user.
   b. On the User Details page, for each non-failed app in the Apps list, choose Action.
   c. From the dropdown list, choose Delete.
   d. On the Delete app dialog box, choose Yes, delete app. Then enter delete in the confirmation field, and choose Delete.
   e. When Status shows as Deleted for all apps, choose Edit.
   f. On the Edit User page, choose Delete user.
   g. On the Delete user dialog box, choose Yes, delete user. Then enter delete in the confirmation field, and choose Delete.

⚠️ Important
When a user is deleted, they lose access to the Amazon EFS volume that contains their data, including notebooks and other artifacts. The data is not deleted and can be accessed by an administrator.

6. When all users are deleted, choose the Space management tab.

7. Repeat the following steps for each shared space in the Spaces list.
   a. Select the name of the shared space.
   b. Choose Delete app for every app.
   c. On the Delete app dialog box, choose Yes, delete app. Then enter delete in the confirmation field, and choose Delete.
   d. Choose Cancel.
   e. Select the shared space.
   f. Choose Delete.
   g. On the Delete space dialog box, choose Yes, delete space. Then enter delete in the confirmation field, and choose Delete space.

8. When all users and shared spaces are deleted, choose the domain settings tab.

9. Choose Edit.

11. On the **Delete domain** dialog box, choose **Yes, delete domain**. Then enter *delete* in the confirmation field, and choose **Delete**.

**Delete an Amazon SageMaker domain (AWS CLI)**

**To delete a domain**

1. Retrieve the list of domains in your account.

   ```
   aws --region Region sagemaker list-domains
   ```

2. Retrieve the list of applications for the domain to be deleted.

   ```
   aws --region Region sagemaker list-apps \ 
   --domain-id-equals DomainId
   ```

3. Delete each application in the list.

   ```
   aws --region Region sagemaker delete-app \ 
   --domain-id DomainId \ 
   --app-name AppName \ 
   --app-type AppType \ 
   --user-profile-name UserProfileName
   ```

4. Retrieve the list of user profiles in the domain.

   ```
   aws --region Region sagemaker list-user-profiles \ 
   --domain-id-equals DomainId
   ```

5. Delete each user profile in the list.

   ```
   aws --region Region sagemaker delete-user-profile \ 
   --domain-id DomainId \ 
   --user-profile-name UserProfileName
   ```

6. Retrieve the list of shared spaces in the domain.

   ```
   aws --region Region sagemaker list-spaces \ 
   --domain-id DomainId
   ```

7. Delete each shared space in the list.
Domain user profiles

A user profile represents a single user within an Amazon SageMaker domain. The user profile is the main way to reference a user for the purposes of sharing, reporting, and other user-oriented features. This entity is created when a user onboards to the Amazon SageMaker domain. A user profile can have (at most) a single JupyterServer application outside the context of a shared space. The user profile's Studio Classic application is directly associated with the user profile and has an isolated Amazon EFS directory, an execution role associated with the user profile, and Kernel Gateway applications. A user profile can also create other applications from the console or from Amazon SageMaker Studio.

Topics

- Add and Remove User Profiles
- View User Profiles and User Profile Details

Add and Remove User Profiles

The following sections demonstrate how to add and remove user profiles from an Amazon SageMaker domain using the SageMaker console or the AWS Command Line Interface (AWS CLI).

Topics

- Add user profiles
- Remove user profiles
Add user profiles

The following section shows how to add user profiles to a domain using the SageMaker console or the AWS CLI.

After you add a user profile to the domain, users can login using a URL. If the domain uses AWS IAM Identity Center for authentication, users receive an email that contains the URL to sign in to the domain. If the domain uses AWS Identity and Access Management, you can create a URL for a user profile using CreatePresignedDomainUrl

Add user profiles from the console

You can add user profiles to a domain from the SageMaker console by following this procedure.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to add a user profile to.
5. On the domain details page, choose the User profiles tab.
6. Choose Add user. This opens a new page.
7. Use the default name for your user profile or add a custom name.
8. For Execution role, choose an option from the role selector. If you choose Enter a custom IAM role ARN, the role must have, at a minimum, an attached trust policy that grants SageMaker permission to assume the role. For more information, see SageMaker Roles.

   If you choose Create a new role, the Create an IAM role dialog box opens:

   a. For S3 buckets you specify, specify additional Amazon S3 buckets that users of your notebooks can access. If you don't want to add access to more buckets, choose None.

   b. Choose Create role. SageMaker creates a new IAM role, AmazonSageMaker-ExecutionPolicy, with the AmazonSageMakerFullAccess policy attached.

9. (Optional) Add tags to the user profile. All resources that the user profile creates will have a domain ARN tag and a user profile ARN tag. The domain ARN tag is based on domain ID, while the user profile ARN tag is based on the user profile name.
10. Choose Next.
11. Under **Default JupyterLab version**, select a JupyterLab version from the dropdown to use as the default for your user profile. For information about selecting a JupyterLab version, see [JupyterLab Versioning](#).

12. In the **SageMaker Projects and JumpStart** section, you have two options. You can accept the default Project and JumpStart settings, or you can customize whether the user profile can create projects and use JumpStart. For more information, see [SageMaker Studio Permissions Required to Use Projects](#).

13. Choose **Next**.

14. (Optional) If the domain has an RStudio license associated, select whether you want to create the user with one of the following authorizations:

   - Unauthorized
   - RStudio Admin
   - RStudio User

15. Choose **Next**.

16. For the **Canvas base permissions configuration**, select whether to establish the minimum required permissions to use the SageMaker Canvas application.

17. (Optional) For the **Time series forecasting configuration**: To grant user permissions for time series forecasting in SageMaker Canvas, leave the **Enable time series forecasting** option turned on. It is turned on by default.

18. (Optional) If you left **Enable time series forecasting** turned on, select **Create and use a new execution role**. Alternatively, if you already have an IAM role with the required Amazon Forecast permissions attached, select **Use an existing execution role**. For more information, see the [IAM role setup method](#).

19. Choose **Submit**.

### Create user profiles from the AWS CLI

To create a user profile in a domain from the AWS CLI, run the following command from the terminal of your local machine. For information about the available JupyterLab version ARNs, see [Setting a default JupyterLab version](#).

```bash
aws --region region \
  sagemaker create-user-profile \
  --domain-id domain-id \
  --user-profile-name user-name \
```

**Create user profiles from the AWS CLI**

To create a user profile in a domain from the AWS CLI, run the following command from the terminal of your local machine. For information about the available JupyterLab version ARNs, see [Setting a default JupyterLab version](#).

```bash
aws --region region \
  sagemaker create-user-profile \
  --domain-id domain-id \
  --user-profile-name user-name \
```
Remove user profiles

All apps launched by a user profile must be deleted to delete the user profile. The following section shows how to remove user profiles from a domain using the SageMaker console or AWS CLI.

Remove user profiles from the console

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to remove a user profile from.
5. On the domain details page, choose the User profiles tab.
6. Select the user profile that you want to delete.
7. On the User Details page, for each non-failed app in the Apps list, choose Action.
8. From the dropdown list, choose Delete.
9. On the Delete app dialog box, choose Yes, delete app. Then enter delete in the confirmation field, and choose Delete.
10. When Status shows as Deleted for all apps, choose Edit.
12. On the Delete user pop-up, choose Yes, delete user.
13. Enter delete in the field to confirm deletion.

Remove user profiles from the AWS CLI

To delete a user profile from the AWS CLI, run the following command from the terminal of your local machine.
View User Profiles and User Profile Details

This topic shows how to view a list of user profiles in an Amazon SageMaker domain, and view details for a user profile from the SageMaker console or the AWS Command Line Interface (AWS CLI).

Topics

- View user profiles
- View user profile details

View user profiles

The following section describes how to view a list of user profiles in a domain from the SageMaker console or the AWS CLI.

View user profiles from the console

Complete the following procedure to view a list of user profiles in the domain from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to view a list of user profiles for.
5. On the domain details page, choose the User profiles tab.

View user profiles from the AWS CLI

To view the user profiles in a domain from the AWS CLI, run the following command from the terminal of your local machine.

```
aws sagemaker list-user-profiles
```
View user profile details

The following section describes how to view the details of a user profile from the SageMaker console or the AWS CLI.

View user profile details from the console

Complete the following procedure to view the details of a user profile from the SageMaker console.

2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to view a list of user profiles for.
5. On the domain details page, choose the User profiles tab.
6. Select the user profile that you want to view details for.

View user profile details from the AWS CLI

To describe a user profile from the AWS CLI, run the following command from the terminal of your local machine.

```
aws sagemaker describe-user-profile \
  --region region \
  --domain-id domain-id \
  --user-profile-name user-name
```

IAM Identity Center Groups in a domain

If you use AWS IAM Identity Center authentication for your Amazon SageMaker domain, you can add and edit group and user access to a domain. For more information about IAM Identity Center authentication, see [What is IAM Identity Center?](#). The following topics show how to manage IAM Identity Center users and groups that have access to a domain.

Topics
• View groups and users
• Add groups and users
• Remove groups

View groups and users

Complete the following procedure to view a list of IAM Identity Center groups and users from the Amazon SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to open the domain settings page for.
5. On the domain details page, choose the Groups tab.

Add groups and users

The following sections show how to add groups and users to a domain from the SageMaker console or AWS CLI.

⚠️ Note
If the domain was created before October 1st, 2023, you can only add groups and users to the domain from the SageMaker console.

SageMaker console

Complete the following procedure to add groups and users to your domain from the SageMaker console.

1. On the Groups tab, choose Assign users and groups.
2. On the Assign users and groups page, select the users and groups that you want to add.
3. Choose Assign users and groups.
**AWS CLI**

Complete the following procedure to add groups and users to your domain from the AWS CLI.

1. **Fetch the SingleSignOnApplicationArn of the domain with a call to `describe-domain`**.
   SingleSignOnApplicationArn is the ARN of the application managed in IAM Identity Center.

   ```bash
   aws sagemaker describe-domain \
   --region region \
   --domain-id domain-id
   ```

2. **Associate the user or group with the domain**. To accomplish this, pass the SingleSignOnApplicationArn value returned from the `describe-domain` command as the application-arn parameter in a call to `create-application-assignment`. You must also pass the type and ID of the entity to associate.

   ```bash
   aws sso-admin create-application-assignment \
   --application-arn application-arn \
   --principal-id principal-id \
   --principal-type principal-type
   ```

**Remove groups**

Complete the following procedure to remove groups from your domain from the SageMaker console. For information about deleting a user, see [Remove user profiles](#).

1. On the **Groups** tab, choose the group that you want to remove.
2. Choose **Unassign groups**.
3. On the pop-up window, choose **Yes, unassign groups**.
4. Enter `unassign` in the field.
5. Choose **Unassign groups**.

**Quick onboard to Amazon SageMaker domain**

This topic describes how to onboard to Amazon SageMaker domain by using the **Set up for single users** procedure from the SageMaker console. This procedure sets up your domain in a single click and uses AWS Identity and Access Management (IAM) authentication.
For information on how to set up your domain with advanced settings, see Custom onboarding using IAM for the custom IAM authentication set up and Custom onboarding to Amazon SageMaker domain using IAM Identity Center for the custom IAM Identity Center authentication set up.

RStudio support is not currently available when onboarding using the Set up for single users procedure. To use RStudio, you must onboard using the Set up for organizations procedure.

**Topics**

- Quick setup for single users
- Default settings

**Quick setup for single users**

After satisfying the prerequisites (create an AWS account, create an administrative user, and set up your AWS CLI) in Set Up Amazon SageMaker Prerequisites, expand one of the following instruction options and follow the steps.

**Onboard to the domain quickly using Set up for single user from the SageMaker landing page**

1. Open the SageMaker console.
2. On the landing page you will see New to SageMaker? on the right. If you do not see this option, open the the left navigation pane and choose Amazon SageMaker at the top of the navigation pane.
3. Under New to SageMaker?, choose Set up for single user. Your domain and user profile are created automatically.

**Onboard to the domain quickly using Set up for single user from the domain page**

1. Open the SageMaker console.
2. Open the left navigation pane.
3. Under Admin configurations, choose Domains.
4. Choose Create domain.
5. Choose Set up for single user (Quick setup). Your domain and user profile are created automatically.
Now that you've onboarded to the domain, you can launch an application by opening the SageMaker console and choosing the environment from the left navigation pane.

For information about adding users to your domain, see Add and Remove User Profiles.

Default settings

When you onboard to Amazon SageMaker domain using the Set up for single user procedure, SageMaker uses the following defaults as part of the domain and user profile creation. You can't modify these defaults using the Set up for single user procedure.

- **Domain name**: SageMaker automatically assigns the name of the domain with a timestamp in the following format.

  QuickSetupDomain-YYYYMMDDTHHMMSS

- **User profile name**: SageMaker automatically assigns the name of the user profile with a timestamp in the following format.

  default-YYYYMMDDTHHMMSS

- **Domain execution role**: SageMaker creates a new IAM role and attaches the AmazonSageMakerFullAccess policy. When using the quick setup and the updated Amazon SageMaker Studio is your default experience, your IAM role also includes the AmazonSageMakerCanvasFullAccess, AmazonSageMakerCanvasAIServicesAccess, AmazonS3FullAccess policies.

- **User profile execution role**: SageMaker sets the user profile execution role to the same IAM role used for the domain execution role.

- **Shared space execution role**: SageMaker sets the shared space execution role to the same IAM role used for the domain execution role.

- **SageMaker Canvas time series forecasting role**: SageMaker creates a new IAM role with the permissions required to use the SageMaker Canvas time series forecasting feature.

- **Amazon S3 bucket**: SageMaker creates an Amazon S3 bucket named with the following format for notebook sharing.

  sagemaker-studio-XXXXXXXXXXXXXXXX

- **Amazon VPC**: SageMaker selects a public VPC with the following logic.
1. If there is a default VPC with associated subnets in the Region, SageMaker uses it.

2. If there is no default VPC or the default VPC has no associated subnets, then SageMaker uses any existing VPC with associated subnets. If there are multiple existing VPCs, SageMaker can select any of them.

**Custom onboarding to Amazon SageMaker domain using IAM Identity Center**

This topic describes how to onboard to Amazon SageMaker domain using authentication using AWS IAM Identity Center from the SageMaker console or the AWS CLI. For information about setting up IAM Identity Center for use with a domain, see [Set up IAM Identity Center for use with Amazon SageMaker domain](#). For information on how to onboard using AWS Identity and Access Management (IAM) authentication, see [Quick onboarding](#) or [Custom onboarding using IAM](#).

**Topics**

- Onboard from the console
- Onboard from the AWS CLI
- Access the domain after onboarding
- Set up IAM Identity Center for use with Amazon SageMaker domain

**Onboard from the console**

After satisfying the prerequisites (create an AWS account, create an administrative user, and set up your AWS CLI) in [Set Up Amazon SageMaker Prerequisites](#), follow the steps.

**To onboard to domain using IAM Identity Center**

1. Open the [SageMaker console](#).
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the domains page, choose Create domain.
5. On the Set up SageMaker domain page, choose Set up for organizations.
6. Select Configure.
Step 1: Domain details

1. For **domain Name**, enter a unique name for your domain. For example, this can be your project or team name.
2. Choose **Next**.

Step 2: Users and ML Activities

Select the group or create the users for the domain and grant them permissions for the ML activities to which you want them to have access.

In these setup instructions we use the IAM Identity Center option. To use authentication in IAM Identity Center, you must belong to an organization in AWS Organizations. For information on setting up an IAM Identity Center, see [Set up IAM Identity Center for use with Amazon SageMaker domain](#).

The IAM role you configure in this step is assigned to the selected IAM Identity Center group and all users who are part of that group.

1. Under **How do you want to access Studio?**, choose **AWS Identity Center**.
2. Under **Who will use Studio?** select the IAM Identity Center users or groups, then choose **Select**. You need to set up Amazon SageMaker Studio within the same Region in which your IAM Identity Center is configured. You can change the Region of your domain by choosing the Region from the dropdown list on the top right of the console or you can change your IAM Identity Center Region by navigating to the [AWS access portal](#).
3. Under **What ML activities do they perform?** you can use an existing role by choosing **Use an existing role** or you can create a new role by choosing **Create a new role** and checking the ML activities to which you wish the role to have access. You can select at most 10 ML activities.
4. While selecting ML activities, you may need to satisfy requirements. To satisfy a requirement, choose **Add** and complete the requirement.
5. After all requirements are satisfied, choose **Next**.

Step 3: Applications

In this step, you can configure the applications you have enabled in the previous step. For more information on the ML activities, see [ML activity reference](#).
If the application has not been enabled, you receive a warning for that application. To enable an application that has not been enabled, return to the previous step by choosing **Back** and follow the previous instructions.

**Studio** configuration:

Under **Studio**, you have the option to choose between the new and classic version of Studio as your default experience. This means choosing which ML environment you will interact with after opening Studio.

- **Studio - New** includes multiple integrated development environments (IDEs) and applications, including Amazon SageMaker Studio Classic. If chosen, the Studio Classic IDE has default settings. For information on the default settings, see [Default settings](#).

- **Studio Classic** includes the Jupyter IDE. If chosen, you may configure your Studio Classic configuration.

For information on Studio Classic, see [Amazon SageMaker Studio Classic](#).

**SageMaker Canvas** configuration:

If you have Amazon SageMaker Canvas enabled, see [Getting started with using Amazon SageMaker Canvas](#) for the instructions and configuration details for onboarding.

**Studio Classic** configuration:

If you chose **Studio - New** (recommended) as your default experience, the Studio Classic IDE has default settings. For information on the default settings, see [Default settings](#).

If you chose Studio Classic as your default experience, you can choose to enable or disable notebook resource sharing. Notebook resources include artifacts such as cell output and Git repositories. For more information on Notebook resources, see [Share and Use an Amazon SageMaker Studio Classic Notebook](#).

If you enabled notebook resource sharing:

1. Under **S3 location for shareable notebook resources**, input your Amazon S3 location.
2. Under **Encryption key - optional**, leave as **No Custom Encryption** or choose an existing AWS KMS key or choose **Enter a KMS key ARN** and enter your AWS KMS key’s ARN.
3. Under **Notebook cell output sharing preference**, choose **Allow users to share cell output** or **Disable cell output sharing**.
RStudio configuration:

To enable RStudio, you need an RStudio license. To set that up, see RStudio license.

1. Under **RStudio Workbench**, verify that your RStudio license is automatically detected. For more information about getting an RStudio license and activating it with SageMaker, see RStudio license.

2. Select an instance type to launch your RStudio Server on. For more information, see RStudioServerPro instance type.

3. Under **Permission**, create your role or select an existing role. The role must have the following permissions policy. This policy allows the RStudioServerPro application to access necessary resources. It also allows Amazon SageMaker to automatically launch an RStudioServerPro application when the existing RStudioServerPro application is in a Deleted or Failed status. For information about adding permissions to a role, see Modifying a role permissions policy (console).

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "VisualEditor0",
            "Effect": "Allow",
            "Action": [
                "license-manager:ExtendLicenseConsumption",
                "license-manager:ListReceivedLicenses",
                "license-manager:GetLicense",
                "license-manager:CheckoutLicense",
                "license-manager:CheckInLicense",
                "logs:CreateLogDelivery",
                "logs:CreateLogGroup",
                "logs:CreateLogStream",
                "logs:DeleteLogDelivery",
                "logs:Describe*",
                "logs:GetLogDelivery",
                "logs:GetLogEvents",
                "logs:ListLogDeliveries",
                "logs:PutLogEvents",
                "logs:PutResourcePolicy",
                "logs:UpdateLogDelivery",
                "sagemaker:CreateApp"
            ]
        }
    ]
}
```
4. Under **RStudio Connect**, add the URL for your RStudio Connect server. RStudio Connect is a publishing platform for Shiny applications, R Markdown reports, dashboards, plots, and more. When you onboard to RStudio on SageMaker, an RStudio Connect server is not created. For more information, see [RStudio Connect URL](#).

5. Under **RStudio Package Manager**, add the URL for your RStudio Package Manager. SageMaker creates a default package repository for the Package Manager when you onboard RStudio. For more information about RStudio Package Manager, see [RStudio Package Manager](#).

6. Select **Next**.

**Code Editor** configuration:

If you have Code Editor enabled, see [Code Editor](#) for an overview and the configuration details.

**Step 4: Network**

Choose how you want Studio to connect to other AWS services.

You can choose to disable internet access to your Studio by specifying using **Virtual Private Cloud (VPC) Only** network access type. If you choose this option, you cannot run a Studio notebook unless your VPC has an interface endpoint to the SageMaker API and runtime, or a Network Address Translation (NAT) gateway with internet access, and your security groups allow outbound connections. For more information on Amazon VPCs, see [Choose an Amazon VPC](#).

If you choose Virtual Private Cloud (VPC) Only the following steps are required. If you choose **Public internet access**, the first two of the following steps are required.

1. Under **VPC**, choose the Amazon VPC ID.
2. Under **Subnet**, choose one or more subnets. If you don't choose any subnets, SageMaker uses all the subnets in the Amazon VPC. We recommend that you use multiple subnets that are not created in constrained Availability Zones. Using subnets in these constrained Availability Zones can result in insufficient capacity errors and longer application creation times. For more information about constrained Availability Zones, see [Availability Zones](#).
3. Under **Security group(s)**, choose one or more subnets.
If VPC only is selected, SageMaker automatically applies the security group settings defined for the domain to all shared spaces created in the domain. If Public internet only is selected, SageMaker does not apply the security group settings to shared spaces created in the domain.

**Step 5: Storage**

You have the option to encrypt your data. The Amazon Elastic File System (Amazon EFS) and Amazon Elastic Block Store (Amazon EBS) file systems that are created for you when you create a domain. Amazon EBS sizes are used by both Code Editor and JupyterLab spaces.

You cannot change the encryption key after encrypt your Amazon EFS and Amazon EBS file systems. To encrypt your Amazon EFS and Amazon EBS file systems, you can use the following configurations.

- Under **Encryption key - optional**, leave as No Custom Encryption or choose an existing KMS key or choose Enter a KMS key ARN and enter the ARN of your KMS key.
- Under **Default space size - optional**, enter the default space size.
- Under **Maximum space size - optional**, enter the maximum space size.

**Step 6: Review and create**

Review your domain settings. If you need to change the settings, choose Edit next to the relevant step. Once you confirm that your domain settings are accurate, choose Submit and the domain is created for you. This process may take a few minutes.

**Onboard from the AWS CLI**

Use the following commands to onboard to a domain using authentication using IAM Identity Center from the AWS CLI.

After satisfying the prerequisites (create an AWS account, create an administrative user, and set up your AWS CLI) in Set Up Amazon SageMaker Prerequisites, use the following the steps.

1. Create an execution role that is used to create a domain and attach the AmazonSageMakerFullAccess policy. You can also use an existing role that has, at a minimum, an attached trust policy that grants SageMaker permission to assume the role. For more information, see SageMaker Roles.

   ```
   aws iam create-role --role-name execution-role-name --assume-role-policy-document file://execution-role-trust-policy.json
   ```
2. Get the default Amazon Virtual Private Cloud (Amazon VPC) of your account.

```bash
aws --region region ec2 describe-vpcs --filters Name=isDefault,Values=true --query "Vpcs[0].VpcId" --output text
```

3. Get the list of subnets in the default Amazon VPC.

```bash
aws --region region ec2 describe-subnets --filters Name=vpc-id,Values=default-vpc-id --query "Subnets[*].SubnetId" --output json
```

4. Create a domain by passing the default Amazon VPC ID, subnets, and execution role ARN. You must also pass a SageMaker image ARN. For information on the available JupyterLab version ARNs, see Setting a default JupyterLab version.

```bash
aws --region region sagemaker create-domain --domain-name domain-name --vpc-id default-vpc-id --subnet-ids subnet-ids --auth-mode SSO --default-user-settings "ExecutionRole=arn:aws:iam::account-number:role/execution-role-name,JupyterServerAppSettings={DefaultResourceSpec={InstanceType=system,SageMakerImageArn=image-arn}}" \ --query DomainArn --output text
```

5. Verify that the domain has been created.

```bash
aws --region region sagemaker list-domains
```

**Access the domain after onboarding**

After you are given access to the domain, you are sent an email inviting you to create a password and use IAM Identity Center. The email also contains the URL to sign in to the domain. For more information about signing in and session duration, see How to sign in to the user portal.

After you activate your account, go to the domain URL, sign in, and wait for your user profile to be created. On subsequent visits, you only need to wait for the Studio app to load.

Bookmark the URL. The URL is also available on the domain settings page.
Set up IAM Identity Center for use with Amazon SageMaker domain

To use authentication in IAM Identity Center, you must belong to an organization in AWS Organizations. If you don't belong to an organization, you can create one by following the steps in Tutorial: Creating and configuring an organization.

Multi-factor authentication (MFA) is enabled by default when you create an IAM Identity Center instance. Users are prompted to sign in with MFA when their device, browser, or location changes. As a security best practice, we strongly recommend enabling MFA for your workforce identities. For more information, see Manage MFA devices in IAM Identity Center.

After you have created your organization and user, you can create a SageMaker user profile for that user in IAM Identity Center as follows.

1. From the Amazon SageMaker console: – You can use the Amazon SageMaker console to create a user profile for the user in IAM Identity Center. If the user in IAM Identity Center hasn’t already been associated with the domain, it is automatically associated.

2. Using the AWS CLI or AWS CloudFormation – A user in IAM Identity Center assigned to the domain can create a user profile using the SageMaker console, the AWS CLI or AWS CloudFormation.
   - First, you must use the IAM Identity Center console to assign the user or the user’s group to the domain. For more information about application assignment, see Assign user access.
   - Then, use the AWS CLI or AWS CloudFormation to create a SageMaker user profile for the user.

Note
To simplify administration of access permissions, we recommend using IAM Identity Center to assign groups to the domain (instead of assigning users). Groups allow permissions to be granted or denied to multiple users at once. A user can be moved out of a group or to a different group if needed. When assigning user access to applications, IAM Identity Center does not currently support users being added to nested groups. If a user is added to a nested group, they might receive a "You do not have any applications" error message during sign-in. Assignments must be made to the immediate group the user is a member of.

Return to the Domains page to continue to onboard using IAM Identity Center authentication.
Custom onboarding to Amazon SageMaker domain using IAM

This topic describes how to onboard to Amazon SageMaker domain using the **Set up for organizations** procedure for AWS Identity and Access Management (IAM) authentication from the SageMaker console or the AWS CLI. To onboard faster using IAM, see [Quick onboarding](#).

For information on how to onboard using AWS IAM Identity Center (IAM Identity Center), see [Custom onboarding using IAM Identity Center](#).

### Onboard using console

After satisfying the prerequisites (create an AWS account, create an administrative user, and set up your AWS CLI) in [Set Up Amazon SageMaker Prerequisites](#), follow the steps.

**To onboard to domain using IAM**

1. Open the [SageMaker console](#).
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the **domains** page, choose **Create domain**.
5. On the **Setup SageMaker domain** page, choose **Set up for organizations**.
6. Select **Configure**.

#### Step 1: domain details

1. For **domain Name**, enter a unique name for your domain. For example, this can be your project or team name.
2. Choose **Next**.

#### Step 2: Users and ML Activities

Select the group or create the users for the domain and grant permissions to which ML activities they will have access.

In these setup instructions, we use the **Login through IAM** option.

The IAM role you configure in this step is assigned to all of the users you add in this step.
Step 1: Access

1. Under **How do you want to access Studio?**, choose **Login through IAM**.

2. Under **Who will use Studio?** add the user profile names. To add a user profile name, choose **Add** user, enter a user profile name, then choose **Select**.

3. Under **What ML activities do they perform?** you can use an existing role by choosing **Use an existing role** or you can create a new role by choosing **Create a new role** and checking the ML activities to which you wish the role to have access. You can select at most 10 ML activities.

4. While selecting ML activities, you may need to satisfy requirements. To satisfy a requirement, choose **Add** and complete the requirement.

5. After all requirements are satisfied, choose **Next**.

**Step 3: Applications**

In this step, you can configure the applications you have enabled in the previous step. For more information on the ML activities, see [ML activity reference](#).

If the application has not been enabled, you receive a warning for that application. To enable an application that has not been enabled, return to the previous step by choosing **Back** and follow the previous instructions.

**Studio** configuration:

Under **Studio**, you have the option to choose between the new and classic version of Studio as your default experience. This means choosing which ML environment you will interact with after opening Studio.

- **Studio - New** includes multiple integrated development environments (IDEs) and applications, including Amazon SageMaker Studio Classic. If chosen, the Studio Classic IDE has default settings. For information on the default settings, see [Default settings](#).

- **Studio Classic** includes the Jupyter IDE. If chosen, you may configure your Studio Classic configuration.

  For information on Studio Classic, see [Amazon SageMaker Studio Classic](#).

**SageMaker Canvas** configuration:

If you have Amazon SageMaker Canvas enabled, see [Getting started with using Amazon SageMaker Canvas](#) for the instructions and configuration details for onboarding.
Studio Classic configuration:

If you chose **Studio - New** (recommended) as your default experience, the Studio Classic IDE has default settings. For information on the default settings, see Default settings.

If you chose Studio Classic as your default experience, you can choose to enable or disable notebook resource sharing. Notebook resources include artifacts such as cell output and Git repositories. For more information on Notebook resources, see Share and Use an Amazon SageMaker Studio Classic Notebook.

If you enabled notebook resource sharing:

1. Under **S3 location for shareable notebook resources**, input your Amazon S3 location.
2. Under **Encryption key - optional**, leave as No Custom Encryption or choose an existing AWS KMS key or choose Enter a KMS key ARN and enter your AWS KMS key’s ARN.
3. Under **Notebook cell output sharing preference**, choose Allow users to share cell output or Disable cell output sharing.

RStudio configuration:

To enable RStudio you will need an RStudio license. To set that up, see RStudio license.

1. Under **RStudio Workbench**, verify that your RStudio license is automatically detected. For more information about getting an RStudio license and activating it with SageMaker, see RStudio license.
2. Select an instance type to launch your RStudio Server on. For more information, see RStudioServerPro instance type.
3. Under **Permission**, create your role or select an existing role. The role must have the following permissions policy. This policy allows the RStudioServerPro application to access necessary resources. It also allows Amazon SageMaker to automatically launch an RStudioServerPro app when the existing RStudioServerPro application is in a Deleted or Failed status. For information about adding permissions to a role, see Modifying a role permissions policy (console).

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": "sagemaker:CreateRStudioServerProInstance",
            "Resource": "arn:aws:sagemaker:region:account-id:instance-id"
        },
        {
            "Effect": "Allow",
            "Action": "sagemaker:DeleteRStudioServerProInstance",
            "Resource": "arn:aws:sagemaker:region:account-id:instance-id"
        },
        {
            "Effect": "Allow",
            "Action": "sagemaker:DescribeRStudioServerProInstance",
            "Resource": "arn:aws:sagemaker:region:account-id:instance-id"
        },
        {
            "Effect": "Allow",
            "Action": "sagemaker:StopRStudioServerProInstance",
            "Resource": "arn:aws:sagemaker:region:account-id:instance-id"
        },
        {
            "Effect": "Allow",
            "Action": "sagemaker:StartRStudioServerProInstance",
            "Resource": "arn:aws:sagemaker:region:account-id:instance-id"
        },
        {
            "Effect": "Allow",
            "Action": "sagemaker:UpdateRStudioServerProInstance",
            "Resource": "arn:aws:sagemaker:region:account-id:instance-id"
        }
    ]
}
```
4. Under **RStudio Connect**, add the URL for your RStudio Connect server. RStudio Connect is a publishing platform for Shiny applications, R Markdown reports, dashboards, plots, and more. When you onboard to RStudio on SageMaker, an RStudio Connect server is not created. For more information, see [RStudio Connect URL](#).

5. Under **RStudio Package Manager**, add the URL for your RStudio Package Manager. SageMaker creates a default package repository for the Package Manager when you onboard RStudio. For more information about RStudio Package Manager, see [RStudio Package Manager](#).

6. Select **Next**.

**Code Editor** configuration:

If you have Code Editor enabled, see [Code Editor](#) for an overview and the configuration details.

**Step 4: Network**

Choose how you want Studio to connect to other AWS services.
You can choose to disable internet access to your Studio by specifying using **Virtual Private Cloud (VPC) Only** network access type. If you choose this option, you cannot run a Studio notebook unless your VPC has an interface endpoint to the SageMaker API and runtime, or a Network Address Translation (NAT) gateway with internet access, and your security groups allow outbound connections. For more information on Amazon VPCs, see [Choose an Amazon VPC](#).

If you choose Virtual Private Cloud (VPC) Only the following steps are required. If you choose **Public internet access**, the first two of the following steps are required.

1. **Under VPC**, choose the Amazon VPC ID.

2. **Under Subnet**, choose one or more subnets. If you don't choose any subnets, SageMaker uses all the subnets in the Amazon VPC. We recommend that you use multiple subnets that are not created in constrained Availability Zones. Using subnets in these constrained Availability Zones can result in insufficient capacity errors and longer application creation times. For more information about constrained Availability Zones, see [Availability Zones](#).

3. **Under Security group(s)**, choose one or more subnets.

If **VPC only** is selected, SageMaker automatically applies the security group settings defined for the domain to all shared spaces created in the domain. If **Public internet only** is selected, SageMaker does not apply the security group settings to shared spaces created in the domain.

**Step 5: Storage**

You have the option to encrypt your data. The [Amazon Elastic File System (Amazon EFS)](#) and [Amazon Elastic Block Store (Amazon EBS)](#) file systems that are created for you when you create a domain. Amazon EBS sizes are used by both Code Editor and JupyterLab spaces.

You cannot change the encryption key after encrypt your Amazon EFS and Amazon EBS file systems. To encrypt your Amazon EFS and Amazon EBS file systems, you can use the following configurations.

- **Under Encryption key - optional**, leave as **No Custom Encryption** or choose an existing KMS key or choose **Enter a KMS key ARN** and enter the ARN of your KMS key.
- **Under Default space size - optional**, enter the default space size.
- **Under Maximum space size - optional**, enter the maximum space size.

**Step 6: Review and create**
Review your domain settings. If you need to change the settings, choose **Edit** next to the relevant step. Once you confirm that your domain settings are accurate, choose **Submit** and the domain is created for you. This process may take a few minutes.

### Onboard using the AWS CLI

After satisfying the prerequisites (create an AWS account, create an administrative user, and set up your AWS CLI) in [Set Up Amazon SageMaker Prerequisites](#), use the following commands to onboard to a domain using authentication using IAM from the AWS CLI.

1. Create an execution role that is used to create a domain and attach the [AmazonSageMakerFullAccess](#) policy. You can also use an existing role that has, at a minimum, an attached trust policy that grants SageMaker permission to assume the role. For more information, see [SageMaker Roles](#).

   ```bash
   aws iam create-role --role-name execution-role-name
   aws iam attach-role-policy --role-name execution-role-name --policy-arn arn:aws:iam::aws:policy/AmazonSageMakerFullAccess
   ```

2. Get the default Amazon Virtual Private Cloud (Amazon VPC) of your account.

   ```bash
   aws --region region ec2 describe-vpcs --filters Name=isDefault,Values=true --query "Vpcs[0].VpcId" --output text
   ```

3. Get the list of subnets in the default Amazon VPC.

   ```bash
   aws --region region ec2 describe-subnets --filters Name=vpc-id,Values=default-vpc-id --query "Subnets[*].SubnetId" --output json
   ```

4. Create a domain by passing the default Amazon VPC ID, subnets, and execution role ARN. You must also pass a SageMaker image ARN. For information on the available JupyterLab version ARNs, see [Setting a default JupyterLab version](#).

   ```bash
   aws --region region sagemaker create-domain --domain-name domain-name --vpc-id default-vpc-id --subnet-ids subnet-ids --auth-mode IAM --default-user-settings "ExecutionRole=arn:aws:iam::account-number:role/execution-role-name,JupyterServerAppSettings={DefaultResourceSpec={InstanceType=system,SageMakerImageArn=image-arn}}" \
   --query DomainArn --output text
   ```

5. Verify that the domain has been created.
Choose an Amazon VPC

This topic provides detailed information about choosing an Amazon Virtual Private Cloud (Amazon VPC) when you onboard to Amazon SageMaker domain. For more information about onboarding to SageMaker domain, see Amazon SageMaker domain overview.

By default, SageMaker domain uses two Amazon VPCs. One Amazon VPC is managed by Amazon SageMaker and provides direct internet access. You specify the other Amazon VPC, which provides encrypted traffic between the domain and your Amazon Elastic File System (Amazon EFS) volume.

You can change this behavior so that SageMaker sends all traffic over your specified Amazon VPC. When you choose this option, you must provide the subnets, security groups, and interface endpoints that are necessary to communicate with the SageMaker API and SageMaker runtime, and various AWS services, such as Amazon Simple Storage Service (Amazon S3) and Amazon CloudWatch, that are used by Studio.

When you onboard to SageMaker domain, you tell SageMaker to send all traffic over your Amazon VPC by setting the network access type to VPC only.

To specify the Amazon VPC information

When you specify the Amazon VPC entities (that is, the Amazon VPC, subnet, or security group) in the following procedure, one of three options is presented based on the number of entities you have in the current AWS Region. The behavior is as follows:

- One entity – SageMaker uses that entity. This can't be changed.
- Multiple entities – You must choose the entities from the dropdown list.
- No entities – You must create one or more entities in order to use domain. Choose Create <entity> to open the VPC console in a new browser tab. After you create the entities, return to the domain Get started page to continue the onboarding process.

This procedure is part of the Amazon SageMaker domain onboarding process when you choose Set up for organizations. Your Amazon VPC information is specified under the Network section.

1. Select the network access type.
Note

If **VPC only** is selected, SageMaker automatically applies the security group settings defined for the domain to all shared spaces created in the domain. If **Public internet only** is selected, SageMaker does not apply the security group settings to shared spaces created in the domain.

- **Public internet only** – Non-Amazon EFS traffic goes through a SageMaker managed Amazon VPC, which allows internet access. Traffic between the domain and your Amazon EFS volume is through the specified Amazon VPC.
- **VPC only** – All SageMaker traffic is through the specified Amazon VPC and subnets. You must use a subnet that does not have direct internet access in **VPC only** mode. Internet access is disabled by default.

2. Choose the Amazon VPC.

3. Choose one or more subnets. If you don't choose any subnets, SageMaker uses all the subnets in the Amazon VPC. We recommend that you use multiple subnets that are not created in constrained Availability Zones. Using subnets in these constrained Availability Zones can result in insufficient capacity errors and longer application creation times. For more information about constrained Availability Zones, see [Availability Zones](https://docs.aws.amazon.com/en_us/sexualtheValue.html).

4. Choose the security groups. If you chose **Public internet only**, this step is optional. If you chose **VPC only**, this step is required.

Note

For the maximum number of allowed security groups, see [UserSettings](https://docs.aws.amazon.com/en_us/sexualtheValue.html).

For Amazon VPC requirements in **VPC only** mode, see [Connect SageMaker Studio Notebooks in a VPC to External Resources](https://docs.aws.amazon.com/en_us/sexualtheValue.html).

**Supported Regions and Quotas**

For the AWS Regions supported by Amazon SageMaker and the Amazon Elastic Compute Cloud (Amazon EC2) instance types that are available in each Region, see [Amazon SageMaker Pricing](https://docs.aws.amazon.com/en_us/sexualtheValue.html).
For a list of the SageMaker service endpoints for each Region, see Amazon SageMaker endpoints and quotas in the AWS General Reference.

**Quotas**

For a list of SageMaker quotas, see Amazon SageMaker endpoints and quotas in the AWS General Reference.

The Service Quotas console provides information about your service quotas. You can use the Service Quotas console to view your default service quotas or to request quota increases. To request a quota increase for adjustable quotas, see Requesting a quota increase.

You can set up a quota request template for your AWS Organization that automatically requests quota increases during account creation. For more information, see Using Service Quotas request templates.
Use automated ML, no-code, or low-code

Amazon SageMaker offers the following features to automate key machine learning tasks and use no-code or low-code solutions.

- **Amazon SageMaker Autopilot** is an automated machine learning (AutoML) feature-set that automates the end-to-end process of building, training, tuning, and deploying machine learning models. Amazon SageMaker Autopilot analyzes your data, selects algorithms suitable for your problem type, preprocesses the data to prepare it for training, handles automatic model training, and performs hyperparameter optimization to find the best performing model for your dataset.

- **SageMaker JumpStart** provides pretrained, open-source models for a wide range of problem types to help you get started with machine learning. You can incrementally train and tune these models before deployment. JumpStart also provides solution templates that set up infrastructure for common use cases, and executable example notebooks for machine learning with SageMaker.

**Topics**

- [SageMaker Autopilot](#)
- [SageMaker JumpStart](#)

## SageMaker Autopilot

**Important**

As of November 30, 2023, Autopilot's UI is migrating to [Amazon SageMaker Canvas](#) as part of the updated [Amazon SageMaker Studio](#) experience. SageMaker Canvas provides data scientists with no-code capabilities for tasks such as data preparation, feature engineering, algorithm selection, training and tuning, inference, continuous model monitoring, and more. SageMaker Canvas supports a variety of use cases, including computer vision, demand forecasting, intelligent search, and generative AI.

Users of [Amazon SageMaker Studio Classic](#), the previous experience of [Studio](#), can continue using the Autopilot UI in Studio Classic. Users with coding experience can continue using all [API references](#) in any supported SDK for technical implementation.

If you have been using Autopilot in Studio Classic until now and want to migrate to SageMaker Canvas, you might have to grant additional permissions to your user profile
Amazon SageMaker Autopilot is a feature set that simplifies and accelerates various stages of the machine learning workflow by automating the process of building and deploying machine learning models (AutoML).

Autopilot performs the following key tasks that you can use on autopilot or with various degrees of human guidance:

- **Data analysis and preprocessing:** Autopilot identifies your specific problem type, handles missing values, normalizes your data, selects features, and overall prepares the data for model training.

- **Model selection:** Autopilot explores a variety of algorithms and uses a cross-validation resampling technique to generate metrics that evaluate the predictive quality of the algorithms based on predefined objective metrics.

- **Hyperparameter optimization:** Autopilot automates the search for optimal hyperparameter configurations.

- **Model training and evaluation:** Autopilot automates the process of training and evaluating various model candidates. It splits the data into training and validation sets, trains the selected model candidates using the training data, and evaluates their performance on the unseen data of the validation set. Lastly, it ranks the optimized model candidates based on their performance and identifies the best performing model.

- **Model deployment:** Once Autopilot has identified the best performing model, it provides the option to deploy the model automatically by generating the model artifacts and the endpoint exposing an API. External applications can send data to the endpoint and receive the corresponding predictions or inferences.

Autopilot supports building machine learning models on large datasets up to hundreds of GBs.

The following diagram outlines the tasks of this AutoML process managed by Autopilot.
Depending on your comfort level with the machine learning process and coding experience, you can use Autopilot in different ways:

- **Using the Studio Classic UI**, users can choose between a no-code experience or have some level of human input.

  - **Note**
    
    Only experiments created from tabular data for problem types such as regression or classification are available via the Studio Classic UI.

- **Using the AutoML API**, users with coding experience can use available SDKs to create AutoML jobs. This approach provides greater flexibility and customization options and is available for all problem types.

Autopilot currently supports the following problem types:

  - **Note**
    
    For regression or classification problems involving tabular data, users can choose between two options: using the Studio Classic user interface or the API Reference. Tasks such as text and image classification, time-series forecasting, and fine-tuning of large language models are exclusively available through the version 2 of the Autopilot API. For Python users, we recommend using the AWS SDK for Python (Boto3) as the Amazon SageMaker Python SDK is not currently supported for the Autopilot API version 2.

    Users who prefer the convenience of a user interface can use Amazon SageMaker Canvas to access pre-trained models and generative AI foundation models, or create custom models tailored for specific text, image classification, forecasting needs, or generative AI.
• **Regression, binary, and multiclass classification** with tabular data formatted as CSV or Parquet files in which each column contains a feature with a specific data type and each row contains an observation. The column data types accepted include numerical, categorical, text, and time series that consists of strings of comma-separated numbers.
  
  • To create an Autopilot job as a pilot experiment using the SageMaker API reference, see [Create a regression or classification job for tabular data using the AutoML API](#).
  
  • To create an Autopilot job as a pilot experiment using the Studio Classic UI, see [Create a Regression or Classification Autopilot experiment for tabular data using the Studio Classic UI](#).
  
  • If you are an administrator looking to pre-configure default infrastructure, networking, or security parameters of Autopilot experiments in Studio Classic UI, see [Configure the default parameters of an Autopilot experiment (for administrators)](#).
  
  • **Text classification** with data formatted as CSV or Parquet files in which a column provides the sentences to be classified, while another column should provide the corresponding class label. See [Create an AutoML job for text classification using the API](#).
  
  • **Image classification** with image formats such as PNG, JPEG, or a combination of both. See [Create an AutoML job for image classification using the API](#).
  
  • **Time-series forecasting** with time-series data formatted as CSV or Parquet files. See [Create an AutoML job for time-series forecasting using the API](#).
  
  • Fine-tuning of large language models (LLMs) for **text generation** with data formatted as CSV or Parquet files. See [Create an AutoML job to fine-tune text generation models using the API](#).

Additionally, Autopilot helps users understand how models make predictions by automatically generating reports that show the importance of each individual feature. This provides transparency and insights into the factors influencing the predictions, which can be used by risk and compliance teams and external regulators. Autopilot also provides a model performance report, which encompasses a summary of evaluation metrics, a confusion matrix, various visualizations such as receiver operating characteristic curves and precision-recall curves, and more. The specific content of each report vary depending on the problem type of the Autopilot experiment.

The explainability and performance reports for the best model candidate in an Autopilot experiment are available for text, image, and tabular data classification problem types.

For tabular data use cases such as regression or classification, Autopilot offers additional visibility into how the data was wrangled and how the model candidates were selected, trained, and tuned by generating notebooks that contain the code used to explore the data and find the best performing model. These notebooks provide an interactive and exploratory environment to help
you learn about the impact of various inputs or the trade-offs made in the experiments. You can experiment further with the higher performing model candidate by making your own modifications to the data exploration and candidate definition notebooks provided by Autopilot.

With Amazon SageMaker, you pay only for what you use. You pay for the underlying compute and storage resources within SageMaker or other AWS services, based on your usage. For more information about the cost of using SageMaker, see Amazon SageMaker Pricing.

Topics

- Create a regression or classification job for tabular data using the AutoML API
- Create an AutoML job for image classification using the API
- Create an AutoML job for text classification using the API
- Create an AutoML job for time-series forecasting using the API
- Create an AutoML job to fine-tune text generation models using the API
- Create a Regression or Classification Autopilot experiment for tabular data using the Studio Classic UI
- Amazon SageMaker Autopilot example notebooks
- Amazon SageMaker Autopilot quotas
- API Reference guide for Amazon SageMaker Autopilot

Create a regression or classification job for tabular data using the AutoML API

You can create an Autopilot experiment for tabular data programmatically by calling the CreateAutoMLJobV2 API action in any language supported by Autopilot or the AWS CLI.

For information on how this API action translates into a function in the language of your choice, see the See Also section of CreateAutoMLJobV2 and choose an SDK. As an example, for Python users, see the full request syntax of create_auto_ml_job_v2 in AWS SDK for Python (Boto3).

⚠️ Note

CreateAutoMLJobV2 and DescribeAutoMLJobV2 are new versions of CreateAutoMLJob and DescribeAutoMLJob which offer backward compatibility. We recommend using the CreateAutoMLJobV2. CreateAutoMLJobV2 can manage tabular problem types identical to those of its previous version CreateAutoMLJob, as
well as non-tabular problem types such as image or text classification, or time-series forecasting.

At a minimum, all experiments on tabular data require the specification of the experiment name, providing locations for the input and output data, and specifying which target data to predict. Optionally, you can also specify the type of problem that you want to solve (regression, classification, multiclass classification), choose your modeling strategy (stacked ensembles or hyperparameters optimization), select the list of algorithms used by the Autopilot job to train the data, and more.

After the experiment runs, you can compare trials and delve into the details of the pre-processing steps, algorithms, and hyperparameter ranges of each model. You also have the option to download their explainability and performance reports. Use the provided notebooks to see the results of the automated data exploration or the candidate model definitions.

The following is a collection of mandatory and optional input request parameters for the CreateAutoMLJobV2 API action. You can find the alternative information for the previous version of this action, CreateAutoMLJob. However, we recommend using CreateAutoMLJobV2.

Find guidelines on how to migrate a CreateAutoMLJob to CreateAutoMLJobV2 in Migrate a CreateAutoMLJob to CreateAutoMLJobV2.

**Required parameters**

**CreateAutoMLJobV2**

When calling CreateAutoMLJobV2 to create an Autopilot experiment for tabular data, you must provide the following values:

- An AutoMLJobName to specify the name of your job.
- At least one AutoMLJobChannel in AutoMLJobInputDataConfig to specify your data source.
- Both an AutoMLJobObjective metric and your chosen type of supervised learning problem (binary classification, multiclass classification, regression) in AutoMLProblemTypeConfig, or none at all. For tabular data, you must choose TabularJobConfig as the type of AutoMLProblemTypeConfig. You set the supervised learning problem in the ProblemType attribute of TabularJobConfig.
• An **OutputDataConfig** to specify the Amazon S3 output path to store the artifacts of your AutoML job.

• A **RoleArn** to specify the ARN of the role used to access your data.

**CreateAutoMLJob**

When calling **CreateAutoMLJob** to create an AutoML experiment, you must provide the following four values:

• An **AutoMLJobName** to specify the name of your job.

• At least one **AutoMLChannel** in **InputDataConfig** to specify your data source.

• An **OutputDataConfig** to specify the Amazon S3 output path to store the artifacts of your AutoML job.

• A **RoleArn** to specify the ARN of the role used to access your data.

All other parameters are optional.

**Optional parameters**

The following sections provide details of some optional parameters that you can pass to your **CreateAutoMLJobV2** API action when using tabular data. You can find the alternative information for the previous version of this action, **CreateAutoMLJob**. However, we recommend using **CreateAutoMLJobV2**.

**How to set the training mode of an AutoML job**

For tabular data, the set of algorithms run on your data to train your model candidates is dependent on your modeling strategy (ENSEMBLING or HYPERPARAMETER_TUNING). The following details how to set this training mode.

If you keep blank (or null), the Mode is inferred based on the size of your dataset.

For information on Autopilot’s *stacked ensembles* and *hyperparameters optimization* training methods, see [Training modes and algorithm support](#).

**CreateAutoMLJobV2**

For tabular data, you must choose **TabularJobConfig** as the type of **AutoMLProblemTypeConfig**.
You can set the training method of an AutoML job V2 with the `TabularJobConfig.Mode` parameter.

CreateAutoMLJob

You can set the training method of an AutoML job with the `AutoMLJobConfig.Mode` parameter.

How to select features and algorithms for training an AutoML job

Features selection

Autopilot provides automatic data-preprocessing steps including feature selection and feature extraction. However, you can manually provide the features to be used in training with the `FeatureSpecificationS3Uri` attribute.

Selected features should be contained within a JSON file in the following format:

```
{ "FeatureAttributeNames": ["col1", "col2", ...] }
```

The values listed in `["col1", "col2", ...]` are case sensitive. They should be a list of strings containing unique values that are subsets of the column names in the input data.

Note

The list of columns provided as features cannot include the target column.

CreateAutoMLJobV2

For tabular data, you must choose `TabularJobConfig` as the type of `AutoMLProblemTypeConfig`.

You can set the URL to your selected features with the `TabularJobConfig.FeatureSpecificationS3Uri` parameter.

CreateAutoMLJob

You can set the `FeatureSpecificationS3Uri` attribute of `AutoMLCandidateGenerationConfig` within the `CreateAutoMLJob` API with the following format:
Algorithms selection

By default, your Autopilot job runs a pre-defined list of algorithms on your dataset to train model candidates. The list of algorithms depends on the training mode (ENSEMBLING or HYPERPARAMETER_TUNING) used by the job.

You can provide a subset of the default selection of algorithms.

CreateAutoMLJobV2

For tabular data, you must choose `TabularJobConfig` as the type of `AutoMLProblemTypeConfig`.

You can specify an array of selected AutoMLAlgorithms in the `AlgorithmsConfig` attribute of `CandidateGenerationConfig`.

The following is an example of an `AlgorithmsConfig` attribute listing exactly three algorithms ("xgboost", "fastai", "catboost") in its `AutoMLAlgorithms` field for the ensembling training mode.

```
{
    "AutoMLProblemTypeConfig": {
        "TabularJobConfig": {
            "Mode": "ENSEMBLING",
            "CandidateGenerationConfig": {
                "AlgorithmsConfig": [
                    {"AutoMLAlgorithms": ["xgboost", "fastai", "catboost"]}
                ]
            },
            },
        },
    },
}
```
CreateAutoMLJob

You can specify an array of selected AutoMLAlgorithms in the AlgorithmsConfig attribute of `AutoMLCandidateGenerationConfig`.

The following is an example of an AlgorithmsConfig attribute listing exactly three algorithms ("xgboost", "fastai", "catboost") in its AutoMLAlgorithms field for the ensembling training mode.

```json
{
   "AutoMLJobConfig": {
      "CandidateGenerationConfig": {
         "AlgorithmsConfig": [
            {
               "AutoMLAlgorithms": ["xgboost", "fastai", "catboost"]
            }
         ],
         "Mode": "ENSEMBLING"
      }
   }
}
```

For the list of available algorithms per training Mode, see `AutoMLAlgorithms`. For details on each algorithm, see Training modes and algorithm support.

How to specify the training and validation datasets of an AutoML job

You can provide your own validation dataset and custom data split ratio, or let Autopilot split the dataset automatically.

CreateAutoMLJobV2

Each `AutoMLJobChannel` object (see the required parameter `AutoMLJobInputDataConfig`) has a ChannelType, which can be set to either training or validation values that specify how the data is to be used when building a machine learning model. At least one data source must be provided and a maximum of two data sources is allowed: one for training data and one for validation data.

How you split the data into training and validation datasets depends on whether you have one or two data sources.

- If you only have one data source, the ChannelType is set to training by default and must have this value.
- If the ValidationFraction value in AutoMLDataSplitConfig is not set, 0.2 (20%) of the data from this source is used for validation by default.

- If the ValidationFraction is set to a value between 0 and 1, the dataset is split based on the value specified, where the value specifies the fraction of the dataset used for validation.

- If you have two data sources, the ChannelType of one of the AutoMLJobChannel objects must be set to training, the default value. The ChannelType of the other data source must be set to validation. The two data sources must have the same format, either CSV or Parquet, and the same schema. You must not set the value for the ValidationFraction in this case because all of the data from each source is used for either training or validation. Setting this value causes an error.

CreateAutoMLJob

Each AutoMLChannel object (see the required parameter InputDataConfig) has a ChannelType, which can be set to either training or validation values that specify how the data is to be used when building a machine learning model. At least one data source must be provided and a maximum of two data sources is allowed: one for training data and one for validation data.

How you split the data into training and validation datasets depends on whether you have one or two data sources.

- If you only have one data source, the ChannelType is set to training by default and must have this value.
  - If the ValidationFraction value in AutoMLDataSplitConfig is not set, 0.2 (20%) of the data from this source is used for validation by default.
  - If the ValidationFraction is set to a value between 0 and 1, the dataset is split based on the value specified, where the value specifies the fraction of the dataset used for validation.

- If you have two data sources, the ChannelType of one of the AutoMLChannel objects must be set to training, the default value. The ChannelType of the other data source must be set to validation. The two data sources must have the same format, either CSV or Parquet, and the same schema. You must not set the value for the ValidationFraction in this case because all of the data from each source is used for either training or validation. Setting this value causes an error.
For information on split and cross-validation in Autopilot see [Cross-validation in Autopilot](#).

**How to set the problem type of an AutoML job**

CreateAutoMLJobV2

For tabular data, you must choose `TabularJobConfig` as the type of `AutoMLProblemTypeConfig`.

You can further specify the type of supervised learning problem (binary classification, multiclass classification, regression) available for the model candidates of your AutoML job V2 with the `TabularJobConfig.ProblemType` parameter.

CreateAutoMLJob

You can set the type of problem on an AutoML job with the `CreateAutoPilot.ProblemType` parameter. This limits the kind of preprocessing and algorithms that Autopilot tries. After the job is finished, if you had set the `CreateAutoPilot.ProblemType`, then the `ResolvedAttribute.ProblemType` matches the `ProblemType` you set. If you keep it blank (or null), the `ProblemType` is inferred on your behalf.

**Note**

In some cases, Autopilot is unable to infer the `ProblemType` with high enough confidence, in which case you must provide the value for the job to succeed.

**How to add sample weights to an AutoML job**

You can add a sample weights column to your tabular dataset and then pass it to your AutoML job to request dataset rows to be weighted during training and evaluation.

Support for sample weights is available in [ensembling mode](#) only. Your weights should be numeric and non-negative. Data points with invalid or no weight value are excluded. For more information on the available objective metrics, see [Autopilot weighted metrics](#).

CreateAutoMLJobV2

For tabular data, you must choose `TabularJobConfig` as the type of `AutoMLProblemTypeConfig`. 

---

Create a Regression or Classification Job Using the AutoML API

81
To set sample weights when creating an experiment (see CreateAutoMLJobV2), you can pass the name of your sample weights column in the SampleWeightAttributeName attribute of the TabularJobConfig object. This ensures that your objective metric uses the weights for the training, evaluation, and selection of model candidates.

CreateAutoMLJob

To set sample weights when creating an experiment (see CreateAutoMLJob), you can pass the name of your sample weights column in the SampleWeightAttributeName attribute of the AutoMLChannel object. This ensures that your objective metric uses the weights for the training, evaluation, and selection of model candidates.

Migrate a CreateAutoMLJob to CreateAutoMLJobV2

We recommend users of CreateAutoMLJob to migrate to CreateAutoMLJobV2. This section explains the differences in the input parameters between CreateAutoMLJob and CreateAutoMLJobV2 by highlighting the changes in the position, name, or structure of the objects and attributes of the input request between the two versions.

- Request attributes that did not change between versions.

```json
{
    "AutoMLJobName": "string",
    "AutoMLJobObjective": {
        "MetricName": "string"
    },
    "ModelDeployConfig": {
        "AutoGenerateEndpointName": boolean,
        "EndpointName": "string"
    },
    "OutputDataConfig": {
        "KmsKeyId": "string",
        "S3OutputPath": "string"
    },
    "RoleArn": "string",
    "Tags": [
        {
            "Key": "string",
            "Value": "string"
        }
    ]
}
```
• Request attributes that changed position and structure between versions.

The following attributes changed position: DataSplitConfig, SecurityConfig, CompletionCriteria, Mode, FeatureSpecificationS3Uri, SampleWeightAttributeName, TargetAttributeName.

CreateAutoMLJob

```json
{
    "AutoMLJobConfig": {
        "Mode": "string",
        "CompletionCriteria": {
            "MaxAutoMLJobRuntimeInSeconds": number,
            "MaxCandidates": number,
            "MaxRuntimePerTrainingJobInSeconds": number
        },
        "DataSplitConfig": {
            "ValidationFraction": number
        },
        "SecurityConfig": {
            "EnableInterContainerTrafficEncryption": boolean,
            "VolumeKmsKeyId": "string",
            "VpcConfig": {
                "SecurityGroupIds": [ "string" ],
                "Subnets": [ "string" ]
            }
        },
        "CandidateGenerationConfig": {
            "FeatureSpecificationS3Uri": "string"
        },
        "GenerateCandidateDefinitionsOnly": boolean,
        "ProblemType": "string"
    }
}
```

CreateAutoMLJobV2

```json
{
    "AutoMLProblemTypeConfig": {
        "TabularJobConfig": {
            "Mode": "string",
            "CandidateGenerationConfig": {
                "FeatureSpecificationS3Uri": "string"
            }
        }
    }
}
```
The following attributes changed position and structure between versions.

The following JSON illustrates how `AutoMLJobConfig.CandidateGenerationConfig` of type `AutoMLCandidateGenerationConfig` moved to `AutoMLProblemTypeConfig.TabularJobConfig.CandidateGenerationConfig` of type `CandidateGenerationConfig` in V2.

CreateAutoMLJob

```json
{
    "AutoMLJobConfig": {
        "CandidateGenerationConfig": {
            "AlgorithmsConfig": [
                {
                    "AutoMLAlgorithms": [ "string" ]
                }
            ],
            "FeatureSpecificationS3Uri": "string"
        }
    }
}
```
CreateAutoMLJobV2

{  
    "AutoMLProblemTypeConfig": {  
        "TabularJobConfig": {  
            "CandidateGenerationConfig": {  
                "AlgorithmsConfig": [  
                    {  
                        "AutoMLAlgorithms": [ "string" ]  
                    }  
                ],  
            },  
        },  
    },  
},

- **Request attributes that changed name and structure.**

The following JSON illustrates how `InputDataConfig` (An array of `AutoMLChannel`) changed to `AutoMLJobInputDataConfig` (An array of `AutoMLJobChannel`) in V2. Note that the attributes `SampleWeightAttributeName` and `TargetAttributeName` move out of `InputDataConfig` and into `AutoMLProblemTypeConfig`.

CreateAutoMLJob

{  
    "InputDataConfig": [  
        {  
            "ChannelType": "string",  
            "CompressionType": "string",  
            "ContentType": "string",  
            "DataSource": {  
                "S3DataSource": {  
                    "S3DataFormat": "string",  
                    "S3Uri": "string"  
                }  
            },  
            "SampleWeightAttributeName": "string",  
            "TargetAttributeName": "string"  
        }  
    ]  
}
Autopilot datasets and problem types

For tabular data (that is data in which each column contains a feature with a specific data type and each row contains an observation), Autopilot gives you the option of specifying the type of supervised learning problem available for the model candidates of the AutoML job, such as binary classification or regression, or of detecting it on your behalf based on the data you provide.

Topics

- Autopilot datasets, data types, and formats
- Autopilot problem types

Autopilot datasets, data types, and formats

Autopilot supports tabular data formatted as CSV files or as Parquet files: each column contains a feature with a specific data type and each row contains an observation. The properties of these two file formats differ considerably.
• **CSV** (comma-separated-values) is a row-based file format that stores data in human readable plaintext which is a popular choice for data exchange as they are supported by a wide range of applications.

• **Parquet** is a column-based file format where the data is stored and processed more efficiently than row-based file formats. This makes them a better option for big data problems.

The **data types** accepted for columns include numerical, categorical, text, and time series that consists of strings of comma-separated numbers. If Autopilot detects it is dealing with **time series** sequences, it processes them through specialized feature transformers provided by the **tsfresh** library. This library takes the time series as an input and outputs a feature such as the highest absolute value of the time series or descriptive statistics on autocorrelation. These outputted features are then used as inputs to one of the three problem types.

Autopilot supports building machine learning models on large datasets up to hundreds of GBs. For details on the default resource limits for input datasets and how to increase them, see **Autopilot quotas**.

**Autopilot problem types**

For the tabular data, you further specify the type of supervised learning problems available for the model candidates as follows:

**Regression**

Regression estimates the values of a dependent target variable based on one or more other variables or attributes that are correlated with it. An example is the prediction of house prices using features like the number of bathrooms and bedrooms, square footage of the house and garden. Regression analysis can create a model that takes one or more of these features as an input and predicts the price of a house.

**Binary classification**

Binary classification is a type of supervised learning that assigns an individual to one of two predefined and mutually exclusive classes based on their attributes. It is supervised because the models are trained using examples where the attributes are provided with correctly labeled objects. A medical diagnosis for whether an individual has a disease or not based on the results of diagnostic tests is an example of binary classification.
Multiclass classification

Multiclass classification is a type of supervised learning that assigns an individual to one of several classes based on their attributes. It is supervised because the models are trained using examples where the attributes are provided with correctly labelled objects. An example is the prediction of the topic most relevant to a text document. A document may be classified as being about, say, religion or politics or finance, or about one of several other predefined topic classes.

Training modes and algorithm support

Autopilot supports different training modes and algorithms to address machine learning problems, report on quality and objective metrics, and to use cross-validation automatically, when needed.

Training modes

SageMaker Autopilot can automatically select the training method based on the dataset size, or you can select it manually. The choices are as follows:

- **Ensembling** – Autopilot uses the AutoGluon library to train several base models. To find the best combination for your dataset, ensemble mode runs 10 trials with different model and meta parameter settings. Then Autopilot combines these models using a stacking ensemble method to create an optimal predictive model. For a list of algorithms that Autopilot supports in ensembling mode for tabular data, see the following Algorithms support section.

- **Hyperparameter optimization (HPO)** – Autopilot finds the best version of a model by tuning hyperparameters using Bayesian optimization or multi-fidelity optimization while running training jobs on your dataset. HPO mode selects the algorithms that are most relevant to your dataset and selects the best range of hyperparameters to tune your models. To tune your models, HPO mode runs up to 100 trials (default) to find the optimal hyperparameters settings within the selected range. If your dataset size is less than 100 MB, Autopilot uses Bayesian optimization. Autopilot chooses multi-fidelity optimization if your dataset is larger than 100 MB. In multi-fidelity optimization, metrics are continuously emitted from the training containers. A trial that is performing poorly against a selected objective metric is stopped early. A trial that is performing well is allocated more resources.

For a list of algorithms that Autopilot supports in HPO mode, see the following Algorithm support section.
• **Auto** – Autopilot automatically chooses either ensembling mode or HPO mode based on your dataset size. If your dataset is larger than 100 MB, Autopilot chooses HPO. Otherwise, it chooses ensembling mode. Autopilot can fail to read the size of your dataset in the following cases.

• If you enable Virtual Private Cloud (VPC) mode, for an AutoML job but the S3 bucket containing the dataset only allows access from the VPC.

• The input `S3DataType` of your dataset is a `ManifestFile`.

• The input `S3Uri` contains more than 1000 items.

If Autopilot is unable to read your dataset size, it defaults to choosing HPO mode.

**Note**

For optimal runtime and performance, use ensemble training mode for datasets that are smaller than 100 MB.

**Algorithms support**

In **HPO mode**, Autopilot supports the following types of machine learning algorithms:

• **Linear learner** – A supervised learning algorithm that can solve either classification or regression problems.

• **XGBoost** – A supervised learning algorithm that attempts to accurately predict a target variable by combining an ensemble of estimates from a set of simpler and weaker models.

• Deep learning algorithm – A multilayer perceptron (MLP) and feedforward artificial neural network. This algorithm can handle data that is not linearly separable.

**Note**

You don't need to specify an algorithm to use for your machine learning problem. Autopilot automatically selects the appropriate algorithm to train.

In **ensembling mode**, Autopilot supports the following types of machine learning algorithms:
• **LightGBM** – An optimized framework that uses tree-based algorithms with gradient boosting. This algorithm uses trees that grow in breadth, rather than depth, and is highly optimized for speed.

• **CatBoost** – A framework that uses tree-based algorithms with gradient boosting. Optimized for handling categorical variables.

• **XGBoost** – A framework that uses tree-based algorithms with gradient boosting that grows in depth, rather than breadth.

• **Random Forest** – A tree-based algorithm that uses several decision trees on random sub-samples of the data with replacement. The trees are split into optimal nodes at each level. The decisions of each tree are averaged together to prevent overfitting and improve predictions.

• **Extra Trees** – A tree-based algorithm that uses several decision trees on the entire dataset. The trees are split randomly at each level. The decisions of each tree are averaged to prevent overfitting and to improve predictions. Extra trees add a degree of randomization in comparison to the random forest algorithm.

• **Linear Models** – A framework that uses a linear equation to model the relationship between two variables in observed data.

• Neural network torch – A neural network model that's implemented using Pytorch.

• Neural network fast.ai – A neural network model that's implemented using fast.ai.

---

### Metrics and validation

This guide shows metrics and validation techniques that you can use to measure machine learning model performance. Amazon SageMaker Autopilot produces metrics that measure the predictive quality of machine learning model candidates. The metrics calculated for candidates are specified using an array of `MetricDatum` types.

#### Autopilot metrics

The following list contains the names of the metrics that are currently available to measure model performance within Autopilot.

1. **Note**

   Autopilot supports sample weights. To learn more about sample weights and the available objective metrics, see [Autopilot weighted metrics](#).
The following are the available metrics.

**Accuracy**

The ratio of the number of correctly classified items to the total number of (correctly and incorrectly) classified items. It is used for both binary and multiclass classification. Accuracy measures how close the predicted class values are to the actual values. Values for accuracy metrics vary between zero (0) and one (1). A value of 1 indicates perfect accuracy, and 0 indicates perfect inaccuracy.

**AUC**

The area under the curve (AUC) metric is used to compare and evaluate binary classification by algorithms that return probabilities, such as logistic regression. To map the probabilities into classifications, these are compared against a threshold value.

The relevant curve is the receiver operating characteristic curve. The curve plots the true positive rate (TPR) of predictions (or recall) against the false positive rate (FPR) as a function of the threshold value, above which a prediction is considered positive. Increasing the threshold results in fewer false positives, but more false negatives.

AUC is the area under this receiver operating characteristic curve. Therefore, AUC provides an aggregated measure of the model performance across all possible classification thresholds. AUC scores vary between 0 and 1. A score of 1 indicates perfect accuracy, and a score of one half (0.5) indicates that the prediction is not better than a random classifier.

**BalancedAccuracy**

BalancedAccuracy is a metric that measures the ratio of accurate predictions to all predictions. This ratio is calculated after normalizing true positives (TP) and true negatives (TN) by the total number of positive (P) and negative (N) values. It is used in both binary and multiclass classification and is defined as follows: \(0.5 \times (TP/P + TN/N)\), with values ranging from 0 to 1. BalancedAccuracy gives a better measure of accuracy when the number of positives or negatives differ greatly from each other in an imbalanced dataset, such as when only 1% of email is spam.

**F1**

The F1 score is the harmonic mean of the precision and recall, defined as follows: \(F1 = \frac{2 \times (precision \times recall)}{precision + recall}\). It is used for binary classification into classes traditionally referred to as positive and negative. Predictions are said to be true when they match their actual (correct) class, and false when they do not.
Precision is the ratio of the true positive predictions to all positive predictions, and it includes the false positives in a dataset. Precision measures the quality of the prediction when it predicts the positive class.

Recall (or sensitivity) is the ratio of the true positive predictions to all actual positive instances. Recall measures how completely a model predicts the actual class members in a dataset.

F1 scores vary between 0 and 1. A score of 1 indicates the best possible performance, and 0 indicates the worst.

**F1macro**

The F1macro score applies F1 scoring to multiclass classification problems. It does this by calculating the precision and recall, and then taking their harmonic mean to calculate the F1 score for each class. Lastly, the F1macro averages the individual scores to obtain the F1macro score. F1macro scores vary between 0 and 1. A score of 1 indicates the best possible performance, and 0 indicates the worst.

**InferenceLatency**

Inference latency is the approximate amount of time between making a request for a model prediction to receiving it from a real time endpoint to which the model is deployed. This metric is measured in seconds and only available in ensembling mode.

**LogLoss**

Log loss, also known as cross-entropy loss, is a metric used to evaluate the quality of the probability outputs, rather than the outputs themselves. It is used in both binary and multiclass classification and in neural nets. It is also the cost function for logistic regression. Log loss is an important metric to indicate when a model makes incorrect predictions with high probabilities. Values range from 0 to infinity. A value of 0 represents a model that perfectly predicts the data.

**MAE**

The mean absolute error (MAE) is a measure of how different the predicted and actual values are, when they're averaged over all values. MAE is commonly used in regression analysis to understand model prediction error. If there is linear regression, MAE represents the average distance from a predicted line to the actual value. MAE is defined as the sum of absolute errors divided by the number of observations. Values range from 0 to infinity, with smaller numbers indicating a better model fit to the data.
MSE

The mean squared error (MSE) is the average of the squared differences between the predicted and actual values. It is used for regression. MSE values are always positive. The better a model is at predicting the actual values, the smaller the MSE value is.

Precision

Precision measures how well an algorithm predicts the true positives (TP) out of all of the positives that it identifies. It is defined as follows: Precision = TP/(TP+FP), with values ranging from zero (0) to one (1), and is used in binary classification. Precision is an important metric when the cost of a false positive is high. For example, the cost of a false positive is very high if an airplane safety system is falsely deemed safe to fly. A false positive (FP) reflects a positive prediction that is actually negative in the data.

PrecisionMacro

The precision macro computes precision for multiclass classification problems. It does this by calculating precision for each class and averaging scores to obtain precision for several classes. PrecisionMacro scores range from zero (0) to one (1). Higher scores reflect the model's ability to predict true positives (TP) out of all of the positives that it identifies, averaged across multiple classes.

R2

$R^2$, also known as the coefficient of determination, is used in regression to quantify how much a model can explain the variance of a dependent variable. Values range from one (1) to negative one (-1). Higher numbers indicate a higher fraction of explained variability. $R^2$ values close to zero (0) indicate that very little of the dependent variable can be explained by the model. Negative values indicate a poor fit and that the model is outperformed by a constant function. For linear regression, this is a horizontal line.

Recall

Recall measures how well an algorithm correctly predicts all of the true positives (TP) in a dataset. A true positive is a positive prediction that is also an actual positive value in the data. Recall is defined as follows: Recall = TP/(TP+FN), with values ranging from 0 to 1. Higher scores reflect a better ability of the model to predict true positives (TP) in the data. It is used in binary classification.

Recall is important when testing for cancer because it's used to find all of the true positives. A false positive (FP) reflects a positive prediction that is actually negative in the data. It is often...
insufficient to measure only recall, because predicting every output as a true positive yields a perfect recall score.

RecallMacro

The RecallMacro computes recall for multiclass classification problems by calculating recall for each class and averaging scores to obtain recall for several classes. RecallMacro scores range from 0 to 1. Higher scores reflect the model's ability to predict true positives (TP) in a dataset, whereas a true positive reflects a positive prediction that is also an actual positive value in the data. It is often insufficient to measure only recall, because predicting every output as a true positive will yield a perfect recall score.

RMSE

Root mean squared error (RMSE) measures the square root of the squared difference between predicted and actual values, and is averaged over all values. It is used in regression analysis to understand model prediction error. It's an important metric to indicate the presence of large model errors and outliers. Values range from zero (0) to infinity, with smaller numbers indicating a better model fit to the data. RMSE is dependent on scale, and should not be used to compare datasets of different sizes.

Metrics that are automatically calculated for a model candidate are determined by the type of problem being addressed.

Refer to the Amazon SageMaker API reference documentation for the list of available metrics supported by Autopilot.

Autopilot weighted metrics

Note

Autopilot supports sample weights in ensembling mode only for all available metrics with the exception of Balanced Accuracy and InferenceLatency. BalanceAccuracy comes with its own weighting scheme for imbalanced datasets that does not require sample weights. InferenceLatency does not support sample weights. Both objective Balanced Accuracy and InferenceLatency metrics ignore any existing sample weights when training and evaluating a model.
Users can add a sample weights column to their data to ensure that each observation used to train a machine learning model is given a weight corresponding to its perceived importance to the model. This is especially useful in scenarios in which the observations in the dataset have varying degrees of importance, or in which a dataset contains a disproportionate number of samples from one class compared to others. Assigning a weight to each observation based on its importance or greater importance to a minority class can help a model's overall performance, or ensure that a model is not biased toward the majority class.

For information about how to pass sample weights when creating an experiment in the Studio Classic UI, see Step 7 in Create an Autopilot experiment using Studio Classic.

For information about how to pass sample weights programmatically when creating an Autopilot experiment using the API, see How to add sample weights to an AutoML job in Create an Autopilot experiment programatically.

**Cross-validation in Autopilot**

Cross-validation is used in to reduce overfitting and bias in model selection. It is also used to assess how well a model can predict the values of an unseen validation dataset, if the validation dataset is drawn from the same population. This method is especially important when training on datasets that have a limited number of training instances.

Autopilot uses cross-validation to build models in hyperparameter optimization (HPO) and ensemble training mode. The first step in the Autopilot cross-validation process is to split the data into k-folds.

**K-fold splitting**

K-fold splitting is a method that separates an input training dataset into multiple training and validation datasets. The dataset is split into k equally-sized sub-samples called folds. Models are then trained on k−1 folds and tested against the remaining k\(^{th}\) fold, which is the validation dataset. The process is repeated k times using a different data set for validation.

The following image depicts k-fold splitting with k = 4 folds. Each fold is represented as a row. The dark-toned boxes represent the parts of the data used in training. The remaining light-toned boxes indicate the validation datasets.
Autopilot uses k-fold cross-validation for both hyperparameter optimization (HPO) mode and ensembling mode.

You can deploy Autopilot models that are built using cross-validation like you would with any other Autopilot or SageMaker model.

**HPO mode**

K-fold cross-validation uses the k-fold splitting method for cross-validation. In HPO mode, Autopilot automatically implements k-fold cross-validation for small datasets with 50,000 or fewer training instances. Performing cross-validation is especially important when training on small datasets because it protects against overfitting and selection bias.

HPO mode uses a $k$ value of 5 on each of the candidate algorithms that are used to model the dataset. Multiple models are trained on different splits, and the models are stored separately. When training is complete, validation metrics for each of the models are averaged to produce a single estimation metric. Lastly, Autopilot combines the models from the trial with the best validation metric into an ensemble model. Autopilot uses this ensemble model to make predictions.

The validation metric for the models trained by Autopilot is presented as the objective metric in the model leaderboard. Autopilot uses the default validation metric for each problem type that it
handles, unless you specify otherwise. For the list of all metrics that Autopilot uses, see Autopilot metrics.

For example, the Boston Housing dataset contains only 861 samples. If you build a model to predict house sale prices using this dataset without cross-validation, you risk training on a dataset that is not representative of the Boston housing stock. If you split the data only once into training and validation subsets, the training fold may only contain data mainly from the suburbs. As a result, you would train on data that isn't representative of the rest of the city. In this example, your model would likely overfit on this biased selection. K-fold cross-validation can reduce the risk of this kind of error by making full and randomized use of the available data for both training and validation.

Cross-validation can increase training times by an average of 20%. Training times may also increase significantly for complex datasets.

**Note**

In HPO mode, you can see the training and validation metrics from each fold in your /aws/sagemaker/TrainingJobs CloudWatch Logs. For more information about CloudWatch Logs, see Log Amazon SageMaker Events with Amazon CloudWatch.

**Ensembling mode**

**Note**

Autopilot supports sample weights in ensembling mode. For the list of available metrics supporting sample weights, see Autopilot metrics.

In ensembling mode, cross-validation is performed regardless of dataset size. Customers can either provide their own validation dataset and custom data split ratio, or let Autopilot split the dataset automatically into an 80-20% split ratio. The training data is then split into k-folds for cross-validation, where the value of k is determined by the AutoGluon engine. An ensemble consists of multiple machine learning models, where each model is known as the base model. A single base model is trained on (k-1) folds and makes out-of-fold predictions on the remaining fold. This process is repeated for all k folds, and the out-of-fold (OOF) predictions are concatenated to form a single set of predictions. All base models in the ensemble follow this same process of generating OOF predictions.
The following image depicts k-fold validation with \( k = 4 \) folds. Each fold is represented as a row. The dark-toned boxes represent the parts of the data used in training. The remaining light-toned boxes indicate the validation datasets.

In the upper part of the image, in each fold, the first base model makes predictions on the validation dataset after training on the training datasets. At each subsequent fold, the datasets change roles. A dataset that was previously used for training is now used for validation, and this also applies in reverse. At the end of \( k \) folds, all of the predictions are concatenated to form a single set of predictions called an out-of-fold (OOF) prediction. This process is repeated for each \( n \) base models.

The OOF predictions for each base model are then used as features to train a stacking model. The stacking model learns the importance weights for each base model. These weights are used to combine the OOF predictions to form the final prediction. Performance on the validation dataset determines which base or stacking model is the best, and this model is returned as the final model.

In ensemble mode, you can either provide your own validation dataset or let Autopilot split the input dataset automatically into 80% train and 20% validation datasets. The training data is then split into \( k \)-folds for cross-validation and produces an OOF prediction and a base model for each fold.

These OOF predictions are used as features to train a stacking model, which simultaneously learns weights for each base model. These weights are used to combine the OOF predictions to form the final prediction. The validation datasets for each fold are used for hyperparameter tuning of all
base models and the stacking model. Performance on the validation datasets determines which base or stacking model is the best model, and this model is returned as the final model.

Amazon SageMaker Autopilot model deployment and prediction

This Amazon SageMaker Autopilot guide includes steps for model deployment, setting up real-time inference, and running inference with batch jobs.

After you train your Autopilot models, you can deploy them to get predictions in one of two ways:

1. Use Real-time inferencing to set up an endpoint and obtain predictions interactively.
2. Use Batch inferencing to make predictions in parallel on batches of observations on an entire dataset.

Note

To avoid incurring unnecessary charges: After the endpoints and resources that were created from model deployment are no longer needed, you can delete them. For information about pricing of instances by Region, see Amazon SageMaker Pricing.

Real-time inferencing

Real-time inference is ideal for inference workloads where you have real-time, interactive, low latency requirements. This section shows how you can use real-time inferencing to obtain predictions interactively from your model.

To deploy the model that produced the best validation metric in an Autopilot experiment, you have several options. For example, when using Autopilot in SageMaker Studio Classic, you can deploy the model automatically or manually. You can also use SageMaker APIs to manually deploy an Autopilot model.

The following tabs show three options for deploying your model. These instructions assume that you have already created a model in Autopilot. If you don't have a model, see Create a regression or classification job for tabular data using the AutoML API. To see examples for each option, open each tab.
Deploy using the Autopilot User Interface (UI)

The Autopilot UI contains helpful dropdown menus, toggles, tooltips, and more to help you navigate through model deployment. You can deploy using either one of the following procedures: Automatic or Manual.

- **Automatic Deployment**: To automatically deploy the best model from an Autopilot experiment to an endpoint
  2. Toggle the Auto deploy value to Yes.

  **Note**
  Automatic deployment will fail if either the default resource quota or your customer quota for endpoint instances in a Region is too limited. In hyperparameter optimization (HPO) mode, you are required to have at least two ml.m5.2xlarge instances. In ensembling mode, you are required to have at least one ml.m5.12xlarge instance. If you encounter a failure related to quotas, you can request a service limit increase for SageMaker endpoint instances.

- **Manual Deployment**: To manually deploy the best model from an Autopilot experiment to an endpoint
  2. Toggle the Auto deploy value to No.
  3. Select the model that you want to deploy under Model name.
  4. Select the orange **Deployment and advanced settings** button located on the right of the leaderboard. This opens a new tab.
  5. Configure the endpoint name, instance type, and other optional information.
  6. Select the orange **Deploy model** to deploy to an endpoint.
  7. Check the progress of the endpoint creation process in the [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/) by navigating to the Endpoints section. That section is located in the **Inference** dropdown menu in the navigation panel.
  8. After the endpoint status changes from **Creating** to **InService**, as shown below, return to Studio Classic and invoke the endpoint.

Create a Regression or Classification Job Using the AutoML API

100
Deploy using SageMaker APIs

You can also obtain real-time inference by deploying your model using API calls. This section shows the five steps of this process using AWS Command Line Interface (AWS CLI) code snippets.

For complete code examples for both AWS CLI commands and AWS SDK for Python (boto3), open the tabs directly following these steps.

1. **Obtain candidate definitions**

   Obtain the candidate container definitions from `InferenceContainers`. These candidate definitions are used to create a SageMaker model.

   The following example uses the `DescribeAutoMLJob` API to obtain candidate definitions for the best model candidate. See the following AWS CLI command as an example.

   ```
   aws sagemaker describe-auto-ml-job --auto-ml-job-name <job-name> --region <region>
   ```

2. **List candidates**

   The following example uses the `ListCandidatesForAutoMLJob` API to list all candidates. See the following AWS CLI command as an example.

   ```
   aws sagemaker list-candidates-for-auto-ml-job --auto-ml-job-name <job-name> --region <region>
   ```

3. **Create a SageMaker model**

   Use the container definitions from the previous steps to create a SageMaker model by using the `CreateModel` API. See the following AWS CLI command as an example.
4. Create an endpoint configuration

The following example uses the CreateEndpointConfig API to create an endpoint configuration. See the following AWS CLI command as an example.

```
aws sagemaker create-endpoint-config --endpoint-config-name '<your-custom-endpoint-config-name>' \
  --production-variants '<list-of-production-variants>' \
  --region '<region>'
```

5. Create the endpoint

The following AWS CLI example uses the CreateEndpoint API to create the endpoint.

```
aws sagemaker create-endpoint --endpoint-name '<your-custom-endpoint-name>' \
  --endpoint-config-name '<endpoint-config-name-you-just-created>' \
  --region '<region>'
```

Check the progress of your endpoint deployment by using the DescribeEndpoint API. See the following AWS CLI command as an example.

```
aws sagemaker describe-endpoint --endpoint-name '<endpoint-name>' --region <region>
```

After the EndpointStatus changes to InService, the endpoint is ready to use for real-time inference.

6. Invoke the endpoint

The following command structure invokes the endpoint for real-time inferencing.

```
aws sagemaker invoke-endpoint --endpoint-name '<endpoint-name>' \
  --region '<region>' --body '<your-data>' [--content-type] '<content-type>' <outfile>
```
The following tabs contain complete code examples for deploying a model with AWS SDK for Python (boto3) or the AWS CLI.

AWS SDK for Python (boto3)

1. **Obtain the candidate definitions** by using the following code example.

```python
import sagemaker
import boto3

session = sagemaker.session.Session()

sagemaker_client = boto3.client('sagemaker', region_name='us-west-2')
job_name = 'test-auto-ml-job'

describe_response = sm_client.describe_auto_ml_job(AutoMLJobName=job_name)
# extract the best candidate definition from DescribeAutoMLJob response
best_candidate = describe_response['BestCandidate']
# extract the InferenceContainers definition from the candidate definition
inference_containers = best_candidate['InferenceContainers']
```

2. **Create the model** by using the following the code example.

```python
# Create Model
model_name = 'test-model'
sagemaker_role = 'arn:aws:iam:44445556666:role/sagemaker-execution-role'
create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    Containers = inference_containers
)
```

3. **Create the endpoint configuration** by using the following the code example.

```python
endpoint_config_name = 'test-endpoint-config'

instance_type = 'ml.m5.2xlarge'
# for all supported instance types, see
# https://docs.aws.amazon.com/sagemaker/latest/APIReference/
# API_ProductionVariant.html#sagemaker-Type-ProductionVariant-InstanceType
# Create endpoint config
endpoint_config_response = sagemaker_client.create_endpoint_config(
```
4. Create the endpoint and deploy the model with the following code example.

```python
# create endpoint and deploy the model
deployment_config = {
    "VariantName": "variant1",
    "ModelName": model_name,
    "InstanceType": instance_type,
    "InitialInstanceCount": 1
}

create_endpoint_response = sagemaker_client.create_endpoint(EndpointName=endpoint_name,
                                                            ProductionVariant=deployment_config)

print(f"Created EndpointConfig: {endpoint_config_response['EndpointConfigArn']}")
```

5. Invoke the endpoint for real-time inferencing by using the following command structure.

```bash
# once endpoint status is InService, you can invoke the endpoint for inferencing
if status == "InService":
    sm_runtime = boto3.Session().client('sagemaker-runtime')
    inference_result = sm_runtime.invoke_endpoint(EndpointName='test-endpoint',
                                                  ContentType='text/csv',
                                                  Body='1,2,3,4,class')
```

**AWS Command Line Interface (AWS CLI)**

1. Obtain the candidate definitions by using the following code example.
2. **Create the model** by using the following code example.

```bash
aws sagemaker describe-auto-ml-job --auto-ml-job-name 'test-automl-job' --region us-west-2
```

```bash
aws sagemaker create-model --model-name 'test-sagemaker-model' --containers '[
    {
        "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-sklearn-automl:2.5-1-cpu-py3", DOC-EXAMPLE-BUCKET1
        "ModelDataUrl": "s3://DOC-EXAMPLE-BUCKET/output/model.tar.gz",
        "Environment": {
            "AUTOML_SPARSE_ENCODE_RECORDIO_PROTOBUF": "1",
            "AUTOML_TRANSFORM_MODE": "feature-transform",
            "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "application/x-recordio-protobuf",
            "SAGEMAKER_PROGRAM": "sagemaker_serve",
            "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code"
        }
    }, {
        "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-xgboost:1.3-1-cpu-py3",
        "ModelDataUrl": "s3://DOC-EXAMPLE-BUCKET/output/model.tar.gz",
        "Environment": {
            "MAX_CONTENT_LENGTH": "20971520",
            "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "text/csv",
            "SAGEMAKER_INFERENCE_OUTPUT": "predicted_label",
            "SAGEMAKER_INFERENCE_SUPPORTED": "predicted_label,probability,probabilities"
        }
    }, {
        "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-sklearn-automl:2.5-1-cpu-py3", aws-region
        "ModelDataUrl": "s3://DOC-EXAMPLE-BUCKET/output/model.tar.gz",
        "Environment": {
            "AUTOML_TRANSFORM_MODE": "inverse-label-transform",
            "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "text/csv",
            "SAGEMAKER_INFERENCE_INPUT": "predicted_label",
            "SAGEMAKER_INFERENCE_OUTPUT": "predicted_label",
            "SAGEMAKER_INFERENCE_SUPPORTED": "predicted_label,probability,labels,probabilities",
            "SAGEMAKER_PROGRAM": "sagemaker_serve",
            "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code"
        }
    }
]
```

**Create a Regression or Classification Job Using the AutoML API**
For additional details, see creating a model.

The create model command will return a response in the following format.

```json
{
}
```

3. Create an endpoint configuration by using the following code example.

```bash
aws sagemaker create-endpoint-config --endpoint-config-name 'test-endpoint-config' \
--production-variants '[["VariantName": "variant1", 
    "ModelName": "test-sagemaker-model", 
    "InitialInstanceCount": 1, 
    "InstanceType": "ml.m5.2xlarge" 
}}' \
--region us-west-2
```

The create endpoint configuration command will return a response in the following format.

```json
{
}
```

4. Create an endpoint by using the following code example.

```bash
aws sagemaker create-endpoint --endpoint-name 'test-endpoint' \
--endpoint-config-name 'test-endpoint-config' \
--region us-west-2
```

The create endpoint command will return a response in the following format.

```json
{
}
```
Check the progress of the endpoint deployment by using the following `describe-endpoint` CLI code example.

```bash
aws sagemaker describe-endpoint --endpoint-name 'test-endpoint' --region us-west-2
```

The previous progress check will return a response in the following format.

```json
{
  "EndpointName": "test-endpoint",
  "EndpointConfigName": "test-endpoint-config",
  "EndpointStatus": "Creating",
  "CreationTime": 1660251167.595,
  "LastModifiedTime": 1660251167.595
}
```

After the `EndpointStatus` changes to `InService`, the endpoint is ready for use in real-time inference.

5. **Invoke the endpoint** for real-time inferencing by using the following command structure.

```bash
aws sagemaker-runtime invoke-endpoint --endpoint-name 'test-endpoint' \
  --region 'us-west-2' \
  --body '1,51,3.5,1.4,0.2' \
  --content-type 'text/csv' \
  '/tmp/inference_output'
```

For more options, see [invoking an endpoint](#).

### Deploy models from different accounts

You can deploy an Autopilot model from a different account than the original account that a model was generated in. To implement cross-account model deployment, this section shows how to do the following:

1. **Grant permission to the deploying account**
To assume the role in the generating account, you must grant permission to the deploying account. This allows the deploying account to describe Autopilot jobs in the generating account.

The following example uses a generating account with a trusted sagemaker-role entity. The example shows how to give a deploying account with the ID 111122223333 permission to assume the role of the generating account.

```
"Statement": [

  
  "Effect": "Allow",
  "Principal": {
    "Service": [
      "sagemaker.amazonaws.com"
    ],
    "AWS": [ "111122223333"
  ],
  "Action": "sts:AssumeRole"
]
```

The new account with the ID 111122223333 can now assume the role for the generating account.

Next, call the DescribeAutoMLJob API from the deploying account to obtain a description of the job created by the generating account.

The following code example describes the model from the deploying account.

```
import sagemaker
import boto3
session = sagemaker.session.Session()

sts_client = boto3.client('sts')
sts_client.assume_role

role = 'arn:aws:iam::111122223333:role/sagemaker-role'
role_session_name = "role-session-name"
_assumed_role = sts_client.assume_role(RoleArn=role,
                                     RoleSessionName=role_session_name)

credentials = _assumed_role["Credentials"]
access_key = credentials["AccessKeyId"]
```
```python
secret_key = credentials["SecretAccessKey"]
session_token = credentials["SessionToken"]

session = boto3.session.Session()

sm_client = session.client('sagemaker', region_name='us-west-2',
                             aws_access_key_id=access_key,
                             aws_secret_access_key=secret_key,
                             aws_session_token=session_token)

# now you can call describe automl job created in account A

job_name = "test-job"
response= sm_client.describe_auto_ml_job(AutoMLJobName=job_name)
```

2. **Grant access to the deploying account** to the model artifacts in the generating account.

The deploying account only needs access to the model artifacts in the generating account to deploy it. These are located in the **S3OutputPath** that was specified in the original `CreateAutoMLJob` API call during model generation.

To give the deploying account access to the model artifacts, choose one of the following options:

a. **Give access** to the `ModelDataUrl` from the generating account to the deploying account.

   Next, you need to give the deploying account permission to assume the role. follow the **real-time inferencing steps** to deploy.

b. **Copy model artifacts** from the generating account's original **S3OutputPath** to the generating account.

To grant access to the model artifacts, you must define a `best_candidate` model and reassign model containers to the new account.

The following example shows how to define a `best_candidate` model and reassign the `ModelDataUrl`.

```python
best_candidate = automl.describe_auto_ml_job()["BestCandidate"]

# reassigning ModelDataUrl for best_candidate containers below
new_model_locations = ['new-container-1-ModelDataUrl', 'new-container-2-ModelDataUrl', 'new-container-3-ModelDataUrl']
```
After this assignment of containers, follow the steps in Deploy using SageMaker APIs to deploy.

To build a payload in real-time inferencing, see the notebook example to define a test payload. To create the payload from a CSV file and invoke an endpoint, see the Predict with your model section in Create a machine learning model automatically.

**Batch inferencing**

Batch inferencing, also known as offline inferencing, generates model predictions on a batch of observations. Batch inference is a good option for large datasets or if you don't need an immediate response to a model prediction request.

By contrast, online inference (real-time inferencing) generates predictions in real time.

You can make batch inferences from an Autopilot model using the SageMaker Python SDK, the Autopilot user interface (UI), the AWS SDK for Python (boto3), or the AWS Command Line Interface (AWS CLI).

The following tabs show three options for deploying your model: Using APIs, Autopilot UI, or using APIs to deploy from different accounts. These instructions assume that you have already created a model in Autopilot. If you don't have a model, see Create a regression or classification job for tabular data using the AutoML API. To see examples for each option, open each tab.

**Deploy a model using Autopilot UI**

The Autopilot UI contains helpful dropdown menus, toggles, tooltips, and more to help you navigate through model deployment.

The following steps show how to deploy a model from an Autopilot experiment for batch predictions.

2. On the left navigation pane, choose Studio.
3. Under **Get started**, select the Domain that you want to launch the Studio application in. If your user profile only belongs to one Domain, you do not see the option for selecting a Domain.

4. Select the user profile that you want to launch the Studio Classic application for. If there is no user profile in the domain, choose **Create user profile**. For more information, see Add and Remove User Profiles.

5. Choose **Launch Studio**. If the user profile belongs to a shared space, choose **Open Spaces**.

6. When the SageMaker Studio Classic console opens, choose the **Launch SageMaker Studio** button.

7. Select **AutoML** from the left navigation pane.

8. Under **Name**, select the Autopilot experiment corresponding to the model that you want to deploy. This opens a new **AUTOPILOT JOB** tab.

9. In the **Model name** section, select the model that you want to deploy.

10. Choose **Deploy model**. This opens a new tab.

11. Choose **Make batch predictions** at the top of the page.

12. For **Batch transform job configuration**, input the **Instance type**, **Instance count** and other optional information.

13. In the **Input data configuration** section, open the dropdown menu.

   a. For **S3 data type**, choose **ManifestFile** or **S3Prefix**.

   b. For **Split type**, choose **Line**, **RecordIO**, **TFRecord** or **None**.

   c. For **Compression**, choose **Gzip** or **None**.

14. For **S3 location**, enter the Amazon S3 bucket location of the input data and other optional information.

15. Under **Output data configuration**, enter the S3 bucket for the output data, and choose how to assemble the output of your job.

   a. For **Additional configuration (optional)**, you can enter a MIME type and an **S3 Encryption key**.

16. For **Input/output filtering and data joins (optional)**, you enter a JSONpath expression to filter your input data, join the input source data with your output data, and enter a JSONpath expression to filter your output data.

   a. For examples for each type of filter, see the DataProcessing API.

17. To perform batch predictions on your input dataset, select **Create batch transform job**. A new **Batch Transform Jobs** tab appears.
18 In the **Batch Transform Jobs** tab: Locate the name of your job in **Status** section. Then check the progress of the job.

**Deploy using SageMaker APIs**

To use the SageMaker APIs for batch inferencing, there are three steps:

1. **Obtain candidate definitions**

   Candidate definitions from [InferenceContainers](https://example.com) are used to create a SageMaker model.

   The following example shows how to use the [DescribeAutoMLJob](https://example.com) API to obtain candidate definitions for the best model candidate. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker describe-auto-ml-job --auto-ml-job-name <job-name> --region <region>
   ```

   Use the [ListCandidatesForAutoMLJob](https://example.com) API to list all candidates. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker list-candidates-for-auto-ml-job --auto-ml-job-name <job-name> --region <region>
   ```

2. **Create a SageMaker model**

   To create a SageMaker model using the [CreateModel](https://example.com) API, use the container definitions from the previous steps. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker create-model --model-name '<your-custom-model-name>'
   --containers ['<container-definition1>', '<container-definition2>', '<container-definition3>']
   --execution-role-arn '<execution-role-arn>' --region '<region>'
   ```

3. **Create a SageMaker transform job**

   The following example creates a SageMaker transform job with the [CreateTransformJob](https://example.com) API. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker create-transform-job --transform-job-name '<your-custom-transform-job-name>'
   --model-name '<your-custom-model-name-from-last-step>'
   --transform-input '{
       "DataSource": {
   ```
Check the progress of your transform job using the [DescribeTransformJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_DescribeTransformJob.html) API. See the following AWS CLI command as an example.

```bash
aws sagemaker describe-transform-job --transform-job-name '<your-custom-transform-job-name>' --region '<region>'
```

After the job is finished, the predicted result will be available in `<your-output-path>`.

The output file name has the following format: `<input_data_file_name>.out`. As an example, if your input file is `text_x.csv`, the output name will be `text_x.csv.out`.

The following tabs show code examples for SageMaker Python SDK, AWS SDK for Python (boto3), and the AWS CLI.

### SageMaker Python SDK

The following example uses the [SageMaker Python SDK](https://docs.aws.amazon.com/sagemaker/latest/dg/SDK.html) to make predictions in batches.

```python
from sagemaker import AutoML

sagemaker_session= sagemaker.session.Session()

job_name = 'test-auto-ml-job' # your autopilot job name
automl = AutoML.attach(auto_ml_job_name=job_name)
output_path = 's3://test-auto-ml-job/output'
```

Create a Regression or Classification Job Using the AutoML API
input_data = 's3://test-auto-ml-job/test_X.csv'

# call DescribeAutoMLJob API to get the best candidate definition
best_candidate = automl.describe_auto_ml_job()['BestCandidate']
best_candidate_name = best_candidate['CandidateName']

# create model
model = automl.create_model(name=best_candidate_name,
                             candidate=best_candidate)

# create transformer
transformer = model.transformer(instance_count=1,
                                 instance_type='ml.m5.2xlarge',
                                 assemble_with='Line',
                                 output_path=output_path)

# do batch transform
transformer.transform(data=input_data,
                      split_type='Line',
                      content_type='text/csv',
                      wait=True)

AWS SDK for Python (boto3)

The following example uses **AWS SDK for Python (boto3)** to make predictions in batches.

```python
import sagemaker
import boto3

session = sagemaker.session.Session()

sm_client = boto3.client('sagemaker', region_name='us-west-2')
role = 'arn:aws:iam::1234567890:role/sagemaker-execution-role'
output_path = 's3://test-auto-ml-job/output'
input_data = 's3://test-auto-ml-job/test_X.csv'

best_candidate = sm_client.describe_auto_ml_job(AutoMLJobName=job_name)
['BestCandidate']
best_candidate_containers = best_candidate['InferenceContainers']
best_candidate_name = best_candidate['CandidateName']

# create model
response = sm_client.create_model(
    ModelName = best_candidate_name,
    ...)
```
ExecutionRoleArn = role,
Containers = best_candidate_containers
)

# Launch Transform Job
response = sm_client.create_transform_job(
    TransformJobName=f'{best_candidate_name}-transform-job',
    ModelName=model_name,
    TransformInput={
        'DataSource': {
            'S3DataSource': {
                'S3DataType': 'S3Prefix',
                'S3Uri': input_data
            }
        },
        'ContentType': "text/csv",
        'SplitType': 'Line'
    },
    TransformOutput={
        'S3OutputPath': output_path,
        'AssembleWith': 'Line',
    },
    TransformResources={
        'InstanceType': 'ml.m5.2xlarge',
        'InstanceCount': 1,
    },
)

The batch inference job returns a response in the following format.

```json
 "ResponseMetadata": {'RequestId': '659f97fc-28c4-440b-b957-a49733f7c2f2',
 'HTTPStatusCode': 200,
 'HTTPHeaders': {'x-amzn-requestid': '659f97fc-28c4-440b-b957-a49733f7c2f2',
 'content-type': 'application/x-amz-json-1.1',
 'content-length': '96',
 'date': 'Thu, 11 Aug 2022 22:23:49 GMT'},
 'RetryAttempts': 0}
```

AWS Command Line Interface (AWS CLI)

1. **Obtain the candidate definitions** by using the following the code example.
2. **Create the model** by using the following the code example.

```bash
aws sagemaker create-model --model-name 'test-sagemaker-model'
--containers '[{
  "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-sklearn-automl:2.5-1-cpu-py3",
  "ModelDataUrl": "s3://test-bucket/out/test-job1/data-processor-models/test-job1-dpp0-1-e569ff7ad77f4e55a7e549a/output/model.tar.gz",
  "Environment": {
    "AUTOML_SPARSE_ENCODE_RECORDIO_PROTOBUF": "1",
    "AUTOML_TRANSFORM_MODE": "feature-transform",
    "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "application/x-recordio-protobuf",
    "SAGEMAKER_PROGRAM": "sagemaker_serve",
    "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code"
  }
}, {
  "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-xgboost:1.3-1-cpu-py3",
  "ModelDataUrl": "s3://test-bucket/out/test-job1/tuning/flicdf10v2-dpp0-xgb/test-job1E9-244-7490a1c0/output/model.tar.gz",
  "Environment": {
    "MAX_CONTENT_LENGTH": "20971520",
    "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "text/csv",
    "SAGEMAKER_INFERENCE_INPUT": "predicted_label",
    "SAGEMAKER_INFERENCE_OUTPUT": "predicted_label",
    "SAGEMAKER_INFERENCE_SUPPORTED": "predicted_label,probability,labels,probabilities"
  }
}, {
  "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-sklearn-automl:2.5-1-cpu-py3",
  "ModelDataUrl": "s3://test-bucket/out/test-job1/data-processor-models/test-job1-dpp0-1-e569ff7ad77f4e55a7e549a/output/model.tar.gz",
  "Environment": {
    "AUTOML_TRANSFORM_MODE": "inverse-label-transform",
    "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "text/csv",
    "SAGEMAKER_INFERENCE_INPUT": "predicted_label",
    "SAGEMAKER_INFERENCE_OUTPUT": "predicted_label",
    "SAGEMAKER_INFERENCE_SUPPORTED": "predicted_label,probability,labels,probabilities"
  }
}
]

aws sagemaker describe-auto-ml-job --auto-ml-job-name 'test-automl-job' --region us-west-2

Create a Regression or Classification Job Using the AutoML API
3. **Create the transform job** by using the following the code example.

```bash
aws sagemaker create-transform-job --transform-job-name 'test-transform-job' --model-name 'test-sagemaker-model' --transform-input '{
    "DataSource": {
        "S3DataSource": {
            "S3DataType": "S3Prefix",
            "S3Uri": "s3://test-bucket/data.csv"
        }
    },
    "ContentType": "text/csv",
    "SplitType": "Line"
}'
--transform-output '{
    "S3OutputPath": "s3://test-bucket/output/",
    "AssembleWith": "Line"
}'
--transform-resources '{
    "InstanceType": "ml.m5.2xlarge",
    "InstanceCount": 1
}'
--region 'us-west-2'
```

4. **Check the progress of the transform job** by using the following the code example.

```bash
aws sagemaker describe-transform-job --transform-job-name 'test-transform-job' --region us-west-2
```

The following is the response from the transform job.

```json
{
    "TransformJobName": "test-transform-job",
    "TransformJobStatus": "InProgress",
}
```
"ModelName": "test-model",
"TransformInput": {
    "DataSource": {
        "S3DataSource": {
            "S3DataType": "S3Prefix",
            "S3Uri": "s3://test-bucket/data.csv"
        }
    },
    "ContentType": "text/csv",
    "CompressionType": "None",
    "SplitType": "Line"
},
"TransformOutput": {
    "S3OutputPath": "s3://test-bucket/output/",
    "AssembleWith": "Line",
    "KmsKeyId": ""
},
"TransformResources": {
    "InstanceType": "ml.m5.2xlarge",
    "InstanceCount": 1
},
"CreationTime": 1662495635.679,
"TransformStartTime": 1662495847.496,
"DataProcessing": {
    "InputFilter": "$",
    "OutputFilter": "$",
    "JoinSource": "None"
}
}

After the TransformJobStatus changes to Completed, you can check the inference result in the S3OutputPath.

**Deploy models from different accounts**

To create a batch inferencing job in a different account than the one that the model was generated in, follow the instructions in Deploy models from different accounts. Then you can create models and transform jobs by following the Deploy using SageMaker APIs.
Models generated by Amazon SageMaker Autopilot

This procedure describes how to share a model that you created in Amazon SageMaker Autopilot with another user in SageMaker Canvas. It also shows how to view details about jobs that you've run.

Prerequisites

Before you begin this procedure, you must have created and run an Autopilot experiment. For instructions, see Create a regression or classification job for tabular data using the AutoML API.

Share your Autopilot model

You can share your Autopilot model with another user in SageMaker Canvas. The other user can then import your model and use it to generate predictions.

To share the model in the Autopilot user interface using a button, see the following section View model details. The Share Model button is discussed in Step 6.

For more information about how to share a model, see Bring Your Own Model Into Canvas.

View model details

Autopilot generates details about the candidate models that you can obtain. These details include the following:

- A plot of the aggregated SHAP values that indicate the importance of each feature. This helps explain your models predictions.
- The summary statistics for various training and validation metrics, including the objective metric.
- A list of the hyperparameters used to train and tune the model.

To view model details after running an Autopilot job, follow these steps:

1. Choose the Home icon from the left navigation pane to view the top-level Amazon SageMaker Studio Classic navigation menu.
2. Select the AutoML card from the main working area. This opens a new Autopilot tab.
3. In the Name section, select the Autopilot job that has the details that you want to examine. This opens a new Autopilot job tab.
4. The **Autopilot job** panel lists the metric values including the **Objective** metric for each model under **Model name**. The **Best model** is listed at the top of the list under **Model name** and is also highlighted in the **Models** tab.

   - To review model details, select the model that you are interested in and select **View model details**. This opens a new **Model Details** tab.

5. The **Model Details** tab is divided into four subsections.

   1. The top of the **Explainability** tab contains a plot of aggregated SHAP values that indicate the importance of each feature. Following that are the metrics and hyperparameter values for this model.
   
   2. The **Performance** tab contains metrics statistics a confusion matrix.
   
   3. The **Artifacts** tab contains information about model inputs, outputs, and intermediate results.
   
   4. The **Network** tab summarizes your network isolation and encryption choices.

   ![Note](image)

   Feature importance and information in the **Performance** tab is only generated for the **Best model**.

   For more information about how the SHAP values help explain predictions based on feature importance, see the whitepaper [Understanding the model explainability](#). Additional information is also available in the [Model Explainability](#) topic in the SageMaker Developer Guide.

6. To share your Autopilot model with another SageMaker Canvas user, choose **Share Model**. That button is located at the top right of the **Model Details** tab.

   - In the **Add Canvas users** section, use the down arrow to select a SageMaker Canvas user.

**View an Autopilot Model Performance Report**

An Amazon SageMaker model quality report (also referred to as performance report) provides insights and quality information for the best model candidate generated by an AutoML job. This includes information about the job details, model problem type, objective function, and other
information related to the problem type. This guide shows how to view Amazon SageMaker Autopilot performance metrics graphically, or view metrics as raw data in a JSON file.

For example, in classification problems, the model quality report includes the following:

- Confusion matrix
- Area under the receiver operating characteristic curve (AUC)
- Information to understand false positives and false negatives
- Tradeoffs between true positives and false positives
- Tradeoffs between precision and recall

Autopilot also provides performance metrics for all of your candidate models. These metrics are calculated using all of the training data and are used to estimate model performance. The main working area includes these metrics by default. The type of metric is determined by the type of problem being addressed.

Refer to the [Amazon SageMaker API reference documentation](#) for the list of available metrics supported by Autopilot.

You can sort your model candidates with the relevant metric to help you select and deploy the model that addresses your business needs. For definitions of these metrics, see the [Autopilot candidate metrics](#) topic.

To view a performance report from an Autopilot job, follow these steps:

1. Choose the **Home** icon

   from the left navigation pane to view the top-level [Amazon SageMaker Studio Classic](#) navigation menu.

2. Select the **AutoML** card from the main working area. This opens a new **Autopilot** tab.

3. In the **Name** section, select the Autopilot job that has the details that you want to examine. This opens a new **Autopilot job** tab.

4. The **Autopilot job** panel lists the metric values including the **Objective** metric for each model under **Model name**. The **Best model** is listed at the top of the list under **Model name** and it is highlighted in the **Models** tab.
To review model details, select the model that you are interested in and select **View in model details**. This opens a new **Model Details** tab.

5. Choose the **Performance** tab between the **Explainability** and **Artifacts** tab.

   a. On the top right section of the tab, select the down arrow on the **Download Performance Reports** button.

   b. The down arrow provides two options to view Autopilot performance metrics:
      
      i. You can download a PDF of the performance report to view the metrics graphically.
      
      ii. You can view metrics as raw data and download it as a JSON file.

For instructions on how to create and run an AutoML job in SageMaker Studio Classic, see [Create a regression or classification job for tabular data using the AutoML API](#).

The performance report contains two sections. The first contains details about the Autopilot job that produced the model. The second section contains a model quality report.

**Autopilot Job details**

This first section of the report gives some general information about the Autopilot job that produced the model. These job details include the following information:

- Autopilot candidate name
- Autopilot job name
- Problem type
- Objective metric
- Optimization direction

**Model quality report**

Model quality information is generated by Autopilot model insights. The report's content that is generated depends on the problem type it addressed: regression, binary classification, or multiclass classification. The report specifies the number of rows that were included in the evaluation dataset and the time at which the evaluation occurred.
Metrics tables

The first part of the model quality report contains metrics tables. These are appropriate for the type of problem that the model addressed.

The following image is an example of a metrics table that Autopilot generates for a regression problem. It shows the metric name, value, and standard deviation.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Value</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>mae</td>
<td>5.347324</td>
<td>0.118636</td>
</tr>
<tr>
<td>mse</td>
<td>87.874017</td>
<td>4.346468</td>
</tr>
<tr>
<td>rmse</td>
<td>9.374114</td>
<td>0.232349</td>
</tr>
<tr>
<td>r2</td>
<td>0.924700</td>
<td>0.003710</td>
</tr>
</tbody>
</table>

The following image is an example of a metrics table generated by Autopilot for a multiclass classification problem. It shows the metric name, value, and standard deviation.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Value</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>weighted_recall</td>
<td>0.597104</td>
<td>0.005410</td>
</tr>
<tr>
<td>weighted_precision</td>
<td>0.591693</td>
<td>0.005729</td>
</tr>
<tr>
<td>accuracy</td>
<td>0.597104</td>
<td>0.005410</td>
</tr>
<tr>
<td>weighted_f0_5</td>
<td>0.592155</td>
<td>0.005659</td>
</tr>
<tr>
<td>weighted_f1</td>
<td>0.593423</td>
<td>0.005554</td>
</tr>
<tr>
<td>weighted_f2</td>
<td>0.595392</td>
<td>0.005456</td>
</tr>
<tr>
<td>accuracy_best_constant_classifier</td>
<td>0.200699</td>
<td>0.004422</td>
</tr>
<tr>
<td>weighted_recall_best_constant_classifier</td>
<td>0.200699</td>
<td>0.004422</td>
</tr>
<tr>
<td>weighted_precision_best_constant_classifier</td>
<td>0.040280</td>
<td>0.001753</td>
</tr>
<tr>
<td>weighted_f0_5_best_constant_classifier</td>
<td>0.047944</td>
<td>0.002039</td>
</tr>
<tr>
<td>weighted_f1_best_constant_classifier</td>
<td>0.067094</td>
<td>0.002684</td>
</tr>
<tr>
<td>weighted_f2_best_constant_classifier</td>
<td>0.111716</td>
<td>0.003808</td>
</tr>
</tbody>
</table>

Graphical model performance information

The second part of the model quality report contains graphical information to help you evaluate model performance. The contents of this section depend on the problem type used in modeling.
The area under the receiver operating characteristic curve

The area under the receiver operating characteristic curve represents the trade-off between true positive and false positive rates. It is an industry-standard accuracy metric used for binary classification models. AUC (area under the curve) measures the ability the model to predict a higher score for positive examples, as compared to negative examples. The AUC metric provides an aggregated measure of the model performance across all possible classification thresholds.

The AUC metric returns a decimal value from 0 to 1. AUC values near 1 indicate that the machine learning model is highly accurate. Values near 0.5 indicate that the model is performing no better than guessing at random. AUC values close to 0 indicate that the model has learned the correct patterns, but is making predictions that are as inaccurate as possible. Values near zero can indicate a problem with the data. For more information about the AUC metric, see the Receiver operating characteristic article on Wikipedia.

The following is an example of an area under the receiver operating characteristic curve graph to evaluate predictions made by a binary classification model. The dashed thin line represents the area under the receiver operating characteristic curve that a model which classifies no-better-than-random guessing would score, with an AUC score of 0.5. The curves of more accurate classification models lie above this random baseline, where the rate of true positives exceeds the rate of false positives. The area under the receiver operating characteristic curve representing the performance of the binary classification model is the thicker solid line.
A summary of the graph's components of false positive rate (FPR) and true positive rate (TPR) are defined as follows.

- **Correct predictions**
  - True positive (TP): The predicted value is 1, and the true value is 1.
  - True negative (TN): The predicted value is 0, and the true value is 0.

- **Erroneous predictions**
  - False positive (FP): The predicted value is 1, but the true value is 0.
  - False negative (FN): The predicted value is 0, but the true value is 1.
The **false positive rate** (FPR) measures the fraction of true negatives (TN) that were falsely predicted as positives (FP), over the sum of FP and TN. The range is 0 to 1. A smaller value indicates better predictive accuracy.

- \( \text{FPR} = \frac{FP}{(FP+TN)} \)

The **true positive rate** (TPR) measures the fraction true positives that were correctly predicted as positives (TP) over the sum of TP and false negatives (FN). The range is 0 to 1. A larger value indicates better predictive accuracy.

- \( \text{TPR} = \frac{TP}{(TP+FN)} \)

**Confusion matrix**

A confusion matrix provides a way to visualize the accuracy of the predictions made by a model for binary and multiclass classification for different problems. The confusion matrix in the model quality report contains the following.

- The number and percentage of correct and incorrect predictions for the actual labels
- The number and percentage of accurate predictions on the diagonal from the upper-left to the lower-right corner
- The number and percentage of inaccurate predictions on the diagonal from the upper-right to the lower-left corner

The incorrect predictions on a confusion matrix are the confusion values.

The following diagram is an example of a confusion matrix for a binary classification problem. It contains the following information:

- The vertical axis is divided into two rows containing true and false actual labels.
- The horizontal axis is divided into two columns containing true and false labels that were predicted by the model.
- The color bar assigns a darker tone to a larger number of samples to visually indicate the number of values that were classified in each category.

In this example, the model predicted actual 2817 false values correctly, and 353 actual true values correctly. The model incorrectly predicted 130 actual true values to be false and 33 actual false
values to be true. The difference in tone indicates that the dataset is not balanced. The imbalance is because there are many more actual false labels than actual true labels.

The following diagram is an example of a confusion matrix for a multi-class classification problem. The confusion matrix in the model quality report contains the following.

- The vertical axis is divided into three rows containing three different actual labels.
- The horizontal axis is divided into three columns containing labels that were predicted by the model.
- The color bar assigns a darker tone to a larger number of samples to visually indicate the number of values that were classified in each category.

In the example below, the model correctly predicted actual 354 values for label $f$, 1094 values for label $i$ and 852 values for label $m$. The difference in tone indicates that the dataset is not balanced because there are many more labels for the value $i$ than for $f$ or $m$. 

The confusion matrix in the model quality report provided can accommodate a maximum of 15 labels for multiclass classification problem types. If a row corresponding to a label shows a Nan value, it means that the validation dataset used to check model predictions does not contain data with that label.

**Gain curve**

In binary classification, a gain curve predicts the cumulative benefit of using a percentage of the dataset to find a positive label. The gain value is calculated during training by dividing the cumulative number of positive observations by the total number of positive observations in the data, at each decile. If the classification model created during training is representative of the unseen data, you can use the gain curve to predict the percentage of data that you must target to obtain a percentage of positive labels. The greater the percentage of the dataset used, the higher the percentage of positive labels found.

In the following example graph, the gain curve is the line with changing slope. The straight line is the percentage of positive labels found by selecting a percentage of data from the dataset at
random. Upon targeting 20% of the dataset, you would expect to find larger than 40% of the positive labels. As an example, you might consider using a gain curve to determine your efforts in a marketing campaign. Using our gain curve example, for 83% of people in a neighborhood to purchase cookies, you'd send an advertisement to about 60% of the neighborhood.

**Gain Curve**

Lift curve

In binary classification, the lift curve illustrates the uplift of using a trained model to predict the likelihood of finding a positive label compared to a random guess. The lift value is calculated during training using the ratio of percentage gain to the ratio of positive labels at each decile. If the model created during training is representative of the unseen data, use the lift curve to predict the benefit of using the model over randomly guessing.

In the following example graph, the lift curve is the line with changing slope. The straight line is the lift curve associated with selecting the corresponding percentage randomly from the dataset.
Upon targeting 40% of the dataset with your model's classification labels, you would expect to find about 1.7 times the number of the positive labels that you would have found by randomly selecting 40% of the unseen data.

![Lift Curve](image)

**Precision-recall curve**

The precision-recall curve represents the tradeoff between precision and recall for binary classification problems.

**Precision** measures the fraction of actual positives that are predicted as positive (TP) out of all positive predictions (TP and false positive). The range is 0 to 1. A larger value indicates better accuracy in the predicted values.

- Precision = TP/(TP+FP)
Recall measures the fraction of actual positives that are predicted as positive (TP) out of all actual positive predictions (TP and false negative). This is also known as the sensitivity or as the true positive rate. The range is 0 to 1. A larger value indicates better detection of positive values from the sample.

- Recall = TP/(TP+FN)

The objective of a classification problem is to correctly label as many elements as possible. A system with high recall but low precision returns a high percentage of false positives.

The following graphic depicts a spam filter that marks every email as spam. It has high recall, but low precision, because recall doesn’t measure false positives.

Give more weight to recall over precision if your problem has a low penalty for false positive values, but a high penalty for missing a true positive result. For example, detecting an impending collision in a self-driving vehicle.

By contrast, a system with high precision, but low recall, returns a high percentage of false negatives. A spam filter that marks every email as desirable (not spam) has high precision but low recall because precision doesn’t measure false negatives.

If your problem has a low penalty for false negative values, but a high penalty for missing a true negative results, give more weight to precision over recall. For example, flagging a suspicious filter for a tax audit.
The following graphic depicts a spam filter that has high precision but low recall, because precision doesn't measure false negatives.

A model that makes predictions with both high precision and high recall produces a high number of correctly labeled results. For more information, see [Precision and recall](https://en.wikipedia.org/wiki/Precision_and_recall) article in Wikipedia.

**Area under precision-recall curve (AUPRC)**

For binary classification problems, Amazon SageMaker Autopilot includes a graph of the area under the precision-recall curve (AUPRC). The AUPRC metric provides an aggregated measure of the model performance across all possible classification thresholds and uses both precision and recall. AUPRC does not take the number of true negatives into account. Therefore, it can be useful to evaluate model performance in cases where there's a large number of true negatives in the data. For example, to model a gene containing a rare mutation.

The following graphic is an example of an AUPRC graph. Precision at its highest value is 1, and recall is at 0. In the lower right corner of the graph, recall is its highest value (1) and precision is 0. In between these two points, the AUPRC curve illustrates the tradeoff between precision and recall at different thresholds.
Actual against predicted plot

The actual against predicted plot shows the difference between actual and predicted model values. In the following example graph, the solid line is a linear line of best fit. If the model were 100% accurate, each predicted point would equal its corresponding actual point and lie on this line of best fit. The distance away from the line of best fit is a visual indication of model error. The larger the distance away from the line of best fit, the higher the model error.
Standardized residual plot

A standardized residual plot incorporates the following statistical terms:

**residual**

A (raw) residual shows the difference between actual and values predicted by your model. The larger the difference, the larger the residual value.

**standard deviation**

The standard deviation is a measure of how values vary from an average value. A high standard deviation indicates that many values are very different from their average value. A low standard deviation indicates that many values are close to their average value.
standardized residual

A standardized residual divides the raw residuals by their standard deviation. Standardized residuals have units of standard deviation and are useful in identifying outliers in data regardless of the difference in scale of the raw residuals. If a standardized residual is much smaller or larger than the other standardized residuals, it indicates that the model is not fitting these observations well.

The standardized residual plot measures the strength of the difference between observed and expected values. The actual predicted value is displayed on the x axis. A point with a value larger than an absolute value of 3 is commonly regarded as an outlier.

The following example graph shows that a large number of standardized residuals are clustered around 0 on the horizontal axis. The values close to zero indicate that the model is fitting these points well. The points towards the top and bottom of the plot are not predicted well by the model.
Residual histogram

A residual histogram incorporates the following statistical terms:

**residual**

A (raw) residual shows the difference between actual and values predicted by your model. The larger the difference, the larger the residual value.

**standard deviation**

The standard deviation is a measure of how much values vary from an average value. A high standard deviation indicates that many values are very different from their average value. A low standard deviation indicates that many values are close to their average value.

**standardized residual**

A standardized residual divides the raw residuals by their standard deviation. Standardized residuals have units of standard deviation. These are useful in identifying outliers in data regardless of the difference in scale of the raw residuals. If a standardized residual is much smaller or larger than the other standardized residuals, it would indicate that the model is not fitting these observations well.

**histogram**

A histogram is a graph that shows how often a value occurred.

The residual histogram shows the distribution of standardized residual values. A histogram distributed in a bell shape and centered at zero indicates that the model does not systematically overpredict or underpredict any particular range of target values.

In the following graphic, the standardized residual values indicate that the model is fitting the data well. If the graph showed values far away from the center value, it would indicate that those values don’t fit the model well.
Amazon SageMaker Autopilot notebooks generated to manage AutoML tasks

Amazon SageMaker Autopilot manages the key tasks in an automatic machine learning (AutoML) process using an AutoML job.

The AutoML job creates three notebook-based reports that describe the plan that Autopilot follows to generate candidate models. A candidate model consists of a (pipeline, algorithm) pair. First, there's a data exploration notebook that describes what Autopilot learned about the data that you provided. Second, there's a candidate definition notebook, which uses the information about the data to generate candidates. Third, a model insights report that can help detail the performance characteristics of the best model in the leaderboard of an Autopilot experiment.

Topics
- Amazon SageMaker Autopilot Data exploration report
- Candidate definition notebook
You can run these notebooks in Amazon SageMaker, or locally, if you have installed the Amazon SageMaker Python SDK. You can share the notebooks just like any other SageMaker Studio Classic notebook. The notebooks are created for you to conduct experiments. For example, you could edit the following items in the notebooks:

- Preprocessors used on the data
- Amount of hyperparameter optimization (HPO) runs and their parallelism
- Algorithms to try
- Instance types used for the HPO jobs
- Hyperparameter ranges

Modifications to the candidate definition notebook are encouraged as a learning tool. With this capability, you learn how decisions made during the machine learning process impact your results.

### Note

When you run the notebooks in your default instance, you incur baseline costs. However, when you run HPO jobs from the candidate notebook, these jobs use additional compute resources that incur additional costs.

---

**Amazon SageMaker Autopilot Data exploration report**

Amazon SageMaker Autopilot cleans and pre-processes your dataset automatically. High-quality data improves machine learning efficiency and produces models that make more accurate predictions.

There are issues with customer-provided datasets that cannot be fixed automatically without the benefit of some domain knowledge. Large outlier values in the target column for regression problems, for example, may cause suboptimal predictions for the non-outlier values. Outliers may need to be removed depending on the modeling objective. If a target column is included by accident as one of the input features, the final model will validate well, but be of little value for future predictions.

To help customers discover these sorts of issues, Autopilot provides a data exploration report that contains insights into potential issues with their data. The report also suggests how to handle the issues.
A data exploration notebook containing the report is generated for every Autopilot job. The report is stored in an Amazon S3 bucket and can be accessed from your output path. The path of the data exploration report usually adheres to the following pattern.

```
[s3 output path]/[name of the automl job]/sagemaker-automl-candidates/[name of processing job used for data analysis]/notebooks/
SageMakerAutopilotDataExplorationNotebook.ipynb
```

The location of the data exploration notebook can be obtained from the Autopilot API using the **DescribeAutoMLJob** operation response, which is stored in **DataExplorationNotebookLocation**.

When running Autopilot from SageMaker Studio Classic, you can open the data exploration report using the following steps:

1. Choose the **Home** icon from the left navigation pane to view the top-level Amazon SageMaker Studio Classic navigation menu.
2. Select the **AutoML** card from the main working area. This opens a new **Autopilot** tab.
3. In the **Name** section, select the Autopilot job that has the data exploration notebook that you want to examine. This opens a new **Autopilot job** tab.
4. Select **Open data exploration notebook** from the top right section of the **Autopilot job** tab.

The data exploration report is generated from your data before the training process begins. This allows you to stop Autopilot jobs that might lead to meaningless results. Likewise, you can address any issues or improvements with your dataset before rerunning Autopilot. This way, you can use your domain expertise to improve the data quality manually, before you train a model on a better-curated dataset.

The data report contains only static markdown and can be opened in any Jupyter environment. The notebook that contains the report can be converted to other formats, such as PDF or HTML. For more information about conversions, see Using the nbconvert script to convert Jupyter notebooks to other formats.

**Topics**

- Dataset Summary
- Target Analysis
• **Data Sample**
• **Duplicate rows**
• **Cross column correlations**
• **Anomalous Rows**
• **Missing values, cardinality, and descriptive statistics**

**Dataset Summary**

This **Dataset Summary** provides key statistics characterizing your dataset including the number of rows, columns, percent duplicate rows and missing target values. It is intended to provide you with a quick alert when there are issues with your dataset that Amazon SageMaker Autopilot has detected and that are likely to require your intervention. The insights are surfaced as warnings that are classified as being of either “high” or “low” severity. The classification depends on the level of confidence that the issue will adversely impact the performance of the model.

The high and low severity insights appear in the summary as pop-ups. For most of the insights, recommendations are offered for how to confirm that there is an issue with the dataset that requires your attention. Proposals are also provided for how to resolve the issues.

Autopilot provides additional statistics about missing or not valid target values in our dataset to help you detect other issues that may not be captured by high severity insights. An unexpected number of columns of a particular type might indicate that some columns that you want to use may be missing from the dataset. It could also indicate that there was an issue with how the data was prepared or stored. Fixing these data problems brought to your attention by Autopilot can improve the performance of the machine learning models trained on your data.

High severity insights are shown in the summary section and in other relevant sections in the report. Examples of high and low-severity insights are usually given depending on the section of the data report.

**Target Analysis**

Various high and low-severity insights are shown in this section related to the distribution of values in the target column. Check that target column contains the correct values. Incorrect values in target column will likely result in a machine learning model that doesn't serve the intended business purpose. Several data insights of high and low severity are present in this section. Here are several examples.
• **Outlier target values** - Skewed or unusual target distribution for regression, such as heavy tailed targets.

• **High or low target cardinality** - Infrequent number of class labels or a large number of unique classes for classification.

For both regression and classification problem types, not valid values such as numeric infinity, NaN or empty space in target column are surfaced. Depending on the problem type, different dataset statistics are presented. A distribution of target column values for a regression problem allows you to verify if the distribution is what you expected.

The following screenshot shows an Autopilot data report, which includes statistics such as the mean, median, minimum, maximum, percentage of outliers in your dataset. The screenshot also includes a histogram showing the distribution of labels in the target column. The histogram shows **Target Column Values** on the horizontal axis and **Count** on the vertical axis. A box highlights the **Outliers Percentage** section of the screenshot to indicate where this statistic appears.

<table>
<thead>
<tr>
<th>Mean</th>
<th>Median</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Skew</th>
<th>Kurtosis</th>
<th>Number of Uniques</th>
<th>Outliers Percentage</th>
<th>Invalid Percentage</th>
<th>Missing Percentage</th>
<th>Missing Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>3017.90</td>
<td>2116.24</td>
<td>0.67</td>
<td>121012.25</td>
<td>2.86</td>
<td>16.33</td>
<td>130809</td>
<td>1.30%</td>
<td>0.00%</td>
<td>0.00%</td>
<td>0</td>
</tr>
</tbody>
</table>

Multiple statistics are shown regarding target values and their distribution. If any of the outliers, not valid values, or missing percentages are greater than zero, these values are surfaced so you can investigate why your data contains unusable target values. Some unusable target values are highlighted as a low severity insight warning.

In the following screenshot, a ` symbol was added accidentally to the target column, which prevented the numeric value of the target from being parsed. A **Low severity insight: "Invalid"**
A "target values" warning appears. The warning in this example states "0.14% of the labels in the target column could not be converted to numeric values. The most common non-numeric values are: ["-3.8e-05","-9-05","-4.7e-05","-1.4999999999999999e-05","-4.3e-05"]. That usually indicates that there are problems with data collection or processing. Amazon SageMaker Autopilot ignores all observations with invalid target label."

Autopilot also provides a histogram showing the distribution of labels for classification.

The following screenshot shows an example of statistics given for your target column including the number of classes, missing or not valid values. A histogram with **Target Label** on the horizontal axis and **Frequency** on the vertical axis shows the distribution of each label category.
Note

You can find definitions of all the terms presented in this and other sections in Definitions section at the bottom of the report notebook.

Data Sample

Autopilot presents an actual sample of your data to help you spot issues with your dataset. The sample table scrolls horizontally. Inspect the sample data to verify that all the necessary columns are present in the dataset.
Autopilot also calculates a measure of prediction power, that can be used to identify a linear or nonlinear relationship between a feature and the target variable. A value of 0 indicates that the feature has no predictive value in predicting the target variable. A value of 1 indicates the highest predictive power for the target variable. For more information on predictive power, see the Definitions section.

Note

It is not recommended that you use prediction power as a substitute for feature importance. Only use it if you're certain that prediction power is an appropriate measure for your use case.

The following screenshot shows example data sample. The top row contains the prediction power of each column in your dataset. The second row contains the column data type. Subsequent rows contain the labels. The columns contain the target column followed by each feature column. Each feature column has an associated prediction power, highlighted in this screenshot, with a box. In this example, the column containing the feature x51 has a predictive power of 0.68 for the target variable y. The feature x55 is slightly less predictive with a prediction power of 0.59.
Duplicate rows

If duplicate rows are present in the dataset, Amazon SageMaker Autopilot displays a sample of them.

Note

It is not recommended to balance a dataset by up-sampling before providing it to Autopilot. This may result in inaccurate validation scores for the models trained by Autopilot, and the models that are produced may be unusable.

Cross column correlations

Autopilot uses the Pearson's correlation coefficient, a measure of linear correlation between two features, to populate a correlation matrix. In the correlation matrix, numeric features are plotted on both the horizontal and vertical axes, with the Pearson's correlation coefficient plotted at their intersections. The higher the correlation between two features, the higher the coefficient, with a maximum value of $|1|$.

- A value of $-1$ indicates that the features are perfectly negatively correlated.
- A value of $1$, which occurs when a feature is correlated with itself, indicates perfect positive correlation.

You can use the information in the correlation matrix to remove highly correlated features. A smaller number of features reduces chances of overfitting a model and can reduce the costs of production in two ways. It lessens the Autopilot runtime needed and, for some applications, can make data collection procedures cheaper.

The following screenshot shows an example of a correlation matrix between 7 features. Each feature is displayed in a matrix on both the horizontal and vertical axes. The Pearson's correlation coefficient is displayed at the intersection between two features. Each feature intersection has a color tone associated with it. The higher the correlation, the darker the tone. The darkest tones occupy the diagonal of the matrix, where each feature is correlated with itself, representing perfect correlation.
Anomalous Rows

Amazon SageMaker Autopilot detects which rows in your dataset might be anomalous. It then assigns an anomaly score to each row. Rows with negative anomaly scores are considered anomalous.

The following screenshot shows the output from an Autopilot analysis for rows containing anomalies. A column containing an anomalous score appears next to the dataset columns for each row.
Amazon SageMaker Autopilot examines and reports on properties of the individual columns of your dataset. In each section of the data report that presents this analysis, the content is arranged in order. This is so you can check the most “suspicious” values first. Using these statistics you can improve contents of individual columns, and improve the quality of the model produced by Autopilot.

Autopilot calculates several statistics on the categorical values in columns that contain them. These include the number of unique entries and, for text, the number of unique words.

Autopilot calculates several standard statistics on the numerical values in columns that contain them. The following image depicts these statistics, including the mean, median, minimum and maximum values, and the percentages of numerical types and of outlier values.

### Missing values, cardinality, and descriptive statistics

Amazon SageMaker Autopilot examines and reports on properties of the individual columns of your dataset. In each section of the data report that presents this analysis, the content is arranged in order. This is so you can check the most “suspicious” values first. Using these statistics you can improve contents of individual columns, and improve the quality of the model produced by Autopilot.

Autopilot calculates several statistics on the categorical values in columns that contain them. These include the number of unique entries and, for text, the number of unique words.

Autopilot calculates several standard statistics on the numerical values in columns that contain them. The following image depicts these statistics, including the mean, median, minimum and maximum values, and the percentages of numerical types and of outlier values.
Candidate definition notebook

The candidate definition notebook contains each suggested preprocessing step, algorithm, and hyperparameter ranges.

You can choose which candidate to train and tune in two ways. The first, by running sections of the notebook. The second, by running the entire notebook to optimize all candidates to identify a best candidate. If you run the entire notebook, only the best candidate is displayed after job completion.

To run Autopilot from SageMaker Studio Classic, open the candidate definition notebook by following these steps:

1. Choose the **Home** icon from the left navigation pane to view the top-level *Amazon SageMaker Studio Classic* navigation menu.
2. Select the **AutoML** card from the main working area. This opens a new **Autopilot** tab.
3. In the **Name** section, select the Autopilot job that has the candidate definition notebook that you want to examine. This opens a new **Autopilot job** tab.

<table>
<thead>
<tr>
<th>% of Numerical Values</th>
<th>Mean</th>
<th>Median</th>
<th>Min</th>
<th>Max</th>
<th>% of Outlier Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>y</td>
<td>100.0%</td>
<td>9.93957</td>
<td>9.0</td>
<td>3.0</td>
<td>27.0</td>
</tr>
<tr>
<td>1</td>
<td>100.0%</td>
<td>0.523612</td>
<td>0.545</td>
<td>0.11</td>
<td>0.815</td>
</tr>
<tr>
<td>2</td>
<td>100.0%</td>
<td>0.407799</td>
<td>0.425</td>
<td>0.09</td>
<td>0.65</td>
</tr>
<tr>
<td>3</td>
<td>100.0%</td>
<td>0.13995</td>
<td>0.145</td>
<td>0.015</td>
<td>0.515</td>
</tr>
<tr>
<td>4</td>
<td>100.0%</td>
<td>0.828266</td>
<td>0.81</td>
<td>0.008</td>
<td>2.8255</td>
</tr>
<tr>
<td>5</td>
<td>100.0%</td>
<td>0.358844</td>
<td>0.339</td>
<td>0.0025</td>
<td>1.2395</td>
</tr>
<tr>
<td>6</td>
<td>100.0%</td>
<td>0.180348</td>
<td>0.1725</td>
<td>0.002</td>
<td>0.6415</td>
</tr>
<tr>
<td>7</td>
<td>100.0%</td>
<td>0.238783</td>
<td>0.235</td>
<td>0.003</td>
<td>1.005</td>
</tr>
</tbody>
</table>
4. Choose **Open candidate generation notebook** from the top right section of the **Autopilot job** tab. This opens a new read-only preview of the **Amazon SageMaker Autopilot Candidate Definition Notebook**.

To run the candidate definition notebook, follow these steps:

1. Choose **Import notebook** at the top right of the **Amazon SageMaker Autopilot Candidate Definition Notebook** tab. This opens a tab to set up a new notebook environment to run the notebook.
2. Select an existing SageMaker **Image** or use a **Custom Image**.
3. Select a **Kernel**, an **Instance type**, and an optional **Start-up script**.

You can now run the notebook in this new environment.

**Configure inference output in generated containers**

Autopilot generates an ordered **ContainerDefinition** list. This can be used to build a model to deploy in a machine learning pipeline. This model can be used for online hosting and inference.

Customers can list inference container definitions with the **ListCandidateForAutoMLJob** API. The list of inference container definitions that represent the best candidate is also available in the **DescribeAutoMLJob** response.

**Inference container definitions for regression and classification problem types**

Autopilot generates inference containers specific to the **training mode** and the **problem type** of the job.

**Container definitions for hyperparameter optimization (HPO) mode**

- **Regression**: HPO generates two containers:
  1. A feature engineering container that transforms the original features into features that the regression algorithms can train on.
  2. An algorithm container that transforms features and generates a regression score for the dataset.
- **Classification**: HPO generates three containers:
  1. A feature engineering container that transforms the original features into features that the classification algorithms can train on.
2. An algorithm container that generates the `predicted_label` with the highest probability. This container can also produce the various probabilities associated with the classification outcomes in the inference response.

3. A feature engineering container that performs post-processing of the algorithm prediction. For example, it can perform an inverse transform on the predicted label and change it to the original label.

**Container definitions for ensembling mode**

In ensembling mode, both regression and classification problem types have only one inference container. This inference container transforms the features and generates the predictions based on problem type.

**Inference responses per problem type**

**Inference responses for classification models**

For classification inference containers, you can select the content of the inference response by using four predefined keys:

- **predicted_label**: The label with the highest probability of predicting the correct label, as determined by Autopilot.
- **probability**:
  - **HPO models**: The probability of the True class for binary classification. The probability of the `predicted_label` for multiclass classification.
  - **Ensemble models**: The probability of the `predicted_label` for binary and multiclass classification.
- **probabilities**: The list of probabilities for all corresponding classes.
- **labels**: The list of all labels.

For example, for a binary classification problem, if you pass the inference response keys `['predicted_label', 'probability', 'probabilities', 'labels']` and the output response appears as `[1, 0.1, "[0.9, 0.1]", ["1", "0"]]`, you should interpret it as follows:

1. `predicted_label` equals 1 because label "1" has a higher probability (0.9 in this case).
2. For HPO models, probability equals 0.1 which is the probability of the `positive_class` (0 in this case) selected by Autopilot.

   For Ensemble models, probability equals 0.9 which is the probability of the `predicted_label`.

3. `probabilities` lists the probability of each label in `labels`.

4. `labels` are the unique labels in the dataset, where the second label ("0" in this case) is the `positive_class` selected by Autopilot.

By default, inference containers are configured to generate only the `predicted_label`. To select additional inference content, you can update the `inference_response_keys` parameter to include up to these three environment variables:

- `SAGEMAKER_INFERENCE_SUPPORTED`: This is set to provide hints to you about what content each container supports.
- `SAGEMAKER_INFERENCE_INPUT`: This should be set to the keys that the container expects in input payload.
- `SAGEMAKER_INFERENCE_OUTPUT`: This should be populated with the set of keys that the container outputs.

**Inference responses for classification models in HPO mode**

This section shows how to configure the inference response from classification models using hyperparameter optimization (HPO) mode.

To choose the inference response content in HPO mode: Add the `SAGEMAKER_INFERENCE_INPUT` and `SAGEMAKER_INFERENCE_OUTPUT` variables to the second and third containers that are generated in HPO mode for classification problems.

The keys supported by the second container (algorithm) are `predicted_label`, `probability`, and `probabilities`. Note that `labels` is deliberately not added to `SAGEMAKER_INFERENCE_SUPPORTED`.

The keys supported by the third classification model container are `predicted_label`, `labels`, `probability`, and `probabilities`. Therefore, the `SAGEMAKER_INFERENCE_SUPPORTED` environment includes the names of these keys.

To update the definition of the inference containers to receive `predicted_label` and `probability`, use the following code example.
The following code example updates the definition of the inference containers to receive predicted_label, probabilities, and labels. Do not pass the labels to the second container (the algorithm container), because it is generated by the third container independently.

```python
containers[1]['Environment'].update({'SAGEMAKER_INFERENCE_OUTPUT': 'predicted_label, probabilities'})
containers[2]['Environment'].update({'SAGEMAKER_INFERENCE_INPUT': 'predicted_label, probabilities'})
containers[2]['Environment'].update({'SAGEMAKER_INFERENCE_OUTPUT': 'predicted_label, probabilities, labels'})
```

The following collapsible sections provide code examples for AWS SDK for Python (Boto3) and for SageMaker SDK for Python. Each section shows how to select the content of the inference responses in HPO mode for the respective code example.

**AWS SDK for Python (Boto3)**

```python
import boto3

sm_client = boto3.client('sagemaker', region_name='<Region>')

role = '<IAM role>'
input_data = '<S3 input uri>'
output_path = '<S3 output uri>'

best_candidate = sm_client.describe_auto_ml_job(AutoMLJobName='<AutoML Job Name>')
['BestCandidate']
best_candidate_containers = best_candidate['InferenceContainers']
best_candidate_name = best_candidate['CandidateName']

best_candidate_containers[1]['Environment'].update({'SAGEMAKER_INFERENCE_OUTPUT': 'predicted_label, probability'})
best_candidate_containers[2]['Environment'].update({'SAGEMAKER_INFERENCE_INPUT': 'predicted_label, probability'})
```
```python
best_candidate_containers[2]['Environment'].update({'SAGEMAKER_INFERENCE_OUTPUT': 'predicted_label, probability'})

# create model
response = sm_client.create_model(
    ModelName = '<Model Name>',
    ExecutionRoleArn = role,
    Containers = best_candidate_containers
)

# Launch Transform Job
response = sm_client.create_transform_job(
    TransformJobName='<Transform Job Name>',
    ModelName='<Model Name>',
    TransformInput={
        'DataSource': {
            'S3DataSource': {
                'S3DataType': 'S3Prefix',
                'S3Uri': 'input_data
            }
        },
        'ContentType': "text/CSV",
        'SplitType': 'Line'
    },
    TransformOutput={
        'S3OutputPath': 'output_path',
        'AssembleWith': 'Line',
    },
    TransformResources={
        'InstanceType': 'ml.m4.xlarge',
        'InstanceCount': 1,
    }
)

SageMaker SDK for Python

from sagemaker import AutoML

aml = AutoML.attach(auto_ml_job_name='<AutoML Job Name>')
aml_best_model = aml.create_model(name='<Model Name>',
    candidate=None,
    inference_response_keys**=['probabilities', 'labels'])
```

Create a Regression or Classification Job Using the AutoML API
Inference responses for classification models in ensembling mode

This section shows how to configure the inference response from classification models using ensembling mode.

In **ensembling mode**, to choose the content of the inference response, update the SAGEMAKER_INFERENCE_OUTPUT environment variable.

The keys supported by the classification model container are `predicted_label`, `labels`, `probability`, and `probabilities`. These keys are included in the SAGEMAKER_INFERENCE_SUPPORTED environment.

To update the inference container definition to receive `predicted_label` and `probability`, refer to the following code example.

```python
containers[0]['Environment'].update({'SAGEMAKER_INFERENCE_OUTPUT': 'predicted_label, probability'})
```

The following collapsible section provides a code example for selecting the content of the inference responses in ensembling mode. The example uses AWS SDK for Python (Boto3).

**AWS SDK for Python (Boto3)**

```python
import boto3
sm_client = boto3.client('sagemaker', region_name='<Region>')

role = '<IAM role>'
input_data = '<S3 input uri>'
output_path = '<S3 output uri>'
```
best_candidate = sm_client.describe_auto_ml_job(AutoMLJobName='<AutoML Job Name>')['BestCandidate']
best_candidate_containers = best_candidate['InferenceContainers']
best_candidate_name = best_candidate['CandidateName']

*best_candidate_containers[0]['Environment'].update({'SAGEMAKER_INFERENCE_OUTPUT': 'predicted_label, probability'})
*

# create model
response = sm_client.create_model(
    ModelName = '<Model Name>',
    ExecutionRoleArn = role,
    Containers = best_candidate_containers
)

# Launch Transform Job
response = sm_client.create_transform_job(
    TransformJobName='<Transform Job Name>',
    ModelName='<Model Name>',
    TransformInput={
        'DataSource': {
            'S3DataSource': {
                'S3DataType': 'S3Prefix',
                'S3Uri': input_data
            }
        },
        'ContentType': "text/CSV",
        'SplitType': 'Line'
    },
    TransformOutput={
        'S3OutputPath': output_path,
        'AssembleWith': 'Line',
    },
    TransformResources={
        'InstanceType': 'ml.m4.xlarge',
        'InstanceCount': 1,
    }
)

The following collapsible section provides a code example that is identical to the SageMaker SDK for Python example for HPO. It is included for your convenience.
SageMaker SDK for Python

The following HPO code example uses SageMaker SDK for Python.

```python
from sagemaker import AutoML

aml = AutoML.attach(auto_ml_job_name='<AutoML Job Name>')</n
aml_best_model = aml.create_model(name='<Model Name>',
                                   candidate=None,
                                   *inference_response_keys**=['probabilities',
                                   'labels'])*

aml_transformer = aml_best_model.transformer(accept='text/csv',
                                              assemble_with='Line',
                                              instance_type='ml.m5.xlarge',
                                              instance_count=1,)

aml_transformer.transform('<S3 input uri>',
                           content_type='text/csv',
                           split_type='Line',
                           job_name='<Transform Job Name>',
                           wait=True)
```

Tutorials and example notebooks

Example notebooks, tutorial videos, and walkthroughs to get started with Amazon SageMaker Autopilot.

Topics
- Example notebooks: Explore modeling with Amazon SageMaker Autopilot
- Videos: Use Autopilot to automate and explore the machine learning process
- Tutorials: Get started with Amazon SageMaker Autopilot

Example notebooks: Explore modeling with Amazon SageMaker Autopilot

Amazon SageMaker Autopilot provides the following example notebooks.

- Direct marketing with Amazon SageMaker Autopilot: This notebook demonstrates how uses the Bank Marketing Data Set to predict whether a customer will enroll for a term deposit at a bank. You can use Autopilot on this dataset to get the most accurate ML pipeline by exploring options contained in various candidate pipelines. Autopilot generates each candidate in a two-step process.
procedure. The first step performs automated feature engineering on the dataset. The second step trains and tunes an algorithm to produce a model. The notebook contains instructions on how to train the model and how to deploy the model to perform batch inference using the best candidate.

- **Customer Churn Prediction with Amazon SageMaker Autopilot**: This notebook describes using machine learning for the automated identification of unhappy customers, also known as customer churn prediction. The example shows how to analyze a publicly available dataset and perform feature engineering on it. Next it shows how to tune a model by selecting the best performing pipeline along with the optimal hyperparameters for the training algorithm. Finally, it shows how to deploy the model to a hosted endpoint and how to evaluate its predictions against ground truth. However, ML models rarely give perfect predictions. That's why this notebook also shows how to incorporate the relative costs of prediction mistakes when determining the financial outcome of using ML.

- **Top Candidates Customer Churn Prediction with Amazon SageMaker Autopilot and Batch Transform (Python SDK)**: This notebook also describes using machine learning for the automated identification of unhappy customers, also known as customer churn prediction. This notebook demonstrates how to configure the model to obtain the inference probability, select the top N models, and make Batch Transform on a hold-out test set for evaluation.

  **Note**

  This notebook works with SageMaker Python SDK >= 1.65.1 released on 6/19/2020.

- **Bringing your own data processing code to Amazon SageMaker Autopilot**: This notebook demonstrates how to incorporate and deploy custom data processing code when using Amazon SageMaker Autopilot. It adds a custom feature selection step to remove irrelevant variables to an Autopilot job. It then shows how to deploy both the custom processing code and models generated by Autopilot on a real-time endpoint and, alternatively, for batch processing.

**Videos: Use Autopilot to automate and explore the machine learning process**

Here is a video series that provides a tour of Amazon SageMaker Autopilot capabilities using Studio Classic. They show how to start an AutoML job, analyze and preprocess data, how to do feature engineering and hyperparameter optimization on candidate models, and how to visualize and compare the resulting model metrics.

**Topics**
• **Start an AutoML job with Amazon SageMaker Autopilot**

• **Review data exploration and feature engineering automated in Autopilot.**

• **Tune models to optimize performance**

• **Choose and deploy the best model**

• **Amazon SageMaker Autopilot tutorial**

**Start an AutoML job with Amazon SageMaker Autopilot**

This video shows you to how to start an AutoML job with Autopilot. (Length: 8:41)

[Amazon SageMaker Studio - AutoML with Amazon SageMaker Autopilot (part 1)](https://example.com)

**Review data exploration and feature engineering automated in Autopilot.**

This video shows you how to review the data exploration and candidate definition notebooks generated by Amazon SageMaker Autopilot. (Length: 10:04)

[Amazon SageMaker Studio - AutoML with Amazon SageMaker Autopilot (part 2)](https://example.com)

**Tune models to optimize performance**

This video shows you how to optimize model performance during training using hyperparameter tuning. (Length: 4:59)

[SageMaker Studio - AutoML with Amazon SageMaker Autopilot (part 3)](https://example.com)

**Choose and deploy the best model**

This video shows you how to use job metrics to choose the best model and then how to deploy it. (Length: 5:20)

[SageMaker Studio - AutoML with Amazon SageMaker Autopilot (part 4)](https://example.com)

**Amazon SageMaker Autopilot tutorial**

This video walks you through an end to end demo where we first build a binary classification model automatically with Amazon SageMaker Autopilot. We see how candidate models have been built and optimized using auto-generated notebooks. We also look at the top candidates with Amazon SageMaker Experiments. Finally, we deploy the top candidate (based on XGBoost), and configure data capture with SageMaker Model Monitor.
End to end demo with AutoML on SageMaker

Tutorials: Get started with Amazon SageMaker Autopilot

Get started tutorials for Autopilot demonstrate how to create a machine learning model automatically without writing code. They show you how Autopilot simplifies the machine learning experience by helping you explore your data and try different algorithms. Autopilot builds the best machine learning model for the problem type using AutoML capabilities while allowing full control and visibility.

- **Create a machine learning model automatically with Autopilot**: You assume the role of a developer working at a bank in this tutorial. You have been asked to develop a machine learning model to predict if a customer will enroll for a certificate of deposit (CD). This is a binary classification problem. The model is trained on the marketing dataset that contains information on customer demographics, responses to marketing events, and external factors.

Create an AutoML job for image classification using the API

The following instructions show how to create an Amazon SageMaker Autopilot job as a pilot experiment for image classification problem types using SageMaker API Reference.

**Note**

Tasks such as text and image classification, time-series forecasting, and fine-tuning of large language models are exclusively available through the version 2 of the Autopilot API. For Python users, we recommend using the AWS SDK for Python (Boto3) as the Amazon SageMaker Python SDK is not currently supported for the Autopilot API version 2. Users who prefer the convenience of a user interface can use Amazon SageMaker Canvas to access pre-trained models and generative AI foundation models, or create custom models tailored for specific text, image classification, forecasting needs, or generative AI.

You can create an Autopilot image classification experiment programmatically by calling the CreateAutoMLJobV2 API action in any language supported by Amazon SageMaker Autopilot or the AWS CLI.

For information on how this API action translates into a function in the language of your choice, see the See Also section of CreateAutoMLJobV2 and choose an SDK. As an example, for Python users, see the full request syntax of create_auto_ml_job_v2 in AWS SDK for Python (Boto3).
The following is a collection of mandatory and optional input request parameters for the CreateAutoMLJobV2 API action used in image classification.

**Required parameters**

When calling CreateAutoMLJobV2 to create an Autopilot experiment for image classification, you must provide the following values:

- An AutoMLJobName to specify the name of your job.
- At least one AutoMLJobChannel in AutoMLJobInputDataConfig to specify your data source.
- An AutoMLProblemTypeConfig of type ImageClassificationJobConfig.
- An OutputDataConfig to specify the Amazon S3 output path to store the artifacts of your AutoML job.
- A RoleArn to specify the ARN of the role used to access your data.

All other parameters are optional.

**Optional parameters**

The following sections provide details of some optional parameters that you can pass to your image classification AutoML job.

**How to specify the training and validation datasets of an AutoML job**

You can provide your own validation dataset and custom data split ratio, or let Autopilot split the dataset automatically.

Each AutoMLJobChannel object (see the required parameter AutoMLJobInputDataConfig) has a ChannelType, which can be set to either training or validation values that specify how the data is to be used when building a machine learning model.

At least one data source must be provided and a maximum of two data sources is allowed: one for training data and one for validation data. How you split the data into training and validation datasets depends on whether you have one or two data sources.

How you split the data into training and validation datasets depends on whether you have one or two data sources.

- If you only have one data source, the ChannelType is set to training by default and must have this value.
• If the ValidationFraction value in `AutoMLDataSplitConfig` is not set, 0.2 (20%) of the data from this source is used for validation by default.

• If the ValidationFraction is set to a value between 0 and 1, the dataset is split based on the value specified, where the value specifies the fraction of the dataset used for validation.

• If you have two data sources, the ChannelType of one of the `AutoMLJobChannel` objects must be set to `training`, the default value. The ChannelType of the other data source must be set to `validation`. The two data sources must have the same format, either CSV or Parquet, and the same schema. You must not set the value for the ValidationFraction in this case because all of the data from each source is used for either training or validation. Setting this value causes an error.

**How to specify the automatic model deployment configuration for an AutoML job**

To enable automatic deployment for the best model candidate of an AutoML job, include a `ModelDeployConfig` in the AutoML job request. This will allow the deployment of the best model to a SageMaker endpoint. Below are the available configurations for customization.

• To let Autopilot generate the endpoint name, set `AutoGenerateEndpointName` to True.

• To provide your own name for the endpoint, set `AutoGenerateEndpointName` to False and provide a name of your choice in `EndpointName`.

**Datasets format and objective metric for image classification**

In this section we learn about the available formats for datasets used in image classification as well as the objective metric used to evaluate the predictive quality of machine learning model candidates. The metrics calculated for candidates are specified using an array of `MetricDatum` types.

**Datasets formats**

Autopilot supports .png, .jpg, and .jpeg image formats. If your dataset contains all .png images use `image/png`, if it contains all .jpg or .jpeg images use `image/jpeg`, and if your dataset contains a mix of image formats use `image/*`.

**Objective metric**

The following list contains the names of the metrics that are currently available to measure the performance of models for image classification.
Accuracy

The ratio of the number of correctly classified items to the total number of (correctly and incorrectly) classified items. Accuracy measures how close the predicted class values are to the actual values. Values for accuracy metrics vary between zero (0) and one (1). A value of 1 indicates perfect accuracy, and 0 indicates perfect inaccuracy.

Autopilot model deployment and prediction

This Autopilot guide includes steps for model deployment and setting up real-time inference.

After you train your Autopilot models, you can set up an endpoint and obtain predictions interactively.

Real-time inferencing

Real-time inference is ideal for inference workloads where you have real-time, interactive, low latency requirements. This section shows how you can use real-time inferencing to obtain predictions interactively from your model.

You can use SageMaker APIs to manually deploy the model that produced the best validation metric in an Autopilot experiment as follows.

Alternatively, you can chose the automatic deployment option when creating your Autopilot experiment. For information on setting up the automatic deployment of models, see ModelDeployConfig in the request parameters of CreateAutoMLJobV2. This creates an endpoint automatically.

Note

To avoid incurring unnecessary charges, you can delete unneeded endpoint and resources created from model deployment. For information about pricing of instances by Region, see Amazon SageMaker Pricing.

1. Obtain the candidate container definitions

Obtain the candidate container definitions from InferenceContainers. A container definition for inference refers to the containerized environment designed for deploying and running your trained SageMaker model to make predictions.
The following AWS CLI command example uses the `DescribeAutoMLJobV2` API to obtain candidate definitions for the best model candidate.

```
aws sagemaker describe-auto-ml-job-v2 --auto-ml-job-name job-name --region region
```

2. List candidates

The following AWS CLI command example uses the `ListCandidatesForAutoMLJob` API to list all model candidates.

```
aws sagemaker list-candidates-for-auto-ml-job --auto-ml-job-name <job-name> --region <region>
```

3. Create a SageMaker model

Use the container definitions from the previous steps and a candidate of your choice to create a SageMaker model by using the `CreateModel` API. See the following AWS CLI command as an example.

```
aws sagemaker create-model --model-name '<your-candidate-name>' \
--containers ['<container-definition1>, <container-definition2>, <container-definition3>'] \
--execution-role-arn '<execution-role-arn>' --region '<region>'
```

4. Create an endpoint configuration

The following AWS CLI command example uses the `CreateEndpointConfig` API to create an endpoint configuration.

```
aws sagemaker create-endpoint-config --endpoint-config-name '<your-endpoint-config-name>' \
--production-variants '<list-of-production-variants>' \
--region '<region>'
```

5. Create the endpoint

The following AWS CLI example uses the `CreateEndpoint` API to create the endpoint.

```
aws sagemaker create-endpoint --endpoint-name '<your-endpoint-name>' \
--endpoint-config-name '<endpoint-config-name-you-just-created>'
```
Check the progress of your endpoint deployment by using the `DescribeEndpoint` API. See the following AWS CLI command as an example.

```bash
aws sagemaker describe-endpoint --endpoint-name '<endpoint-name>' --region '<region>'
```

After the `EndpointStatus` changes to `InService`, the endpoint is ready to use for real-time inference.

6. **Invoke the endpoint**

   The following command structure invokes the endpoint for real-time inferencing.

   ```bash
   aws sagemaker invoke-endpoint --endpoint-name '<endpoint-name>' 
   --region '<region>' --body '<your-data>' [--content-type] '<content-type>' <outfile>
   ```

---

**Explainability report**

Amazon SageMaker Autopilot provides an explainability report to help explain how a best model candidate makes predictions for image classification problems. This report can assist ML engineers, product managers, and other internal stakeholders in understanding the characteristics of the model. Both consumers and regulators rely on transparency in machine learning to trust and interpret decisions made on model predictions. You can use these explanations for auditing and meeting regulatory requirements, establishing trust in the model, supporting human decision-making, and debugging and improving model performance.

The Autopilot explanatory functionality for image classification uses a visual class activation map (CAM) approach that produces a heatmap where the distribution and intensity of each color highlights the areas of an image that contribute the most to a specific prediction. This approach relies on principal components derived from an implementation of [Eigen-CAM](#).

Autopilot generates the explainability report as a JSON file. The report includes analysis details that are based on the validation dataset. Each image used to generate the report contains the following information:

- `input_image_uri`: The Amazon S3 URI to the input image taken as input for the heatmap.
- `heatmap_image_uri`: The Amazon S3 URI to the heatmap image generated by Autopilot.
• **predicted_label**: The label class predicted by best model trained by Autopilot.

• **probability**: The confidence with which the predicted_label is predicted.

You can find the Amazon S3 prefix to the explainability artifacts generated for the best candidate in the response to `DescribeAutoMLJobV2` at `BestCandidate.CandidateProperties.CandidateArtifactLocations.Explainability`.

The following examples illustrates what the heatmaps look like on few samples from **Oxford-IIIT Pet Dataset**. The heatmap image displays color gradients that indicate the relative importance of different features within the image. The red color represents regions with greater importance in predicting the "predicted_label" of the input image compared to the features represented by the blue color.

<table>
<thead>
<tr>
<th>Input Image</th>
<th>Heatmap Image</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1.png" alt="Input Image" /></td>
<td><img src="image2.png" alt="Heatmap Image" /></td>
</tr>
</tbody>
</table>
Model performance report

An Amazon SageMaker model quality report (also referred to as performance report) provides insights and quality information for the best model candidate generated by an AutoML job. This includes information about the job details, model problem type, objective function, and various metrics. This section details the content of a performance report for image classification problems and explains how to access the metrics as raw data in a JSON file.

You can find the Amazon S3 prefix to the model quality report artifacts generated for the best candidate in the response to DescribeAutoMLJobV2 at BestCandidate.CandidateProperties.CandidateArtifactLocations.ModelInsights.

The performance report contains two sections:

- The first section contains details about the Autopilot job that produced the model.
- The second section contains a model quality report with various performance metrics.
Autopilot job details

This first section of the report gives some general information about the Autopilot job that produced the model. These details include the following information:

- Autopilot candidate name: The name of the best model candidate.
- Autopilot job name: The name of the job.
- Problem type: The problem type. In our case, image classification.
- Objective metric: The objective metric used to optimize the performance of the model. In our case, Accuracy.
- Optimization direction: Indicates whether to minimize or maximize the objective metric.

Model quality report

Model quality information is generated by Autopilot model insights. The report’s content that is generated depends on the problem type it addressed. The report specifies the number of rows that were included in the evaluation dataset and the time at which the evaluation occurred.

Metrics tables

The first part of the model quality report contains metrics tables. These are appropriate for the type of problem that the model addressed.

The following image is an example of a metrics table generated by Autopilot for an image or text classification problem. It shows the metric name, value, and standard deviation.
The second part of the model quality report contains graphical information to help you evaluate model performance. The contents of this section depend on the selected problem type.

**Confusion matrix**

A confusion matrix provides a way to visualize the accuracy of the predictions made by a model for binary and multiclass classification for different problems.

A summary of the graph's components of **false positive rate** (FPR) and **true positive rate** (TPR) are defined as follows.

- Correct predictions
  - **True positive** (TP): The predicted the value is 1, and the true value is 1.
  - **True negative** (TN): The predicted the value is 0, and the true value is 0.

- Erroneous predictions
  - **False positive** (FP): The predicted the value is 1, but the true value is 0.
  - **False negative** (FN): The predicted the value is 0, but the true value is 1.

The confusion matrix in the model quality report contains the following.

- The number and percentage of correct and incorrect predictions for the actual labels.

### Metrics table

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Value</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>weighted_recall</td>
<td>0.597104</td>
<td>0.005410</td>
</tr>
<tr>
<td>weighted_precision</td>
<td>0.591693</td>
<td>0.005729</td>
</tr>
<tr>
<td>accuracy</td>
<td>0.597104</td>
<td>0.005410</td>
</tr>
<tr>
<td>weighted_f0_5</td>
<td>0.592155</td>
<td>0.005659</td>
</tr>
<tr>
<td>weighted_f1</td>
<td>0.593423</td>
<td>0.005554</td>
</tr>
<tr>
<td>weighted_f2</td>
<td>0.595392</td>
<td>0.005456</td>
</tr>
<tr>
<td>accuracy_best_constant_classifier</td>
<td>0.200699</td>
<td>0.004422</td>
</tr>
<tr>
<td>weighted_recall_best_constant_classifier</td>
<td>0.200699</td>
<td>0.004422</td>
</tr>
<tr>
<td>weighted_precision_best_constant_classifier</td>
<td>0.040280</td>
<td>0.001753</td>
</tr>
<tr>
<td>weighted_f0_5_best_constant_classifier</td>
<td>0.047944</td>
<td>0.002039</td>
</tr>
<tr>
<td>weighted_f1_best_constant_classifier</td>
<td>0.067094</td>
<td>0.002684</td>
</tr>
<tr>
<td>weighted_f2_best_constant_classifier</td>
<td>0.111716</td>
<td>0.003808</td>
</tr>
</tbody>
</table>
• The number and percentage of accurate predictions on the diagonal from the upper-left to the lower-right corner
• The number and percentage of inaccurate predictions on the diagonal from the upper-right to the lower-left corner

The incorrect predictions on a confusion matrix are the confusion values.

The following diagram is an example of a confusion matrix for a multi-class classification problem. The confusion matrix in the model quality report contains the following.

• The vertical axis is divided into three rows containing three different actual labels.
• The horizontal axis is divided into three columns containing labels that were predicted by the model.
• The color bar assigns a darker tone to a larger number of samples to visually indicate the number of values that were classified in each category.

In the example below, the model correctly predicted actual 354 values for label f, 1094 values for label i and 852 values for label m. The difference in tone indicates that the dataset is not balanced because there are many more labels for the value i than for f or m.
The confusion matrix in the model quality report provided can accommodate a maximum of 15 labels for multiclass classification problem types. If a row corresponding to a label shows a NaN value, it means that the validation dataset used to check model predictions does not contain data with that label.

Create an AutoML job for text classification using the API

The following instructions show how to create an Amazon SageMaker Autopilot job as a pilot experiment for text classification problem types using SageMaker API Reference.

Note

Tasks such as text and image classification, time-series forecasting, and fine-tuning of large language models are exclusively available through the version 2 of the Autopilot API. For Python users, we recommend using the AWS SDK for Python (Boto3) as the Amazon SageMaker Python SDK is not currently supported for the Autopilot API version 2.
Users who prefer the convenience of a user interface can use Amazon SageMaker Canvas to access pre-trained models and generative AI foundation models, or create custom models tailored for specific text, image classification, forecasting needs, or generative AI.

You can create an Autopilot text classification experiment programmatically by calling the CreateAutoMLJobV2 API action in any language supported by Amazon SageMaker Autopilot or the AWS CLI.

For information on how this API action translates into a function in the language of your choice, see the See Also section of CreateAutoMLJobV2 and choose an SDK. As an example, for Python users, see the full request syntax of create_auto_ml_job_v2 in AWS SDK for Python (Boto3).

The following is a collection of mandatory and optional input request parameters for the CreateAutoMLJobV2 API action used in text classification.

**Required parameters**

When calling CreateAutoMLJobV2 to create an Autopilot experiment for text classification, you must provide the following values:

- An AutoMLJobName to specify the name of your job.
- At least one AutoMLJobChannel in AutoMLJobInputDataConfig to specify your data source.
- An AutoMLProblemTypeConfig of type TextClassificationJobConfig.
- An OutputDataConfig to specify the Amazon S3 output path to store the artifacts of your AutoML job.
- A RoleArn to specify the ARN of the role used to access your data.

All other parameters are optional.

**Optional parameters**

The following sections provide details of some optional parameters that you can pass to your text classification AutoML job.

**How to specify the training and validation datasets of an AutoML job**

You can provide your own validation dataset and custom data split ratio, or let Autopilot split the dataset automatically.
Each `AutoMLJobChannel` object (see the required parameter `AutoMLJobInputDataConfig`) has a `ChannelType`, which can be set to either `training` or `validation` values that specify how the data is to be used when building a machine learning model.

At least one data source must be provided and a maximum of two data sources is allowed: one for training data and one for validation data. How you split the data into training and validation datasets depends on whether you have one or two data sources.

How you split the data into training and validation datasets depends on whether you have one or two data sources.

- If you only have **one data source**, the `ChannelType` is set to `training` by default and must have this value.
  - If the `ValidationFraction` value in `AutoMLDataSplitConfig` is not set, 0.2 (20%) of the data from this source is used for validation by default.
  - If the `ValidationFraction` is set to a value between 0 and 1, the dataset is split based on the value specified, where the value specifies the fraction of the dataset used for validation.

- If you have **two data sources**, the `ChannelType` of one of the `AutoMLJobChannel` objects must be set to `training`, the default value. The `ChannelType` of the other data source must be set to `validation`. The two data sources must have the same format, either CSV or Parquet, and the same schema. You must not set the value for the `ValidationFraction` in this case because all of the data from each source is used for either training or validation. Setting this value causes an error.

**How to specify the automatic model deployment configuration for an AutoML job**

To enable automatic deployment for the best model candidate of an AutoML job, include a `ModelDeployConfig` in the AutoML job request. This will allow the deployment of the best model to a SageMaker endpoint. Below are the available configurations for customization.

- To let Autopilot generate the endpoint name, set `AutoGenerateEndpointName` to `True`.
- To provide your own name for the endpoint, set `AutoGenerateEndpointName` to `False` and provide a name of your choice in `EndpointName`. 
Datasets format and objective metric for text classification

In this section we learn about the available formats for datasets used in text classification as well as the metric used to evaluate the predictive quality of machine learning model candidates. The metrics calculated for candidates are specified using an array of `MetricDatum` types.

Datasets formats

Autopilot supports tabular data formatted as CSV files or as Parquet files. For tabular data, each column contains a feature with a specific data type and each row contains an observation. The properties of these two file formats differ considerably.

- **CSV** (comma-separated-values) is a row-based file format that stores data in human readable plaintext which a popular choice for data exchange as they are supported by a wide range of applications.
- **Parquet** is a column-based file format where the data is stored and processed more efficiently than row-based file formats. This makes them a better option for big data problems.

The **data types** accepted for columns include numerical, categorical, text.

Autopilot supports building machine learning models on large datasets up to hundreds of GBs. For details on the default resource limits for input datasets and how to increase them, see Amazon SageMaker Autopilot quotas.

Objective metric

The following list contains the names of the metrics that are currently available to measure the performance of models for text classification.

**Accuracy**

The ratio of the number of correctly classified items to the total number of (correctly and incorrectly) classified items. Accuracy measures how close the predicted class values are to the actual values. Values for accuracy metrics vary between zero (0) and one (1). A value of 1 indicates perfect accuracy, and 0 indicates perfect inaccuracy.

Autopilot model deployment and prediction

This Autopilot guide includes steps for model deployment and setting up real-time inference.
After you train your Autopilot models, you can set up an endpoint and obtain predictions interactively.

**Real-time inferencing**

Real-time inference is ideal for inference workloads where you have real-time, interactive, low latency requirements. This section shows how you can use real-time inferencing to obtain predictions interactively from your model.

You can use SageMaker APIs to manually deploy the model that produced the best validation metric in an Autopilot experiment as follows.

Alternatively, you can choose the automatic deployment option when creating your Autopilot experiment. For information on setting up the automatic deployment of models, see **ModelDeployConfig** in the request parameters of **CreateAutoMLJobV2**. This creates an endpoint automatically.

---

**Note**

To avoid incurring unnecessary charges, you can delete unneeded endpoint and resources created from model deployment. For information about pricing of instances by Region, see [Amazon SageMaker Pricing](https://aws.amazon.com/sagemaker/pricing/).

---

1. **Obtain the candidate container definitions**

   Obtain the candidate container definitions from **InferenceContainers**. A container definition for inference refers to the containerized environment designed for deploying and running your trained SageMaker model to make predictions.

   The following AWS CLI command example uses the **DescribeAutoMLJobV2** API to obtain candidate definitions for the best model candidate.

   ```bash
   aws sagemaker describe-auto-ml-job-v2 --auto-ml-job-name job-name --region region
   ```

2. **List candidates**

   The following AWS CLI command example uses the **ListCandidatesForAutoMLJob** API to list all model candidates.

---

Create a Text Classification job using the AutoML API
3. **Create a SageMaker model**

Use the container definitions from the previous steps and a candidate of your choice to create a SageMaker model by using the CreateModel API. See the following AWS CLI command as an example.

```bash
aws sagemaker create-model --model-name '<your-candidate-name>' \
    --containers ['<container-definition1>, <container-definition2>, <container-definition3>'] \
    --execution-role-arn '<execution-role-arn>' --region '<region>'
```

4. **Create an endpoint configuration**

The following AWS CLI command example uses the CreateEndpointConfig API to create an endpoint configuration.

```bash
aws sagemaker create-endpoint-config --endpoint-config-name '<your-endpoint-config-name>' \
    --production-variants '<list-of-production-variants>' \
    --region '<region>'
```

5. **Create the endpoint**

The following AWS CLI example uses the CreateEndpoint API to create the endpoint.

```bash
aws sagemaker create-endpoint --endpoint-name '<your-endpoint-name>' \
    --endpoint-config-name '<endpoint-config-name-you-just-created>' \
    --region '<region>'
```

Check the progress of your endpoint deployment by using the DescribeEndpoint API. See the following AWS CLI command as an example.

```bash
aws sagemaker describe-endpoint --endpoint-name '<endpoint-name>' --region '<region>'
```

After the EndpointStatus changes to InService, the endpoint is ready to use for real-time inference.
6. Invoke the endpoint

The following command structure invokes the endpoint for real-time inferencing.

```bash
aws sagemaker invoke-endpoint --endpoint-name '<endpoint-name>' \
--region '<region>' --body '<your-data>' [--content-type] '<content-type>' <outfile>
```

Explainability report

Amazon SageMaker Autopilot provides an explainability report to help explain how a best model candidate makes predictions for text classification problems. This report can assist ML engineers, product managers, and other internal stakeholders in understanding the characteristics of the model. Both consumers and regulators rely on transparency in machine learning to trust and interpret decisions made on model predictions. You can use these explanations for auditing and meeting regulatory requirements, establishing trust in the model, supporting human decision-making, and debugging and improving model performance.

The Autopilot explanatory functionality for text classification uses the axiomatic attribution method *Integrated Gradients*. This approach relies on an implementation of *Axiomatic Attribution for Deep Network*.

Autopilot generates the explainability report as a JSON file. The report includes analysis details that are based on the validation dataset. Each sample used to generate the report contains the following information:

- **text**: The input text content explained.
- **token_scores**: The list of scores for every token in the text.
- **attribution**: The score depicting the importance of the token.
  - **description.partial_text**: The partial substring that represents the token.
- **predicted_label**: The label class predicted by the best model candidate.
- **probability**: The confidence with which the predicted_label was predicted.

You can find the Amazon S3 prefix to the explainability artifacts generated for the best candidate in the response to `DescribeAutoMLJobV2` at `BestCandidate.CandidateProperties.CandidateArtifactLocations.Explainability`. 

Create a Text Classification job using the AutoML API
The following is an example of analysis content that you could find in the explainability artifacts.

```json
{
  "text": "It was a fantastic movie!",
  "predicted_label": 2,
  "probability": 0.9984835,
  "token_scores": [
    {
      "attribution": 0,
      "description": {
        "partial_text": "It"
      }
    },
    {
      "attribution": -0.022447118861679088,
      "description": {
        "partial_text": "was"
      }
    },
    {
      "attribution": -0.2164326456817965,
      "description": {
        "partial_text": "a"
      }
    },
    {
      "attribution": 0.675,
      "description": {
        "partial_text": "fantastic"
      }
    },
    {
      "attribution": 0.416,
      "description": {
        "partial_text": "movie!"
      }
    }
  ]
}
```

In this sample of the JSON report, the explanatory functionality evaluates the text *It was a fantastic movie!* and scores the contribution of each of its token to the overall predicted label. The predicted label is 2, which is a strong positive sentiment, with a probability of 99.85%.
The JSON sample then details the contribution of each individual token to this prediction. For example, the token fantastic has a stronger attribution than the token was. It is the token that contributed the most to the final prediction.

**Model performance report**

An Amazon SageMaker model quality report (also referred to as performance report) provides insights and quality information for the best model candidate generated by an AutoML job. This includes information about the job details, model problem type, objective function, and various metrics. This section details the content of a performance report for text classification problems and explains how to access the metrics as raw data in a JSON file.

You can find the Amazon S3 prefix to the model quality report artifacts generated for the best candidate in the response to `DescribeAutoMLJobV2` at `BestCandidate.CandidateProperties.CandidateArtifactLocations.ModelInsights`.

The performance report contains two sections:

- The first section contains details about the Autopilot job that produced the model.
- The second section contains a model quality report with various performance metrics.

**Autopilot job details**

This first section of the report gives some general information about the Autopilot job that produced the model. These details include the following information:

- Autopilot candidate name: The name of the best model candidate.
- Autopilot job name: The name of the job.
- Problem type: The problem type. In our case, text classification.
- Objective metric: The objective metric used to optimize the performance of the model. In our case, Accuracy.
- Optimization direction: Indicates whether to minimize or maximize the objective metric.

**Model quality report**

Model quality information is generated by Autopilot model insights. The report's content that is generated depends on the problem type it addressed. The report specifies the number of rows that were included in the evaluation dataset and the time at which the evaluation occurred.
Metrics tables

The first part of the model quality report contains metrics tables. These are appropriate for the type of problem that the model addressed.

The following image is an example of a metrics table generated by Autopilot for an image or text classification problem. It shows the metric name, value, and standard deviation.

### Metrics table

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Value</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>weighted_recall</td>
<td>0.597104</td>
<td>0.005410</td>
</tr>
<tr>
<td>weighted_precision</td>
<td>0.591693</td>
<td>0.005729</td>
</tr>
<tr>
<td>accuracy</td>
<td>0.597104</td>
<td>0.005410</td>
</tr>
<tr>
<td>weighted_f0.5</td>
<td>0.592155</td>
<td>0.005659</td>
</tr>
<tr>
<td>weighted_f1</td>
<td>0.593423</td>
<td>0.005554</td>
</tr>
<tr>
<td>weighted_f2</td>
<td>0.595392</td>
<td>0.005456</td>
</tr>
<tr>
<td>accuracy_best_constant_classifier</td>
<td>0.200699</td>
<td>0.004422</td>
</tr>
<tr>
<td>weighted_recall_best_constant_classifier</td>
<td>0.200699</td>
<td>0.004422</td>
</tr>
<tr>
<td>weighted_precision_best_constant_classifier</td>
<td>0.040280</td>
<td>0.001753</td>
</tr>
<tr>
<td>weighted_f0.5_best_constant_classifier</td>
<td>0.047944</td>
<td>0.002039</td>
</tr>
<tr>
<td>weighted_f1_best_constant_classifier</td>
<td>0.067094</td>
<td>0.002684</td>
</tr>
<tr>
<td>weighted_f2_best_constant_classifier</td>
<td>0.111716</td>
<td>0.003808</td>
</tr>
</tbody>
</table>

Graphical model performance information

The second part of the model quality report contains graphical information to help you evaluate model performance. The contents of this section depend on the selected problem type.

Confusion matrix

A confusion matrix provides a way to visualize the accuracy of the predictions made by a model for binary and multiclass classification for different problems.

A summary of the graph's components of **false positive rate** (FPR) and **true positive rate** (TPR) are defined as follows.

- Correct predictions
  - **True positive** (TP): The predicted is 1, and the true value is 1.
  - **True negative** (TN): The predicted is 0, and the true value is 0.
- Erroneous predictions
• **False positive** (FP): The predicted value is 1, but the true value is 0.

• **False negative** (FN): The predicted value is 0, but the true value is 1.

The confusion matrix in the model quality report contains the following.

• The number and percentage of correct and incorrect predictions for the actual labels

• The number and percentage of accurate predictions on the diagonal from the upper-left to the lower-right corner

• The number and percentage of inaccurate predictions on the diagonal from the upper-right to the lower-left corner

The incorrect predictions on a confusion matrix are the confusion values.

The following diagram is an example of a confusion matrix for a multi-class classification problem. The confusion matrix in the model quality report contains the following.

• The vertical axis is divided into three rows containing three different actual labels.

• The horizontal axis is divided into three columns containing labels that were predicted by the model.

• The color bar assigns a darker tone to a larger number of samples to visually indicate the number of values that were classified in each category.

In the example below, the model correctly predicted actual 354 values for label \( f \), 1094 values for label \( i \) and 852 values for label \( m \). The difference in tone indicates that the dataset is not balanced because there are many more labels for the value \( i \) than for \( f \) or \( m \).
The confusion matrix in the model quality report provided can accommodate a maximum of 15 labels for multiclass classification problem types. If a row corresponding to a label shows a Nan value, it means that the validation dataset used to check model predictions does not contain data with that label.

Create an AutoML job for time-series forecasting using the API

Forecasting in machine learning refers to the process of predicting future outcomes or trends based on historical data and patterns. By analyzing past time-series data and identifying underlying patterns, machine learning algorithms can make predictions and provide valuable insights into future behavior. In forecasting, the goal is to develop models that can accurately capture the relationship between input variables and the target variable over time. This involves examining various factors such as trends, seasonality, and other relevant patterns within the data. The collected information is then used to train a machine learning model. The trained model is capable of generating predictions by taking new input data and applying the learned patterns and
relationships. It can provide forecasts for a wide range of use cases, such as sales projections, stock market trends, weather forecasts, demand forecasting, and many more.

The following instructions show how to create an Amazon SageMaker Autopilot job as a pilot experiment for time-series forecasting problem types using SageMaker API Reference.

Note

Tasks such as text and image classification, time-series forecasting, and fine-tuning of large language models are exclusively available through the version 2 of the Autopilot API. For Python users, we recommend using the AWS SDK for Python (Boto3) as the Amazon SageMaker Python SDK is not currently supported for the Autopilot API version 2. Users who prefer the convenience of a user interface can use Amazon SageMaker Canvas to access pre-trained models and generative AI foundation models, or create custom models tailored for specific text, image classification, forecasting needs, or generative AI.

You can create an Autopilot time-series forecasting experiment programmatically by calling the CreateAutoMLJobV2 API in any language supported by Amazon SageMaker Autopilot or the AWS CLI.

For information on how this API action translates into a function in the language of your choice, see the See Also section of CreateAutoMLJobV2 and choose an SDK. As an example, for Python users, see the full request syntax of create_auto_ml_job_v2 in AWS SDK for Python (Boto3).

Autopilot trains several model candidates with your target time-series, then selects an optimal forecasting model for a given objective metric. When your model candidates have been trained, you can find the best candidate metrics in the response to DescribeAutoMLJobV2 at BestCandidate.

The following sections define the mandatory and optional input request parameters for the CreateAutoMLJobV2 API used in time-series forecasting.

Note

Refer to the notebook Time-Series Forecasting with Amazon SageMaker Autopilot for a practical, hands-on time-series forecasting example. In this notebook, you use Amazon SageMaker Autopilot to train a time-series model and produce predictions using the trained
The notebook provides instructions for retrieving a ready-made dataset of tabular historical data on Amazon S3.

**Prerequisites**

Before using Autopilot to create a time-series forecasting experiment in SageMaker, make sure to:

- Prepare your time-series dataset. Dataset preparation involves collecting relevant data from various sources, cleaning and filtering it to remove noise and inconsistencies, and organizing it into a structured format. See [Time-series datasets format and missing values filling methods](#) to learn more about time-series formats requirements in Autopilot. Optionally, you can supplement your dataset with the public holiday calendar of the country of your choice to capture associated patterns. For more information on holiday calendars, see [National holiday calendars](#).

  **Note**
  
  We recommend providing at least 3-5 historical data points for each 1 future data point you want to predict. For example, to forecast 7 days ahead (horizon of 1 week) based on daily data, train your model on a minimum of 21-35 days of historical data. Make sure to provide enough data to capture seasonal and recurrent patterns.

- Place your time-series data in an Amazon S3 bucket.
- Grant full access to the Amazon S3 bucket containing your input data for the SageMaker execution role used to run your experiment. Once this is done, you can use the ARN of this execution role in Autopilot API requests.
- For information on retrieving your SageMaker execution role, see [Get execution role](#).
- For information on granting your SageMaker execution role permissions to access one or more specific buckets in Amazon S3, see [Add Additional Amazon S3 Permissions to a SageMaker Execution Role](#) in [Create execution role](#).

**Required parameters**

When calling [CreateAutoMLJobV2](#) to create an Autopilot experiment for time-series forecasting, you must provide the following values:
• An **AutoMLJobName** to specify the name of your job. The name should be of type `string`, and should have a minimum length of 1 character and a maximum length of 32.

• At least one **AutoMLJobChannel** in **AutoMLJobInputDataConfig** in which you specify the name of the Amazon S3 bucket that contains your data. Optionally, you can specify the content (CSV or Parquet files) and compression (GZip) types.

• An **AutoMLProblemTypeConfig** of type **TimeSeriesForecastingJobConfig** to configure the settings of your time-series forecasting job. In particular, you must specify:

  • The **frequency** of predictions, which refers to the desired granularity (hourly, daily, monthly, etc) of your forecast.

    Valid intervals are an integer followed by **Y** (Year), **M** (Month), **W** (Week), **D** (Day), **H** (Hour), and **min** (Minute). For example, 1D indicates every day and 15min indicates every 15 minutes. The value of a frequency must not overlap with the next larger frequency. For example, you must use a frequency of 1H instead of 60min.

    The valid values for each frequency are the following:

    • Minute - 1-59
    • Hour - 1-23
    • Day - 1-6
    • Week - 1-4
    • Month - 1-11
    • Year - 1

  • The **horizon** of predictions in your forecast, which refers to the number of time-steps that the model predicts. The forecast horizon is also called the prediction length. The maximum forecast horizon is the lesser of 500 time-steps or 1/4 of the time-steps in the dataset.

  • A **TimeSeriesConfig** in which you define the schema of your dataset to map the column headers to your forecast by specifying:

    • A **TargetAttributeName**: The column that contains historical data of the target field to forecast.

    • A **TimestampAttributeName**: The column that contains a point in time at which the target value of a given item is recorded.

    • A **ItemIdentifierAttributeName**: The column that contains the item identifiers for which you want to predict the target value.
The following is an example of those request parameters. In this example, you are setting up a daily forecast for the expected quantity or level of demand of specific items over a period of 20 days.

```json
"AutoMLProblemTypeConfig": {
   "ForecastFrequency": "D",
   "ForecastHorizon": 20,
   "TimeSeriesConfig": {
      "TargetAttributeName": "demand",
      "TimestampAttributeName": "timestamp",
      "ItemIdentifierAttributeName": "item_id"
   }
},
```

- An **OutputDataConfig** to specify the Amazon S3 output path to store the artifacts of your AutoML job.
- A **RoleArn** to specify the ARN of the role used to access your data. You can use the ARN of the execution role to which you have granted access to your data.

All other parameters are optional. For example, you can set specific forecast quantiles, choose a filling method for missing values in the dataset, or define how to aggregate data that does not align with forecast frequency. To learn how to set those additional parameters, see **Optional parameters**.

### Optional parameters

The following sections provide details of some optional parameters that you can pass to your time-series forecasting AutoML job.

#### How to specify custom quantiles

Autopilot trains 6 models candidates with your target time-series, then combines these models using a stacking ensemble method to create an optimal forecasting model for a given objective metric. Each Autopilot forecasting model generates a probabilistic forecast by producing forecasts at quantiles between P1 and P99. These quantiles are used to account for forecast uncertainty. By default, forecasts will be generated for the 0.1 (p10), 0.5 (p50), and 0.9 (p90). You can choose to specify your own quantiles.
In Autopilot, you can specify up to five forecast quantiles from 0.01 (p1) to 0.99 (p99), by increments of 0.01 or higher in the ForecastQuantiles attribute of `TimeSeriesForecastingJobConfig`.

In the following example, you are setting up a daily 10th, 25th, 50th, 75th, and 90th percentile forecast for the expected quantity or level of demand of specific items over a period of 20 days.

```
"AutoMLProblemTypeConfig": {
  "ForecastFrequency": "D",
  "ForecastHorizon": 20,
  "ForecastQuantiles": ["p10", "p25", "p50", "p75", "p90"],
  "TimeSeriesConfig": {
    "TargetAttributeName": "demand",
    "TimestampAttributeName": "timestamp",
    "ItemIdentifierAttributeName": "item_id"
  }
},
```

**How to aggregate data for different forecast frequencies**

To create a forecast model (also referred to as the best model candidate from your experiment), you must specify a forecast frequency. The forecast frequency determines the frequency of predictions in your forecasts. For example, monthly sales forecasts. Autopilot best model can generate forecasts for data frequencies that are higher than the frequency at which your data is recorded.

During training, Autopilot aggregates any data that does not align with the forecast frequency you specify. For example, you might have some daily data but specify a weekly forecast frequency. Autopilot aligns the daily data based on the week that it belongs in. Autopilot then combines it into single record for each week.

During aggregation, the default transformation method is to sum the data. You can configure the aggregation when you create your AutoML job in the `Transformations` attribute of `TimeSeriesForecastingJobConfig`. The supported aggregation methods are sum (default), avg, first, min, max. Aggregation is only supported for the target column.

In the following example, you configure the aggregation to calculate the average of the individual promo forecasts to provide the final aggregated forecast values.

```
"Transformations": {
  "Aggregation": {
  ```
How to handle missing values in your input datasets

Autopilot provides a number of filling methods to handle missing values in the target and other numeric columns of your time-series datasets. For information on the list of supported filling methods and their available filling logic, see Handle missing values.

You configure your filling strategy in the Transformations attribute of TimeSeriesForecastingJobConfig when creating your AutoML job.

To set a filling method, you need to provide a key-value pair:

- The key is the name of the column for which you want to specify the filling method.
- The value associated with the key is an object that defines the filling strategy for that column.

You can specify multiple filling methods for a single column.

To set a specific value for the filling method, you should set the fill parameter to the desired filling method value (for example "backfill" : "value"), and define the actual filling value in an additional parameter suffixed with "_value". For example, to set backfill to a value of 2, you must include two parameters: "backfill": "value" and "backfill_value": "2".

In the following example, you specify the filling strategy for the incomplete data column, "price" as follows: All missing values between the first data point of an item and the last are set to 0 after which all missing values are filled with the value 2 until the end date of the dataset.

```
"Transformations": {

  "Filling": {

    "price": {

      "middlefill": "zero",
      "backfill": "value",
      "backfill_value": "2"

    }

  }

}
```
How to specify an objective metric

Autopilot produces accuracy metrics to evaluate the model candidates and help you choose which to use to generate forecasts. When you run a time-series forecasting experiment, you can either choose AutoML to let Autopilot optimize the predictor for you, or you can manually choose an algorithm for your predictor.

By default, Autopilot uses the Average Weighted Quantile Loss. However, you can configure the objective metric when you create your AutoML job in the MetricName attribute of AutoMLJobObjective.

For the list of available algorithms, see Algorithms support for time-series forecasting.

How to incorporate national holiday information to your dataset

In Autopilot, you can incorporate a feature-engineered dataset of national holiday information to your time-series. Autopilot provide native support for the holiday calendars of over 250 countries. After you choose a country, Autopilot applies that country's holiday calendar to every item in your dataset during training. This allows the model to identify patterns associated with specific holidays.

You can enable the holiday featurization when you create your AutoML job by passing an HolidayConfigAttributes object to the HolidayConfig attribute of TimeSeriesForecastingJobConfig. The HolidayConfigAttributes object contains the two letters CountryCode attribute that determines the country of the public national holiday calendar used to augment your time-series dataset.

Refer to Country Codes for the list of supported calendars and their corresponding country code.

How to enable automatic deployment

Autopilot allows you to automatically deploy your forecast model to an endpoint. To enable automatic deployment for the best model candidate of an AutoML job, include a ModelDeployConfig in the AutoML job request. This allows the deployment of the best model to a SageMaker endpoint. Below are the available configurations for customization.

- To let Autopilot generate the endpoint name, set AutoGenerateEndpointName to True.
- To provide your own name for the endpoint, set AutoGenerateEndpointName to False and provide a name of your choice in EndpointName.
**Time-series datasets format and missing values filling methods**

Time-series data refers to a collection of observations or measurements recorded over regular intervals of time. In this type of data, each observation is associated with a specific timestamp or time period, creating a sequence of data points ordered chronologically.

The specific columns you include in your time-series dataset depend on the goals of your analysis and the data available to you. At a minimum, the time-series data is composed of a 3-column table where:

- One column contains unique identifiers assigned to individual items to refer to their value at a specific moment.
- Another column represents the point-in-time value or `target` to log the value of a given item at a specific moment. After the model is trained on those target values, this target column contains the values that the model predicts at a specified frequency within a defined horizon.
- And a timestamp column is included to record the date and time when the value was measured.
- Additional columns can contain other factors that may influence the forecast performance. For example, in a time-series dataset for retail where the target is the sales or revenue, you might include features that provide information about units sold, product ID, store location, customer count, inventory levels, as well as covariate indicators such as weather data or demographic information.

**Note**

You can add a feature-engineered dataset of national holiday information to your time-series. By including holidays in your time series model, you can capture the periodic patterns that holidays create. This helps your forecasts better reflect the underlying seasonality of your data. For information on the available calendars per country, see [National holiday calendars](#).

**Datasets format for time-series forecasting**

Autopilot supports numeric, categorical, text, and datetime data types. The data type of the target column must be numeric.

Autopilot supports time-series data formatted as CSV (default) files or as Parquet files.
• **CSV** (comma-separated-values) is a row-based file format that stores data in human readable plaintext which a popular choice for data exchange as they are supported by a wide range of applications.

• **Parquet** is a column-based file format where the data is stored and processed more efficiently than row-based file formats. This makes them a better option for big data problems.

For more information about the resource limits on time-series datasets for forecasting in Autopilot, see [Amazon SageMaker Autopilot time-series forecasting resource limits](#).

### Handle missing values

A common issue in time-series forecasting data is the presence of missing values. Your data might contain missing values for a number of reasons, including measurement failures, formatting problems, human errors, or a lack of information to record. For instance, if you are forecasting product demand for a retail store and an item is sold out or unavailable, there would be no sales data to record while that item is out of stock. If prevalent enough, missing values can significantly impact a model's accuracy.

Autopilot provides a number of filling methods to handle missing values, with distinct approaches for the target column and other additional columns. Filling is the process of adding standardized values to missing entries in your dataset.

Refer to [How to handle missing values in your input datasets](#) to learn how to set the method for filling missing values in your time-series dataset.

Autopilot supports the following filling methods:

• **Front filling**: Fills any missing values between the earliest recorded data point among all items and the starting point of each item (each item can start at a different time). This ensures that the data for each item is complete and spans from the earliest recorded data point to its respective starting point.

• **Middle filling**: Fills any missing values between the start and end dates of the items in the dataset.

• **Back filling**: Fills any missing values between the last data point of each item (each item can stop at a different time) and the last recorded data point among all items.

• **Future filling**: Fills any missing values between the last recorded data point among all items and the end of the forecast horizon.
The following image provides a visual representation of the different filling methods.

<table>
<thead>
<tr>
<th>Start date of dataset (First record)</th>
<th>First recorded data of item A</th>
<th>Last recorded data of item A</th>
<th>End date of dataset (Last record)</th>
<th>End of horizon</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Front</td>
<td>Middle</td>
<td>Back</td>
<td>Future</td>
</tr>
</tbody>
</table>

Choose a filling logic

When choosing a filling logic, you should consider how the logic will be interpreted by your model. For instance, in a retail scenario, recording 0 sales of an available item is different from recording 0 sales of an unavailable item, as the latter does not imply a lack of customer interest in the item. Because of this, 0 filling in the target column of the time-series might cause the predictor to be under-biased in its predictions, while NaN filling might ignore actual occurrences of 0 available items being sold and cause the predictor to be over-biased.

Filling logic

You can perform filling on the target column and other numeric columns in your datasets. Target columns have different filling guidelines and restrictions than the rest of the numeric columns.

Filling Guidelines

<table>
<thead>
<tr>
<th>Column type</th>
<th>Filling by default?</th>
<th>Supported filling methods</th>
<th>Default filling logic</th>
<th>Accepted filling logic</th>
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<tbody>
<tr>
<td>Target column</td>
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<td>• zero - 0 filling.</td>
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<td>• value - an integer or float number.</td>
</tr>
<tr>
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<td>• nan - not a number.</td>
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<tr>
<td></td>
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<td></td>
<td></td>
<td>• mean - the mean value from the data series.</td>
</tr>
<tr>
<td>Column type</td>
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<td>Supported filling methods</td>
<td>Default filling logic</td>
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<td>• median - the median value from the data series.</td>
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<td></td>
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<td>• max - the maximum value from the data series.</td>
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</table>

**Note**

For both the target and other numeric columns, mean, median, min, and max are calculated based on a rolling window of the 64 most recent data entries before the missing values.
# National holiday calendars

Autopilot supports a feature-engineered dataset of national holiday information that provides access to the holiday calendars of over 250 countries.

Holiday calendar features are especially useful in the retail domain, where public holidays can significantly affect demand.

Refer to [How to incorporate national holiday information to your dataset](#) to learn how to add a calendar to your dataset.

## Country Codes

Autopilot provides native support for the public holiday calendars of the following countries. Use the **Country Code** when specifying a country with the API.

### Supported Countries

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Create a Time-series Forecasting job using the AutoML API
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Create a Time-series Forecasting job using the AutoML API
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<tr>
<td>Zimbabwe</td>
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</table>
Objective metrics

Autopilot produces accuracy metrics to evaluate the model candidates and help you choose which to use to generate forecasts. You can either let Autopilot optimize the predictor for you, or you can manually choose an algorithm for your predictor. By default, Autopilot uses the Average Weighted Quantile Loss.

The following list contains the names of the metrics that are currently available to measure the performance of models for time-series forecasting.

**RMSE**

Root mean squared error (RMSE) – Measures the square root of the squared difference between predicted and actual values, and is averaged over all values. It's an important metric to indicate the presence of large model errors and outliers. Values range from zero (0) to infinity, with smaller numbers indicating a better model fit to the data. RMSE is dependent on scale, and should not be used to compare datasets of different sizes.

**wQL**

Weighted Quantile Loss (wQL) – Assess the accuracy of the forecast by measuring the weighted absolute differences between predicted and actual P10, P50, and P90 quantiles with lower values indicating better performance.

**Average wQL (default)**

Average Weighted Quantile Loss (Average wQL) – Evaluates the forecast by averaging the accuracy at the P10, P50, and P90 quantiles. A lower value indicates a more accurate model.

**MASE**

Mean Absolute Scaled Error (MASE) – The mean absolute error of the forecast normalized by the mean absolute error of a simple baseline forecasting method. A lower value indicates a more accurate model, where MASE < 1 is estimated to be better than the baseline and MASE > 1 is estimated to be worse than the baseline.

**MAPE**

Mean Absolute Percent Error (MAPE) – The percentage error (percent difference of the mean forecasted value versus the actual value) averaged over all time points. A lower value indicates a more accurate model, where MAPE = 0 is a model with no errors.
WAPE

Weighted Absolute Percent Error (WAPE) – The sum of the absolute error normalized by the sum of the absolute target, which measure the overall deviation of forecasted values from observed values. A lower value indicates a more accurate model.

Algorithms support for time-series forecasting

Autopilot trains the following six built-in algorithms with your target time-series. Then, using a stacking ensemble method, it combines these model candidates to create an optimal forecasting model for a given objective metric.


- **Prophet** – Prophet is a popular local Bayesian structural time series model based on an additive model where non-linear trends are fit with yearly, weekly, and daily seasonality. The Autopilot Prophet algorithm uses the *Prophet class* of the Python implementation of Prophet. It works best with time-series with strong seasonal effects and several seasons of historical data.

- **Non-Parametric Time Series (NPTS)** – The NPTS proprietary algorithm is a scalable, probabilistic baseline forecaster. It predicts the future value distribution of a given time-series by sampling from past observations. NPTS is especially useful when working with sparse or intermittent time series.

- **Autoregressive Integrated Moving Average (ARIMA)** – ARIMA is a commonly used statistical algorithm for time-series forecasting. The algorithm captures standard temporal structures (patterned organizations of time) in the input dataset. It is especially useful for simple datasets with under 100 time series.

- **Exponential Smoothing (ETS)** – ETS is a commonly used statistical algorithm for time-series forecasting. The algorithm is especially useful for simple datasets with under 100 time series, and datasets with seasonality patterns. ETS computes a weighted average over all observations in the time series dataset as its prediction, with exponentially decreasing weights over time.
Autopilot model deployment and forecasts

After you train your Autopilot predictor (best model), you can deploy a model to get predictions in one of two ways:

1. Use **Real-time forecasting** to set up an endpoint and obtain predictions interactively.
2. Use **Batch forecasting** to make predictions in parallel on batches of observations on an entire dataset.

When providing input data for forecasting, the schema of your data should remain the same as the one used to train your model, including the number of columns, column headers, and data types. You can forecast for existing or new item IDs within the same or different timestamp range to predict for a different time period.

Forecasting models predict for the forecast horizon points in the future specified in the input request at training, which is from the *target end date* to the *target end date + forecast horizon*. To use the model for predicting specific dates, you should provide the data in the same format as the original input data, extending up to a specified *target end date*. In this scenario, the model will start predicting from the new target end date.

For example, if your dataset had monthly data from January to June with a Forecast horizon of 2, then the model would predict the target value for the next 2 months, which would be July and August. If in August, you want to predict for the next 2 months, this time your input data should be from January to August and the model will predict for the next 2 months (September, October).

When forecasting future data points, there is no set minimum for the amount of historical data to provide. Include enough data to capture seasonal and recurrent patterns in your time-series.

**Note**

We recommend using the following instance types for forecasting:

- For real-time forecasting, use **m5.12xlarge** instances.
- For batch forecasting, use m5.12xlarge instances for general-purpose workloads and m5.24xlarge instances for big data forecasting tasks.
Real-time forecasting

You can use real-time forecasting for inference workloads where you have real-time, interactive, low latency requirements.

Note

For real time forecasting, the dataset should be a subset of the input dataset. The real time endpoint has an input data size of approximately 6MB and a response timeout limitation of 60 seconds. We recommend bringing in one or few items at a time.

You can use SageMaker APIs to manually deploy the model that produced the best validation metric in an Autopilot experiment as follows.

Alternatively, you can chose the automatic deployment option when creating your Autopilot experiment. For information on setting up the automatic deployment of models, see How to enable automatic deployment.

1. Obtain the candidate container definitions

Obtain the candidate container definitions from InferenceContainers. A container definition for inference refers to the containerized environment designed for deploying and running your trained SageMaker model to make predictions.

The following AWS CLI command example uses the DescribeAutoMLJobV2 API to obtain candidate definitions for the best model candidate.

```bash
aws sagemaker describe-auto-ml-job-v2 --auto-ml-job-name job-name --region region
```

2. List candidates

The following AWS CLI command example uses the ListCandidatesForAutoMLJob API to list all model candidates.

```bash
aws sagemaker list-candidates-for-auto-ml-job --auto-ml-job-name <job-name> --region <region>
```

3. Create a SageMaker model
Use the container definitions from the previous steps and a candidate of your choice to create a SageMaker model by using the `create-model` API. See the following AWS CLI command as an example.

```bash
aws sagemaker create-model --model-name '<your-candidate-name>' \
    --containers ['<container-definition1>, <container-definition2>, <container-definition3>'] \
    --execution-role-arn '<execution-role-arn>' --region '<region>'
```

4. Create an endpoint configuration

The following AWS CLI command example uses the `create-endpoint-config` API to create an endpoint configuration.

```bash
aws sagemaker create-endpoint-config --endpoint-config-name '<your-endpoint-config-name>' \
    --production-variants '<list-of-production-variants>' \
    --region '<region>'
```

5. Create the endpoint

The following AWS CLI example uses the `create-endpoint` API to create the endpoint.

```bash
aws sagemaker create-endpoint --endpoint-name '<your-endpoint-name>' \
    --endpoint-config-name '<endpoint-config-name-you-just-created>' \
    --region '<region>'
```

Check the progress of your endpoint deployment by using the `describe-endpoint` API. See the following AWS CLI command as an example.

```bash
aws sagemaker describe-endpoint --endpoint-name '<endpoint-name>' --region '<region>'
```

After the EndpointStatus changes to InService, the endpoint is ready to use for real-time inference.

6. Invoke the endpoint

The following command structure invokes the endpoint for real-time inferencing.
Batch forecasting

Batch forecasting, also known as offline inferencing, generates model predictions on a batch of observations. Batch inference is a good option for large datasets or if you don't need an immediate response to a model prediction request.

By contrast, online inference (real-time inferencing) generates predictions in real time.

You can make batch inferences from an Autopilot model using the API reference.

To use the SageMaker APIs for batch inferencing:

1. Obtain candidate definitions

   Candidate definitions from `InferenceContainers` are used to create a SageMaker model.

   The following example shows how to use the `DescribeAutoMLJobV2` API to obtain candidate definitions for the best model candidate. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker describe-auto-ml-job-v2 --auto-ml-job-name <job-name> --region <region>
   ```

   Use the `ListCandidatesForAutoMLJob` API to list all candidates. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker list-candidates-for-auto-ml-job --auto-ml-job-name <job-name> --region <region>
   ```

2. Create a SageMaker model

   To create a SageMaker model using the `CreateModel` API, use the container definitions from the previous steps. See the following AWS CLI command as an example.

   ```bash
   aws sagemaker create-model --model-name 'your-custom-model-name' --containers ["<container-definition1>, <container-definition2>, <container-definition3>]
   ```
3. Create a SageMaker transform job

The following example creates a SageMaker transform job with the CreateTransformJob API. See the following AWS CLI command as an example.

```
aws sagemaker create-transform-job --transform-job-name '<your-custom-transform-job-name>' --model-name '<your-custom-model-name-from-last-step>'
   --transform-input '{
       "DataSource": {
           "S3DataSource": {
               "S3DataType": "S3Prefix",
               "S3Uri": "<your-input-data>"
           },
           "ContentType": "text/csv",
           "SplitType": "None"
       }
   }
   --transform-output '{
       "S3OutputPath": "<your-output-path>",
       "AssembleWith": "Line"
   }
   --transform-resources '{
       "InstanceType": "<instance-type>",
       "InstanceCount": 1
   }' --region '<region>'
```

Check the progress of your transform job using the DescribeTransformJob API. See the following AWS CLI command as an example.

```
aws sagemaker describe-transform-job --transform-job-name '<your-custom-transform-job-name>' --region <region>
```

After the job is finished, the predicted result is available in <your-output-path>.

The output file name has the following format: <input_data_file_name>.out. As an example, if your input file is text_x.csv, the output name will be text_x.csv.out.

The following tabs show code examples for the AWS SDK for Python (boto3), and the AWS CLI.
The following example uses **AWS SDK for Python (boto3)** to make predictions in batches.

```python
import sagemaker
import boto3

session = sagemaker.session.Session()

sm_client = boto3.client('sagemaker', region_name='us-west-2')
role = 'arn:aws:iam::1234567890:role/sagemaker-execution-role'
output_path = 's3://test-auto-ml-job/output'
input_data = 's3://test-auto-ml-job/test_X.csv'

best_candidate = sm_client.describe_auto_ml_job_v2(AutoMLJobName=job_name)['BestCandidate']
best_candidate_containers = best_candidate['InferenceContainers']
best_candidate_name = best_candidate['CandidateName']

# create model
response = sm_client.create_model(
    ModelName = best_candidate_name,
    ExecutionRoleArn = role,
    Containers = best_candidate_containers
)

# Launch Transform Job
response = sm_client.create_transform_job(
    TransformJobName=f'{best_candidate_name}-transform-job',
    ModelName=model_name,
    TransformInput={
        'DataSource': {
            'S3DataSource': {
                'S3DataType': 'S3Prefix',
                'S3Uri': input_data
            }
        },
        'ContentType': "text/csv",
        'SplitType': 'None'
    },
    TransformOutput={
        'S3OutputPath': output_path,
        'AssembleWith': 'Line',
    },
)
```

Create a Time-series Forecasting job using the AutoML API
The batch inference job returns a response in the following format.

```json
{
    'ResponseMetadata': {
        'RequestId': '659f97fc-28c4-440b-b957-a49733f7c2f2',
        'HTTPStatusCode': 200,
        'HTTPHeaders': {
            'x-amzn-requestid': '659f97fc-28c4-440b-b957-a49733f7c2f2',
            'content-type': 'application/x-amz-json-1.1',
            'content-length': '96',
            'date': 'Thu, 11 Aug 2022 22:23:49 GMT'},
    'RetryAttempts': 0
}
```

AWS Command Line Interface (AWS CLI)

1. **Obtain the candidate definitions** by using the following the code example.

   ```bash
   aws sagemaker describe-auto-ml-job-v2 --auto-ml-job-name 'test-automl-job' --region us-west-2
   ```

2. **Create the model** by using the following the code example.

   ```bash
   aws sagemaker create-model --model-name 'test-sagemaker-model' --containers '[[
       "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-sklearn-automl:2.5-1-cpu-py3",
       "ModelDataUrl": "s3://test-bucket/out/test-job1/data-processor-models/test-job1-dpp0-1-e569ff7ad77f4e55a7e549a/output/model.tar.gz",
       "Environment": {
           "AUTOML_SPARSE_ENCODE_RECORDIO_PROTOBUF": "1",
           "AUTOML_TRANSFORM_MODE": "feature-transform",
           "SAGEMAKER_DEFAULT_INVOCATIONS_ACCEPT": "application/x-recordio-protobuf",
           "SAGEMAKER_PROGRAM": "sagemaker_serve",
           "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code"
       }
    }, {
       "Image": "348316444620.dkr.ecr.us-west-2.amazonaws.com/sagemaker-xgboost:1.3-1-cpu-py3",
    ]]
   ```
Create the transform job by using the following the code example.

```bash
aws sagemaker create-transform-job --transform-job-name 'test-transform-job'
   --model-name 'test-sagemaker-model'
   --transform-input '{
       "DataSource": {
           "S3DataSource": {
               "S3DataType": "S3Prefix",
               "S3Uri": "s3://test-bucket/data.csv"
           },
       "ContentType": "text/csv",
       "SplitType": "None"
   }
   --transform-output '{
       "S3OutputPath": "s3://test-bucket/output/"
   }'
```

Create a Time-series Forecasting job using the AutoML API
4. **Check the progress of the transform job** by using the following the code example.

```bash
aws sagemaker describe-transform-job --transform-job-name 'test-transform-job' --region us-west-2
```

The following is the response from the transform job.

```json
{
    "TransformJobName": "test-transform-job",
    "TransformJobStatus": "InProgress",
    "ModelName": "test-model",
    "TransformInput": {
        "DataSource": {
            "S3DataSource": {
                "S3DataType": "S3Prefix",
                "S3Uri": "s3://test-bucket/data.csv"
            }
        },
        "ContentType": "text/csv",
        "CompressionType": "None",
        "SplitType": "None"
    },
    "TransformOutput": {
        "S3OutputPath": "s3://test-bucket/output/",
        "AssembleWith": "Line",
        "KmsKeyId": ""
    },
    "TransformResources": {
        "InstanceType": "ml.m5.2xlarge",
        "InstanceCount": 1
    },
    "CreationTime": 1662495635.679,
    "TransformStartTime": 1662495847.496,
}
```
After the `TransformJobStatus` changes to `Completed`, you can check the inference result in the `S3OutputPath`.

**Amazon SageMaker Autopilot data exploration notebook**

Amazon SageMaker Autopilot cleans and pre-processes your dataset automatically. To help users understand their data, uncover patterns, relationships, and anomalies about the time-series, Amazon SageMaker Autopilot generates a **data exploration** static report in the form of a notebook for users to reference.

The data exploration notebook is generated for every Autopilot job. The report is stored in an Amazon S3 bucket and can be accessed from the job output path.

You can find the Amazon S3 prefix to the data exploration notebook in the response to `DescribeAutoMLJobV2` at `AutoMLJobArtifacts.DataExplorationNotebookLocation`.

**Reports generated by Amazon SageMaker Autopilot**

In addition to the data exploration notebook, Autopilot generates various reports for the best model candidate of each experiment.

- An explainability report provides insights into how the model makes forecasts.
- A performance report provides a quantitative assessment of the model's forecasting capabilities.
- A backtest results report is generated after testing the model's performance on historical data.

**Explainability report**

Autopilot explainability report helps you better understand how the attributes in your datasets impact forecasts for specific time-series (item and dimension combinations) and time points. Autopilot uses a metric called **Impact scores** to quantify the relative impact of each attribute and determine whether they increase or decrease forecast values.
For example, consider a forecasting scenario where the target is sales and there are two related attributes: price and color. Autopilot may find that the item’s color has a high impact on sales for certain items, but a negligible effect for other items. It may also find that a promotion in the summer has a high impact on sales, but a promotion in the winter has little effect.

The explainability report is generated only when:

- The time series dataset includes additional feature columns or is associated with a holiday calendar.
- The base models CNN-QR and DeepAR+ are included in the final ensemble.

**Interpret Impact scores**

Impact scores measure the relative impact attributes have on forecast values. For example, if the price attribute has an impact score that is twice as large as the store location attribute, you can conclude that the price of an item has twice the impact on forecast values than the store location.

Impact scores also provide information on whether attributes increase or decrease forecast values.

The Impact scores range from -1 to 1, where the sign denotes the direction of the impact. A score of 0 indicates no impact, while scores close to 1 or -1 indicate a significant impact.

It is important to note that Impact scores measure the relative impact of attributes, not the absolute impact. Therefore, Impact scores cannot be used to determine whether particular attributes improve model accuracy. If an attribute has a low Impact score, that does not necessarily mean that it has a low impact on forecast values; it means that it has a lower impact on forecast values than other attributes used by the predictor.

**Find the explainability report**

You can find the Amazon S3 prefix to the explainability artifacts generated for the best candidate in the response to `DescribeAutoMLJobV2` at `BestCandidate.CandidateProperties.CandidateArtifactLocations.Explainability`.

**Model performance report**

Autopilot model quality report (also referred to as performance report) provides insights and quality information for the best model candidate (best predictor) generated by an AutoML job. This
includes information about the job details, objective function, and accuracy metrics (wQL, MAPE, WAPE, RMSE, MASE).

You can find the Amazon S3 prefix to the model quality report artifacts generated for the best candidate in the response to DescribeAutoMLJobV2 at BestCandidate.CandidateProperties.CandidateArtifactLocations.ModelInsights.

Backtests results report

Backtests results provide insights into the performance of a time-series forecasting model by evaluating its predictive accuracy and reliability. It helps analysts and data scientists assess its performance on historical data and assists in understanding its potential performance on future, unseen data.

Autopilot uses backtesting to tune parameters and produce accuracy metrics. During backtesting, Autopilot automatically splits your time-series data into two sets, a training set and a testing set. The training set is used to train a model which is then used to generate forecasts for data points in the testing set. Autopilot uses this testing dataset to evaluate the model's accuracy by comparing forecasted values with observed values in the testing set.

You can find the Amazon S3 prefix to the model quality report artifacts generated for the best candidate in the response to DescribeAutoMLJobV2 at BestCandidate.CandidateProperties.CandidateArtifactLocations.BacktestResults.

Amazon SageMaker Autopilot time-series forecasting resource limits

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### Resource limits

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<th>Resource limits</th>
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### Create an AutoML job to fine-tune text generation models using the API

Large language models (LLMs) excel in multiple generative tasks, including text generation, summarization, completion, question answering, and more. Their performance can be attributed to their significant size and extensive training on diverse datasets and various tasks. However, specific domains, such as healthcare and financial services, may require customized fine-tuning to adapt to unique data and use cases. By tailoring their training to their particular domain, LLMs can improve their performance and provide more accurate outputs for targeted applications.

Autopilot offers the capability to fine-tune a selection of pre-trained generative text models. In particular, Autopilot supports the **instruction-based fine tuning** of a selection of general-purpose large language models (LLMs) powered by SageMaker JumpStart.

#### Note

The text generation models that support fine-tuning in Autopilot are currently accessible exclusively in Regions supported by SageMaker Canvas. See the documentation of SageMaker Canvas for the [full list of its supported Regions](#).

Fine-tuning a pre-trained model requires a specific dataset of clear instructions that guide the model on how to generate output or behave for that task. The model learns from the dataset, adjusting its parameters to conform to the provided instructions. Instruction-based fine-tuning involves using labeled examples formatted as prompt-response pairs and phrased as instructions. For more information about fine-tuning, see [Fine-tune a foundation model](#).
The following guidelines outline the process of creating an Amazon SageMaker Autopilot job as a pilot experiment to fine-tune text generation LLMs using the SageMaker API Reference.

**Note**
Tasks such as text and image classification, time-series forecasting, and fine-tuning of large language models are exclusively available through the version 2 of the Autopilot API. For Python users, we recommend using the AWS SDK for Python (Boto3) as the Amazon SageMaker Python SDK is not currently supported for the Autopilot API version 2.

Users who prefer the convenience of a user interface can use Amazon SageMaker Canvas to access pre-trained models and generative AI foundation models, or create custom models tailored for specific text, image classification, forecasting needs, or generative AI.

To create an Autopilot experiment programmatically for fine-tuning an LLM, you can call the CreateAutoMLJobV2 API in any language supported by Amazon SageMaker Autopilot or the AWS CLI.

For information about how this API action translates into a function in the language of your choice, see the See Also section of CreateAutoMLJobV2 and choose an SDK. As an example, for Python users, see the full request syntax of create_auto_ml_job_v2 in AWS SDK for Python (Boto3).

**Note**
Autopilot fine-tunes large language models without requiring multiple candidates to be trained and evaluated. Instead, using your dataset, Autopilot directly fine-tunes your target model to enhance a default objective metric, the cross-entropy loss. Fine-tuning language models in Autopilot does not require setting the AutoMLJobObjective field.

Once your LLM is fine-tuned, you can evaluate its performance by accessing various ROUGE scores through the BestCandidate when making a DescribeAutoMLJobV2 API call. The model also provides information about its training and validation loss as well as perplexity. For a comprehensive list of metrics for evaluating the quality of the text generated by the fine-tuned models, see Metrics for fine-tuning large language models in Autopilot.
Prerequisites

Before using Autopilot to create a fine-tuning experiment in SageMaker, make sure to take the following steps:

- (Optional) Choose the pre-trained model you want to fine-tune.

  For the list of pre-trained models available for fine-tuning in Amazon SageMaker Autopilot, see Supported large language models for fine-tuning. The selection of a model is not mandatory; if no model is specified, Autopilot automatically defaults to the model *Falcon7BInstruct*.

- Create a dataset of instructions. See Dataset file types and input data format to learn about the format requirements for your instruction-based dataset.

- Place your dataset in an Amazon S3 bucket.

- Grant full access to the Amazon S3 bucket containing your input data for the SageMaker execution role used to run your experiment.

  - For information on retrieving your SageMaker execution role, see Get execution role.

  - For information on granting your SageMaker execution role permissions to access one or more specific buckets in Amazon S3, see Add Additional Amazon S3 Permissions to a SageMaker Execution Role in Create execution role.

- Additionally, you should provide your execution role with the necessary permissions to access the default storage Amazon S3 bucket used by SageMaker JumpStart. This access is required for storing and retrieving pre-trained model artifacts in SageMaker JumpStart. To grant access to this Amazon S3 bucket, you must create a new inline custom policy on your execution role.

  Here’s an example policy that you can use in your JSON editor when configuring AutoML fine-tuning jobs in us-west-2:

  *SageMaker JumpStart’s bucket names follow a predetermined pattern that depends on the AWS Regions. You must adjust the name of the bucket accordingly.*

```json
{
    "Sid": "Statement1",
    "Effect": "Allow",
    "Action": [
        "s3:GetObject",
        "s3:PutObject",
        "s3:ListBucket"
    ],
}
```
Once this is done, you can use the ARN of this execution role in Autopilot API requests.

**Required parameters**

When calling `CreateAutoMLJobV2` to create an Autopilot experiment for LLM fine-tuning, you must provide the following values:

- An **AutoMLJobName** to specify the name of your job. The name should be of type string, and should have a minimum length of 1 character and a maximum length of 32.

- At least one **AutoMLJobChannel** of the training type within the **AutoMLJobInputDataConfig**. This channel specifies the name of the Amazon S3 bucket where your fine-tuning dataset is located. You have the option to define a validation channel. If no validation channel is provided, and a `ValidationFraction` is configured in the **AutoMLDataSplitConfig**, this fraction is utilized to randomly divide the training dataset into training and validation sets. Additionally, you can specify the type of content (CSV or Parquet files) for the dataset.

- An **AutoMLProblemTypeConfig** of type `TextGenerationJobConfig` to configure the settings of your training job.

  In particular, you can specify the name of the base model to fine-tune in the `BaseModelName` field. For the list of pre-trained models available for fine-tuning in Amazon SageMaker Autopilot, see [Supported large language models for fine-tuning](https://docs.aws.amazon.com/sagemaker/latest/dg/llm-fine-tuning-supported-models.html).

- An **OutputDataConfig** to specify the Amazon S3 output path to store the artifacts of your AutoML job.

- A **RoleArn** to specify the ARN of the role used to access your data.

The following is an example of the full request format used when making an API call to `CreateAutoMLJobV2` for fine-tuning a (Falcon7BInstruct) model.

```json
{
    "AutoMLJobName": "<job_name>",
```
"AutoMLJobInputDataConfig": [  
  {  
    "ChannelType": "training",
    "CompressionType": "None",
    "ContentType": "text/csv",
    "DataSource": {  
      "S3DataSource": {  
        "S3DataType": "S3Prefix",
        "S3Uri": "s3://<bucket_name>/<input_data>.csv"
      }
    }
  },  
  
  "OutputDataConfig": {  
    "S3OutputPath": "s3://<bucket_name>/output",
    "KmsKeyId": "arn:aws:kms:<region>:<account_id>:key/<key_value>"
  },  
  "RoleArn": "arn:aws:iam::<account_id>:role/<sagemaker_execution_role_name>",
  "AutoMLProblemTypeConfig": {  
    "TextGenerationJobConfig": {
      "BaseModelName": "Falcon7BInstruct"
    }
  }
}

All other parameters are optional.

Optional parameters

The following sections provide details of some optional parameters that you can pass to your fine-tuning AutoML job.

How to specify the training and validation datasets of an AutoML job

You can provide your own validation dataset and custom data split ratio, or let Autopilot split the dataset automatically.

Each AutoMLJobChannel object (see the required parameter AutoMLJobInputDataConfig) has a ChannelType, which can be set to either training or validation values that specify how the data is to be used when building a machine learning model.
At least one data source must be provided and a maximum of two data sources is allowed: one for training data and one for validation data. How you split the data into training and validation datasets depends on whether you have one or two data sources.

- If you only have **one data source**, the ChannelType is set to training by default and must have this value.
  - If the ValidationFraction value in `AutoMLDataSplitConfig` is not set, 0.2 (20%) of the data from this source is used for validation by default.
  - If the ValidationFraction is set to a value between 0 and 1, the dataset is split based on the value specified, where the value specifies the fraction of the dataset used for validation.
- If you have **two data sources**, the ChannelType of one of the `AutoMLJobChannel` objects must be set to training, the default value. The ChannelType of the other data source must be set to validation. The two data sources must have the same format, either CSV or Parquet, and the same schema. You must not set the value for the ValidationFraction in this case because all of the data from each source is used for either training or validation. Setting this value causes an error.

**How to enable automatic deployment**

With Autopilot, you can automatically deploy your fine-tuned model to an endpoint. To enable automatic deployment for your fine-tuned model, include a `ModelDeployConfig` in the AutoML job request. This allows the deployment of your fine-tuned model to a SageMaker endpoint. Below are the available configurations for customization.

- To let Autopilot generate the endpoint name, set `AutoGenerateEndpointName` to True.
- To provide your own name for the endpoint, set `AutoGenerateEndpointName` to False and provide a name of your choice in `EndpointName`.

**How to set the EULA acceptance when fine-tuning a model using the AutoML API**

For models requiring the acceptance of an end-user license agreement before fine-tuning, you can accept the EULA by setting the AcceptEula attribute of the `ModelAccessConfig` to True in `TextGenerationJobConfig` when configuring your `AutoMLProblemTypeConfig`.
**How to set hyperparameters to optimize the learning process of a model**

You can optimize the learning process of your text generation model by setting hyperparameter values in the `TextGenerationHyperParameters` attribute of `TextGenerationJobConfig` when configuring your `AutoMLProblemTypeConfig`.

Autopilot allows for the setting of four common hyperparameters across all models.

- `epochCount`: Its value should be a string containing an integer value within the range of 1 to 10.
- `batchSize`: Its value should be a string containing an integer value within the range of 1 to 64.
- `learningRate`: Its value should be a string containing a floating-point value within the range of 0 to 1.
- `learningRateWarmupSteps`: Its value should be a string containing an integer value within the range of 0 to 250.

For more details on each hyperparameter, see [Optimize the learning process of your text generation models with hyperparameters](#).

The following JSON example shows a `TextGenerationHyperParameters` field passed to the `TextGenerationJobConfig` where all four hyperparameters are configured.

```
"AutoMLProblemTypeConfig": {
  "TextGenerationJobConfig": {
    "BaseModelName": "Falcon7B",
    "TextGenerationHyperParameters": {
      "epochCount": "5",
      "learningRate": "0.000001",
      "batchSize": "32",
      "learningRateWarmupSteps": "10"
    }
  }
}
```

**Supported large language models for fine-tuning**

Using Autopilot API, users can fine-tune the following large language models (LLMs). Those models are powered by Amazon SageMaker JumpStart.

Note for fine-tuning models that require the acceptance of an end-user license agreement, you must explicitly declare EULA acceptance when creating your AutoML job. Note that after...
fine-tuning a pretrained model, the weights of the original model are changed, so you do not need to later accept a EULA when deploying the fine-tuned model. For information on how to accept the EULA when creating a fine-tuning job using the AutoML API, see the section called “Set EULA”.

You can find the full details of each model by searching for your SageMaker JumpStart Model ID in the following model table, and then following the link in the Source column. These details might include the languages supported by the model, biases it may exhibit, the datasets employed for fine-tuning, and more.

<table>
<thead>
<tr>
<th>SageMaker JumpStart Model ID</th>
<th>BaseModelName in API</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>huggingface-textgeneration-dolly-v2-3b-bf16</td>
<td>Dolly3B</td>
<td>Dolly 3B is a 2.8 billion parameter instruction-following large language model based on pythia-2.8b. It is trained on the instruction/response fine tuning dataset databricks-dolly-15k and can perform tasks including brainstorming, classification, questions and answers, text generation, information extraction, and summarization.</td>
</tr>
<tr>
<td>huggingface-textgeneration-dolly-v2-7b-bf16</td>
<td>Dolly7B</td>
<td>Dolly 7B is a 6.9 billion parameter instruction-following large language model based on pythia-6.9b. It is trained on the instruction/response fine tuning dataset databricks-dolly-15k and can perform tasks including brainstorming, classification,</td>
</tr>
<tr>
<td><strong>SageMaker JumpStart Model ID</strong></td>
<td><strong>BaseModelName in API request</strong></td>
<td><strong>Description</strong></td>
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<tr>
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<td>----------------</td>
</tr>
<tr>
<td>huggingface-textgeneration-dolly-v2-12b-bf16</td>
<td>Dolly12B</td>
<td>Dolly 12B is a 12 billion parameter instruction-following large language model based on <strong>pythia-12b</strong>. It is trained on the instruction/response fine tuning dataset <strong>databricks-dolly-15k</strong> and can perform tasks including brainstorming, classification, questions and answers, text generation, information extraction, and summarization.</td>
</tr>
<tr>
<td>huggingface-llm-falcon-7b-bf16</td>
<td>Falcon7B</td>
<td>Falcon 7B is a 7 billion parameter causal large language model trained on 1,500 billion tokens enhanced with curated corpora. Falcon-7B is trained on English and French data only, and does not generalize appropriately to other languages. Because the model was trained on large amounts of web data, it carries the stereotypes and biases commonly found online.</td>
</tr>
<tr>
<td>SageMaker JumpStart Model ID</td>
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</tr>
<tr>
<td>huggingface-llm-falcon-7b-instruct-bf16</td>
<td>Falcon7BInstruct</td>
<td>Falcon 7B Instruct is a 7 billion parameter causal large language model built on Falcon 7B and fine-tuned on a 250 million tokens mixture of chat/instruct datasets. Falcon 7B Instruct is mostly trained on English data, and does not generalize appropriately to other languages. Furthermore, as it is trained on a large-scale corpora representative of the web, it carries the stereotypes and biases commonly encountered online.</td>
</tr>
<tr>
<td>SageMaker JumpStart Model ID</td>
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</tr>
<tr>
<td>huggingface-llm-falcon-40b-bf16</td>
<td>Falcon40B</td>
<td>Falcon 40B is a 40 billion parameter causal large language model trained on 1,000 billion tokens enhanced with curated corpora. It is trained mostly on English, German, Spanish, and French, with limited capabilities in Italian, Portuguese, Polish, Dutch, Romanian, Czech, and Swedish. It does not generalize appropriately to other languages. Furthermore, as it is trained on a large-scale corpora representative of the web, it carries the stereotypes and biases commonly encountered online.</td>
</tr>
<tr>
<td>huggingface-llm-falcon-40b-instruct-bf16</td>
<td>Falcon40BInstruct</td>
<td>Falcon 40B Instruct is a 40 billion parameter causal large language model built on Falcon40B and fine-tuned on a mixture of Baize. It is mostly trained on English and French data, and does not generalize appropriately to other languages. Furthermore, as it is trained on a large-scale corpora representative of the web, it carries the stereotypes and biases commonly encountered online.</td>
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<td>-------------</td>
</tr>
<tr>
<td>huggingface-text2text-flan-t5-large</td>
<td>FlanT5L</td>
<td>The <strong>Flan-T5</strong> model family is a set of large language models that are fine-tuned on multiple tasks and can be further trained. These models are well-suited for tasks such as language translation, text generation, sentence completion, word sense disambiguation, summarization, or question answering. Flan T5 L is a 780 million parameter large language model trained on numerous languages. You can find the list of the languages supported by Flan T5 L in the details of the model retrieved from your search by model ID in SageMaker JumpStart's <a href="#">model table</a>.</td>
</tr>
<tr>
<td><strong>SageMaker JumpStart Model ID</strong></td>
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</tr>
<tr>
<td>huggingface-text2text-flan-t5-xl</td>
<td>FlanT5XL</td>
<td>The Flan-T5 model family is a set of large language models that are fine-tuned on multiple tasks and can be further trained. These models are well-suited for tasks such as language translation, text generation, sentence completion, word sense disambiguation, summarization, or question answering. Flan T5 XL is a 3 billion parameter large language model trained on numerous languages. You can find the list of the languages supported by Flan T5 XL in the details of the model retrieved from your search by model ID in SageMaker JumpStart's model table.</td>
</tr>
<tr>
<td>SageMaker JumpStart Model ID</td>
<td>BaseModelName in API request</td>
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<tr>
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<td>-------------</td>
</tr>
<tr>
<td>huggingface-text2text-flan-t5-xxll</td>
<td>FlanT5XXL</td>
<td>The Flan-T5 model family is a set of large language models that are fine-tuned on multiple tasks and can be further trained. These models are well-suited for tasks such as language translation, text generation, sentence completion, word sense disambiguation, summarization, or question answering. Flan T5 XXL is a 11 billion parameter model. You can find the list of the languages supported by Flan T5 XXL in the details of the model retrieved from your search by model ID in SageMaker JumpStart's model table.</td>
</tr>
<tr>
<td>meta-textgeneration-llama-2-7b</td>
<td>Llama2-7B</td>
<td>Llama 2 is a collection of pretrained and fine-tuned generative text models, ranging in scale from 7 billion to 70 billion parameters. Llama2-7B is the 7 billion parameter model that is intended for English use and can be adapted for a variety of natural language generation tasks.</td>
</tr>
<tr>
<td>SageMaker JumpStart Model ID</td>
<td>BaseModelName in API request</td>
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</tr>
<tr>
<td>meta-textgeneration-llama-2-7b-f</td>
<td>Llama2-7BChat</td>
<td>Llama 2 is a collection of pretrained and fine-tuned generative text models, ranging in scale from 7 billion to 70 billion parameters. Llama2-7B is the 7 billion parameter chat model that is optimized for dialogue use cases.</td>
</tr>
<tr>
<td>meta-textgeneration-llama-2-13b</td>
<td>Llama2-13B</td>
<td>Llama 2 is a collection of pretrained and fine-tuned generative text models, ranging in scale from 7 billion to 70 billion parameters. Llama2-13B is the 13 billion parameter model that is intended for English use and can be adapted for a variety of natural language generation tasks.</td>
</tr>
<tr>
<td>meta-textgeneration-llama-2-13b-f</td>
<td>Llama2-13BChat</td>
<td>Llama 2 is a collection of pretrained and fine-tuned generative text models, ranging in scale from 7 billion to 70 billion parameters. Llama2-13B is the 13 billion parameter chat model that is optimized for dialogue use cases.</td>
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<td>-----------------------------</td>
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<td>-------------</td>
</tr>
<tr>
<td>huggingface-llm-mistral-7b</td>
<td>Mistral7B</td>
<td>Mistral 7B is a seven billion parameters code and general purpose English text generation model. It can be used in a variety of use cases including text summarization, classification, text completion, or code completion.</td>
</tr>
<tr>
<td>huggingface-llm-mistral-7b-instruct</td>
<td>Mistral7BInstruct</td>
<td>Mistral 7B Instruct is the fine-tuned version of Mistral 7B for conversational use cases. It was specialized using a variety of publicly available conversation datasets in English.</td>
</tr>
<tr>
<td>huggingface-textgeneration1-mpt-7b-bf16</td>
<td>MPT7B</td>
<td>MPT 7B is a decoder-style transformer large language model with 6.7 billion parameters, pre-trained from scratch on 1 trillion tokens of English text and code. It is prepared to handle long context lengths.</td>
</tr>
<tr>
<td>huggingface-textgeneration1-mpt-7b-instruct-bf16</td>
<td>MPT7BInstruct</td>
<td>MPT 7B Instruct is a model for short-form instruction following tasks. It is built by fine-tuning MPT 7B on a dataset derived from databricks-dolly-15k and the Anthropic Helpful and Harmless (HH-RLHF) datasets.</td>
</tr>
</tbody>
</table>
Dataset file types and input data format

Instruction-based fine-tuning uses labeled datasets to improve the performance of pre-trained LLMs on specific natural language processing (NLP) tasks. The labeled examples are formatted as prompt-response pairs and phrased as instructions.

To learn about the supported dataset file types, see Supported dataset file types.

To learn about input data format, see Input data format for instruction-based fine-tuning.

Supported dataset file types

Autopilot supports instruction-based fine-tuning datasets formatted as CSV files (default) or as Parquet files.

• CSV (comma separated values) is a row-based file format that stores data in human readable plaintext, which is a popular choice for data exchange as it is supported by a wide range of applications.

• Parquet is a binary, column-based file format where the data is stored and processed more efficiently than in human readable file formats such as CSV. This makes it a better option for big data problems.

Note

The dataset may consist of multiple files, each of which must adhere to a specific template. For information on how to format your input data, see Input data format for instruction-based fine-tuning.

Input data format for instruction-based fine-tuning

Each file in the dataset must adhere to the following format:

• The dataset must contain exactly two comma-separated and named columns, `input` and `output`. Autopilot does not allow any additional columns.

• The `input` columns contain the prompts, and their corresponding `output` contains the expected answer. Both the `input` and `output` are in string format.
The following example illustrates the input data format for instruction-based fine-tuning in Autopilot.

```
input,output
"<prompt text>","<expected generated text>"
```

**Note**
We recommend using datasets with a minimum of 1000 rows to ensure optimal learning and performance of the model.

Additionally, Autopilot sets a maximum limit on the number of rows in the dataset and the context length based on the type of model being used.

- The limits on the number of rows in a dataset apply to the cumulative count of rows across all files within the dataset, including multiple files. If there are two channel types defined (one for training and one for validation), the limit applies to the total number of rows across all datasets within both channels. When the number of rows exceeds the threshold, the job fails with a validation error.
- When the length of the input or output of a row in the dataset exceeds the limit set on the context of the language model, it is automatically truncated. If more than 60% of the rows in the dataset are truncated, whether in their input or output, Autopilot fails the job with a validation error.

The following table presents those limits for each model.

<table>
<thead>
<tr>
<th>SageMaker JumpStart Model ID</th>
<th>BaseModelName in API request</th>
<th>Row Limit</th>
<th>Context Length Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>huggingface-textgeneration-dolly-v2-3b-bf16</td>
<td>Dolly3B</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-textgeneration-dolly-v2-7b-bf16</td>
<td>Dolly7B</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>JumpStart Model ID</td>
<td>BaseModelName in API request</td>
<td>Row Limit</td>
<td>Context Length Limit</td>
</tr>
<tr>
<td>--------------------</td>
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</tr>
<tr>
<td>huggingface-textgeneration-dolly-v2-12b-bf16</td>
<td>Dolly12B</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-llm-falcon-7b-bf16</td>
<td>Falcon7B</td>
<td>1,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-llm-falcon-7b-instruct-bf16</td>
<td>Falcon7BI</td>
<td>1,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-llm-falcon-40b-bf16</td>
<td>Falcon40B</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-llm-falcon-40b-instruct-bf16</td>
<td>Falcon40B</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-text2text-flan-t5-large</td>
<td>FlanT5L</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-text2text-flan-t5-xl</td>
<td>FlanT5XL</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-text2text-flan-t5-xxl</td>
<td>FlanT5XXL</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>meta-textgeneration-llama-2-7b</td>
<td>Llama2-7B</td>
<td>10,000 rows</td>
<td>2048 tokens</td>
</tr>
<tr>
<td>meta-textgeneration-llama-2-7b-f</td>
<td>Llama2-7BChat</td>
<td>10,000 rows</td>
<td>2048 tokens</td>
</tr>
<tr>
<td>meta-textgeneration-llama-2-13b</td>
<td>Llama2-13B</td>
<td>7,000 rows</td>
<td>2048 tokens</td>
</tr>
</tbody>
</table>
Optimize the learning process of your text generation models with hyperparameters

You can optimize the learning process of your base model by adjusting any combination of the following hyperparameters. These parameters are available for all models.

- **Epoch Count**: The epochCount hyperparameter determines how many times the model goes through the entire training dataset. It influences the training duration and can prevent overfitting when set appropriately. Large number of epochs may increase the overall runtime of fine-tuning jobs. We recommend setting a large MaxAutoMLJobRuntimeInSeconds within the CompletionCriteria of the `TextGenerationJobConfig` to avoid fine-tuning jobs from stopping prematurely.

- **Batch Size**: The batchSize hyperparameter defines the number of data samples used in each iteration of training. It can affect the convergence speed and memory usage. With large batch size, the risk of out of memory (OOM) errors increases, which may surface as an internal server error in Autopilot. To check for such error, check the `/aws/sagemaker/TrainingJobs` log group for the training jobs launched by your Autopilot job. You can access those logs in

<table>
<thead>
<tr>
<th>SageMaker JumpStart Model ID</th>
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<td>Llama2-13BChat</td>
<td>7,000 rows</td>
<td>2048 tokens</td>
</tr>
<tr>
<td>huggingface-llm-mistral-7b</td>
<td>Mistral7B</td>
<td>10,000 rows</td>
<td>2048 tokens</td>
</tr>
<tr>
<td>huggingface-llm-mistral-7b-instruct</td>
<td>Mistral7B Instruct</td>
<td>10,000 rows</td>
<td>2048 tokens</td>
</tr>
<tr>
<td>huggingface-textgeneration1-mpt-7b-bf16</td>
<td>MPT7B</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
<tr>
<td>huggingface-textgeneration1-mpt-7b-instruct-bf16</td>
<td>MPT7BInstruct</td>
<td>10,000 rows</td>
<td>1024 tokens</td>
</tr>
</tbody>
</table>
CloudWatch from the AWS management console. Choose Logs, and then choose the /aws/sagemaker/TrainingJobs log group. To remedy OOM errors, reduce the batch size.

We recommend starting with a batch size of 1, then incrementally increase it until an out of memory error occurs. As a reference, 10 epochs typically takes up to 72h to complete.

- **Learning Rate**: The learningRate hyperparameter controls the step size at which a model's parameters are updated during training. It determines how quickly or slowly the model's parameters are updated during training. A high learning rate means that the parameters are updated by a large step size, which can lead to faster convergence but may also cause the optimization process to overshoot the optimal solution and become unstable. A low learning rate means that the parameters are updated by a small step size, which can lead to more stable convergence but at the cost of slower learning.

- **Learning Rate Warmup Steps**: The learningRateWarmupSteps hyperparameter specifies the number of training steps during which the learning rate gradually increases before reaching its target or maximum value. This helps the model converge more effectively and avoid issues like divergence or slow convergence that can occur with an initially high learning rate.

To learn about how to adjust hyperparameters for your fine-tuning experiment in Autopilot and discover their possible values, see [How to set hyperparameters to optimize the learning process of a model](#).

**Metrics for fine-tuning large language models in Autopilot**

Using your dataset, Autopilot directly fine-tunes a target language model (LLM) to enhance a default objective metric, the cross-entropy loss.

Cross-entropy loss is a widely used metric to assess the dissimilarity between the predicted probability distribution and the actual distribution of words in the training data. By minimizing cross-entropy loss, the model learns to make more accurate and contextually relevant predictions, particularly in tasks related to text generation.

After fine-tuning an LLM you can evaluate the quality of its generated text using a range of ROUGE scores. Additionally, you can analyze the perplexity and cross-entropy training and validation losses as part of the evaluation process.

- Perplexity loss measures how well the model can predict the next word in a sequence of text, with lower values indicating a better understanding of the language and context.
Recall-Oriented Understudy for Gisting Evaluation (ROUGE) is a set of metrics used in the field of natural language processing (NLP) and machine learning to evaluate the quality of machine-generated text, such as text summarization or text generation. It primarily assesses the similarities between the generated text and the ground truth reference (human-written) text of a validation dataset. ROUGE measures are designed to assess various aspects of text similarity, including the precision and recall of n-grams (contiguous sequences of words) in the system-generated and reference texts. The goal is to assess how well a model captures the information present in the reference text.

There are several variants of ROUGE metrics, depending on the type of n-grams used and the specific aspects of text quality being evaluated.

The following list contains the name and description of the ROUGE metrics available after the fine-tuning of large language models in Autopilot.

**ROUGE-1, ROUGE-2**

ROUGE-N, the primary ROUGE metric, measures the overlap of n-grams between the system-generated and reference texts. ROUGE-N can be adjusted to different values of n (here 1 or 2) to evaluate how well the system-generated text captures the n-grams from the reference text.

**ROUGE-L**

ROUGE-L (ROUGE-Longest Common Subsequence) calculates the longest common subsequence between the system-generated text and the reference text. This variant considers word order in addition to content overlap.

**ROUGE-L-Sum**

ROUGE-L-SUM (Longest Common Subsequence for Summarization) is designed for the evaluation of text summarization systems. It focuses on measuring the longest common subsequence between the machine-generated summary and the reference summary. ROUGE-L-SUM takes into account the order of words in the text, which is important in text summarization tasks.

**Autopilot model deployment and predictions**

After fine-tuning a large language model (LLM), you can deploy the model for real-time text generation by setting up an endpoint to obtain interactive predictions.
Amazon SageMaker Developer Guide

**Note**

We recommend running real-time inference jobs on `ml.g5.12xlarge` for better performances. Alternatively, `ml.g5.8xlarge` instances are suitable for Falcon-7B-Instruct and MPT-7B-Instruct text generation tasks.

You can find the specifics of these instances within the [Accelerated Computing](https://aws.amazon.com/ec2/instance-types/) category in the selection of instance types provided by Amazon EC2.

---

### Real-time text generation

You can use SageMaker APIs to manually deploy your fine-tuned model to a SageMaker Hosting [real-time inference endpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/hosting-endpoints.html), then begin making predictions by invoking the endpoint as follows.

**Note**

Alternatively, you can chose the automatic deployment option when creating your fine-tuning experiment in Autopilot. For information on setting up the automatic deployment of models, see [How to enable automatic deployment](https://docs.aws.amazon.com/sagemaker/latest/dg/enable-automatic-deployment.html).

You can also use the SageMaker Python SDK and the JumpStartModel class to perform inferences with models fine-tuned by Autopilot. This can be done by specifying a custom location for the model's artifact in Amazon S3. For information on defining your model as a SageMaker JumpStart model and deploying your model for inference, see [Low-code deployment with the JumpStartModel class](https://docs.aws.amazon.com/sagemaker/latest/dg/jumpstart-deployment.html).

---

1. **Obtain the candidate inference container definitions**

   You can find the `InferenceContainerDefinitions` within the `BestCandidate` object retrieved from the response to the [DescribeAutoMLJobV2](https://docs.aws.amazon.com/sagemaker/latest/dg/api-model-describe-auto-ml.html) API call. A container definition for inference refers to the containerized environment designed for deploying and running your trained model to make predictions.

   The following AWS CLI command example uses the [DescribeAutoMLJobV2](https://docs.aws.amazon.com/cli/latest/reference/sagemaker/describe-auto-ml-job-v2.html) API to obtain recommended container definitions for your job name.

   ```bash
   aws sagemaker describe-auto-ml-job-v2 --auto-ml-job-name job-name --region region
   ```

2. **Create a SageMaker model**

---

Create an LLM fine-tuning job using the AutoML API
Use the container definitions from the previous step to create a SageMaker model by using the CreateModel API. See the following AWS CLI command as an example. Use the CandidateName for your model name.

```bash
aws sagemaker create-model --model-name '"<your-candidate-name>"' \
--primary-container '"<container-definition>"' \
--execution-role-arn '"<execution-role-arn>"' --region '"<region>"
```

3. Create an endpoint configuration

The following AWS CLI command example uses the CreateEndpointConfig API to create an endpoint configuration.

```
aws sagemaker create-endpoint-config --endpoint-config-name '"<your-endpoint-config-name>"' \
--production-variants '"<list-of-production-variants>"' ModelDataDownloadTimeoutInSeconds=3600 ContainerStartupHealthCheckTimeoutInSeconds=3600 \
--region '"<region>"
```

**Note**

To prevent the endpoint creation from timing out due to a lengthy model download, we recommend setting ModelDataDownloadTimeoutInSeconds = 3600 and ContainerStartupHealthCheckTimeoutInSeconds = 3600.

```
aws sagemaker create-endpoint-config --endpoint-config-name '"<your-endpoint-config-name>"' \
--primary-container '"<container-definition>"' \
--execution-role-arn '"<execution-role-arn>"' --region '"<region>"
```

4. Create the endpoint

The following AWS CLI example uses the CreateEndpoint API to create the endpoint.

```
aws sagemaker create-endpoint --endpoint-name '"<your-endpoint-name>"' \
--endpoint-config-name '"<endpoint-config-name-you-just-created>"' \
--region '"<region>"
```

Check the progress of your endpoint deployment by using the DescribeEndpoint API. See the following AWS CLI command as an example.
After the EndpointStatus changes to InService, the endpoint is ready to use for real-time inference.

5. Invoke the endpoint

The following command invokes the endpoint for real-time inferencing. Your prompt needs to be encoded in bytes.

```bash
aws sagemaker invoke-endpoint --endpoint-name '<endpoint-name>' \
   --region '<region>' --body '<your-prompt-in-bytes>' [--content-type] 'application/json' <outfile>
```

**Note**

The format of your input prompt depends on the language model. For more information on the format of text generation prompts, see [Request format for text generation models real-time inference](#).

**Request format for text generation models real-time inference**

Different large language models (LLMs) may have specific software dependencies, runtime environments, and hardware requirements influencing Autopilot's recommended container to host the model for inference. Additionally, each model dictates the required input data format and the expected format for predictions and outputs.

Here are example inputs for some models and recommended containers.

- For Falcon models with the recommended container `huggingface-pytorch-tgi-inference:2.0.1-tgi1.0.3-gpu-py39-cu118-ubuntu20.04`:

  ```json
  payload = {
      "inputs": "Large language model fine-tuning is defined as",
      "parameters": {
        "do_sample": false,
        "top_p": 0.9,
  ```
Create a Regression or Classification Autopilot experiment for tabular data using the Studio Classic UI

Note

All UI-related instructions in this guide pertain to Autopilot's standalone features before migrating to Amazon SageMaker Canvas. Users following these instructions should use Studio Classic.

You can use the Amazon SageMaker Studio Classic UI to create Autopilot experiments for classification or regression problems on tabular data. The UI helps you specify the name of your experiment, provide locations for the input and output data, and specify which target data to predict. Optionally, you can also specify the type of problem that you want to solve (regression, classification, multiclass classification), choose your modeling strategy (stacked ensembles or hyperparameters optimization), select the list of algorithms used by the Autopilot job to train the data, and more.

The UI has descriptions, toggle switches, dropdown menus, radio buttons, and more to help you navigate creating your model candidates. After the experiment runs, you can compare trials and delve into the details of the pre-processing steps, algorithms, and hyperparameter ranges of each model. Optionally, you can download their explainability and performance reports. Use the provided notebooks to see the results of the automated data exploration or the candidate model definitions.
Alternatively, you can use Autopilot AutoML API in Create a regression or classification job for tabular data using the AutoML API.

**Configure the default parameters of an Autopilot experiment (for administrators)**

Autopilot supports setting default values to simplify the configuration of Amazon SageMaker Autopilot when you create an Autopilot experiment using the Studio Classic UI. Administrators can use Studio Classic lifecycle configurations (LCC) to set infrastructure, networking, and security values in configuration files and pre-populate the advanced settings of AutoML jobs.

By doing so, they can fully control network connectivity and access permissions for the resources associated with Amazon SageMaker Studio Classic, including SageMaker instances, data sources, output data, and other related services. Specifically, administrators can configure a desired network architecture, such as Amazon VPC, subnets, and security groups, for a Studio Classic domain or individual user profiles. Data scientists can focus on data science specific parameters when creating their Autopilot experiments using the Studio Classic UI. Furthermore, administrators can manage the encryption of data on the instance in which Autopilot experiments run by setting default encryption keys.

---

**Note**

This feature is currently not available in the Asia Pacific (Hong Kong) and Middle East (Bahrain) opt-in Regions.

---

In the following sections, you can find the full list of parameters supporting the setting of defaults when creating an Autopilot experiment using the Studio Classic UI, and learn how to set those default values.

**Topics**

- List of default parameters supported
- Set default Autopilot experiment parameters

**List of default parameters supported**

The following parameters support setting default values with a configuration file for creating an Autopilot experiment using the Studio Classic UI. Once set, the values automatically fill in their
corresponding field in the Autopilot' **Create Experiment** tab in the Studio Classic UI. See [Advanced settings (optional)](https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_CreateAutoMLJob.html) for a full description of each field.

- **Security:** Amazon VPC, subnets, and security groups.
- **Access:** AWS IAM role ARNs.
- **Encryption:** AWS KMS key IDs.
- **Tags:** Key-value pairs used to label and organize SageMaker resources.

**Set default Autopilot experiment parameters**

Administrators can set default values in a configuration file, then manually place the file in a recommended location within the Studio Classic environment of specific users, or they can pass the file to a lifecycle configuration script (LCC) to automate the customization of the Studio Classic environment for a given domain or user profile.

- To set up the configuration file, start by filling in its default parameters.

To configure any or all default values listed in [List of default parameters supported](https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_CreateAutoMLJob.html), administrators can create a configuration file named `config.yaml`, the structure of which should adhere to this [sample configuration file](https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_CreateAutoMLJob.html). The following snippet shows a sample configuration file with all the supported AutoML parameters. For more information on the format of this file, refer to the [full schema](https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_CreateAutoMLJob.html).

```yaml
SchemaVersion: '1.0'
SageMaker:
  AutoMLJob:
    # https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_CreateAutoMLJob.html
    AutoMLJobConfig:
      SecurityConfig:
        EnableInterContainerTrafficEncryption: true
        VolumeKmsKeyId: 'kms-key-id'
      VpcConfig:
        SecurityGroupIds:
          - 'security-group-id-1'
          - 'security-group-id-2'
        Subnets:
          - 'subnet-1'
          - 'subnet-2'
    OutputDataConfig:
```

Create a Regression or Classification Job Using the Studio Classic UI
• Then, place the configuration file in the recommended location by either manually copying the file to its recommended paths or using a lifecycle configuration (LCC).

The configuration file needs to be present in at least one of the following locations in the user's Studio Classic environment. By default, SageMaker searches for a configuration file in two locations:

• First, in /etc/xdg/sagemaker/config.yaml. We refer to this file as the administrator configuration file.

• Then, in /root/.config/sagemaker/config.yaml. We refer to this file as the user configuration file.

Using the administrator configuration file, administrators can define a set of default values. Optionally, they can use the user configuration file to override values set in the administrator configuration file, or set additional default parameter values.

The following snippet shows a sample script which writes the default parameters configuration file to the administrator location in the user's Studio Classic environment. You can replace /etc/xdg/sagemaker with /root/.config/sagemaker to write the file to the user location.

```bash
#!/bin/bash
sudo mkdir -p /etc/xdg/sagemaker

echo "SchemaVersion: '1.0'
CustomParameters: 
  AnyStringKey: 'AnyStringValue'
SageMaker: 
  AutoMLJob: 
    # https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_CreateAutoMLJob.html
    AutoMLJobConfig: 
      SecurityConfig: 
        EnableInterContainerTrafficEncryption: true
      VolumeKmsKeyId: 'kms-key-id'
```

Create a Regression or Classification Job Using the Studio Classic UI

249
VpcConfig:
    SecurityGroupIds:
        - 'security-group-id-1'
        - 'security-group-id-2'
    Subnets:
        - 'subnet-1'
        - 'subnet-2'
OutputDataConfig:
    KmsKeyId: 'kms-key-id'
    RoleArn: 'arn:aws:iam::111222333444:role/Admin'
    Tags:
        - Key: 'tag_key'
          Value: 'tag_value'
" | sudo tee /etc/xdg/sagemaker/config.yaml

- **Copy the files manually** – To copy the configuration files manually, run the `script` created in the previous step from a Studio Classic terminal. In this case, the user profile that executed the script can create Autopilot experiments with the default values applicable only to them.

- **Create a SageMaker lifecycle configuration** – Alternatively, you can use a lifecycle configuration (LCC) to automate the customization of your Studio Classic environment. LCC are shell scripts triggered by Amazon SageMaker Studio Classic lifecycle events such as starting a Studio Classic application. This customization includes installing custom packages, configuring notebook extensions, pre-loading datasets, setting up source code repositories, or, in our case, pre-populating default parameters. Administrators can attach the LCC to a Studio Classic domain to automate the configuration of default values for each user profile within that domain.

The following sections detail how to create a lifecycle configuration so users can load Autopilot default parameters automatically when launching Studio Classic. You can choose to create an LCC using the SageMaker Console or the AWS CLI.

**Create a LCC from the SageMaker Console**

Use the following steps to create an LCC containing your default parameters, attach the LCC to a domain or a user profile, then launch a Studio Classic application pre-populated with the default parameters set by the LCC using the SageMaker Console.

- **To create a lifecycle configuration that runs the script containing your default values using the SageMaker Console**
On the left side, navigate to Admin configurations, then Lifecycle configurations.

From the Lifecycle configurations page, navigate to the Studio Classic tab, then choose Create configuration.

For Name, type a name using alphanumeric characters and ".-", but no spaces. The name can have a maximum of 63 characters.

Paste your script in the Scripts section.

Choose Create configuration to create the lifecycle configuration. This creates an LCC of type Kernel gateway app.

To attach the lifecycle configuration to a Studio Classic domain, a space, or a user profile

Follow the steps in Attach the lifecycle configuration to Studio Classic domain or user profile to attach your LCC to a Studio Classic domain or a specific user profile.

To launch your Studio Classic application with the lifecycle configuration

Once the LCC is attached to a domain or a user profile, impacted users can start a Studio Classic application from the landing page of Studio Classic in Studio to pick up the defaults set by the LCC automatically. This auto-populates the Studio Classic UI when creating an Autopilot experiment.

Create a LCC from the AWS CLI

Use the following snippets to launch a Studio Classic application that runs your script using the AWS CLI. Note that lifecycle_config.sh is the name given to your script in this example.

Before getting started:

- Ensure that you have updated and configured AWS CLI by completing the prerequisites described in Create a lifecycle configuration from the AWS CLI.

- Install OpenSSL documentation. The AWS CLI command uses the open-source library OpenSSL to encode your script in Base64 format. This requirement prevents errors that occur from spacing and line break encoding.

You can now follow these three steps:

- Create a new lifecycle configuration referencing the configuration script lifecycle_config.sh
LCC_CONTENT=`openssl base64 -A -in lifecycle_config.sh`

## Create a new lifecycle config

```bash
aws sagemaker create-studio-lifecycle-config --region region \
--studio-lifecycle-config-name lcc-name \
--studio-lifecycle-config-content $LCC_CONTENT \
--studio-lifecycle-config-app-type default
```

Note the ARN of the newly created lifecycle configuration that is returned. This ARN is required to attach the lifecycle configuration to your application.

- **Attach the lifecycle configuration to your JupyterServerApp**

The following example shows how to create a new user profile with a lifecycle configuration attached. To update an existing user profile, use the AWS CLI `update-user-profile` command. To create or update a domain, see `create-domain` and `update-domain`. Add the lifecycle configuration ARN from the previous step to the settings of the JupyterServerAppSettings application type. You can add multiple lifecycle configurations at the same time by using a list of lifecycle configurations.

```bash
# Create a new UserProfile
aws sagemaker create-user-profile --domain-id domain-id \
--user-profile-name user-profile-name \
--region region \
--user-settings '{
"JupyterServerAppSettings": {
  "LifecycleConfigArns": [
"lifecycle-configuration-arn"
  ]
}
}
```

Once the LCC is attached to a domain or a user profile, impacted users can shut down and update their existing Studio Classic application by following the steps in [Shut down and Update Amazon SageMaker Studio Classic](#), or start a new Studio Classic application from the AWS Console to pick up the defaults set by the LCC automatically. This autopoulates the Studio Classic UI when creating an Autopilot experiment. Alternatively, they can launch a new Studio Classic application using the AWS CLI as follows.

- **Launch your Studio Classic application with the lifecycle configuration using the AWS CLI**
# Create a Jupyter Server application

```
# Create a Jupyter Server application
aws sagemaker create-app --domain-id domain-id \ 
--user-profile-name user-profile-name \ 
--region region \ 
--app-type JupyterServer \ 
--resource-spec LifecycleConfigArn=lifecycle-configuration-arn \ 
--app-name default
```

For more information on creating a lifecycle configuration using the AWS CLI, see [Create a Lifecycle Configuration from the AWS CLI](#).

---

**To create an Autopilot experiment using Studio Classic UI**

1. Sign in at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/), choose **Studio** from the left navigation pane, select your Domain and user profile, then **Open Studio**.

2. In Studio, choose the Studio Classic icon in the top left navigation pane. This opens a Studio Classic app.

3. Run or open a Studio Classic application from the space of your choice, or **Create Studio Classic space**. On the **Home** tab, choose the **AutoML** card. This opens a new **AutoML** tab.

4. Choose **Create an AutoML experiment**. This opens a new **Create experiment** tab.

5. In the **Experiment and data details** section, enter the following information:

   a. **Experiment name** – Must be unique to your account in the current AWS Region and contain a maximum of 63 alphanumeric characters. Can include hyphens (-) but not spaces.

   b. **Input data** – Provide the Amazon Simple Storage Service (Amazon S3) bucket location of your input data. This S3 bucket must be in your current AWS Region. The URL must be in an `s3://` format where Amazon SageMaker has write permissions. The file must be in CSV or Parquet format and contain at least 500 rows. Select **Browse** to scroll through available paths and **Preview** to see a sample of your input data.

   c. **Is your S3 input a manifest file?** – A manifest file includes metadata with your input data. The metadata specifies the location of your data in Amazon S3. It also specifies how the data is formatted and which attributes from the dataset to use when training your model. You can use a manifest file as an alternative to preprocessing when your labeled data is being streamed in Pipe mode.
d. **Auto split data?** – Autopilot can split your data into an 80-20% split for training and validation data. If you prefer a custom split, you can choose the **Specify split ratio**. To use a custom dataset for validation, choose **Provide a validation set**.

e. **Output data location (S3 bucket)** – The name of the S3 bucket location where you want to store the output data. The URL for this bucket must be in an Amazon S3 format where Amazon SageMaker has write permissions. The S3 bucket must be in the current AWS Region. Autopilot can also create this for you in the same location as your input data.

6. Choose **Next: Target and features**. The **Target and features** tab opens.

7. In the **Target and features** section:
   - Select a column to set as a target for model predictions.
   - Optionally, you can pass the name of a sample weights column in the **Sample weight** section to request your dataset rows to be weighted during training and evaluation. For more information on the available objective metrics, see **Autopilot weighted metrics**.

   ![](image)
   **Note**
   Support for sample weights is available in **ensembling mode** only.

   • You can also select features for training and change their data type. The following data types are available: Text, Numerical, Categorical, Datetime, Sequence, and Auto. All features are selected by default.


9. In the **Training method** section, select your training option: **Ensembling**, **Hyperparameter optimization (HPO)**, or **Auto** to let Autopilot choose the training method automatically based on the dataset size. Each training mode runs a pre-defined set of algorithms on your dataset to train model candidates. By default, Autopilot pre-selects all the available algorithms for the given training mode. You can run an Autopilot training experiment with all the algorithms or choose your own subset.

   For more information on the training modes and the available algorithms, see the **Autopilot training modes** section in the **Training modes and algorithms** page.

10. Choose **Next: Deployment and advanced settings** to open the **Deployment and advanced settings** tab. Settings include the auto-display endpoint name, machine learning problem type, and additional choices for running your experiment.
a. **Deployment settings** – Autopilot can automatically create an endpoint and deploy your model for you.

To auto-deploy to an automatically generated endpoint, or to provide an endpoint name for custom deployment, set the toggle to **Yes** under **Auto deploy?** If you are importing data from Amazon SageMaker Data Wrangler, you have additional options to auto-deploy the best model with or without the transforms from Data Wrangler.

**Note**

If your Data Wrangler flow contains multi-row operations such as groupby, join, or concatenate, you can't auto-deploy with these transforms. For more information, see [Automatically Train Models on Your Data Flow](#).

b. **Advanced settings (optional)** – Autopilot provides additional controls to manually set experimental parameters such as defining your problem type, time constraints on your Autopilot job and trials, security, and encryption settings.

**Note**

Autopilot supports the setting of default values to simplify the configuration of Autopilot experiments using Studio Classic UI. Administrators can use Studio Classic lifecycle configurations (LCC) to set infrastructure, networking, and security values in configuration files and pre-populate the **advanced settings** of AutoML jobs.

To learn about how administrators can automate the customization of an Autopilot experiment, see [Configure the default parameters of an Autopilot experiment (for administrators)](#).

i. **Machine learning problem type** – Autopilot can automatically infer the type of supervised learning problem from your dataset. If you prefer to choose it manually, you can use the **Select the machine learning problem type** dropdown menu. Note that it defaults to **Auto**. In some cases, SageMaker is unable to infer accurately. When that happens, you must provide the value for the job to succeed. In particular, you can choose from the following types:
• **Binary classification** – Binary classification assigns input data to one of two predefined and mutually exclusive classes, based on their attributes, such as medical diagnosis based on results of diagnostic tests that determine if someone has a disease.

• **Regression** – Regression establishes a relationship between the input variables (also known as independent variables or features) and the target variable (also known as the dependent variable). This relationship is captured through a mathematical function or model that maps the input variables to a continuous output. It is commonly used for tasks such as predicting house prices based on features like square footage and the number of bathrooms, stock market trends, or estimating sales figures.

• **Multiclass classification** – Multiclass classification assigns input data to one of several classes based on their attributes, like the prediction of the topic most relevant to a text document, such as politics, finance, or philosophy.

  ii. **Runtime** – You can define a maximum time limit. Upon reaching the time limit, trials and jobs that exceed the time constraint automatically stop.

  iii. **Access** – You can choose the role that Amazon SageMaker Studio Classic assumes to gain temporary access to AWS services (in particular, SageMaker and Amazon S3) on your behalf. If no role is explicitly defined, Studio Classic automatically uses the default SageMaker execution role attached to your user profile.

  iv. **Encryption** – To enhance the security of your data at rest and protect it against unauthorized access, you can specify encryption keys to encrypt data in your Amazon S3 buckets and in the Amazon Elastic Block Store (Amazon EBS) volume attached to your Studio Classic domain.

  v. **Security** – You can choose the virtual private cloud (Amazon VPC) in which your SageMaker job runs. Ensure that the Amazon VPC has access to your input and output Amazon S3 buckets.

  vi. **Project** – Specify the name of the SageMaker project to associate with this Autopilot experiment and model outputs. When you specify a project, Autopilot tags the project to an experiment. This lets you know which model outputs are associated with this project.

  vii. **Tags** – Tags are an array of key-value pairs. Use tags to categorize your resources from AWS services, such as their purpose, owner, or environment.
c. Choose **Next: Review and create** to get a summary of your Autopilot experiment before you create it.

11. Select **Create experiment**. The creation of the experiment starts an Autopilot job in SageMaker. Autopilot provides the status of the experiment, information on the data exploration process and model candidates in notebooks, a list of generated models and their reports, and the job profile used to create them.

For information on the notebooks generated by an Autopilot job, see [Amazon SageMaker Autopilot notebooks generated to manage AutoML tasks](https://docs.aws.amazon.com/autopilot/latest/dg/autopilot-generated-notebooks.html). For information on the details of each model candidate and their reports, see [Models generated by Amazon SageMaker Autopilot](https://docs.aws.amazon.com/autopilot/latest/dg/autopilot-generated-models.html).

### Note

To avoid incurring unnecessary charges: If you deploy a model that is no longer needed, delete the endpoints and resources that were created during that deployment. Information about pricing instances by Region is available at [Amazon SageMaker Pricing](https://aws.amazon.com/sagemaker/pricing/).

### Amazon SageMaker Autopilot example notebooks

The following notebooks serve as practical, hands-on examples that address various use cases of Autopilot.

You can find all of Autopilot's notebooks in the `autopilot` directory of SageMaker GitHub examples repository.

We recommend cloning the full Git repository within Studio Classic to access and run the notebooks directly. For information on how to clone a Git repository in Studio Classic, see [Clone a Git Repository in SageMaker Studio Classic](https://docs.aws.amazon.com/sagemaker/latest/dg/clone-git-repository-studio-classic.html).

<table>
<thead>
<tr>
<th>Use case</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Serverless inference</strong></td>
<td>By default, Autopilot allows deploying generated models to real-time inference endpoints. In this repository, the notebook illustrates how to deploy Autopilot models...</td>
</tr>
<tr>
<td>Use case</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>trained with ENSEMBLING and HYPERPARAMETER OPTIMIZATION (HPO) modes to serverless endpoints. Serverless endpoints automatically launch compute resources and scale them in and out depending on traffic, eliminating the need to choose instance types or manage scaling policies.</td>
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<tr>
<td>Custom feature selection</td>
<td>Autopilot inspects your data set, and runs a number of candidates to figure out the optimal combination of data preprocessing steps, machine learning algorithms, and hyperparameters. You can easily deploy either on a real-time endpoint or for batch processing.</td>
</tr>
<tr>
<td></td>
<td>In some cases, you might want to have the flexibility to bring custom data processing code to Autopilot. For example, your datasets might contain a large number of independent variables, and you may wish to incorporate a custom feature selection step to remove irrelevant variables first. The resulting smaller dataset can then be used to launch an Autopilot job. Ultimately, you would also want to include both the custom processing code and models from Autopilot for real-time or batch processing.</td>
</tr>
<tr>
<td>Use case</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Pipeline example</td>
<td>While Autopilot streamlines the process of building ML models, MLOps engineers are still responsible for creating, automating, and managing end-to-end ML workflows in production. SageMaker Pipelines can assist in automating various steps of the ML lifecycle, such as data preprocessing, model training, hyperparameter tuning, model evaluation, and deployment. This notebook serves as a demonstration of how to incorporate Autopilot into a SageMaker Pipelines end-to-end AutoML training workflow. To launch an Autopilot experiment within Pipelines, you must create a model-building workflow by writing custom integration code using Pipelines Lambda or Processing steps. For more information, refer to <a href="https://docs.aws.amazon.com/sagemaker/latest/dg/autopilot-move-to-pipelines.html">Move Amazon SageMaker Autopilot ML models from experimentation to production using Amazon SageMaker Pipelines</a>. Alternatively, when using Autopilot in Ensembling mode, you can refer to the notebook example that demonstrates how to use native AutoML step in SageMaker Pipeline's native AutoML step. With Autopilot supported as a native step within Pipelines, you can now add an automated training step (AutoMLStep) to your Pipelines and invoke an Autopilot experiment in Ensembling mode.</td>
</tr>
<tr>
<td>More notebooks</td>
<td>You can find more notebooks illustrating other use cases such as batch transform, time-series forecasting and more in the root directory.</td>
</tr>
</tbody>
</table>
Amazon SageMaker Autopilot quotas

There are quotas that limit the resources available to you when using Amazon SageMaker Autopilot. Some of these limits are increasable and some are not.

Note

The resource quotas documented in the following sections are valid for versions of Amazon SageMaker Studio Classic 3.22.2 and higher. For information on updating your version of SageMaker Studio Classic, see Shut Down and Update SageMaker Studio Classic and Studio Classic Apps.

Topics

- Quotas that you can increase
- Resource quotas

Quotas that you can increase

Resource limits

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<tr>
<th>Resource</th>
<th>Regions</th>
<th>Default limits</th>
<th>Can be increased up to</th>
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<td>Hundreds of GBs</td>
</tr>
<tr>
<td>Size of a single Parquet file*</td>
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<td>Target dataset size for subsampling**</td>
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<td>Number of concurrent Autopilot jobs</td>
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<td>4</td>
<td>Hundreds</td>
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</table>
Resource limits per Autopilot job

<table>
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<tr>
<th>Resource</th>
<th>Limit per Autopilot job</th>
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</thead>
<tbody>
<tr>
<td>Maximum runtime for an Autopilot job</td>
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</table>
API Reference guide for Amazon SageMaker Autopilot

This section provides a subset of the HTTP service APIs for creating and managing Amazon SageMaker Autopilot resources (AutoML jobs) programmatically.

For information on the entire SageMaker REST APIs and the available SDKs, see API and SDK Reference.

If your language of choice is Python, you can refer to AWS SDK for Python (Boto3) directly.

Actions

This list details the operations available in the Reference API to manage AutoML jobs programmatically.

- `CreateAutoMLJob`
- `CreateAutoMLJobV2`
- `DescribeAutoMLJob`
- `DescribeAutoMLJobV2`
- `ListAutoMLJobs`
- `ListCandidatesForAutoMLJob`
- `StopAutoMLJob`

**Note**

`CreateAutoMLJobV2` and `DescribeAutoMLJobV2` are new versions of `CreateAutoMLJob` and `DescribeAutoMLJob` which offer backward compatibility. We recommend using the `CreateAutoMLJobV2`. `CreateAutoMLJobV2` can manage tabular problem types identical to those of its previous version `CreateAutoMLJob`, as well as non-tabular problem types such as image or text classification, or time-series forecasting.

Find guidelines about how to migrate a `CreateAutoMLJob` to `CreateAutoMLJobV2` in Migrate a `CreateAutoMLJob` to `CreateAutoMLJobV2`.

Data Types
This list details the API AutoML objects used by the actions above as inbound requests or outbound responses.

- `AutoMLAlgorithmConfig`
- `AutoMLCandidate`
- `AutoMLCandidateGenerationConfig`
- `AutoMLCandidateStep`
- `AutoMLChannel`
- `AutoMLContainerDefinition`
- `AutoMLDataSource`
- `AutoMLDataSplitConfig`
- `AutoMLInferenceContainerDefinitions`
- `AutoMLJobArtifacts`
- `AutoMLJobChannel`
- `AutoMLJobCompletionCriteria`
- `AutoMLJobInputDataConfig`
- `AutoMLJobConfig`
- `AutoMLJobObjective`
- `AutoMLJobStepMetadata`
- `AutoMLJobSummary`
- `AutoMLOutputDataConfig`
- `AutoMLProblemTypeConfig`
- `AutoMLJobCompletionCriteria`
- `AutoMLJobSummary`
- `AutoMLOutputDataConfig`
- `AutoMLPartialFailureReason`
- `AutoMLProblemTypeConfig`
- `AutoMLProblemTypeResolvedAttributes`
- `AutoMLResolvedAttributes`
SageMaker JumpStart

SageMaker JumpStart provides pretrained, open-source models for a wide range of problem types to help you get started with machine learning. You can incrementally train and tune these models before deployment. JumpStart also provides solution templates that set up infrastructure for common use cases, and executable example notebooks for machine learning with SageMaker.

You can deploy, fine-tune, and evaluate pretrained models from popular models hubs through the JumpStart landing page in the updated Studio experience.
You can also access pretrained models, solution templates, and examples through the JumpStart landing page in Amazon SageMaker Studio Classic.

The following steps show how to access JumpStart models using Amazon SageMaker Studio and Amazon SageMaker Studio Classic.

You can also access JumpStart models using the SageMaker Python SDK. For information about how to use JumpStart models programmatically, see Use SageMaker JumpStart Algorithms with Pretrained Models.

Open and use JumpStart in Studio

The following sections give information on how to open, use, and manage JumpStart from the Studio UI.

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

Open JumpStart in Studio

In Amazon SageMaker Studio, open the JumpStart landing page either through the Home page or the Home menu on the left-side panel. This opens the SageMaker JumpStart landing page where you can explore model hubs and search for models.

- From the Home page, choose JumpStart in the Prebuilt and automated solutions pane.
- From the Home menu in the left panel, navigate to the SageMaker JumpStart node.

For more information on getting started with Amazon SageMaker Studio, see Amazon SageMaker Studio.
Important

Before downloading or using third-party content: You are responsible for reviewing and complying with any applicable license terms and making sure that they are acceptable for your use case.

Use JumpStart in Studio

From the SageMaker JumpStart landing page in Studio, you can explore model hubs from providers of both proprietary and publicly available models.
You can find specific hubs or models using the search bar. Within each model hub, you can search directly for models, sort by provided attributes, or filter based on a list of provided model tasks.

**Manage JumpStart in Studio**

Choose a model to see its model detail card. In the upper right-hand corner of the model detail card, choose **Fine-tune**, **Deploy**, or **Evaluate** to start working through the fine-tuning, deployment, or evaluation workflows, respectively. Note that not all models are available for fine-tuning or evaluation. For more information on each of these options, see [Use foundation models in Studio](#).

**Open and use JumpStart in Studio Classic**

The following sections give information on how to open, use, and manage JumpStart from the Amazon SageMaker Studio Classic UI.
Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Open JumpStart in Studio Classic

In Amazon SageMaker Studio Classic, open the JumpStart landing page either through the Home page or the Home menu on the left-side panel.

- From the Home page you can either:
  - Choose JumpStart in the Prebuilt and automated solutions pane. This opens the SageMaker JumpStart landing page.
  - Choose a model directly in the SageMaker JumpStart landing page, or choose the Explore All option to see available solutions or models of a specific type.
- From the Home menu in the left panel you can either:
  - Navigate to the SageMaker JumpStart node, then choose Models, notebooks, solutions. This opens the SageMaker JumpStart landing page.
  - Navigate to the JumpStart node, then choose Launched JumpStart assets.

The Launched JumpStart assets page lists your currently launched solutions, deployed model endpoints, and training jobs created with JumpStart. You can access the JumpStart landing page from this tab by clicking on the Browse JumpStart button at the top right of the tab.

The JumpStart landing page lists available end-to-end machine learning solutions, pretrained models, and example notebooks. From any individual solution or model page, you can choose the Browse JumpStart button at the top right of the tab to return to the SageMaker JumpStart page.
Important

Before downloading or using third-party content: You are responsible for reviewing and complying with any applicable license terms and making sure that they are acceptable for your use case.

Use JumpStart in Studio Classic

From the SageMaker JumpStart landing page, you can browse for solutions, models, notebooks, and other resources.
You can find JumpStart resources by using the search bar, or by browsing each category. Use the tabs to filter the available solutions by categories:

- **Solutions** – In one step, launch comprehensive machine learning solutions that tie SageMaker to other AWS services. Select **Explore All Solutions** to view all available solutions.

- **Resources** – Use example notebooks, blogs, and video tutorials to learn and head start your problem types.
  
  - **Blogs** – Read details and solutions from machine learning experts.
  
  - **Video tutorials** – Watch video tutorials for SageMaker features and machine learning use cases from machine learning experts.
  
  - **Example notebooks** – Run example notebooks that use SageMaker features like Spot Instance training and experiments over a large variety of model types and use cases.

- **Data types** – Find a model by data type (e.g., Vision, Text, Tabular, Audio, Text Generation). Select **Explore All Models** to view all available models.

- **ML tasks** – Find a model by problem type (e.g., Image Classification, Image Embedding, Object Detection, Text Generation). Select **Explore All Models** to view all available models.

- **Notebooks** – Find example notebooks that use SageMaker features across multiple model types and use cases. Select **Explore All Notebooks** to view all available example notebooks.

- **Frameworks** – Find a model by framework (e.g., PyTorch, TensorFlow, Hugging Face).
Manage JumpStart in Studio Classic

From the Home menu in the left panel, navigate to SageMaker JumpStart, then choose Launched JumpStart assets to list your currently launched solutions, deployed model endpoints, and training jobs created with JumpStart.

Topics

- JumpStart Foundation Models
- Task-Specific Models
- Shared Models and Notebooks
- Solution Templates
- Amazon SageMaker JumpStart Industry: Financial

JumpStart Foundation Models

Amazon SageMaker JumpStart offers state-of-the-art foundation models for use cases such as content writing, code generation, question answering, copywriting, summarization, classification, information retrieval, and more. Use JumpStart foundation models to build your own generative AI solutions and integrate custom solutions with additional SageMaker features. For more information, see Getting started with Amazon SageMaker JumpStart.

A foundation model is a large pre-trained model that is adaptable to many downstream tasks and often serves as the starting point for developing more specialized models. Examples of foundation models include LLaMa-2-7b, BLOOM 176B, FLAN-T5 XL, or GPT-J 6B, which are pre-trained on massive amounts of text data and can be fine-tuned for specific language tasks.

Amazon SageMaker JumpStart onboards and maintains publicly available foundation models for you to access, customize, and integrate into your machine learning lifecycles. For more information, see Publicly available foundation models. Amazon SageMaker JumpStart also includes proprietary foundation models from third-party providers. For more information, see Proprietary foundation models.

To get started exploring and experimenting with available models, see How to use JumpStart foundation models. All foundation models are available to use programatically with the SageMaker Python SDK. For more information, see Use foundation models with the SageMaker Python SDK.
For more information on considerations to make when choosing a model, see Model sources and license agreements.

For specifics about customization and fine-tuning foundation models, see Customize a foundation model.

For more general information on foundation models, see the paper On the Opportunities and Risks of Foundation Models.

Topics

- Explore the latest foundation models
- How to use JumpStart foundation models
- Model sources and license agreements
- Customize a foundation model
- Evaluate a text generation foundation model in Studio
- Example notebooks

Explore the latest foundation models

Amazon SageMaker JumpStart offers state-of-the-art, built-in publicly available and proprietary foundation models to customize and integrate into your generative AI workflows.

Publicly available foundation models

Amazon SageMaker JumpStart onboards and maintains open source foundation models from third-party sources. To get started with one of these publicly available models, see How to use JumpStart foundation models or explore one of the available Example notebooks. In a given example notebook for a publicly available model, try switching out the model ID to experiment with different models within the same model family.

For more information on model IDs and resources on deploying publicly available JumpStart foundation models with the SageMaker Python SDK, see Use foundation models with the SageMaker Python SDK.

By definition, foundation models are adaptable to many downstream tasks. Foundation models are trained on massive amounts of general domain data and the same model can be implemented or customized for multiple use cases. When choosing your foundation model, start with defining a specific task, such as text generation or image generation.
Publicly available text generation models

Text generation foundation models can be used for a variety of downstream tasks, including text summarization, text classification, question answering, long-form content generation, short-form copywriting, information extraction, and more.

Publicly available text generation model table

<table>
<thead>
<tr>
<th>Model Name</th>
<th>Model ID</th>
<th>Model Source</th>
<th>Fine-tunable</th>
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To explore the latest text generation JumpStart foundation models, use the **Text Generation** filter on the [Getting started with Amazon SageMaker JumpStart](#) product description page. You can also explore foundation models based on tasks directly in the Amazon SageMaker Studio UI or SageMaker Studio Classic UI. Only a subset of publicly available text generation models are available for fine-tuning in JumpStart. For more information, see [Use foundation models in Amazon SageMaker Studio Classic](#).

**Publicly available image generation models**

JumpStart provides a wide variety of Stable Diffusion image generation foundation models including base models from Stability AI as well as pre-trained models for specific text-to-image tasks from Hugging Face. If you need to fine-tune your text-to-image foundation model, you can use Stable Diffusion 2.1 base from Stability AI. If you want to explore models that are already
trained on specific art styles, you can explore one of the many third-party models from Hugging Face directly in the Amazon SageMaker Studio UI or SageMaker Studio Classic UI.

To explore the latest image generation JumpStart foundation models, use the Text to Image filter on the Getting started with Amazon SageMaker JumpStart product description page. To get started with your chosen text-to-image foundation model, see How to use JumpStart foundation models.

Proprietary foundation models

Amazon SageMaker JumpStart provides access to proprietary foundation models from third-party providers such as AI21 Labs, Cohere, and LightOn.

To get started with one of these proprietary models, see How to use JumpStart foundation models. To use a proprietary foundation model, you must first subscribe to the model in AWS Marketplace. After subscribing to the model, locate the foundation model in Studio or SageMaker Studio Classic. For more information, see SageMaker JumpStart.

To explore the latest proprietary foundation models for a variety of use cases, see Getting started with Amazon SageMaker JumpStart.

How to use JumpStart foundation models

Choose, train, or deploy foundation models through Amazon SageMaker Studio or Amazon SageMaker Studio Classic, use JumpStart foundation models programmatically with the SageMaker Python SDK, or discover JumpStart foundation models directly through the SageMaker console.

Topics

- Use foundation models in Studio
- Use foundation models in Amazon SageMaker Studio Classic
- Use foundation models with the SageMaker Python SDK
- Discover foundation models in the SageMaker Console

Use foundation models in Studio

You can fine-tune, deploy, and evaluate both publicly available and proprietary JumpStart foundation models directly through the Amazon SageMaker Studio UI.
Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

In Amazon SageMaker Studio, open the JumpStart landing page either through the Home page or the Home menu on the left-side panel. This opens the SageMaker JumpStart landing page where you can explore model hubs and search for models.

- From the Home page, choose JumpStart in the Prebuilt and automated solutions pane.
- From the Home menu in the left panel, navigate to the JumpStart node.

For more information on getting started with Amazon SageMaker Studio, see Amazon SageMaker Studio.

From the SageMaker JumpStart landing page in Studio, you can explore model hubs from providers of both publicly available and proprietary models. You can find specific hubs or models using the search bar. Within each model hub, you can search directly for models, sort by Most likes, Most downloads, or Recently updated, or filter based on a list of provided model tasks. Choose a model to see its model detail card. In the upper right corner of the model detail card, choose Fine-tune, Deploy, or Evaluate to start working through the fine-tuning, deployment, or evaluation workflows, respectively. Note that not all models are available for fine-tuning or evaluation.

Fine-tune foundation models in Studio

Fine-tuning trains a pre-trained model on a new dataset without training from scratch. This process, also known as transfer learning, can produce accurate models with smaller datasets and less training time. To fine-tune JumpStart foundation models, navigate to a model detail card in the Studio UI. For more information on how to open JumpStart in Studio, see Open and use JumpStart in Studio. After navigating to the model detail card of your choice, choose Train in the upper right corner. Note that not all models have fine-tuning available.
Important

Some foundation models require explicit acceptance of an end-user license agreement (EULA) before fine-tuning. For more information, see EULA acceptance in Amazon SageMaker Studio.

Model settings

When using a pre-trained JumpStart foundation model in Amazon SageMaker Studio, the Model artifact location (Amazon S3 URI) is populated by default. To edit the default Amazon S3 URI, choose Enter model artifact location. Not all models support changing the model artifact location.

Data settings

In the Data field, provide an Amazon S3 URI point to your training dataset location. The default Amazon S3 URI points to an example training dataset. To edit the default Amazon S3 URI, choose Enter training dataset and change the URI. Be sure to review the model detail card in Amazon SageMaker Studio for information on formatting training data.

Hyperparameters

You can customize the hyperparameters of the training job that are used to fine-tune the model. The hyperparameters available for each fine-tunable model differ depending on the model.

The following hyperparameters are common among models:

- **Epochs** – One epoch is one cycle through the entire dataset. Multiple intervals complete a batch, and multiple batches eventually complete an epoch. Multiple epochs are run until the accuracy of the model reaches an acceptable level, or when the error rate drops below an acceptable level.

- **Learning rate** – The amount that values should be changed between epochs. As the model is refined, its internal weights are being nudged and error rates are checked to see if the model improves. A typical learning rate is 0.1 or 0.01, where 0.01 is a much smaller adjustment and could cause the training to take a long time to converge, whereas 0.1 is much larger and can cause the training to overshoot. It is one of the primary hyperparameters that you might adjust for training your model. Note that for text models, a much smaller learning rate (5e-5 for BERT) can result in a more accurate model.

- **Batch size** – The number of records from the dataset that is to be selected for each interval to send to the GPUs for training.
Review the tool tip prompts and additional information in the model detail card in the Studio UI to learn more about hyperparameters specific to the model of your choice.

For more information on available hyperparameters, see Commonly supported fine-tuning hyperparameters.

**Deployment**

Specify the training instance type and output artifact location for your training job. You can only choose from instances that are compatible with the model of your choice within the fine-tuning the Studio UI. The default output artifact location is the SageMaker default bucket. To change the output artifact location, choose **Enter output artifact location** and change the Amazon S3 URI.

**Security**

Specify the security settings to use for your training job, including the IAM role that SageMaker uses to train your model, whether your training job should connect to a virtual private cloud (VPC), and any encryption keys to secure your data.

**Additional information**

In the **Additional Information** field you can edit the training job name. You can also add and remove tags in the form of key-value pairs to help organize and categorize your fine-tuning training jobs.

After providing information for your fine-tuning configuration, choose **Submit**. If the pre-trained foundation model that you chose to fine-tune requires explicit agreement of an end-user license agreement (EULA) before training, the EULA is provided in a pop-up window. To accept the terms of the EULA, choose **Accept**. You are responsible for reviewing and complying with any applicable license terms and making sure they are acceptable for your use case before downloading or using a model.

**Deploy foundation models in Studio**

To deploy JumpStart foundation models, navigate to a model detail card in the Studio UI. For more information on how to open JumpStart in Studio, see Open and use JumpStart in Studio. After navigating to the model detail page of your choice, choose **Deploy** in the upper right corner of the Studio UI. Then, follow the steps in Deploy models with SageMaker Studio.
Important

Some foundation models require explicit acceptance of an end-user license agreement (EULA) before deployment. For more information, see EULA acceptance in Amazon SageMaker Studio.

Evaluate foundation models in Studio

Amazon SageMaker JumpStart has integrations with SageMaker Clarify foundation model evaluations (FME) in Studio. If a JumpStart model has built-in evaluation capabilities available, you can choose Evaluate in the upper right corner of the model detail page in the JumpStart Studio UI. For more information, see Evaluate a foundation model.

Use foundation models in Amazon SageMaker Studio Classic

You can fine-tune and deploy both publicly available and proprietary JumpStart foundation models through the Studio Classic UI.

Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

To get started with Studio Classic, see Launch Amazon SageMaker Studio Classic.
After opening Amazon SageMaker Studio Classic, choose **Models, notebooks, solutions** in the SageMaker JumpStart section of the navigation pane. Then, scroll down to find either the **Foundation Models: Text Generation** or **Foundation Models: Image Generation** section depending on your use case.

You can choose **View model** on a suggested foundation model card, or choose **Explore All Models** to see all available foundation models for either text generation or image generation. If you choose to see all available models, you can further filter available models by task, data type, content type, or framework. You can also search for a model name directly in the **Search** bar. If you need guidance on selecting a model, see **Explore the latest foundation models**.

⚠️ **Important**

Some foundation models require explicit acceptance of an end-user license agreement (EULA). For more information, see **EULA acceptance in Amazon SageMaker Studio**.

After you choose **View model** for the foundation model of your choice in Studio Classic, you can deploy the model. For more information, see **Deploy a Model**.
You can also choose **Open notebook** in the **Run in notebook** section to run an example notebook for the foundation model directly in Studio Classic.

**Note**

To deploy a proprietary foundation model in Studio Classic, you must first subscribe to the model in AWS Marketplace. The AWS Marketplace link is provided in the associated example notebook within Studio Classic.

If the model is fine-tunable, you can also fine-tune the model. For more information, see [Fine-Tune a Model](#). For a list of which JumpStart foundation models are fine-tunable, see [Fine-tune a foundation model](#).

**Use foundation models with the SageMaker Python SDK**

All JumpStart foundation models are available to deploy programmatically using the SageMaker Python SDK. Publicly available text generation foundation models can be deployed using the model ID in the **Publicly available text generation model table**. Proprietary models must be deployed using the model package information after subscribing to the model in AWS Marketplace.

The following sections show how to fine-tune foundation models using the `JumpStartEstimator` class and how to deploy models using the `JumpStartModel` class, along with additional Python SDK utilities.

**Important**

Some foundation models require explicit acceptance of an end-user license agreement (EULA). For more information, see [EULA acceptance with the SageMaker Python SDK](#).

To reference available model IDs for all publicly available foundation models, see the **Built-in Algorithms with pre-trained Model Table**. Search for the name of the foundation model of your choice in the **Search** bar, change the number of entries shown using the **Show entries** dropdown menu, or choose the **Next** text highlighted in blue on the left side of the page to navigate through the available models.
Fine-tune publicly available foundation models with the JumpStartEstimator class

You can fine-tune a built-in algorithm or pre-trained model in just a few lines of code using the SageMaker Python SDK.

1. First, find the model ID for the model of your choice in the **Built-in Algorithms with pre-trained Model Table**.

2. Using the model ID, define your training job as a JumpStart estimator.

   ```python
   from sagemaker.jumpstart.estimator import JumpStartEstimator
   model_id = "huggingface-textgeneration1-gpt-j-6b"
   estimator = JumpStartEstimator(model_id=model_id)
   ```

3. Run `estimator.fit()` on your model, pointing to the training data to use for fine-tuning.

   ```python
   estimator.fit(
       {"train": training_dataset_s3_path, "validation": validation_dataset_s3_path}
   )
   ```

4. Then, use the deploy method to automatically deploy your model for inference. In this example, we use the GPT-J 6B model from Hugging Face.

   ```python
   predictor = estimator.deploy()
   ```

5. You can then run inference with the deployed model using the predict method.

   ```python
   question = "What is Southern California often abbreviated as?"
   response = predictor.predict(question)
   print(response)
   ```

**Note**

This example uses the foundation model GPT-J 6B, which is suitable for a wide range of text generation use cases including question answering, named entity recognition, summarization, and more. For more information about model use cases, see [Explore the latest foundation models](#).
You can optionally specify model versions or instance types when creating your JumpStartEstimator. For more information about the JumpStartEstimator class and its parameters, see `JumpStartEstimator`.

**Check default instance types**

You can optionally include specific model versions or instance types when fine-tuning a pre-trained model using the JumpStartEstimator class. All JumpStart models have a default instance type. Retrieve the default training instance type using the following code:

```python
from sagemaker import instance_types

instance_type = instance_types.retrieve_default(
    model_id=model_id,
    model_version=model_version,
    scope="training")

print(instance_type)
```

You can see all supported instance types for a given JumpStart model with the `instance_types.retrieve()` method.

**Check default hyperparameters**

To check the default hyperparameters used for training, you can use the `retrieve_default()` method from the `hyperparameters` class.

```python
from sagemaker import hyperparameters

my_hyperparameters = hyperparameters.retrieve_default(model_id=model_id,
    model_version=model_version)

print(my_hyperparameters)

# Optionally override default hyperparameters for fine-tuning
my_hyperparameters["epoch"] = "3"
my_hyperparameters["per_device_train_batch_size"] = "4"

# Optionally validate hyperparameters for the model
hyperparameters.validate(model_id=model_id, model_version=model_version, hyperparameters=my_hyperparameters)
```

For more information on available hyperparameters, see [Commonly supported fine-tuning hyperparameters](#).
Check default metric definitions

You can also check the default metric definitions:

```python
print(metric_definitions.retrieve_default(model_id=model_id,
    model_version=model_version))
```

Deploy publicly available foundation models with the JumpStartModel class

You can deploy a built-in algorithm or pre-trained model to a SageMaker endpoint in just a few lines of code using the SageMaker Python SDK.

1. First, find the model ID for the model of your choice in the Built-in Algorithms with pre-trained Model Table.
2. Using the model ID, define your model as a JumpStart model.

```python
from sagemaker.jumpstart.model import JumpStartModel

model_id = "huggingface-text2text-flan-t5-xl"
my_model = JumpStartModel(model_id=model_id)
```

3. Use the deploy method to automatically deploy your model for inference. In this example, we use the FLAN-T5 XL model from Hugging Face.

```python
predictor = my_model.deploy()
```

4. You can then run inference with the deployed model using the predict method.

```python
question = "What is Southern California often abbreviated as?"
response = predictor.predict(question)
print(response)
```

### Note

This example uses the foundation model FLAN-T5 XL, which is suitable for a wide range of text generation use cases including question answering, summarization, chatbot creation, and more. For more information about model use cases, see Explore the latest foundation models.
For more information about the JumpStartModel class and its parameters, see JumpStartModel.

Check default instance types

You can optionally include specific model versions or instance types when deploying a pre-trained model using the JumpStartModel class. All JumpStart models have a default instance type. Retrieve the default deployment instance type using the following code:

```python
from sagemaker import instance_types

instance_type = instance_types.retrieve_default(
    model_id=model_id,
    model_version=model_version,
    scope="inference")
print(instance_type)
```

See all supported instance types for a given JumpStart model with the instance_types.retrieve() method.

Use inference components to deploy multiple models to a shared endpoint

An inference component is a SageMaker hosting object that you can use to deploy one or more models to an endpoint for increased flexibility and scalability. You must change the endpoint_type for your JumpStart model to be inference-component-based rather than the default model-based endpoint.

```python
predictor = my_model.deploy(
    endpoint_name = 'jumpstart-model-id-123456789012',
    endpoint_type = EndpointType.INFERENCE_COMPONENT_BASED
)
```

For more information on creating endpoints with inference components and deploying SageMaker models, see Shared resource utilization with multiple models.

Check valid input and output inference formats

To check valid data input and output formats for inference, you can use the retrieve_options() method from the Serializers and Deserializers classes.

```python
print(sagemaker.serializers.retrieve_options(model_id=model_id,
                                           model_version=model_version))
```
Check supported content and accept types

Similarly, you can use the `retrieve_options()` method to check the supported content and accept types for a model.

```python
print(sagemaker.deserializers.retrieve_options(model_id=model_id,
                                             model_version=model_version))
print(sagemaker.content_types.retrieve_options(model_id=model_id,
                                             model_version=model_version))
print(sagemaker.accept_types.retrieve_options(model_id=model_id,
                                             model_version=model_version))
```

For more information about utilities, see Utility APIs.

Use proprietary foundation models with the SageMaker Python SDK

Proprietary models must be deployed using the model package information after subscribing to the model in AWS Marketplace. For more information about SageMaker and AWS Marketplace, see Buy and Sell Amazon SageMaker Algorithms and Models in AWS Marketplace. To find AWS Marketplace links for the latest proprietary models, see Getting started with Amazon SageMaker JumpStart.

After subscribing to the model of your choice in AWS Marketplace, you can deploy the foundation model using the SageMaker Python SDK and the SDK associated with the model provider. For example, AI21 Labs, Cohere, and LightOn use the "ai21[SM]", cohere-sagemaker, and lightonsage packages, respectively.

For example, to define a JumpStart model using Jurassic-2 Jumbo Instruct from AI21 Labs, use the following code:

```python
import sagemaker
import ai21

role = get_execution_role()

sagemaker_session = sagemaker.Session()

model_package_arn = "arn:aws:sagemaker:us-east-1:865070037744:model-package/j2-jumbo-instruct-v1-1-43-4e47c49e61743066b9d95efed6882f35"

my_model = ModelPackage(
    role=role, model_package_arn=model_package_arn, sagemaker_session=sagemaker_session)
```
For step-by-step examples, find and run the notebook associated with the proprietary foundation model of your choice in SageMaker Studio Classic. See Use foundation models in Amazon SageMaker Studio Classic for more information. For more information on the SageMaker Python SDK, see ModelPackage.

Discover foundation models in the SageMaker Console

You can explore JumpStart foundation models directly through the Amazon SageMaker Console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Find JumpStart on the left navigation panel and choose Foundation models.
3. Browse models or search for a specific model. If you need guidance for model selection, see Explore the latest foundation models. Choose View model to view the model detail page for the foundation model of your choice.
4. If the model is a proprietary model, choose Subscribe in the upper right corner of the model detail page to subscribe to the model in AWS Marketplace. You should receive an email confirming your subscription to the model of your choice. For more information about SageMaker and AWS Marketplace, see Buy and Sell Amazon SageMaker Algorithms and Models in AWS Marketplace. Publicly available foundation models do not require a subscription.
5. To view an example notebook in GitHub, choose View code in the upper right corner of the model detail page.
6. To view and run an example notebook directly in Amazon SageMaker Studio Classic, choose Open notebook in Studio in the upper right corner of the model detail page.

Model sources and license agreements

Amazon SageMaker JumpStart provides access to hundreds of publicly available and proprietary foundation models from third-party sources and partners. You can explore the JumpStart foundation model selection directly in the SageMaker console, Studio, or Studio Classic.

 Licenses and model sources

Amazon SageMaker JumpStart provides access to both publicly available and proprietary foundation models. Foundation models are onboarded and maintained from third-party open source and proprietary providers. As such, they are released under different licenses as designated by the model source. Be sure to review the license for any foundation model that you use. You
are responsible for reviewing and complying with any applicable license terms and making sure they are acceptable for your use case before downloading or using the content. Some examples of common foundation model licenses include:

- Alexa Teacher Model
- Apache 2.0
- BigScience Responsible AI License v1.0
- CreativeML Open RAIL++-M license

Similarly, for any proprietary foundation models, be sure to review and comply with any terms of use and usage guidelines from the model provider. If you have questions about license information for a specific proprietary model, reach out to model provider directly. You can find model provider contact information in the Support tab of each model page in AWS Marketplace.

**End-user license agreements**

Some JumpStart foundation models require explicit acceptance of an end-user license agreement (EULA) before use.

**EULA acceptance in Amazon SageMaker Studio**

You may be prompted to accept an end-user license agreement before fine-tuning, deploying, or evaluating a JumpStart foundation model in Studio. To get started with JumpStart foundation models in Studio, see [Use foundation models in Studio](#).

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).

Some JumpStart foundation models require acceptance of an end-user license agreement before deployment. If this applies to the foundation model that you choose to use, Studio prompts you with a window containing the EULA content. You are responsible for reviewing and complying with any applicable license terms and making sure they are acceptable for your use case before downloading or using a model.
EULA acceptance in Amazon SageMaker Studio Classic

You may be prompted to accept an end-user license agreement before deploying a JumpStart foundation model or opening a JumpStart foundation model notebook in Studio Classic. To get started with JumpStart foundation models in Studio Classic, see Use foundation models in Amazon SageMaker Studio Classic.

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Some JumpStart foundation models require acceptance of an end-user license agreement before deployment. If this applies to the foundation model that you choose to use, Studio Classic prompts you with a window titled Review the End User License Agreement (EULA) and Acceptable Use Policy (AUP) below after you choose either Deploy or Open notebook. You are responsible for reviewing and complying with any applicable license terms and making sure they are acceptable for your use case before downloading or using a model.

EULA acceptance with the SageMaker Python SDK

The following sections show you how to explicitly declare EULA acceptance when deploying or fine-tuning a JumpStart model with the SageMaker Python SDK. For more information on getting started with JumpStart foundation models using the SageMaker Python SDK, see Use foundation models with the SageMaker Python SDK.

Before you begin, make sure that you do the following:

- Upgrade to the latest version of the model that you use.
- Install the latest version of the SageMaker Python SDK.

⚠️ Important
To use the following workflow you must have v2.198.0 or later of the SageMaker Python SDK installed.
EULA acceptance when deploying a JumpStart model

For models that require the acceptance of an end-user license agreement, you must explicitly declare EULA acceptance when deploying your JumpStart model.

```python
from sagemaker.jumpstart.model import JumpStartModel
model_id = "meta-textgeneration-llama-2-13b"
my_model = JumpStartModel(model_id=model_id)

# Declare EULA acceptance when deploying your JumpStart model
predictor = my_model.deploy(accept_eula=True)
```

The `accept_eula` value is `None` by default and must be explicitly redefined as `True` in order to accept the end-user license agreement. For more information, see JumpStartModel.

EULA acceptance when fine-tuning a JumpStart model

For fine-tuning models that require the acceptance of an end-user license agreement, you must explicitly declare EULA acceptance when defining your JumpStart estimator. After fine-tuning a pre-trained model, the weights of the original model are changed. Therefore, when you deploy the fine-tuned model later, you do not need to accept a EULA.

```python
from sagemaker.jumpstart.estimator import JumpStartEstimator
model_id = "meta-textgeneration-llama-2-13b"

# Declare EULA acceptance when defining your JumpStart estimator
estimator = JumpStartEstimator(model_id=model_id, environment={"accept_eula": "true"})
estimator.fit(
    {"train": training_dataset_s3_path, "validation": validation_dataset_s3_path}
)
```

The `accept_eula` value is `None` by default and must be explicitly redefined as "true" within the estimator environment in order to accept the end-user license agreement. For more information, see JumpStartEstimator.

EULA acceptance SageMaker Python SDK versions earlier than 2.198.0

⚠️ Important

When using versions earlier than 2.198.0 of the SageMaker Python SDK, you must use the SageMaker Predictor class to accept a model EULA.
After deploying a JumpStart foundation model programmatically using the SageMaker Python SDK, you can run inference against your deployed endpoint with the SageMaker `Predictor` class. For models that require the acceptance of an end-user license agreement, you must explicitly declare EULA acceptance in your call to the `Predictor` class:

```python
predictor.predict(payload, custom_attributes="accept_eula=true")
```

The `accept_eula` value is false by default and must be explicitly redefined as true in order to accept the end-user license agreement. The predictor returns an error if you try to run inference while `accept_eula` is set to false. For more information on getting started with JumpStart foundation models using the SageMaker Python SDK, see Use foundation models with the SageMaker Python SDK.

⚠️ Important

The `custom_attributes` parameter accepts key-value pairs in the format "key1=value1;key2=value2". If you use the same key multiple times, the inference server uses the last value associated with the key. For example, if you pass "accept_eula=false;accept_eula=true" to the `custom_attributes` parameter, then the inference server associates the value true with the `accept_eula` key.

Customize a foundation model

Foundation models are extremely powerful models able to solve a wide array of tasks. To solve most tasks effectively, these models require some form of customization.

The recommended way to first customize a foundation model to a specific use case is through prompt engineering. Providing your foundation model with well-engineered, context-rich prompts can help achieve desired results without any fine-tuning or changing of model weights. For more information, see Prompt engineering for foundation models.

If prompt engineering alone is not enough to customize your foundation model to a specific task, you can fine-tune a foundation model on additional domain-specific data. For more information, see Fine-tune a foundation model. The fine-tuning process involves changing model weights.

If you want to customize your model with information from a knowledge library without any retraining, see Retrieval Augmented Generation (RAG).
Prompt engineering for foundation models

Prompt engineering is the process of designing and refining the prompts or input stimuli for a language model to generate specific types of output. Prompt engineering involves selecting appropriate keywords, providing context, and shaping the input in a way that encourages the model to produce the desired response and is a vital technique to actively shape the behavior and output of foundation models.

Effective prompt engineering is crucial for directing model behavior and achieving desired responses. Through prompt engineering, you can control a model's tone, style, and domain expertise without more involved customization measures like fine-tuning. We recommend dedicating time to prompt engineering before you consider fine-tuning a model on additional data. The goal is to provide sufficient context and guidance to the model so that it can generalize and perform well on unseen or limited data scenarios.

Zero-shot learning

Zero-shot learning involves training a model to generalize and make predictions on unseen classes or tasks. To perform prompt engineering in zero-shot learning environments, we recommend constructing prompts that explicitly provide information about the target task and the desired output format. For example, if you want to use a foundation model for zero-shot text classification on a set of classes that the model did not see during training, a well-engineered prompt could be: "Classify the following text as either sports, politics, or entertainment: [input text]." By explicitly specifying the target classes and the expected output format, you can guide the model to make accurate predictions even on unseen classes.

Few-shot learning

Few-shot learning involves training a model with a limited amount of data for new classes or tasks. Prompt engineering in few-shot learning environments focuses on designing prompts that effectively use the limited available training data. For example, if you use a foundation model for an image classification task and only have a few examples of a new image class, you can engineer a prompt that includes the available labeled examples with a placeholder for the target class. For example, the prompt could be: "[image 1], [image 2], and [image 3] are examples of [target class]. Classify the following image as [target class]." By incorporating the limited labeled examples and explicitly specifying the target class, you can guide the model to generalize and make accurate predictions even with minimal training data.
Supported inference parameters

Changing inference parameters might also affect the responses to your prompts. While you can try to add as much specificity and context as possible to your prompts, you can also experiment with supported inference parameters. The following are examples of some commonly supported inference parameters:

<table>
<thead>
<tr>
<th>Inference Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>max_new_tokens</td>
<td>The maximum output length of a foundation model response. Valid values: integer, range: Positive integer.</td>
</tr>
<tr>
<td>temperature</td>
<td>Controls the randomness in the output. Higher temperature results in an output sequence with low-probability words and lower temperature results in output sequence with high-probability words. If $temperature=0$, the response is made up of only the highest probability words (greedy decoding). Valid values: float, range: Positive float.</td>
</tr>
<tr>
<td>top_p</td>
<td>In each step of text generation, the model samples from the smallest possible set of words with a cumulative probability of $top_p$. Valid values: float, range: 0.0, 1.0.</td>
</tr>
<tr>
<td>return_full_text</td>
<td>If True, then the input text is part of the generated output text. Valid values: boolean, default: False.</td>
</tr>
</tbody>
</table>

For more information on foundation model inference, see [Deploy publicly available foundation models with the JumpStartModel class](#).

If prompt engineering is not sufficient to adapt your foundation model to specific business needs, domain-specific language, target tasks, or other requirements, you can consider fine-tuning your model on additional data or using Retrieval Augmented Generation (RAG) to augment your model architecture with enhanced context from archived knowledge sources. For more information, see [Fine-tune a foundation model](#) or [Retrieval Augmented Generation (RAG)](#).

Fine-tune a foundation model

Foundation models are computationally expensive and trained on a large, unlabeled corpus. Fine-tuning a pre-trained foundation model is an affordable way to take advantage of their broad
capabilities while customizing a model on your own small, corpus. Fine-tuning is a customization method that involved further training and does change the weights of your model.

Fine-tuning might be useful to you if you need:

- to customize your model to specific business needs
- your model to successfully work with domain-specific language, such as industry jargon, technical terms, or other specialized vocabulary
- enhanced performance for specific tasks
- accurate, relative, and context-aware responses in applications
- responses that are more factual, less toxic, and better-aligned to specific requirements

There are two main approaches that you can take for fine-tuning depending on your use case and chosen foundation model.

1. If you're interested in fine-tuning your model on domain-specific data, see [Domain adaptation fine-tuning](#).
2. If you're interested in instruction-based fine-tuning using prompt and response examples, see [Instruction-based fine-tuning](#).

**Foundation models available for fine-tuning**

You can fine-tune any of the following JumpStart foundation models:

- Bloom 3B
- Bloom 7B1
- BloomZ 3B FP16
- BloomZ 7B1 FP16
- Code Llama 13B
- Code Llama 13B Python
- Code Llama 34B
- Code Llama 34B Python
- Code Llama 70B
- Code Llama 70B Python
- Code Llama 7B
• Code Llama 7B Python
• CyberAgentLM2-7B-Chat (CALM2-7B-Chat)
• Falcon 40B BF16
• Falcon 40B Instruct BF16
• Falcon 7B BF16
• Falcon 7B Instruct BF16
• Flan-T5 Base
• Flan-T5 Large
• Flan-T5 Small
• Flan-T5 XL
• Flan-T5 XXL
• Gemma 2B
• Gemma 2B Instruct
• Gemma 7B
• Gemma 7B Instruct
• GPT-2 XL
• GPT-J 6B
• GPT-Neo 1.3B
• GPT-Neo 125M
• GPT-NEO 2.7B
• LightGPT Instruct 6B
• Llama 2 13B
• Llama 2 13B Chat
• Llama 2 13B Neuron
• Llama 2 70B
• Llama 2 70B Chat
• Llama 2 7B
• Llama 2 7B Chat
• Llama 2 7B Neuron
• Mistral 7B
• Mixtral 8x7B
• Mixtral 8x7B Instruct
• RedPajama INCITE Base 3B V1
• RedPajama INCITE Base 7B V1
• RedPajama INCITE Chat 3B V1
• RedPajama INCITE Chat 7B V1
• RedPajama INCITE Instruct 3B V1
• RedPajama INCITE Instruct 7B V1
• Stable Diffusion 2.1

**Commonly supported fine-tuning hyperparameters**

Different foundation models support different hyperparameters when fine-tuning. The following are commonly-supported hyperparameters that can further customize your model during training:

<table>
<thead>
<tr>
<th>Inference Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>epoch</td>
<td>The number of passes that the model takes through the fine-tuning dataset during training. Must be an integer greater than 1.</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The rate at which the model weights are updated after working through each batch of fine-tuning training examples. Must be a positive float greater than 0.</td>
</tr>
<tr>
<td>instruction_tuned</td>
<td>Whether to instruction-train the model or not. Must be 'True' or 'False'.</td>
</tr>
<tr>
<td>per_device_train_batch_size</td>
<td>The batch size per GPU core or CPU for training. Must be a positive integer.</td>
</tr>
<tr>
<td>per_device_eval_batch_size</td>
<td>The batch size per GPU core or CPU for evaluation. Must be a positive integer.</td>
</tr>
<tr>
<td>max_train_samples</td>
<td>For debugging purposes or quicker training, truncate the number of training examples to this value. Value -1 means that</td>
</tr>
<tr>
<td>Inference Parameter</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>max_val_samples</td>
<td>For debugging purposes or quicker training, truncate the number of validation examples to this value. Value -1 means that the model uses all of the validation samples. Must be a positive integer or -1.</td>
</tr>
<tr>
<td>max_input_length</td>
<td>Maximum total input sequence length after tokenization. Sequences longer than this will be truncated. If -1, max_input_length is set to the minimum of 1024 and the model_max_length defined by the tokenizer. If set to a positive value, max_input_length is set to the minimum of the provided value and the model_max_length defined by the tokenizer. Must be a positive integer or -1.</td>
</tr>
<tr>
<td>validation_split_ratio</td>
<td>If there is no validation channel, ratio of train-validation split from the training data. Must be between 0 and 1.</td>
</tr>
<tr>
<td>train_data_split_seed</td>
<td>If validation data is not present, this fixes the random splitting of the input training data to training and validation data used by the model. Must be an integer.</td>
</tr>
<tr>
<td>preprocessing_num_workers</td>
<td>The number of processes to use for the pre-processing. If None, main process is used for pre-processing.</td>
</tr>
<tr>
<td>lora_r</td>
<td>Low-rank adaptation (LoRA) r value, which acts as the scaling factor for weight updates. Must be a positive integer.</td>
</tr>
<tr>
<td>lora_alpha</td>
<td>Low-rank adaptation (LoRA) alpha value, which acts as the scaling factor for weight updates. Generally 2 to 4 times the size of lora_r. Must be a positive integer.</td>
</tr>
<tr>
<td>lora_dropout</td>
<td>Dropout value for low-rank adaptation (LoRA) layers Must be a positive float between 0 and 1.</td>
</tr>
<tr>
<td>int8_quantization</td>
<td>If True, model is loaded with 8 bit precision for training.</td>
</tr>
<tr>
<td>Inference Parameter</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>enable_fsdp</td>
<td>If True, training uses Fully Sharded Data Parallelism.</td>
</tr>
</tbody>
</table>

You can specify hyperparameter values when you fine-tune your model in Studio. For more information, see [Fine-tune foundation models in Studio](#).

You can also override default hyperparameter values when fine-tuning your model using the SageMaker Python SDK. For more information, see [Fine-tune publicly available foundation models with the JumpStartEstimator class](#).

**Domain adaptation fine-tuning**

Domain adaptation fine-tuning allows you to leverage pre-trained foundation models and adapt them to specific tasks using limited domain-specific data. If prompt engineering efforts do not provide enough customization, you can use domain adaption fine-tuning to get your model working with domain-specific language, such as industry jargon, technical terms, or other specialized data. This fine-tuning process modifies the weights of the model.

Domain adaptation fine-tuning is available with the following foundation models:

- Bloom 3B
- Bloom 7B1
- BloomZ 3B FP16
- BloomZ 7B1 FP16
- GPT-2 XL
- GPT-J 6B
- GPT-Neo 1.3B
- GPT-Neo 125M
- GPT-NEO 2.7B

---

**Note**

Some JumpStart foundation models, such as Llama 2 7B, require acceptance of an end-user license agreement before fine-tuning and performing inference. For more information, see [End-user license agreements](#).
Prepare and upload training data for domain adaptation fine-tuning

Training data for domain adaptation fine-tuning can be provided in CSV, JSON, or TXT file format. All training data must be in a single file within a single folder.

The training data is taken from the **Text** column for CSV or JSON training data files. If no column is labeled **Text**, then the training data is taken from the first column for CSV or JSON training data files.

The following is an example body of a TXT file to be used for fine-tuning:

```
This report includes estimates, projections, statements relating to our business plans, objectives, and expected operating results that are “forward-looking statements” within the meaning of the Private Securities Litigation Reform Act of 1995, Section 27A of the Securities Act of 1933, and Section 21E of ....
```

Split data for training and testing

You can optionally provide another folder containing validation data. This folder should also include one CSV, JSON, or TXT file. If no validation dataset is provided, then a set amount of the training data is set aside for validation purposes. You can adjust the percentage of training data used for validation when you choose the hyperparameters for fine-tuning your model.

Upload fine-tuning data to Amazon S3

Upload your prepared data to Amazon Simple Storage Service (Amazon S3) to use when fine-tuning a JumpStart foundation model. You can use the following commands to upload your data:

```
from sagemaker.s3 import S3Uploader
```
import sagemaker
import random

output_bucket = sagemaker.Session().default_bucket()
local_data_file = "train.txt"
train_data_location = f"s3://{output_bucket}/training_folder"
S3Uploader.upload(local_data_file, train_data_location)
S3Uploader.upload("template.json", train_data_location)
print(f"Training data: {train_data_location}"

Create a training job for instruction-based fine-tuning

After your data is uploaded to Amazon S3, you can fine-tune and deploy your JumpStart foundation model. To fine-tune your model in Studio, see Fine-tune foundation models in Studio. To fine-tune your model using the SageMaker Python SDK, see Fine-tune publicly available foundation models with the JumpStartEstimator class.

Example notebooks

For more information on domain adaptation fine-tuning, see the following example notebooks:

- SageMaker JumpStart Foundation Models - Fine-tuning text generation GPT-J 6B model on domain specific dataset
- Fine-tune LLaMA 2 models on SageMaker JumpStart

Instruction-based fine-tuning

Instruction-based fine-tuning uses labeled examples to improve the performance of a pre-trained foundation model on a specific task. The labeled examples are formatted as prompt, response pairs and phrased as instructions. This fine-tuning process modifies the weights of the model. For more information on instruction-based fine-tuning, see the papers Introducing FLAN: More generalizable Language Models with Instruction Fine-Tuning and Scaling Instruction-Finetuned Language Models.

Fine-tuned LAnguage Net (FLAN) models use instruction tuning to make models more amenable to solving general downstream NLP tasks. Amazon SageMaker JumpStart provides a number of foundation models in the FLAN model family. For example, FLAN-T5 models are instruction fine-tuned on a wide range of tasks to increase zero-shot performance for a variety of common use cases. With additional data and fine-tuning, instruction-based models can be further adapted to more specific tasks that weren’t considered during pre-training.
Models compatible with instruction-based fine-tuning

Only a subset of JumpStart foundation models are compatible with instruction-based fine-tuning. Instruction-based fine-tuning is available with the following foundation models:

- Flan-T5 Base
- Flan-T5 Large
- Flan-T5 Small
- Flan-T5 XL
- Flan-T5 XXL
- Llama 2 13B
- Llama 2 13B Chat
- Llama 2 13B Neuron
- Llama 2 70B
- Llama 2 70B Chat
- Llama 2 7B
- Llama 2 7B Chat
- Llama 2 7B Neuron
- Mistral 7B
- RedPajama INCITE Base 3B V1
- RedPajama INCITE Base 7B V1
- RedPajama INCITE Chat 3B V1
- RedPajama INCITE Chat 7B V1
- RedPajama INCITE Instruct 3B V1
- RedPajama INCITE Instruct 7B V1

Note

Some JumpStart foundation models, such as Llama 2 7B, require acceptance of an end-user license agreement before fine-tuning and performing inference. For more information, see End-user license agreements.
Prepare and upload training data for instruction-based fine-tuning

Training data for instruction-based fine-tuning must be provided in JSON Lines text file format, where each line is a dictionary. All training data must be in a single folder. The folder can include multiple .jsonl files.

The training folder can also include a template JSON file (template.json) that describes the input and output formats of your data. If no template file is provided, the following template file is used:

```json
{
    "prompt": "Below is an instruction that describes a task, paired with an input that provides further context. Write a response that appropriately completes the request.\n\n### Instruction:\n
### Input:\n
### completion": "{response}"
}
```

According to the `template.json` file, each .jsonl entry of the training data must include {instruction}, {context}, and {response} fields.

If you provide a custom template JSON file, use the "prompt" and "completion" keys to define your own required fields. According to the following custom template JSON file, each .jsonl entry of the training data must include {question}, {context}, and {answer} fields:

```json
{
    "prompt": "question: {question} context: {context}\n
### completion": "{answer}"
}
```

Split data for training and testing

You can optionally provide another folder containing validation data. This folder should also include one or more .jsonl files. If no validation dataset is provided, then a set amount of the training data is set aside for validation purposes. You can adjust the percentage of training data used for validation when you choose the hyperparameters for fine-tuning your model.

Upload fine-tuning data to Amazon S3

Upload your prepared data to Amazon Simple Storage Service (Amazon S3) to use when fine-tuning a JumpStart foundation model. You can use the following commands to upload your data:
from sagemaker.s3 import S3Uploader
import sagemaker
import random

output_bucket = sagemaker.Session().default_bucket()
local_data_file = "train.jsonl"
train_data_location = f"s3://{output_bucket}/dolly_dataset"
S3Uploader.upload(local_data_file, train_data_location)
S3Uploader.upload("template.json", train_data_location)
print(f"Training data: {train_data_location}"

Create a training job for instruction-based fine-tuning

After your data is uploaded to Amazon S3, you can fine-tune and deploy your JumpStart foundation model. To fine-tune your model in Studio, see Fine-tune foundation models in Studio. To fine-tune your model using the SageMaker Python SDK, see Fine-tune publicly available foundation models with the JumpStartEstimator class.

Example notebooks

For more information on instruction-based fine-tuning, see the following example notebooks:

- Fine-tune LLaMA 2 models on SageMaker JumpStart
- Introduction to SageMaker JumpStart - Text Generation with Mistral models
- Introduction to SageMaker JumpStart - Text Generation with Falcon models
- SageMaker JumpStart Foundation Models - HuggingFace Text2Text Instruction Fine-Tuning

Retrieval Augmented Generation (RAG)

Foundation models are usually trained offline, making the model agnostic to any data that is created after the model was trained. Additionally, foundation models are trained on very general domain corpora, making them less effective for domain-specific tasks. You can use Retrieval Augmented Generation (RAG) to retrieve data from outside a foundation model and augment your prompts by adding the relevant retrieved data in context. For more information about RAG model architectures, see Retrieval-Augmented Generation for Knowledge-Intensive NLP Tasks.

With RAG, the external data used to augment your prompts can come from multiple data sources, such as a document repositories, databases, or APIs. The first step is to convert your documents and any user queries into a compatible format to perform relevancy search. To make the formats
compatible, a document collection, or knowledge library, and user-submitted queries are converted to numerical representations using embedding language models. *Embedding* is the process by which text is given numerical representation in a vector space. RAG model architectures compare the embeddings of user queries within the vector of the knowledge library. The original user prompt is then appended with relevant context from similar documents within the knowledge library. This augmented prompt is then sent to the foundation model. You can update knowledge libraries and their relevant embeddings asynchronously.

The retrieved document should be large enough to contain useful context to help augment the prompt, but small enough to fit into the maximum sequence length of the prompt. You can use task-specific JumpStart models, such as the General Text Embeddings (GTE) model from Hugging Face, to provide the embeddings for your prompts and knowledge library documents. After comparing the prompt and document embeddings to find the most relevant documents, construct a new prompt with the supplemental context. Then, pass the augmented prompt to a text generation model of your choosing.

**Example notebooks**

For more information on RAG foundation model solutions, see the following example notebooks:
• Retrieval-Augmented Generation: Question Answering using LangChain and Cohere’s Generate and Embedding Models from SageMaker JumpStart

• Retrieval-Augmented Generation: Question Answering using Llama-2, Pinecone and Custom Dataset

• Retrieval-Augmented Generation: Question Answering based on Custom Dataset with Open-sourced LangChain Library

• Retrieval-Augmented Generation: Question Answering based on Custom Dataset

• Retrieval-Augmented Generation: Question Answering using Llama-2 and Text Embedding Models

• Amazon SageMaker JumpStart - Text Embedding and Sentence Similarity

You can clone the Amazon SageMaker examples repository to run the available JumpStart foundation model examples in the Jupyter environment of your choice within Studio. For more information on applications that you can use to create and access Jupyter in SageMaker, see Applications supported in Amazon SageMaker Studio.

Evaluate a text generation foundation model in Studio

Foundation Model Evaluations (FMEval) is in preview release for Amazon SageMaker Clarify and is subject to change.

⚠️ Important

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see Migrating from Amazon SageMaker Studio Classic. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

Amazon SageMaker JumpStart has integrations with SageMaker Clarify Foundation Model Evaluations (FMEval) in Studio. If a JumpStart model has built-in evaluation capabilities available, you can choose Evaluate in the upper right corner of the model detail page in the JumpStart
Studio UI. For more information on navigating the JumpStart Studio UI, see Open and use JumpStart in Studio,

Use Amazon SageMaker JumpStart to evaluate text-based foundation models with FMEval. You can use these model evaluations to compare model quality and responsibility metrics for one model, between two models, or between different versions of the same model, to help you quantify model risks. FMEval can evaluate text-based models that perform the following tasks:

- **Open-ended generation** – The production of natural human responses to text that does not have a pre-defined structure.
- **Text summarization** – The generation of a concise and condensed summary while retaining the meaning and key information contained in larger text.
- **Question Answering** – The generation of an answer in natural language to a question.
- **Classification** – The assignment of a class, such as positive versus negative to a text passage based on its content.

You can use FMEval to automatically evaluate model responses based on specific benchmarks. You can also evaluate model responses against your own criteria by bringing your own prompt datasets. FMEval provides a user interface (UI) that guides you through the setup and configuration of an evaluation job. You can also use the FMEval library inside your own code.

Every evaluation requires quota for two instances:

- **Hosting instance** – An instance that hosts and deploys an LLM.
- **Evaluation instance** – An instance that is used to prompt and perform an evaluation of an LLM on the hosting instance.

If your LLM is already deployed, provide the endpoint, and SageMaker will use your **hosting instance** to host and deploy the LLM.

If you are evaluating a SageMaker JumpStart model that is not yet deployed to your account, FMEval creates a temporary **hosting instance** for you in your account, and keeps it deployed only for the length of your evaluation. FMEval uses the default instance that SageMaker JumpStart recommends for the chosen LLM as your hosting instance. You must have sufficient quota for this recommended instance.

Every evaluation also uses an evaluation instance to provide prompts to and score the responses from the LLM. You must also have sufficient quota and memory to run the evaluation algorithms.
The quota and memory requirements of the evaluation instance are generally smaller than those required for a hosting instance. We recommend selecting the ml.m5.2xlarge instance. For more information about quota and memory, see FMEval Troubleshooting guide.

Automatic evaluations can be used to score LLMs across the following dimensions:

- **Accuracy** – For text summarization, question answering, and text classification
- **Semantic robustness** – For open-ended generation, text summarization and text classification tasks
- **Factual knowledge** – For open-ended generation
- **Prompt stereotyping** – For open-ended generation
- **Toxicity** – For open-ended generation, text summarization, and question answering

You can also use human evaluations to manually evaluate model responses. The FMEval UI guides you through a workflow of selecting one or more models, provisioning resources, and writing instructions for and contacting your human workforce. After the human evaluation is complete, the results are displayed in FMEval.

You can access model evaluation through the JumpStart landing page in Studio by selecting a model to evaluate and then choosing **Evaluate**. Note that not all JumpStart models have evaluation capabilities available. For more information about how to configure, provision and run FMEval, see What are Foundation Model Evaluations?

**Example notebooks**

For step-by-step examples on how to use publicly available JumpStart foundation models with the SageMaker Python SDK, refer to the following notebooks on text generation, image generation, and model customization.

**Tip**

Proprietary and publicly available JumpStart foundation models have different SageMaker Python SDK deployment workflows. Discover proprietary foundation model example notebooks through Amazon SageMaker Studio Classic or the SageMaker console. For more information, see How to use JumpStart foundation models.
You can clone the Amazon SageMaker examples repository to run the available JumpStart foundation model examples in the Jupyter environment of your choice within Studio. For more information on applications that you can use to create and access Jupyter in SageMaker, see Applications supported in Amazon SageMaker Studio.

Text generation

Explore text generation example notebooks, including guidance on general text generation workflows, multilingual text classification, real-time batch inference, few-shot learning, chatbot interactions, and more.

- SageMaker JumpStart Foundation Models - HuggingFace Text2Text Generation with FLAN-T5 XL as an example
- SageMaker JumpStart Foundation Models - BloomZ: Multilingual Text Classification, Question and Answering, Code Generation, Paragraph rephrase, and More
- SageMaker JumpStart Foundation Models - HuggingFace Text2Text Generation Batch Transform and Real-Time Batch Inference
- SageMaker JumpStart Foundation Models - GPT-J, GPT-Neo Few-shot learning
- SageMaker JumpStart Foundation Models - Chatbots
- Introduction to SageMaker JumpStart - Text Generation with Mistral models
- Introduction to SageMaker JumpStart - Text Generation with Falcon models

Image generation

Get started with text-to-image Stable Diffusion models, learn how to deploy an inpainting model, and experiment with a simple workflow to generate images of your dog.

- Introduction to JumpStart - Text to Image
- Introduction to JumpStart Image editing - Stable Diffusion Inpainting
- Generate fun images of your dog

Model customization

Sometimes your use case requires greater foundation model customization for specific tasks. For more information on model customization approaches, see Customize a foundation model or explore one of the following example notebooks.
• SageMaker JumpStart Foundation Models - Fine-tuning text generation GPT-J 6B model on domain specific dataset
• SageMaker JumpStart Foundation Models - HuggingFace Text2Text Instruction Fine-Tuning
• Retrieval-Augmented Generation: Question Answering using LangChain and Cohere's Generate and Embedding Models from SageMaker JumpStart
• Retrieval-Augmented Generation: Question Answering using LLama-2, Pinecone and Custom Dataset
• Retrieval-Augmented Generation: Question Answering based on Custom Dataset with Open-sourced LangChain Library
• Retrieval-Augmented Generation: Question Answering based on Custom Dataset
• Retrieval-Augmented Generation: Question Answering using Llama-2 and Text Embedding Models
• Amazon SageMaker JumpStart - Text Embedding and Sentence Similarity

Task-Specific Models

JumpStart supports task-specific models across fifteen of the most popular problem types. Of the supported problem types, Vision and NLP-related types total thirteen. There are eight problem types that support incremental training and fine-tuning. For more information about incremental training and hyper-parameter tuning, see SageMaker Automatic Model Tuning. JumpStart also supports four popular algorithms for tabular data modeling.

You can search and browse models from the JumpStart landing page in Studio or Studio Classic. When you select a model, the model detail page provides information about the model, and you can train and deploy your model in a few steps. The description section describes what you can do with the model, the expected types of inputs and outputs, and the data type needed for fine-tuning your model.

You can also programmatically utilize models with the SageMaker Python SDK. For a list of all available models, see the JumpStart Available Model Table.

The list of problem types and links to their example Jupyter notebooks are summarized in the following table.
<table>
<thead>
<tr>
<th>Problem types</th>
<th>Supports inference with pre-trained models</th>
<th>Trainable on a custom dataset</th>
<th>Supported frameworks</th>
<th>Example Notebooks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image classification</td>
<td>Yes</td>
<td>Yes</td>
<td>PyTorch, TensorFlow</td>
<td>[Introduction to JumpStart - Image Classification]</td>
</tr>
<tr>
<td>Object detection</td>
<td>Yes</td>
<td>Yes</td>
<td>PyTorch, TensorFlow, MXNet</td>
<td>[Introduction to JumpStart - Object Detection]</td>
</tr>
<tr>
<td>Semantic segmentation</td>
<td>Yes</td>
<td>Yes</td>
<td>MXNet</td>
<td>[Introduction to JumpStart - Semantic Segmentation]</td>
</tr>
<tr>
<td>Instance segmentation</td>
<td>Yes</td>
<td>Yes</td>
<td>MXNet</td>
<td>[Introduction to JumpStart - Instance Segmentation]</td>
</tr>
<tr>
<td>Image embedding</td>
<td>Yes</td>
<td>No</td>
<td>TensorFlow, MXNet</td>
<td>[Introduction to JumpStart - Image Embedding]</td>
</tr>
<tr>
<td>Text classification</td>
<td>Yes</td>
<td>Yes</td>
<td>TensorFlow</td>
<td>[Introduction to JumpStart - Text Classification]</td>
</tr>
<tr>
<td>Sentence pair classification</td>
<td>Yes</td>
<td>Yes</td>
<td>TensorFlow, Hugging Face</td>
<td>[Introduction to JumpStart - Sentence Pair Classification]</td>
</tr>
<tr>
<td>Problem types</td>
<td>Supports inference with pre-trained models</td>
<td>Trainable on a custom dataset</td>
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<td>-------------------------------</td>
<td>--------------------------------------------</td>
<td>------------------------------</td>
<td>----------------------</td>
<td>--------------------------------------------------------</td>
</tr>
<tr>
<td>Question answering</td>
<td>Yes</td>
<td>Yes</td>
<td>PyTorch, Hugging Face</td>
<td>[Introduction to JumpStart - Question Answering]</td>
</tr>
<tr>
<td>Named entity recognition</td>
<td>Yes</td>
<td>No</td>
<td>Hugging Face</td>
<td>[Introduction to JumpStart - Named Entity Recognition]</td>
</tr>
<tr>
<td>Text summarization</td>
<td>Yes</td>
<td>No</td>
<td>Hugging Face</td>
<td>[Introduction to JumpStart - Text Summarization]</td>
</tr>
<tr>
<td>Text generation</td>
<td>Yes</td>
<td>No</td>
<td>Hugging Face</td>
<td>[Introduction to JumpStart - Text Generation]</td>
</tr>
<tr>
<td>Machine translation</td>
<td>Yes</td>
<td>No</td>
<td>Hugging Face</td>
<td>[Introduction to JumpStart - Machine Translation]</td>
</tr>
<tr>
<td>Text embedding</td>
<td>Yes</td>
<td>No</td>
<td>TensorFlow, MXNet</td>
<td>[Introduction to JumpStart - Text Embedding]</td>
</tr>
<tr>
<td>Problem types</td>
<td>Supports inference with pre-trained models</td>
<td>Trainable on a custom dataset</td>
<td>Supported frameworks</td>
<td>Example Notebooks</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------------------------------</td>
<td>-------------------------------</td>
<td>----------------------------------------------------------</td>
<td>--------------------------------------------------------</td>
</tr>
</tbody>
</table>
| Tabular classification | Yes                                         | Yes                           | LightGBM, CatBoost, XGBoost, AutoGluon - Tabular, TabTransformer, Linear Learner | [Introduction to JumpStart - Tabular Classification - LightGBM, CatBoost](#)  
[Introduction to JumpStart - Tabular Classification - XGBoost, Linear Learner](#)  
[Introduction to JumpStart - Tabular Classification - AutoGluon Learner](#)  
[Introduction to JumpStart - Tabular Classification - TabTransformer Learner](#) |
<table>
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</tr>
</thead>
<tbody>
<tr>
<td>Tabular regression</td>
<td>Yes</td>
<td>Yes</td>
<td>LightGBM, CatBoost, XGBoost, AutoGluon -Tabular, TabTransformer, Linear Learner</td>
<td>Introduction to JumpStart - Tabular Regression - LightGBM, CatBoost</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Introduction to JumpStart - Tabular Regression - XGBoost, Linear Learner</td>
<td>Introduction to JumpStart - Tabular Regression - AutoGluon Learner</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Introduction to JumpStart - Tabular Regression - TabTransformer Learner</td>
<td>Introduction to JumpStart - Tabular Regression - TabTransformer Learner</td>
</tr>
</tbody>
</table>
Deploy a Model

When you deploy a model from JumpStart, SageMaker hosts the model and deploys an endpoint that you can use for inference. JumpStart also provides an example notebook that you can use to access the model after it's deployed.

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

ℹ️ Note

For more information on JumpStart model deployment in Studio, see Deploy foundation models in Studio.

Model deployment configuration

After you choose a model, the model's tab opens. In the Deploy Model pane, choose Deployment Configuration to configure your model deployment.

The default instance type for deploying a model depends on the model. The instance type is the hardware that the training job runs on. In the following example, the ml.p2.xlarge instance is the default for this particular BERT model.
You can also change the endpoint name, add key:value resource tags, activate or deactivate the jumpstart- prefix for any JumpStart resources related to the model, and specify an Amazon S3 bucket for storing model artifacts used by your SageMaker endpoint.

Choose **Security Settings** to specify the AWS Identity and Access Management (IAM) role, Amazon Virtual Private Cloud (Amazon VPC), and encryption keys for the model.
Model deployment security

When you deploy a model with JumpStart, you can specify an IAM role, Amazon VPC, and encryption keys for the model. If you don't specify any values for these entries: The default IAM role is your Studio Classic runtime role; default encryption is used; no Amazon VPC is used.

IAM role

You can select an IAM role that is passed as part of training jobs and hosting jobs. SageMaker uses this role to access training data and model artifacts. If you don't select an IAM role, SageMaker deploys the model using your Studio Classic runtime role. For more information about IAM roles, see [Identity and Access Management for Amazon SageMaker](#).

The role that you pass must have access to the resources that the model needs, and must include all of the following.

- For training jobs: [CreateTrainingJob API: Execution Role Permissions](#).
- For hosting jobs: [CreateModel API: Execution Role Permissions](#).
**Note**

You can scope down the Amazon S3 permissions granted in each of the following roles. Do this by using the ARN of your Amazon Simple Storage Service (Amazon S3) bucket and the JumpStart Amazon S3 bucket.

```
{
    "Effect": "Allow",
    "Action": [
        "s3:GetObject",
        "s3:PutObject",
        "s3:ListMultipartUploadParts",
        "s3:ListBucket"
    ],
    "Resources": [
        "arn:aws:s3:::jumpstart-cache-prod-<region>/*",
        "arn:aws:s3:::jumpstart-cache-prod-<region>",
        "arn:aws:s3:::bucket/*"
    ]
}
```

**Find IAM role**

If you select this option, you must select an existing IAM role from the dropdown list.

**Specify the IAM role that Amazon SageMaker should use to deploy your model. Learn more.**

- Default IAM role
- Find IAM role
- Input IAM role

Amazon SageMaker will deploy your model using the IAM role you select below.

**Execution role**

**Input IAM role**
If you select this option, you must manually enter the ARN for an existing IAM role. If your Studio Classic runtime role or Amazon VPC block the `iam:list*` call, you must use this option to use an existing IAM role.

**Specify the IAM role that Amazon SageMaker should use to deploy your model. Learn more.**

- Default IAM role
- Find IAM role
- Input IAM role

Amazon SageMaker will deploy your model using the IAM role you type below.

**Execution role arn**

```
arn:aws:iam::account-id:role/role-name
```

---

**Amazon VPC**

All JumpStart models run in network isolation mode. After the model container is created, no more calls can be made. You can select an Amazon VPC that is passed as part of training jobs and hosting jobs. SageMaker uses this Amazon VPC to push and pull resources from your Amazon S3 bucket. This Amazon VPC is different from the Amazon VPC that limits access to the public internet from your Studio Classic instance. For more information about the Studio Classic Amazon VPC, see [Connect SageMaker Studio Notebooks in a VPC to External Resources](#).

The Amazon VPC that you pass does not need access to the public internet, but it does need access to Amazon S3. The Amazon VPC endpoint for Amazon S3 must allow access to at least the following resources that the model needs.

```json
{
  "Effect": "Allow",
  "Action": [
    "s3:GetObject",
    "s3:PutObject",
    "s3:ListMultipartUploadParts",
    "s3:ListBucket"
  ],
  "Resources": [
    "arn:aws:s3:::jumpstart-cache-prod-<region>/*",
    "arn:aws:s3:::jumpstart-cache-prod-<region>",
    "arn:aws:s3:::bucket/*"
  ]
}
```
If you do not select an Amazon VPC, no Amazon VPC is used.

**Find VPC**

If you select this option, you must select an existing Amazon VPC from the dropdown list. After you select an Amazon VPC, you must select a subnet and security group for your Amazon VPC. For more information about subnets and security groups, see [Overview of VPCs and subnets](#).

**Specify whether your model should connect to a virtual private cloud (VPC).** Learn more.

- No VPC
- Find VPC
- Input VPC

The VPC you select below will control access to and from your model container.

**VPC ID**

Select...

**Input VPC**

If you select this option, you must manually select the subnet and security group that compose your Amazon VPC. If your Studio Classic runtime role or Amazon VPC blocks the `ec2: list*` call, you must use this option to select the subnet and security group.

**Specify whether your model should connect to a virtual private cloud (VPC).** Learn more.

- No VPC
- Find VPC
- Input VPC

The subnets and security groups you type below will control access to and from your model container.

**Subnet(s)**

Type subnet ID

Add

**Security group(s)**

Type security group ID

Add
Encryption keys

You can select an AWS KMS key that is passed as part of training jobs and hosting jobs. SageMaker uses this key to encrypt the Amazon EBS volume for the container, and the repackaged model in Amazon S3 for hosting jobs and the output for training jobs. For more information about AWS KMS keys, see [AWS KMS keys](#).

The key that you pass must trust the IAM role that you pass. If you do not specify an IAM role, the AWS KMS key must trust your Studio Classic runtime role.

If you do not select an AWS KMS key, SageMaker provides default encryption for the data in the Amazon EBS volume and the Amazon S3 artifacts.

Find encryption keys

If you select this option, you must select existing AWS KMS keys from the dropdown list.

![Specify the encryption keys to secure your data. Learn more.](image)

- Default encryption keys
- Find encryption keys
- Input encryption keys

Encrypt your data in the storage volume attached to your ML compute instance and at rest in S3.

**Volume encryption key**

Select...

**Model encryption key**

Select...

Input encryption keys

If you select this option, you must manually enter the AWS KMS keys. If your Studio Classic execution role or Amazon VPC block the `kms: list*` call, you must use this option to select existing AWS KMS keys.
Configure default values for JumpStart models

You can configure default values for parameters such as IAM roles, VPCs, and KMS keys to pre-populate for JumpStart model deployment and training. After configuring default values, the Studio Classic UI automatically provides your specified security settings and tags to SageMaker JumpStart models to simplify deployment and training workflows. Administrators and end-users can initialize default values specified in a configuration file in YAML format.

By default, the SageMaker Python SDK uses two configuration files: one for the administrator and one for the user. Using the administrator configuration file, administrators can define a set of default values. End-users can override values set in the administrator configuration file and set additional default values using the end-user configuration file. For more information, see Default configuration file location.

The following code sample lists the default locations of the configuration files when using the SageMaker Python SDK in Amazon SageMaker Studio Classic.

```
# Location of the admin config file
/etc/xdg/sagemaker/config.yaml

# Location of the user config file
/root/.config/sagemaker/config.yaml
```

Values specified in the user configuration file override values set in the administrator configuration file. The configuration file is unique to each user profile within an Amazon SageMaker domain.
The user profile's Studio Classic application is directly associated with the user profile. For more information, see Domain user profiles.

Administrators can optionally set configuration defaults for JumpStart model training and deployment through JupyterServer lifecycle configurations. For more information, see Create and associate a lifecycle configuration.

**Default value configuration YAML file**

Your configuration file should adhere to the SageMaker Python SDK configuration file structure. Note that specific fields in the TrainingJob, Model, and EndpointConfig configurations apply to JumpStart model training and deployment default values.

```yaml
SchemaVersion: '1.0'
SageMaker:
  TrainingJob:
    OutputDataConfig:
      KmsKeyId: example-key-id
    ResourceConfig:
      # Training configuration - Volume encryption key
      VolumeKmsKeyId: example-key-id
      # Training configuration form - IAM role
      RoleArn: arn:aws:iam::123456789012:role/SageMakerExecutionRole
    VpcConfig:
      # Training configuration - Security groups
      SecurityGroupIds:
      - sg-1
      - sg-2
      # Training configuration - Subnets
      Subnets:
      - subnet-1
      - subnet-2
      # Training configuration - Custom resource tags
      Tags:
      - Key: Example-key
        Value: Example-value
    Model:
      EnableNetworkIsolation: true
      # Deployment configuration - IAM role
      ExecutionRoleArn: arn:aws:iam::123456789012:role/SageMakerExecutionRole
    VpcConfig:
      # Deployment configuration - Security groups
      SecurityGroupIds:
```

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- *sg-1*
- *sg-2*

# Deployment configuration - Subnets
Subnets:
- *subnet-1*
- *subnet-2*

EndpointConfig:
  AsyncInferenceConfig:
  OutputConfig:
    KmsKeyId: *example-key-id*

DataCaptureConfig:
  # Deployment configuration - Volume encryption key
  KmsKeyId: *example-key-id*

KmsKeyId: *example-key-id*

# Deployment configuration - Custom resource tags
Tags:
- Key: *Example-key*
  Value: *Example-value*

---

**Fine-Tune a Model**

Fine-tuning trains a pretrained model on a new dataset without training from scratch. This process, also known as transfer learning, can produce accurate models with smaller datasets and less training time. You can fine-tune a model if its card shows a **fine-tunable** attribute set to **Yes**.
Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Note
For more information on JumpStart model fine-tuning in Studio, see Fine-tune foundation models in Studio.

Fine-Tuning data source

When you fine-tune a model, you can use the default dataset or choose your own data, which is located in an Amazon S3 bucket.

To browse the buckets available to you, choose Find S3 bucket. These buckets are limited by the permissions used to set up your Studio Classic account. You can also specify an Amazon S3 URI by choosing Enter Amazon S3 bucket location.
Tip

To find out how to format the data in your bucket, choose Learn more. The description section for the model has detailed information about inputs and outputs.

For text models:

- The bucket must have a data.csv file.
- The first column must be a unique integer for the class label. For example: 1, 2, 3, 4, n
- The second column must be a string.
- The second column should have the corresponding text that matches the type and language for the model.

For vision models:

- The bucket must have as many subdirectories as the number of classes.
- Each subdirectory should contain images that belong to that class in .jpg format.
Note
The Amazon S3 bucket must be in the same AWS Region where you're running SageMaker Studio Classic because SageMaker doesn't allow cross-Region requests.

Fine-Tuning deployment configuration

The p3 family is recommended as the fastest for deep learning training, and this is recommended for fine-tuning a model. The following chart shows the number of GPUs in each instance type. There are other available options that you can choose from, including p2 and g4 instance types.

<table>
<thead>
<tr>
<th>Instance type</th>
<th>GPUs</th>
</tr>
</thead>
<tbody>
<tr>
<td>p3.2xlarge</td>
<td>1</td>
</tr>
<tr>
<td>p3.8xlarge</td>
<td>4</td>
</tr>
<tr>
<td>p3.16xlarge</td>
<td>8</td>
</tr>
<tr>
<td>p3dn.24xlarge</td>
<td>8</td>
</tr>
</tbody>
</table>

Hyperparameters

You can customize the hyperparameters of the training job that are used to fine-tune the model. The hyperparameters available for each fine-tunable model differ depending on the model. For information on each available hyperparameter, reference the hyperparameters documentation for the model of your choosing in Use Amazon SageMaker Built-in Algorithms or Pre-trained Models. For example, see Image Classification - TensorFlow Hyperparameters for details on the fine-tunable Image Classification - TensorFlow hyperparameters.

If you use the default dataset for text models without changing the hyperparameters, you get a nearly identical model as a result. For vision models, the default dataset is different from the dataset used to train the pretrained models, so your model is different as a result.

The following hyperparameters are common among models:
• **Epochs** – One epoch is one cycle through the entire dataset. Multiple intervals complete a batch, and multiple batches eventually complete an epoch. Multiple epochs are run until the accuracy of the model reaches an acceptable level, or when the error rate drops below an acceptable level.

• **Learning rate** – The amount that values should be changed between epochs. As the model is refined, its internal weights are being nudged and error rates are checked to see if the model improves. A typical learning rate is 0.1 or 0.01, where 0.01 is a much smaller adjustment and could cause the training to take a long time to converge, whereas 0.1 is much larger and can cause the training to overshoot. It is one of the primary hyperparameters that you might adjust for training your model. Note that for text models, a much smaller learning rate (5e-5 for BERT) can result in a more accurate model.

• **Batch size** – The number of records from the dataset that is to be selected for each interval to send to the GPUs for training.

In an image example, you might send out 32 images per GPU, so 32 would be your batch size. If you choose an instance type with more than one GPU, the batch is divided by the number of GPUs. Suggested batch size varies depending on the data and the model that you are using. For example, how you optimize for image data differs from how you handle language data.

In the instance type chart in the deployment configuration section, you can see the number of GPUs per instance type. Start with a standard recommended batch size (for example, 32 for a vision model). Then, multiply this by the number of GPUs in the instance type that you selected. For example, if you're using a p3.8xlarge, this would be 32(batch size) multiplied by 4 (GPUs), for a total of 128, as your batch size adjusts for the number of GPUs. For a text model like BERT, try starting with a batch size of 64, and then reduce as needed.

**Training output**

When the fine-tuning process is complete, JumpStart provides information about the model: parent model, training job name, training job ARN, training time, and output path. The output path is where you can find your new model in an Amazon S3 bucket. The folder structure uses the model name that you provided and the model file is in an /output subfolder and it's always named model.tar.gz.

Example: s3://bucket/model-name/output/model.tar.gz
Configure default values for model training

You can configure default values for parameters such as IAM roles, VPCs, and KMS keys to pre-populate for JumpStart model deployment and training. For more information, see, Configure default values for JumpStart models.

Share Models

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You can share JumpStart models through the Studio Classic UI directly from the Launched JumpStart assets page using the following procedure:

1. Open Amazon SageMaker Studio Classic and choose Launched JumpStart assets in the JumpStart section of the lefthand navigation pane.
2. Select the Training jobs tab to view the list of your model training jobs.
3. Under the Training jobs list, select the training job that you want to share. This opens the training job details page. You cannot share more than one training job at a time.
4. In the header for the training job, choose Share, and select either Share to Canvas or Share with my organization.

For more information about how to share a model with a SageMaker Canvas user, see Bring Your Own Model Into Canvas.

💡 Note
Only tabular models can be shared to SageMaker Canvas. Trying to share a non-tabular model to SageMaker Canvas throws the error Unsupported Data Type.

For more information about sharing models with your organization, see Shared Models and Notebooks.
Shared Models and Notebooks

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Share your models and notebooks to centralize model artifacts, facilitate discoverability, and increase the reuse of models within your organization. When sharing your models, you can provide training and inference environment information and allow collaborators to use these environments for their own training and inference jobs.

All models that you share and models that are shared with you are searchable in a centralized location directly in Amazon SageMaker Studio Classic. For information on the onboarding steps to sign into Amazon SageMaker Studio Classic, see Onboard to Amazon SageMaker Domain.

Access shared models and notebooks

To access your shared content, choose Shared models in the left navigation pane of the Amazon SageMaker Studio Classic UI.

Add shared content

You can share models or notebooks through the Shared models section of the Studio Classic UI. For details about each step, see Share models and notebooks through the Studio Classic UI.

Filter shared content

There are three main options for filtering shared models and notebooks:

1. **Shared by me** – Models and notebooks that you shared to either JumpStart or SageMaker Canvas.
2. **Shared with me** – Models and notebooks shared with you
3. **Shared by my organization** – All models and notebooks that are shared to anyone in your organization
You can also sort your models and notebooks based on the time they were last updated or by ascending or descending alphabetical order. Choose the filter icon to further sort your selections.

**Share tabular models with SageMaker Canvas users**

In addition to sharing models with your organization, you can also share models with collaborators that use SageMaker Canvas. If you share models to SageMaker Canvas, your collaborators can import those models into SageMaker Canvas and use them to generate predictions.

⚠️ **Important**

Important: You can only share tabular models to SageMaker Canvas.

You can filter for models and notebooks shared to and from SageMaker Canvas by selecting the filter icon in the **Shared by me** or **Shared with me** tabs. For more information about how to share a model to SageMaker Canvas, see [Bring Your Own Model Into Canvas](#).

**Share models and notebooks through the Studio Classic UI**

To share models and notebooks, navigate to the **Shared models** section in Amazon SageMaker Studio Classic, choose **Shared by my organization**, and then
select the **Add** dropdown list. Choose to either add a model or add a notebook.

![Add dropdown list](image)

**Add a model**

To add a model, choose **Shared by my organization**, and then select **Add model** from the **Add** dropdown list. Enter the basic information for your model, and add any training or inference information you want to share with collaborators to train or deploy your model. After you enter all the necessary information, choose **Add model** in the lower right corner.

**Basic information**

First, add the basic descriptive information about your model. This information is used to improve the searchability of your model.

1. Add a title for this model. Adding a title automatically populates a unique identifier in the ID field based on the model title.
2. Add a description of the model.
3. Select a data type from the options: *text*, *vision*, *tabular*, or *audio*. 
4. Select a machine learning task from the list of available tasks, such as *image classification* or *text generation*.

5. Select a machine learning framework.

6. Add metadata information with keywords or phrases to use when searching for a model. Use commas to separate keywords. Any spaces are automatically replaced with commas.

### Enable training

When adding a model to share, you can optionally provide a training environment and allow collaborators in your organization to train the shared model.

#### Note

If you are adding a tabular model, you also need to specify a column format and target column to enable training. For more information, see [Amazon SageMaker Canvas](https://aws.amazon.com/sagemaker/canvas) in the *Amazon SageMaker Developer Guide*.

1. Add a container to use for model training. You can select a container used for an existing training job, bring your own container in Amazon ECR, or use an Amazon SageMaker Deep Learning Container.

2. Add environment variables.

3. Provide a training script location.

4. Provide a script mode entry point.

5. Provide an Amazon S3 URI for model artifacts generated during training.

6. Provide the Amazon S3 URI to the default training dataset.

7. Provide a model output path. The model output path should be the Amazon S3 URI path for any model artifacts generated from training. SageMaker saves the model artifacts as a single compressed TAR file in Amazon S3.

8. Provide a validation dataset to use for evaluating your model during training. Validation datasets must contain the same number of columns and the same feature headers as the training dataset.

9. Turn on network isolation. Network isolation isolates the model container so that no inbound or outbound network calls can be made to or from the model container.
10. Provide training channels through which SageMaker can access your data. For example, you might specify input channels named `train` or `test`. For each channel, specify a channel name and a URI to the location of your data. Choose **Browse** to search for Amazon S3 locations.

11. Provide hyperparameters. Add any hyperparameters with which collaborators should experiment during training. Provide a range of valid values for these hyperparameters. This range is used for training job hyperparameter validation. You can define ranges based on the datatype of the hyperparameter.

12. Select an instance type. We recommend a GPU instance with more memory for training with large batch sizes. For a comprehensive list of SageMaker training instances across AWS Regions, see the **On-Demand Pricing** table in Amazon SageMaker Pricing.

13. Provide metrics. Define metrics for a training job by specifying a name and a regular expression for each metric that your training monitors. Design the regular expressions to capture the values of metrics that your algorithm emits. For example, the metric `loss` might have the regular expression "Loss =(.*)\;".

**Enable deployment**

When adding a model to share, you can optionally provide an inference environment in which collaborators in your organization can deploy the shared model for inference.

1. Add a container to use for inference. You can bring your own container in Amazon ECR or use an Amazon SageMaker Deep Learning Container.

2. Provide the Amazon S3 URI to an inference script. Custom inference scripts run inside your chosen container. Your inference script should include a function for model loading, and optionally functions generating predictions, and input and output processing. For more information on creating inference scripts for the framework of your choice, see Frameworks in the SageMaker Python SDK documentation. For example, for TensorFlow, see How to implement the pre- and/or post-processing handler(s).

3. Provide an Amazon S3 URI for model artifacts. Model artifacts are the output that results from training a model, and typically consist of trained parameters, a model definition that describes how to compute inferences, and other metadata. If you trained your model in SageMaker, the model artifacts are saved as a single compressed TAR file in Amazon S3. If you trained your model outside SageMaker, you need to create this single compressed TAR file and save it in an Amazon S3 location.
4. Select an instance type. We recommend a GPU instance with more memory for training with large batch sizes. For a comprehensive list of SageMaker training instances across AWS Regions, see the On-Demand Pricing table in Amazon SageMaker Pricing.

Add a notebook

To add a notebook, choose Shared by my organization, and then select Add notebook from the the Add dropdown list. Enter the basic information for your notebook and provide an Amazon S3 URI for the location of that notebook.

Basic information

First, add the basic descriptive information about your notebook. This information is used to improve the searchability of your notebook.

1. Add a title for this notebook. Adding a title automatically populates a unique identifier in the ID field based on the notebook title.
2. Add a description of the notebook.
3. Select a data type from the options: text, vision, tabular, or audio.
4. Select an ML task from the list of available tasks, such as image classification or text generation.
5. Select an ML framework.
6. Add metadata information with keywords or phrases to use when searching for a notebook. Use commas to separate keywords. Any spaces are automatically replaced with commas.

Add notebook

Provide an Amazon S3 URI for the location of that notebook. You can choose Browse to search through your Amazon S3 buckets for your notebook file location. After you find your notebook, copy the Amazon S3 URI, choose Cancel, and then add the Amazon S3 URI to the Notebook Location field.

After you enter all the necessary information, choose Add notebook in the lower right corner.
Solution Templates

Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Note

JumpStart Solutions are only available in Studio Classic.

SageMaker JumpStart provides one-click, end-to-end solutions for many common machine learning use cases. Explore the following use cases for more information on available solution templates.

- Demand forecasting
- Credit rating prediction
- Fraud detection
- Computer vision
- Extract and analyze data from documents
- Predictive maintenance
- Churn prediction
- Personalized recommendations
- Reinforcement learning
- Healthcare and life sciences
- Financial pricing
- Causal inference

Choose the solution template that best fits your use case from the JumpStart landing page. When you choose a solution template, JumpStart opens a new tab showing a description of the solution and a Launch button. When you select Launch, JumpStart creates all of the resources that you
need to run the solution, including training and model hosting instances. For more information on launching a JumpStart solution, see the section called “Launch a Solution”.

After launching the solution, you can explore solution features and any generated artifacts in JumpStart. Use the Launched JumpStart assets menu to find your solution. In your solution’s tab, select Open Notebook to use provided notebooks and explore the solution’s features. When artifacts are generated during launch or after running the provided notebooks, they’re listed in the Generated Artifacts table. You can delete individual artifacts with the trash icon.

You can delete all of the solution’s resources by choosing Delete solution resources.

**Demand forecasting**

Demand forecasting uses historical time series data in order to make future estimations in relation to customer demand over a specific period and streamline the supply-demand decision-making process across businesses.

Demand forecasting use cases include predicting ticket sales in the transportation industry, stock prices, number of hospital visits, number of customer representatives to hire for multiple locations in the next month, product sales across multiple regions in the next quarter, cloud server usage for the next day for a video streaming service, electricity consumption for multiple regions over the next week, number of IoT devices and sensors such as energy consumption, and more.

Time series data is categorized as **univariate** and **multi-variate**. For example, the total electricity consumption for a single household is a univariate time series over a period of time. When multiple univariate time series are stacked on each other, it’s called a multi-variate time series. For example, the total electricity consumption of 10 different (but correlated) households in a single neighborhood make up a multi-variate time series dataset.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demand forecasting</td>
<td>Demand forecasting for multivariate time series data using three state-of-the-art time series forecasting algorithms: LSTNet, Prophet, and SageMaker DeepAR.</td>
<td>GitHub »</td>
</tr>
</tbody>
</table>

Solution Templates
Credit rating prediction

Use JumpStart's credit rating prediction solutions to predict corporate credit ratings or to explain credit prediction decisions made by machine learning models. Compared to traditional credit rating modeling methods, machine learning models can automate and improve the accuracy of credit prediction.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corporate credit rating prediction</td>
<td>Multimodal (long text and tabular) machine learning for quality credit predictions using AWS AutoGluon Tabular.</td>
<td>GitHub »</td>
</tr>
<tr>
<td>Graph-based credit scoring</td>
<td>Predict corporate credit ratings using tabular data and a corporate network by training a Graph Neural Network GraphSAGE and AWS AutoGluon Tabular model.</td>
<td>Find in Amazon SageMaker Studio Classic.</td>
</tr>
<tr>
<td>Explain credit decisions</td>
<td>Predict credit default in credit applications and provide explanations using LightGBM and SHAP (SHapley Additive exPlanations).</td>
<td>GitHub »</td>
</tr>
</tbody>
</table>

Fraud detection

Many businesses lose billions annually to fraud. Machine learning based fraud detection models can help systematically identify likely fraudulent activities from a tremendous amount of data. The following solutions use transaction and user identity datasets to identify fraudulent transactions.
<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Detect malicious users and transactions</td>
<td>Automatically detect potentially fraudulent activity in transactions using <strong>SageMaker XGBoost</strong> with the over-sampling technique <strong>Synthetic Minority Over-sampling</strong> (SMOTE).</td>
<td><a href="#">GitHub »</a></td>
</tr>
<tr>
<td>Fraud detection in financial transactions using deep graph library</td>
<td>Detect fraud in financial transactions by training a <strong>graph convolutional network</strong> with the <strong>deep graph library</strong> and a <strong>SageMaker XGBoost</strong> model.</td>
<td><a href="#">GitHub »</a></td>
</tr>
<tr>
<td>Financial payment classification</td>
<td>Classify financial payments based on transaction information using <strong>SageMaker XGBoost</strong>. Use this solution template as an intermediate step in fraud detection, personalization, or anomaly detection.</td>
<td>Find in Amazon SageMaker Studio Classic.</td>
</tr>
</tbody>
</table>

**Computer vision**

With the rise of business use cases such as autonomous vehicles, smart video surveillance, healthcare monitoring and various object counting tasks, fast and accurate object detection systems are rising in demand. These systems involve not only recognizing and classifying every object in an image, but localizing each one by drawing the appropriate bounding box around it. In the last decade, the rapid advances of deep learning techniques greatly accelerated the momentum of object detection.
<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visual product defect detection</td>
<td>Identify defective regions in product images either by training an object detection model from scratch or fine-tuning pretrained SageMaker models.</td>
<td>GitHub »</td>
</tr>
<tr>
<td>Handwriting recognition</td>
<td>Recognize handwritten text in images by training an object detection model and handwriting recognition model. Label your own data using SageMaker Ground Truth.</td>
<td>GitHub »</td>
</tr>
<tr>
<td>Object detection for bird species</td>
<td>Identify birds species in a scene using a SageMaker object detection model.</td>
<td>Find in Amazon SageMaker Studio Classic.</td>
</tr>
</tbody>
</table>

**Extract and analyze data from documents**

JumpStart provides solutions for you to uncover valuable insights and connections in business-critical documents. Use cases include text classification, document summarization, handwriting recognition, relationship extraction, question and answering, and filling in missing values in tabular records.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Privacy for sentiment classification</td>
<td>Anonymize text to better preserve user privacy in sentiment classification.</td>
<td>GitHub »</td>
</tr>
<tr>
<td>Document understanding</td>
<td>Document summarization, entity, and relationship extraction using the</td>
<td>GitHub »</td>
</tr>
</tbody>
</table>
**Amazon SageMaker**

**Solution name** | **Description** | **Get started**
---|---|---
transformers library in PyTorch. | | 
**Handwriting recognition** | Recognize handwritten text in images by training an object detection model and handwriting recognition model. Label your own data using SageMaker Ground Truth. | GitHub »
**Filling in missing values in tabular records** | Fill missing values in tabular records by training a SageMaker AutoPilot model. | GitHub »

**Predictive maintenance**

Predictive maintenance aims to optimize the balance between corrective and preventative maintenance by facilitating the timely replacement of components. The following solutions use sensor data from industrial assets to predict machine failures, unplanned downtime, and repair costs.

**Solution name** | **Description** | **Get started**
---|---|---
Predictive maintenance for vehicle fleets | Predict vehicle fleet failures using vehicle sensor and maintenance information with a convolutional neural network model. | GitHub »
Predictive maintenance for manufacturing | Predict the remaining useful life for each sensor by training a stacked Bidirectional LSTM neural network | GitHub »
<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Churn prediction with text</td>
<td>Predict churn using numerical, categorical, and textual features with BERT encoder and RandomForestClassifier.</td>
<td>GitHub »</td>
</tr>
<tr>
<td>Churn prediction for mobile phone customers</td>
<td>Identify unhappy mobile phone customers using SageMaker XGBoost.</td>
<td>Find in Amazon SageMaker Studio Classic.</td>
</tr>
</tbody>
</table>

**Churn prediction**

Customer churn, or rate of attrition, is a costly problem faced by a wide range of companies. In an effort to reduce churn, companies can identify customers that are likely to leave their service in order to focus their efforts on customer retention. Use a JumpStart churn prediction solution to analyze data sources such as user behavior and customer support chat logs to identify customers that are at a high risk of cancelling a subscription or service.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Entity resolution in identity graphs with deep graph library</td>
<td>Perform cross-device entity linking for online advertising by training a graph convolution.</td>
<td>GitHub »</td>
</tr>
</tbody>
</table>
### Solution Templates

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>onal network with deep graph library.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Purchase modeling</td>
<td>Predict whether a customer will make a purchase by training a SageMaker XGBoost model.</td>
<td><a href="#">GitHub »</a></td>
</tr>
<tr>
<td>Customized recommender system</td>
<td>Train and deploy a custom recommender system that generates movie suggestions for a customer based on past behavior using Neural Collaborative Filtering in SageMaker.</td>
<td>Find in Amazon SageMaker Studio Classic.</td>
</tr>
</tbody>
</table>

### Reinforcement learning

Reinforcement learning (RL) is a type of learning that is based on interaction with the environment. This type of learning is used by an agent that must learn behavior through trial-and-error interactions with a dynamic environment in which the goal is to maximize the long-term rewards that the agent receives as a result of its actions. Rewards are maximized by trading off exploring actions that have uncertain rewards with exploiting actions that have known rewards.

RL is well-suited for solving large, complex problems, such as supply chain management, HVAC systems, industrial robotics, game artificial intelligence, dialog systems, and autonomous vehicles.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reinforcement learning for Battlesnake AI competitions</td>
<td>Provide a reinforcement learning workflow for training and inference with the BattleSnake AI competitions.</td>
<td><a href="#">GitHub »</a></td>
</tr>
</tbody>
</table>
### Healthcare and life sciences

Clinicians and researchers can use JumpStart solutions to analyze medical imagery, genomic information, and clinical health records.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lung cancer survival prediction</td>
<td>Predict non-small cell lung cancer patient survival status with 3-dimensional lung computerized tomography (CT) scans, genomic data, and clinical health records using SageMaker XGBoost.</td>
<td>GitHub »</td>
</tr>
</tbody>
</table>

### Financial pricing

Many businesses dynamically adjust pricing on a regular basis in order to maximize their returns. Use the following JumpStart solutions for price optimization, dynamic pricing, option pricing, or portfolio optimization use cases.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Price optimization</td>
<td>Estimate price elasticity using Double Machine Learning (ML) for causal inference and the Prophet forecasting procedure. Use these Find in Amazon SageMaker Studio Classic.</td>
<td></td>
</tr>
</tbody>
</table>
### Causal inference

Researchers can use machine learning models such as Bayesian networks to represent causal dependencies and draw causal conclusions based on data. Use the following JumpStart solution to understand the causal relationship between Nitrogen-based fertilizer application and corn crop yields.

<table>
<thead>
<tr>
<th>Solution name</th>
<th>Description</th>
<th>Get started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crop yield counterfactuals</td>
<td>Generate a counterfactual analysis of corn response to nitrogen. This solution learns the crop phenology cycle in its entirety using multi-spectral satellite imagery and ground-level observations.</td>
<td>Find in Amazon SageMaker Studio Classic.</td>
</tr>
</tbody>
</table>

### Launch a Solution

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

⚠️ **Note**

JumpStart Solutions are only available in Studio Classic.
First, choose a solution through the SageMaker JumpStart landing page in the Amazon SageMaker Studio Classic UI. For information on the onboarding steps to sign in to Amazon SageMaker Studio Classic, see [Onboard to Amazon SageMaker domain](#). For details on getting to the SageMaker JumpStart landing page, see [Open and use JumpStart in Studio Classic](#).

After you choose a solution, a solution's tab opens showing a description of the solution and a Launch button. To launch a solution, select Launch in the Launch Solution section. JumpStart then creates all of the resources needed to run the solution. This includes training and model hosting instances.

### Advanced parameters

The solution that you choose may have advanced parameters that you can select. Choose Advanced Parameters to specify the AWS Identity and Access Management role for the solution.

Solutions are able to launch resources across 9 AWS services that interact with each other. For the solution to work as expected, newly created components from one service must be able to act on newly created components from another service. We recommend that you use the default IAM role to ensure that all needed permissions are added. For more information about IAM roles, see [Identity and Access Management for Amazon SageMaker](#).

#### Default IAM role

If you select this option, the default IAM roles that are required by this solution are used. Each solution requires different resources. The following list describes the default roles that are used for the solutions based on the service needed. For a description of the permissions required for each service, see [AWS Managed Policies for SageMaker projects and JumpStart](#).

- **API Gateway** – AmazonSageMakerServiceCatalogProductsApiGatewayRole
- **CloudFormation** – AmazonSageMakerServiceCatalogProductsCloudformationRole
- **CodeBuild** – AmazonSageMakerServiceCatalogProductsCodeBuildRole
- **CodePipeline** – AmazonSageMakerServiceCatalogProductsCodePipelineRole
- **Events** – AmazonSageMakerServiceCatalogProductsEventsRole
- **Firehose** – AmazonSageMakerServiceCatalogProductsFirehoseRole
- **Glue** – AmazonSageMakerServiceCatalogProductsGlueRole
- **Lambda** – AmazonSageMakerServiceCatalogProductsLambdaRole
- **SageMaker** – AmazonSageMakerServiceCatalogProductsExecutionRole
If you are using a new SageMaker domain with JumpStart project templates enabled, these roles are automatically created in your account.

If you are using an existing SageMaker domain, these roles may not exist in your account. If this is the case, you will receive the following error when launching the solution:

```
Unable to locate the updated roles required to launch this solution, a general role '/service-role/AmazonSageMakerServiceCatalogProductsUseRole' will be used. Please update your studio domain to generate these roles.
```

You can still launch a solution without the needed role, but the legacy default role AmazonSageMakerServiceCatalogProductsUseRole is used in place of the needed role. The legacy default role has trust relationships with all of the services that JumpStart solutions need to interact with. For the best security, we recommend that you update your domain to have the newly created default roles for each AWS service.

If you have already onboarded to a SageMaker domain, you can update your domain to generate the default roles using the following procedure.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Control Panel at the top left of the page.
3. From the domain page, choose the Settings icon (🌐) to edit the domain settings.
4. On General Settings choose Next.
5. Under SageMaker Projects and JumpStart, select Enable Amazon SageMaker project templates and Amazon SageMaker JumpStart for this account and Enable Amazon SageMaker project templates and Amazon SageMaker JumpStart for Studio Classic users, choose Next.
6. Select Submit.

You should be able to see the default roles listed in Projects - Amazon SageMaker project templates enabled for this account under the Apps - Studio tab.

**Find IAM role**

If you select this option, you must select an existing IAM role from the dropdown list for each of the required services. The selected role must have at least the minimum permissions required for
the corresponding service. For a description of the permissions required for each service, see AWS Managed Policies for SageMaker projects and JumpStart.

Input IAM role

If you select this option, you must manually enter the ARN for an existing IAM role. The selected role must have at least the minimum permissions required for the corresponding service. For a description of the permissions required for each service, see AWS Managed Policies for SageMaker projects and JumpStart.

Amazon SageMaker JumpStart Industry: Financial

Use SageMaker JumpStart Industry: Financial solutions, models, and example notebooks to learn about SageMaker features and capabilities through curated one-step solutions and example notebooks of industry-focused machine learning (ML) problems. The notebooks also walk through how to use the SageMaker JumpStart Industry Python SDK to enhance industry text data and fine-tune pretrained models.

Topics

- Amazon SageMaker JumpStart Industry Python SDK
- Amazon SageMaker JumpStart Industry: Financial Solution
- Amazon SageMaker JumpStart Industry: Financial Models
- Amazon SageMaker JumpStart Industry: Financial Example Notebooks
- Amazon SageMaker JumpStart Industry: Financial Blog Posts
- Amazon SageMaker JumpStart Industry: Financial Related Research
- Amazon SageMaker JumpStart Industry: Financial Additional Resources

Amazon SageMaker JumpStart Industry Python SDK

SageMaker Runtime JumpStart provides processing tools for curating industry datasets and fine-tuning pretrained models through its client library called SageMaker JumpStart Industry Python SDK. For detailed API documentation of the SDK, and to learn more about processing and enhancing industry text datasets for improving the performance of state-of-the-art models on SageMaker JumpStart, see the SageMaker JumpStart Industry Python SDK open source documentation.
Amazon SageMaker JumpStart Industry: Financial Solution

SageMaker JumpStart Industry: Financial provides the following solution notebooks:

- **Corporate Credit Rating Prediction**

  This SageMaker JumpStart Industry: Financial solution provides a template for a text-enhanced corporate credit rating model. It shows how to take a model based on numeric features (in this case, Altman's famous 5 financial ratios) combined with texts from SEC filings to achieve an improvement in the prediction of credit ratings. In addition to the 5 Altman ratios, you can add more variables as needed or set custom variables. This solution notebook shows how SageMaker JumpStart Industry Python SDK helps process Natural Language Processing (NLP) scoring of texts from SEC filings. Furthermore, the solution demonstrates how to train a model using the enhanced dataset to achieve a best-in-class model, deploy the model to a SageMaker endpoint for production, and receive improved predictions in real time.

- **Graph-Based Credit Scoring**

  Credit ratings are traditionally generated using models that use financial statement data and market data, which is tabular only (numeric and categorical). This solution constructs a network of firms using [SEC filings](https://www.sec.gov) and shows how to use the network of firm relationships with tabular data to generate accurate rating predictions. This solution demonstrates a methodology to use data on firm linkages to extend the traditionally tabular-based credit scoring models, which have been used by the ratings industry for decades, to the class of machine learning models on networks.

**Note**
The solution notebooks are for demonstration purposes only. They should not be relied on as financial or investment advice.

You can find these financial services solutions through the SageMaker JumpStart page in Studio Classic.

**Important**
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the SageMaker JumpStart Industry: Financial solution notebooks.
Note

The SageMaker JumpStart Industry: Financial solutions, model cards, and example notebooks are hosted and runnable only through SageMaker Studio Classic. Log in to the SageMaker console, and launch SageMaker Studio Classic. For more information about how to find the solution card, see the previous topic at SageMaker JumpStart.

Amazon SageMaker JumpStart Industry: Financial Models

SageMaker JumpStart Industry: Financial provides the following pretrained Robustly Optimized BERT approach (RoBERTa) models:

- Financial Text Embedding (RoBERTa-SEC-Base)
- RoBERTa-SEC-WIKI-Base
- RoBERTa-SEC-Large
- RoBERTa-SEC-WIKI-Large

The RoBERTa-SEC-Base and RoBERTa-SEC-Large models are the text embedding models based on GluonNLP's RoBERTa model and pretrained on S&P 500 SEC 10-K/10-Q reports of the decade of the 2010's (from 2010 to 2019). In addition to these, SageMaker JumpStart Industry: Financial provides two more RoBERTa variations, RoBERTa-SEC-WIKI-Base and RoBERTa-SEC-WIKI-Large, which are pretrained on the SEC filings and common texts of Wikipedia.

You can find these models in SageMaker JumpStart by navigating to the Text Models node, choosing Explore All Text Models, and then filtering for the ML Task Text Embedding. You can access any corresponding notebooks after selecting the model of your choice. The paired notebooks will walk you through how the pretrained models can be fine-tuned for specific classification tasks on multimodal datasets, which are enhanced by the SageMaker JumpStart Industry Python SDK.
Note
The model notebooks are for demonstration purposes only. They should not be relied on as financial or investment advice.

The following screenshot shows the pretrained model cards provided through the SageMaker JumpStart page on Studio Classic.

Note
The SageMaker JumpStart Industry: Financial solutions, model cards, and example notebooks are hosted and runnable only through SageMaker Studio Classic. Log in to the SageMaker console, and launch SageMaker Studio Classic. For more information about how to find the model cards, see the previous topic at SageMaker JumpStart.
Amazon SageMaker JumpStart Industry: Financial Example Notebooks

SageMaker JumpStart Industry: Financial provides the following example notebooks to demonstrate solutions to industry-focused ML problems:

- **Financial TabText Data Construction** – This example introduces how to use the SageMaker JumpStart Industry Python SDK for processing the SEC filings, such as text summarization and scoring texts based on NLP score types and their corresponding word lists. To preview the content of this notebook, see [Simple Construction of a Multimodal Dataset from SEC Filings and NLP Scores](#).

- **Multimodal ML on TabText Data** – This example shows how to merge different types of datasets into a single dataframe called TabText and perform multimodal ML. To preview the content of this notebook, see [Machine Learning on a TabText Dataframe – An Example Based on the Paycheck Protection Program](#).

- **Multi-category ML on SEC filings data** – This example shows how to train an AutoGluon NLP model over the multimodal (TabText) datasets curated from SEC filings for a multiclass classification task. [Classify SEC 10K/Q Filings to Industry Codes Based on the MDNA Text Column](#).

---

**Note**

The example notebooks are for demonstrative purposes only. They should not be relied on as financial or investment advice.

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**Note**

The SageMaker JumpStart Industry: Financial solutions, model cards, and example notebooks are hosted and runnable only through SageMaker Studio Classic. Log in to the [SageMaker console](#), and launch SageMaker Studio Classic. For more information about how to find the example notebooks, see the previous topic at [SageMaker JumpStart](#).

To preview the content of the example notebooks, see [Tutorials – Finance](#) in the [SageMaker JumpStart Industry Python SDK documentation](#).
Amazon SageMaker JumpStart Industry: Financial Blog Posts

For thorough applications of using SageMaker JumpStart Industry: Financial solutions, models, examples, and the SDK, see the following blog posts:

- Use pre-trained financial language models for transfer learning in Amazon SageMaker JumpStart
- Use SEC text for ratings classification using multimodal ML in Amazon SageMaker JumpStart
- Create a dashboard with SEC text for financial NLP in Amazon SageMaker JumpStart
- Build a corporate credit ratings classifier using graph machine learning in Amazon SageMaker JumpStart
- Domain-adaptation Fine-tuning of Foundation Models in Amazon SageMaker JumpStart on Financial data

Amazon SageMaker JumpStart Industry: Financial Related Research

For research related to SageMaker JumpStart Industry: Financial solutions, see the following papers:

- Context, Language Modeling, and Multimodal Data in Finance
- Multimodal Machine Learning for Credit Modeling
- On the Lack of Robust Interpretability of Neural Text Classifiers
- FinLex: An Effective Use of Word Embeddings for Financial Lexicon Generation

Amazon SageMaker JumpStart Industry: Financial Additional Resources

For additional documentation and tutorials, see the following resources:

- The SageMaker JumpStart Industry: Financial Python SDK
- SageMaker JumpStart Industry: Financial Python SDK Tutorials
- The SageMaker JumpStart Industry: Financial GitHub repository
- Getting started with Amazon SageMaker - Machine Learning Tutorials
Use machine learning environments offered by SageMaker

⚠️ Important

Amazon SageMaker Studio and Amazon SageMaker Studio Classic are two of the machine learning environments that you can use to interact with SageMaker.
If your domain was created after November 30, 2023, Studio is your default experience.
If your domain was created before November 30, 2023, Amazon SageMaker Studio Classic is your default experience. To use Studio if Amazon SageMaker Studio Classic is your default experience, see Migrating from Amazon SageMaker Studio Classic.
When you migrate from Amazon SageMaker Studio Classic to Amazon SageMaker Studio, there is no loss in feature availability. Studio Classic also exists as an IDE within Amazon SageMaker Studio to help you run your legacy machine learning workflows.

SageMaker supports the following machine learning environments:

- **Amazon SageMaker Studio** (Recommended): The latest web-based experience for running ML workflows with a suite of IDEs. Studio supports the following applications:
  - Amazon SageMaker Studio Classic
  - Code Editor, based on Code-OSS, Visual Studio Code - Open Source
  - JupyterLab
  - Amazon SageMaker Canvas
  - RStudio
- **Amazon SageMaker Studio Classic**: Lets you build, train, debug, deploy, and monitor your machine learning models.
- **Amazon SageMaker Notebook Instances**: Lets you prepare and process data, and train and deploy machine learning models from a compute instance running the Jupyter Notebook application.
- **Amazon SageMaker Studio Lab**: Studio Lab is a free service that gives you access to AWS compute resources, in an environment based on open-source JupyterLab, without requiring an AWS account.
- **Amazon SageMaker Canvas**: Gives you the ability to use machine learning to generate predictions without needing to code.
• **Amazon SageMaker geospatial**: Gives you the ability to build, train, and deploy geospatial models.

• **RStudio on Amazon SageMaker**: RStudio is an IDE for R, with a console, syntax-highlighting editor that supports direct code execution, and tools for plotting, history, debugging and workspace management.

• **SageMaker HyperPod**: SageMaker HyperPod lets you provision resilient clusters for running machine learning (ML) workloads and developing state-of-the-art models such as large language models (LLMs), diffusion models, and foundation models (FMs).

To use these machine learning environments, except Studio Lab, SageMaker Notebook Instances, and SageMaker HyperPod, you or your organization's administrator must create an Amazon SageMaker domain. Studio Lab has a separate onboarding process.

**Topics**

- [Amazon SageMaker Studio](#)
- [Amazon SageMaker Studio Classic](#)
- [SageMaker JupyterLab](#)
- [Amazon SageMaker Notebook Instances](#)
- [Amazon SageMaker Studio Lab](#)
- [Amazon SageMaker Canvas](#)
- [Amazon SageMaker geospatial capabilities](#)
- [RStudio on Amazon SageMaker](#)
- [Get started with Code Editor in Amazon SageMaker Studio](#)
- [SageMaker HyperPod](#)
- [Use generative AI in SageMaker notebook environments](#)

**Amazon SageMaker Studio**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).
Amazon SageMaker Studio is the latest web-based experience for running ML workflows. Studio offers a suite of integrated development environments (IDEs). These include Code Editor, based on Code-OSS, Visual Studio Code - Open Source, a new JupyterLab application, RStudio, and Amazon SageMaker Studio Classic. For more information, see Applications supported in Amazon SageMaker Studio.

The new web-based UI in Studio is faster and provides access to all SageMaker resources, including jobs and endpoints, in one interface. ML practitioners can also choose their preferred IDE to accelerate ML development. A data scientist can use JupyterLab to explore data and tune models. In addition, a machine learning operations (MLOps) engineer can use Code Editor with the pipelines tool in Studio to deploy and monitor models in production.

The previous Studio experience is still being supported as Amazon SageMaker Studio Classic. Studio Classic is the default experience for existing customers, and is available as an application in Studio. For more information about Studio Classic, see Amazon SageMaker Studio Classic. For information about how to migrate from Studio Classic to Studio, see Migrating from Amazon SageMaker Studio Classic.

Studio offers the following benefits:

- A new JupyterLab application that has a faster start-up time and is more reliable than the existing Studio Classic application. For more information, see SageMaker JupyterLab.

- A suite of IDEs that open in a separate tab, including the new Code Editor, based on Code-OSS, Visual Studio Code - Open Source application. Users can interact with supported IDEs in a full screen experience. For more information, see Applications supported in Amazon SageMaker Studio.

- Access to all of your SageMaker resources in one place. Studio displays running instances across all of your applications.

- Access to all training jobs in a single view, regardless of whether they were scheduled from notebooks or initiated from Amazon SageMaker JumpStart.

- Simplified model deployment workflows and endpoint management and monitoring directly from Studio. You don't need to access the SageMaker console.

- Automatic creation of all configured applications when you onboard to a domain. For information about onboarding to a domain, see Amazon SageMaker domain overview.

- An improved SageMaker JumpStart experience where you can discover, import, register, fine tune, and deploy a foundation model. For more information, see SageMaker JumpStart.
Migrating from Amazon SageMaker Studio Classic

Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

When you open Studio, the web UI that starts up is based on the chosen default experience. Amazon SageMaker currently supports two different default experiences: the Amazon SageMaker Studio experience and the Amazon SageMaker Studio Classic experience.

- If Studio is your default experience, the UI is similar to the images found in Amazon SageMaker Studio UI overview.
- If Studio Classic is your default experience, the UI is similar to the images found in Amazon SageMaker Studio Classic UI Overview.
When you migrate your default experience from Studio Classic to Studio, you won't lose any features, and you will still access the Studio Classic IDE within Studio. For information about the added benefits of the Studio experience, see Amazon SageMaker Studio.

Note

- For existing customers that created their accounts before November 30, 2023, Amazon SageMaker Studio Classic may be the default experience. You can enable the Amazon SageMaker Studio as your default experience using the AWS Command Line Interface (AWS CLI) or the Amazon SageMaker console. For more information about Studio Classic, see Amazon SageMaker Studio Classic.

- For customers that create their accounts after November 30, 2023, we recommend using Studio as the default experience. This is because Studio contains various integrated development environments (IDEs), including the Studio Classic IDE, and other new features.

The following page outlines the major steps needed when enabling Studio as your default experience, from Studio Classic, for your domain. At the end includes instructions on how to revert your default experience to Studio Classic.

Topics

- Prerequisites
- Set Studio as the default experience for your domain
- Limit access to default applications in Studio
- Update your CORS policy to access Amazon S3 buckets
- Migrate workflows from Studio Classic to JupyterLab
- Migrate from Data Wrangler in Studio Classic to SageMaker Canvas
- Migrate from Autopilot in Studio Classic to SageMaker Canvas
- Revert to Studio Classic experience

Prerequisites

Migration of your default experience from Studio Classic to Studio is managed by the administrator of the domain. If you don't have permissions to set Studio as the default experience for your
domain, contact your administrator. To migrate your default experience, you must have administrator permissions or at least have permissions to update the domain, AWS Identity and Access Management (IAM), and Amazon Simple Storage Service (Amazon S3).

Before you use the AWS CLI to enable Studio as your default experience, follow these prerequisites:

- Install or update the AWS CLI by following the steps in Install or update the latest version of the AWS CLI.
- From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.

**Set Studio as the default experience for your domain**

You can make Studio the default experience when updating a domain from the SageMaker console, the AWS CLI, or AWS CloudFormation. Choose one of the following tabs to view the relevant instructions.

**Note**

The release of Studio includes updates to the AWS managed policies. For more information, see SageMaker Updates to AWS Managed Policies.

When Studio is the default experience for the domain, Studio is the default experience for all users in the domain. However, the user settings take precedence over the domain settings. Therefore, if a user has their user settings default experience set to Studio Classic, then that user will have Studio Classic as their default experience.

**Set Studio as the default experience for your domain using the SageMaker console**

You can set Studio as the default experience for the existing domain by using the SageMaker console.

**To set Studio as the default experience (console)**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the domain where you want to enable Studio as your default experience.
4. On the **Domain details** page, if you do not have Studio set as your default experience you should see the **Enable the new Amazon SageMaker Studio** banner.

If you still do not see the **Enable the new Amazon SageMaker Studio** banner, you may need to choose **Enable the new Studio experience** to view the banner.

5. (Optional) To view the details about the steps involved in enabling Studio as your default experience, choose **View details**. The page shows the following:

- In the **SageMaker Studio Overview** section, you can view the applications that are included or available in the Studio web-based interface.

- In the **Enablement process** section, you can view descriptions of the workflow tasks to enable Studio.

    **Note**

    You will need to migrate your data manually. For instructions about migrating your data, lifecycle configurations, and JupyterLab extensions, see [Migrating from SageMaker Studio Classic to SageMaker Studio](#).

- In the **Revert to Studio Classic experience** section, you can view how to revert to Studio Classic after enabling Studio as your default experience.

6. To begin the process to enable Studio as your default experience, choose **Submit**.

7. In the **Specify and configure role** section, you can view the default applications that are automatically included in Studio.

    To prevent users from running these applications, choose the AWS Identity and Access Management (IAM) Role that has an IAM policy that denies access. For more information, see [Limit access to default applications in Studio](#).

8. In the **Choose default S3 bucket to attach CORS policy** section, you can give Studio access to Amazon S3 buckets. The default Amazon S3 bucket, in this case, is the default S3 bucket for your Studio Classic. In this step, you can do the following:

    - Verify the domain's default S3 bucket to attach the CORS policy to. If your domain does not have a default Amazon S3 bucket, SageMaker creates an Amazon S3 bucket with the correct CORS policy attached.

    - You can include 10 additional S3 buckets to attach the CORS policy to.
To include more than 10 buckets, you must add them manually. For more information about manually attaching the CORS policy to your S3 buckets, see Update your CORS policy to access Amazon S3 buckets.

To proceed, select the check box next to **Do you agree to overriding any existing CORS policy on the chosen S3 buckets?**

9. The **Migrate data** section contains information about the different data storage volumes for Studio Classic and Studio. This process doesn't automatically migrate data. For instructions about migrating your data, lifecycle configurations, and JupyterLab extensions, see Migrating from SageMaker Studio Classic to SageMaker Studio.

10. After you complete the tasks on the page and verify your configuration, choose **Submit**.

**Set Studio as the default experience for your domain using the AWS CLI**

To set Studio as the default experience for the existing domain using the AWS CLI, use the `update-domain` call. You must set ENABLED as the value for StudioWebPortal and studio:: as the value for DefaultLandingUri as part of the default-user-settings parameter.

StudioWebPortal indicates if the Studio experience is the default experience. DefaultLandingUri indicates the default experience that the user is directed to when accessing the domain. In this example, setting these values on a domain level (in `default-user-settings`) makes Studio the default experience for users within the domain.

If a user within the domain has their StudioWebPortal set to DISABLED and DefaultLandingUri set to `app:JupyterServer:` on a user level (in `UserSettings`), this takes precedence over the domain settings. In other words, that user will have Studio Classic as their default experience, regardless of the domain settings.

The following code example shows how to set Studio as the default experience for users within the domain:

```bash
aws sagemaker update-domain \
--domain-id example-domain-id \
--region AWS Region \
--default-user-settings '
{
    "StudioWebPortal": "ENABLED",
}n```
To obtain your `example-domain-id`, use the following instructions:

**To get `example-domain-id`**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand `Admin configurations` and choose `Domains`.
3. Choose the relevant domain.
5. Copy the `Domain ID`.

To obtain your `AWS Region`, use the following instructions to ensure you are using the correct AWS Region for your domain:

**To get `AWS Region`**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand `Admin configurations` and choose `Domains`.
3. Choose the relevant domain.
4. On the `Domain details` page, verify that this is the relevant domain.
5. Expand the region dropdown list from the top right of the SageMaker console, and use the corresponding AWS Region ID to the right of your AWS Region name. For example, us-west-1.

After you migrate your default experience to Studio, you can give Studio access to Amazon S3 buckets. For example, you can include access to your Studio Classic IDE default Amazon S3 bucket and additional S3 buckets. To do so, you must manually attach a **Cross-Origin Resource Sharing** (CORS) configuration to the Amazon S3 buckets. For more information, see [Update your CORS policy to access Amazon S3 buckets](#).

Similarly, you can set Studio as the default experience when you create a domain from the AWS CLI by using the `create-domain` call.
Set Studio as the default experience for your domain using the AWS CloudFormation

You can set the default experience when creating a domain by using the AWS CloudFormation. For more information, see Creating Amazon SageMaker domain using AWS CloudFormation.

For information about the domain resource supported by AWS CloudFormation, see AWS::SageMaker::Domain.

After you migrate your default experience to Studio, you can give Studio access to Amazon S3 buckets. For example, you can include access to your Studio Classic IDE default Amazon S3 bucket and additional S3 buckets. To do so, you must manually attach a Cross-Origin Resource Sharing (CORS) configuration to the Amazon S3 buckets. For more information, see Update your CORS policy to access Amazon S3 buckets.

Limit access to default applications in Studio

Because Studio shows an expanded set of applications, users may have access to applications that weren't displayed before. Administrators can limit access to these default applications by creating an AWS Identity and Access Management (IAM) policy that denies access to them. The policy must then be attached to the execution role of the domain or user profile, as shown in the following procedure.

To limit access to applications in Studio

1. Create a policy with the following content by following the steps in Creating IAM policies:

```json
{
   "Version": "2012-10-17",
   "Statement": [
   {
      "Sid": "AllowSageMakerCreateAppOperations",
      "Effect": "Allow",
      "Action": "sagemaker:CreateApp",
      "Resource": "*"
   },
   {
      "Sid": "DenySageMakerCreateApp",
      "Effect": "Deny",
      "Action": "sagemaker:CreateApp",
      "Resource": "arn:aws:sagemaker:AWS Region:555555555555:app/example-domain-id/user-profile-name/example-app-type/*"
   }
]}
```
To obtain your AWS Region, use the following instructions to ensure you are using the correct AWS Region for your domain:

**To get AWS Region**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
4. On the Domain details page, verify that this is the relevant domain.
5. Expand the region dropdown list from the top right of the SageMaker console, and use the corresponding AWS Region ID to the right of your AWS Region name. For example, us-west-1.

To obtain your account ID, 555555555555, use the following instructions:

**To obtain your account ID**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Expand the account information and options menu from the top right of the SageMaker console.
3. Copy your account ID by clicking on the copy icon next to Account ID.

To obtain your example-domain-id, use the following instructions:

**To get example-domain-id**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
5. Copy the Domain ID.

To obtain the user-profile-name to limit access to, you may want to find the list of user profiles in the domain. To find a list of user profile names for your domain, use the following instructions:
To get **user-profile-name**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.
3. Choose the relevant domain.
4. On the **User profiles** page, search for the relevant user profile and copy the name.

   • For *example-app-type*, use the application type that's relevant to the application that you want to stop. For example, replace *example-app-type* with one of the following application types:

   • JupyterLab application type: JupyterLab. For information about JupyterLab, see [SageMaker JupyterLab](amazon_sagemaker).

   • Code Editor application type: CodeEditor. For information about Code Editor, based on Code-OSS, Visual Studio Code - Open Source, see [Get started with Code Editor in Amazon SageMaker Studio](amazon_sagemaker).

   • To obtain your *example-space-name*, use the following steps:

   **To get **example-space-name****

   1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.
3. Choose the relevant domain.
4. On the **Domain details** page, choose the **Space management** tab.
5. Copy the relevant space name.

2. Attach the policy to the execution role of the domain or user profile that you want to deny access to. For instructions, follow the steps in [Adding IAM identity permissions (console)](amazon_sagemaker).

**Update your CORS policy to access Amazon S3 buckets**

In Studio Classic, users can create, list, and upload files to Amazon Simple Storage Service (Amazon S3) buckets. To support the same experience in Studio, administrators must attach a [Cross-Origin Resource Sharing](amazon_sagemaker) (CORS) configuration to the Amazon S3 buckets. This is required because Studio makes Amazon S3 calls from the internet browser. The browser invokes CORS on behalf of users. As a result, all of the requests to Amazon S3 buckets fail unless the CORS policy is attached to the Amazon S3 buckets.
When an Amazon S3 default bucket doesn't exist, SageMaker creates an Amazon S3 bucket with the correct CORS policy attached.

However, you may need to manually attach the CORS policy to Amazon S3 buckets for the following reasons:

- If there is an existing Amazon S3 default bucket that doesn't have the correct CORS policy attached when you migrate your domain's default experience to Studio.
- If you are using the AWS CLI to migrate your domain's default experience to Studio. For information about using the AWS CLI to migrate, see Set Studio as the default experience for your domain using the AWS CLI.
- If you want to attach the CORS policy to additional Amazon S3 buckets.

**Note**

If you plan to use the SageMaker console to enable Studio as your default experience, the S3 buckets that you attach the CORS policy to will have their existing CORS policies overridden during the migration. For this reason, you can ignore following manual instructions.

However, if you already used the SageMaker console to migrate and want to include more Amazon S3 buckets to attach the CORS policy to, then continue with the following manual instructions.

The following procedure shows how to manually add a CORS configuration to an Amazon S3 bucket.

**To manually add a CORS configuration to an Amazon S3 bucket**

1. Verify that there is an Amazon S3 bucket in the same Region as your domain with the following name. For instructions, see Viewing the properties for an Amazon S3 bucket.

   `<sagemaker-region-accountId>`

2. Add a CORS configuration with the following content to the default Amazon S3 bucket. For instructions, see Configuring cross-origin resource sharing (CORS):
Migrate workflows from Studio Classic to JupyterLab

Migrating your workflows from the Studio Classic application to the JupyterLab application involves two distinct processes:

1. Setting Studio as the default experience.
2. Moving data and workflows from Studio Classic to the JupyterLab application.

Studio Classic and Studio use two different types of storage volumes. Studio Classic uses a single Amazon Elastic File System (Amazon EFS) volume for all data in the application. In Studio, each application gets its own Amazon Elastic Block Store (Amazon EBS) volume.

When you update the default experience of an existing domain, SageMaker doesn't automatically transfer data between these two types of volumes. As a result, data that's stored in an Amazon EBS
or Amazon EFS volume stays in that volume. If a user with data in Studio Classic accesses Studio, when the default experience changes, they won’t see that data in the JupyterLab Amazon EBS.

For more information about migrating workflows from Studio Classic to JupyterLab, see Migrating from SageMaker Studio Classic to SageMaker Studio.

**Migrate from Data Wrangler in Studio Classic to SageMaker Canvas**

Amazon SageMaker Data Wrangler exists as its own feature in the Studio Classic experience. When you enable Studio as your default experience, use the Amazon SageMaker Canvas application to access Data Wrangler functionality. SageMaker Canvas is an application in which you can train and deploy machine learning models without writing any code, and Canvas provides data preparation features powered by Data Wrangler.

The new Studio experience doesn't support the classic Data Wrangler UI, and you must create a Canvas application if you want to continue using Data Wrangler. However, you must have the necessary permissions to create and use Canvas applications.

Complete the following steps to attach the necessary permissions policies to the AWS IAM role for either your SageMaker domain or user.

**To grant permissions for Data Wrangler functionality inside Canvas**

1. Attach the AWS managed policy AmazonSageMakerFullAccess to your user's IAM role. For a procedure that shows you how to attach IAM policies to a role, see Adding IAM identity permissions (console) in the AWS IAM User Guide.

   If this permissions policy is too permissive for your use case, you can create scoped-down policies that include at least the following permissions:

   ```json
   {
       "Sid": "AllowStudioActions",
       "Effect": "Allow",
       "Action": [
           "sagemaker:CreatePresignedDomainUrl",
           "sagemaker:DescribeDomain",
           "sagemaker:ListDomains",
           "sagemaker:DescribeUserProfile",
           "sagemaker:ListUserProfiles",
           "sagemaker:DescribeSpace",
           "sagemaker:ListSpaces",
       ]
   }
   ```
2. Attach the AWS managed policy `AmazonSageMakerCanvasDataPrepFullAccess` to your user's IAM role.

After attaching the necessary permissions, you can create a Canvas application and log in. For more information, see [Getting started with using Amazon SageMaker Canvas](#).

When you've logged into Canvas, you can directly access Data Wrangler and begin creating data flows. For more information, see [Prepare data](#) in the Canvas documentation.

**Migrate from Autopilot in Studio Classic to SageMaker Canvas**

Amazon SageMaker Autopilot exists as its own feature in the Studio Classic experience. When you migrate to using the updated Studio experience, use the Amazon SageMaker Canvas application to continue using the same automated machine learning (AutoML) capabilities via a user interface (UI). SageMaker Canvas is an application in which you can train and deploy machine learning models without writing any code, and Canvas provides a UI to run your AutoML tasks.

The new Studio experience doesn't support the classic Autopilot UI. You must create a Canvas application if you want to continue using Autopilot's AutoML features via a UI.

However, you must have the necessary permissions to create and use Canvas applications.
• If you are accessing SageMaker Canvas from Studio, add those permissions to the execution role of your SageMaker domain or user profile.
• If you are accessing SageMaker Canvas from the Console, add those permissions to your user's AWS IAM role.
• If you are accessing SageMaker Canvas via a presigned URL, add those permissions to the IAM role that you're using for Okta SSO access.

To enable AutoML capabilities in Canvas, add the following policies to your execution role or IAM user role.

• AWS managed policy: CanvasFullAccess.
• Inline policy:

```
{
   "Sid": "AllowAppActionsForUserProfile",
   "Effect": "Allow",
   "Action": [
      "sagemaker:CreateApp",
      "sagemaker:DeleteApp"
   ],
   "Resource": "arn:aws:sagemaker:*:*:app/*/*//*.",
   "Condition": {
      "Null": {
         "sagemaker:OwnerUserProfileArn": "true"
      }
   }
}
```

To attach IAM policies to an execution role

1. **Find the execution role attached to your SageMaker user profile**
   a. In the SageMaker console https://console.aws.amazon.com/sagemaker/, navigate to Domains, then choose your SageMaker domain.
   b. The execution role ARN is listed under Execution role on the User Details page of your user profile. Make note of the execution role name in the ARN.
d. Search for your role by name in the search field.
e. Select the role.

2. Add policies to the role

b. Search for your role by name in the search field.
c. Select the role.
d. In the Permissions tab, navigate to the dropdown menu Add permissions.
e. • For managed policies: Select Attach policies, search for the name of the manage policy you want to attach.

  Select the policy then choose Add permissions.
  • For inline policies: Select Create inline policy, paste your policy in the JSON tab, choose next, name your policy, and choose Create.

For a procedure that shows you how to attach IAM policies to a role, see Adding IAM identity permissions (console) in the AWS IAM User Guide.

After attaching the necessary permissions, you can create a Canvas application and log in. For more information, see Getting started with using Amazon SageMaker Canvas.

**Revert to Studio Classic experience**

Administrators can revert to Studio Classic as the default experience for your domain by updating your domain. This can be done through the SageMaker console or the AWS CLI. Choose one of the following tabs to view the relevant instructions.

When Studio Classic is the default experience for the domain, Studio Classic is the default experience for all users in the domain. However, the user settings take precedence over the domain settings. Therefore, if a user has their default experience set to Studio, then that user will have Studio as their default experience.

**Revert your default experience to Studio Classic using the SageMaker console**

This section shows how to revert to Studio Classic as the default experience using the SageMaker console.
To revert your default experience to Studio Classic (console)


2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.

3. Choose the domain to revert.

4. On the **Domain details** page, scroll to the **Revert to Studio Classic experience** section (bottom of page).

5. Choose **Revert to Studio Classic process**.

6. On the **Revert domain to Studio Classic** page, complete the following tasks and select the corresponding boxes as you do. Go through the following tasks before reverting your domain's default experience to Studio Classic:

   a. **Step 1 - Back up your data** contains information about the different data storage volumes for Studio Classic and Studio. Your data will not be migrated automatically through this process. For instructions about migrating your data, lifecycle configurations, and JupyterLab extensions, see [Migrating from SageMaker Studio Classic to SageMaker Studio](https://docs.aws.amazon.com/sagemaker/latest/dg/migrate-to-studio-classic.html).

   b. **Delete all JupyterLab and Code Editor applications from Studio** reminds you to delete your Studio applications to avoid additional charge. This is not a mandatory step because you can delete your applications and spaces after reverting your domain to Studio Classic. It is recommended to delete your unused applications and spaces to avoid additional costs from them.

      For instructions on how to delete applications and spaces from your domain, see [Delete or stop your Studio running instances, applications, and spaces](https://docs.aws.amazon.com/sagemaker/latest/dg/delete-deregister-stop.html).

   c. **Step 2 - Confirm you want to revert this domain to Studio Classic** asks you to confirm your intention to revert your domain's default experience to Studio Classic.

   d. **Provide feedback** provides the option to leave feedback about why you're reverting your domain to Studio Classic.

7. After all of the steps have been completed and the check boxes are filled, the **Revert domain to Studio Classic** button becomes available.

8. After you complete the tasks on the page and verify your changes, choose **Revert domain to Studio Classic** to revert your domain.
Revert your default experience to Studio Classic using the AWS CLI

To revert to Studio Classic as the default experience for the existing domain using the AWS CLI, use the `update-domain` call. You must set DISABLED as the value for StudioWebPortal and app:JupyterServer: as the value for DefaultLandingUri as part of the default-user-settings parameter.

StudioWebPortal indicates if the Studio experience is the default experience and DefaultLandingUri indicates the default experience that the user is directed to when accessing the domain. In this example, setting these values on a domain level (in default-user-settings) makes Studio Classic the default experience for users within the domain.

If a user within the domain has their StudioWebPortal set to ENABLED and DefaultLandingUri set to studio:: on a user level (in UserSettings), this takes precedence over the domain settings. In other words, that user will have Studio as their default experience, regardless of the domain settings.

The following code example shows how to set Studio Classic as the default experience for users within the domain:

```bash
aws sagemaker update-domain \
--domain-id example-domain-id \
--region AWS Region \
--default-user-settings '{
  "StudioWebPortal": "DISABLED",
  "DefaultLandingUri": "app:JupyterServer:" 
}
'
```

To obtain your `example-domain-id`, use the following instructions:

**To get `example-domain-id`**

2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
5. Copy the Domain ID.
To obtain your **AWS Region**, use the following instructions to ensure you are using the correct AWS Region for your domain:

2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.
3. Choose the relevant domain.
4. On the **Domain details** page, verify that this is the relevant domain.
5. Expand the region dropdown list from the top right of the SageMaker console, and use the corresponding AWS Region ID to the right of your AWS Region name. For example, us-west-1.

### Launch Amazon SageMaker Studio

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

This page's topics demonstrate how to launch Amazon SageMaker Studio from the Amazon SageMaker console and the AWS Command Line Interface (AWS CLI).

### Topics

- **Prerequisites**
- **Launch from the Amazon SageMaker console**
- **Launch using the AWS CLI**

### Prerequisites

Before you begin, complete the following prerequisites:
Onboard to a SageMaker domain with Studio access. If you don't have permissions to set Studio as the default experience for your domain, contact your administrator. For more information, see Amazon SageMaker domain overview.

Update the AWS CLI by following the steps in Installing the current AWS CLI Version.

From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.

Launch from the Amazon SageMaker console

Complete the following procedure to launch Studio from the Amazon SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, choose Studio.
3. From the Studio landing page, select the domain and user profile for launching Studio.
4. Choose Open Studio.
5. To launch Studio, choose Launch personal Studio.

Launch using the AWS CLI

This section demonstrates how to launch Studio using the AWS CLI. The procedure to access Studio using the AWS CLI depends if the domain uses AWS Identity and Access Management (IAM) authentication or AWS IAM Identity Center authentication. You can use the AWS CLI to launch Studio by creating a presigned domain URL when your domain uses IAM authentication. For information about launching Studio with IAM Identity Center authentication, see Custom onboarding to Amazon SageMaker domain using IAM Identity Center.

Launch if Studio is the default experience

The following code snippet demonstrates how to launch Studio from the AWS CLI using a presigned domain URL if Studio is the default experience. For more information, see create-presigned-domain-url.

```
aws sagemaker create-presigned-domain-url \
  --region region \
  --domain-id domain-id \
  --user-profile-name user-profile-name \
```
Launch if Amazon SageMaker Studio Classic is your default experience

The following code snippet demonstrates how to launch Studio from the AWS CLI using a presigned domain URL if Studio Classic is the default experience. For more information, see create-presigned-domain-url.

```bash
aws sagemaker create-presigned-domain-url \
  --region region \n  --domain-id domain-id \n  --user-profile-name user-profile-name \n  --session-expiration-duration-in-seconds 43200 \n  --landing-uri studio:
```

Amazon SageMaker Studio UI overview

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

The Amazon SageMaker Studio user interface is split into three distinct parts.

- **Navigation bar**– This section of the UI includes the URL, breadcrumbs, notifications, and user options.
- **Navigation pane**– This section of the UI includes a list of the applications that are supported in Studio and options for the main workflows in Studio.
- **Content pane**– The main working area that displays the current page of the Studio UI that you have open.
Amazon SageMaker Studio navigation bar

The navigation bar of the Studio UI includes the URL, breadcrumbs, notifications, and user options.

URL Structure

The URL of Studio changes as you navigate the UI. When you navigate to a different page in the UI, the URL changes to reflect that page. With the updated URL, you open any page in the Studio UI directly without navigating to the landing page first.

Breadcrumbs

As you navigate through the Studio UI, the breadcrumbs keep track of the parent pages of the current page. By choosing one of these breadcrumbs, you can navigate to parent pages in the UI.

Notifications
The notifications section of the UI gives information about important changes to Studio, updates to applications, and issues to resolve.

**User options**

The user options give information about the user profile that is currently using Studio, and gives the option to sign out of Studio.

**Amazon SageMaker Studio navigation pane**

**Navigation pane**

The navigation pane of the UI includes a list of the applications that are supported in Studio. It also provides options for the main workflows in Studio.

This section of the UI can be used in an expanded or collapsed state. To change whether the section is expanded or collapsed, select the **Collapse** icon.

**Applications**

The applications section lists the applications that are available in Studio. If you choose one of the application types, you are directed to the landing page for that application.

**Workflows**

The list of workflows includes all of the available actions that you can take in Studio. Choose one of the options to navigate to the landing page for that workflow. If there are multiple workflows available for that option, choosing the option opens a dropdown menu where you can select the desired landing page.

The following list describes the options and provides a link for more information:

- **Home**– The main landing page with an overview, getting started, and what's new.
- **Running instances**– All of the instances that are currently running in Studio. For more information, see [View, stop, or delete your Studio running instances, applications, and spaces](#).
• **Data**– Data preparation options where you can collaborate to store, explore, prepare, transform, and share your data.

  • For more information about Amazon SageMaker Data Wrangler, see [Prepare data](#).
  
  • For more information about Amazon SageMaker Feature Store, see [Create, store, and share features with Amazon SageMaker Feature Store](#).

  • For more information about Amazon EMR clusters, see [Prepare data using Amazon EMR](#).

• **Auto ML**– Automatically build, train, tune, and deploy machine learning (ML) models. For more information, see [Amazon SageMaker Canvas](#).

• **Experiments**– Create, manage, analyze, and compare your machine learning experiments using Amazon SageMaker Experiments. For more information, see [Manage Machine Learning with Amazon SageMaker Experiments](#).

• **Jobs**– View jobs created in Studio.

  • For more information about training, see [Train machine learning models](#).

  • For more information about model evaluation, see [Use SageMaker Clarify to evaluate large language models](#).

• **Pipelines**– Automate your ML workflow with Amazon SageMaker Model Building Pipelines, which provides resources to help you build, track, and manage your pipeline resources. For more information, see [Amazon SageMaker Model Building Pipelines](#).

• **Models**– Organize your models into groups and collections in the model registry, where you can manage model versions, view metadata, and deploy models to production. For more information, see [Register and Deploy Models with Model Registry](#).

• **JumpStart**– Amazon SageMaker JumpStart provides pretrained, open-source models for a wide range of problem types to help you get started with machine learning. For more information, see [SageMaker JumpStart](#).

• **Deployments**– Deploy your machine learning (ML) models for inference.

  • For more information about Amazon SageMaker Inference Recommender, see [Amazon SageMaker Inference Recommender](#).

  • For more information about endpoints, see [Deploy models for inference](#).

---

**Studio content pane**

The main working area is also called the content pane. It displays the current page of the Studio UI that you have open.
Studio home page

The Studio home page is the primary landing page in the main working area. The home page includes two distinct tabs. There is an **Overview** tab and a **Getting started** tab.

Overview

The **Overview** tab includes options to start spaces for popular application types, get started with pre-built and automated solutions for ML workflows, and links to common tasks in the Studio UI.

Getting started

The **Getting started** tab includes information, guidance, and resources on how to begin with Studio. This includes a guided tour of the Studio UI, a link to documentation about Studio, and a selection of quick tips.

Applications supported in Amazon SageMaker Studio

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).

Amazon SageMaker Studio supports the following applications:

- **Code Editor, based on Code-OSS, Visual Studio Code - Open Source**– Code Editor offers a lightweight and powerful integrated development environment (IDE) with familiar shortcuts, terminal, and advanced debugging capabilities and refactoring tools. It is a fully managed, browser-based application in Studio. For more information, see [Get started with Code Editor in Amazon SageMaker Studio](#).

- **Amazon SageMaker Studio Classic**– Amazon SageMaker Studio Classic is a web-based IDE for machine learning. With Studio Classic, you can build, train, debug, deploy, and monitor your machine learning models. For more information, see [Amazon SageMaker Studio Classic](#).

- **JupyterLab**– JupyterLab offers a set of capabilities that augment the fully managed notebook offering. It includes kernels that start in seconds, a pre-configured runtime with popular
data science, machine learning frameworks, and high performance block storage. For more information, see SageMaker JupyterLab.

- **Amazon SageMaker Canvas**– With SageMaker Canvas, you can use machine learning to generate predictions without writing code. With Canvas, you can chat with popular large language models (LLMs), access ready-to-use models, or build a custom model that's trained on your data. For more information, see Amazon SageMaker Canvas.

- **RStudio**– RStudio is an integrated development environment for R. It includes a console and syntax-highlighting editor that supports running code directly. It also includes tools for plotting, history, debugging, and workspace management. For more information, see RStudio on Amazon SageMaker.

### Amazon SageMaker Studio spaces

⚠ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

Spaces are used to manage the storage and resource needs of some Amazon SageMaker Studio applications. Each space has a 1:1 relationship with an instance of an application. Every supported application that is created gets its own space. The following applications in Studio run on spaces:

- Get started with Code Editor in Amazon SageMaker Studio
- SageMaker JupyterLab
- Amazon SageMaker Studio Classic

A space is composed of the following resources:

- A storage volume.
  - For Studio Classic, the space is connected to the shared Amazon Elastic File System (Amazon EFS) volume for the domain.
  - For other applications, a distinct Amazon Elastic Block Store (Amazon EBS) volume is attached to the space. All applications are given their own Amazon EBS volume. Applications do not
have access to the Amazon EBS volume of other applications. For more information about Amazon EBS volumes, see Amazon Elastic Block Store (Amazon EBS).

- The application type of the space.
- The image that the application is based on.

Spaces can be either private or shared:

- **Private**: Private spaces are scoped to a single user in a domain. Private spaces cannot be shared with other users. All applications that support spaces also support private spaces.
- **Shared**: Shared spaces are accessible by all users in the domain. Only Studio Classic supports shared spaces. For more information about shared spaces, see Collaborate with shared spaces.

Spaces can be created in domains that use either AWS IAM Identity Center or AWS Identity and Access Management (IAM) authentication. The following sections give general information about how to access spaces. For specific information about creating and accessing a space, see the documentation for the respective application type of the space that you're creating.

For information about viewing, stopping, or deleting your applications, instances, or spaces, see Delete or stop your Studio running instances, applications, and spaces.

**Topics**

- **Access spaces**

**Access spaces**

The following sections show how to access the list of spaces associated with the user profile in the domain.

**Accessing spaces from the Amazon SageMaker console**

**To access spaces from the Amazon SageMaker console**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Under Admin configurations, choose Domains.
3. From the list of domains, select the domain that contains the spaces.
4. On the Domain details page, select the Space management tab. For more information about managing spaces, see Collaborate with shared spaces.
5. From the list of spaces for that domain, select the space to launch.
6. Choose **Launch Studio** for the space that you want to launch.

### Accessing spaces from Studio

Follow these steps to access spaces from Studio for a specific application type.

**To access spaces from Studio**

1. Open Studio by following the steps in [Launch Amazon SageMaker Studio](#).
2. Select the application type with spaces that you want to access.

### Accessing spaces using the AWS CLI

The following sections show how to access a space from the AWS Command Line Interface (AWS CLI). The procedures are for domains that use AWS Identity and Access Management (IAM) or AWS IAM Identity Center authentication.

#### IAM authentication

The following procedure outlines generally how to access a space using IAM authentication from the AWS CLI.

1. Create a presigned domain URL specifying the name of the space that you want to access.

   ```bash
   aws \\n   --region region \\n   sagemaker \\n   create-presigned-domain-url \\n   --domain-id domain-id \\n   --user-profile-name user-profile-name \\n   --space-name space-name
   ```

2. Navigate to the URL.

#### Accessing a space in IAM Identity Center authentication

The following procedure outlines how to access a space using IAM Identity Center authentication from the AWS CLI.
1. Use the following command to return the URL associated with the space.

   ```bash
   aws \\
   --region region \\
   sagemaker \\
   describe-space \\
   --domain-id domain-id \\
   --space-name space-name
   ```

2. Append the respective redirect parameter for the application type to the URL to be federated through IAM Identity Center. For more information about the redirect parameters, see `describe-space`.

3. Navigate to the URL to be federated through IAM Identity Center.

**Perform common tasks**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

The following sections describe how to perform common tasks in Amazon SageMaker Studio. For an overview of the Studio user interface, see Amazon SageMaker Studio UI overview.

### Set cookie preferences

1. Launch Studio following the steps in Launch Amazon SageMaker Studio.
2. At the bottom of the Studio user interface, choose Cookie Preferences.
3. Select the check box for each type of cookie that you want Amazon SageMaker to use.
4. Choose Save preferences.

### Manage notifications

Notifications give information about important changes to Studio, updates to applications, and issues to resolve.
1. Launch Studio following the steps in Launch Amazon SageMaker Studio.

2. On the top navigation bar, choose the **Notifications** icon ( ).

3. From the list of notifications, select the notification to get information about it.

**Leave feedback**

We take your feedback seriously. We encourage you to provide feedback.

At the top navigation of Studio, choose **Provide feedback**.

**Sign out**

Signing out of the Studio UI is different than closing the browser window. Signing out clears session data from the browser and deletes unsaved changes.

This same behavior also happens when the Studio session times out. This happens after 5 minutes.

1. Launch Studio following the steps in Launch Amazon SageMaker Studio.

2. Choose the **User options** icon ( ).

3. Choose **Sign out**.

4. In the pop-up window, choose **Sign out**.

**Use NVMe stores with Amazon SageMaker Studio**

Amazon SageMaker Studio applications and their associated notebooks run on Amazon Elastic Compute Cloud (Amazon EC2) instances. Some of the Amazon EC2 instance types, such as the m1. m5d instance family, offer non-volatile memory express (NVMe) solid state drives (SSD) instance stores.

NVMe instance stores are local ephemeral disk stores that are physically connected to an instance for fast temporary storage. Studio applications support NVMe instance stores for supported instance types. For more information about instance types and their associated NVMe store volumes, see the Amazon Elastic Compute Cloud Instance Type Details.
This following topic provides information about accessing and using NVMe instance stores, as well as considerations when using NVMe instance stores with Studio.

Considerations

The following considerations apply when using NVMe instance stores with Studio.

- An NVMe instance store is temporary storage. The data stored on the NVMe store is deleted when the instance is terminated, stopped, or hibernated. When using NVMe stores with Studio applications, the data on the NVMe instance store is lost whenever the application is deleted, restarted, or patched. We recommend that you back up valuable data to persistent storage solutions, such as Amazon Elastic Block Store, Amazon Elastic File System, or Amazon Simple Storage Service.

- Studio patches instances periodically to install new security updates. When an instance is patched, the instance is restarted. This restart results in the deletion of data stored in the NVMe instance store. We recommend that you frequently back up necessary data from the NVMe instance store to persistent storage solutions, such as Amazon Elastic Block Store, Amazon Elastic File System, or Amazon Simple Storage Service.

- The following Studio applications support using NVMe storage:
  - JupyterLab
  - Code Editor, based on Code-OSS, Visual Studio Code - Open Source
  - KernelGateway

Access NVMe instance stores

When you select an instance type with attached NVMe instance stores to host a Studio application, the NVMe instance store directory is mounted to the application container at the following location:

```
/mnt/sagemaker-nvme
```

If an instance has more than 1 NVMe instance store attached to it, Studio creates a striped logical volume that spans all of the local disks attached. Studio then mounts this striped logical volume to the `/mnt/sagemaker-nvme` directory. As a result, the directory storage size is the sum of all NVMe instance store volume sizes attached to the instance.
If the `/mnt/sagemaker-nvme` directory does not exist, verify that the instance type hosting your application has an attached NVMe instance store volume.

**Local mode support in Amazon SageMaker Studio**

Amazon SageMaker Studio applications support the use of local mode to create estimators, processors, and pipelines, then deploy them to a local environment. With local mode, you can test machine learning scripts before running them in Amazon SageMaker managed training or hosting environments. Studio supports local mode in the following applications:

- Amazon SageMaker Studio Classic
- JupyterLab
- Code Editor, based on Code-OSS, Visual Studio Code - Open Source

Local mode in Studio applications is invoked using the SageMaker Python SDK. In Studio applications, local mode functions similarly to how it functions in Amazon SageMaker notebook instances, with some differences. For more information about using local mode with the SageMaker Python SDK, see [Local Mode](#).

**Note**

Studio applications do not support multi-container jobs in local mode. Local mode jobs are limited to a single instance for training, inference, and processing jobs. When creating a local mode job, the instance count configuration must be 1.

As part of local mode support, Studio applications support limited Docker access capabilities. With this support, users can interact with the Docker API from Jupyter notebooks or the image terminal of the application. Customers can interact with Docker using one of the following:

- [Docker CLI](#)
- [Docker Compose CLI](#)
- Language specific Docker SDK clients

**Prerequisites**

Complete the following prerequisites to use local mode in Studio applications:
• To pull images from an Amazon Elastic Container Registry repository, the account hosting the Amazon ECR image must provide access permission for the user’s execution role. The domain's execution role must also allow Amazon ECR access.

• Verify that you are using the latest version of the Studio Python SDK by using the following command:

   ```bash
   pip install -U sagemaker
   ```

• To use local mode and Docker capabilities, set the following parameter of the domain's `DockerSettings` using the AWS Command Line Interface (AWS CLI):

   ```bash
   EnableDockerAccess : ENABLED
   ```

• Using `EnableDockerAccess`, you can also control whether users in the domain can use local mode. By default, local mode and Docker capabilities aren't allowed in Studio applications. For more information, see Setting `EnableDockerAccess`.

• Install the Docker CLI in the Studio application by following the steps in Docker installation.

**Setting `EnableDockerAccess`**

The following sections show how to set `EnableDockerAccess` when the domain has public internet access or is in VPC-only mode.

**Note**

Changes to `EnableDockerAccess` only apply to applications created after the domain is updated. You must create a new application after updating the domain.

**Public internet access**

The following example commands show how to set `EnableDockerAccess` when creating a new domain or updating an existing domain with public internet access:

```bash
# create new domain
aws --region region \
    sagemaker create-domain --domain-name domain-name \ 
    --vpc-id vpc-id
```
VPC-only mode

When using a domain in VPC-only mode, Docker image push and pull requests are routed through the service VPC instead of the VPC configured by the customer. Because of this functionality, administrators can configure a list of trusted AWS accounts that users can make Amazon ECR Docker pull and push operations requests to.

If a Docker image push or pull request is made to an AWS account that is not in the list of trusted AWS accounts, the request fails. Docker pull and push operations outside of Amazon Elastic Container Registry (Amazon ECR) aren't supported in VPC-only mode.

The following AWS accounts are trusted by default:

- The account hosting the SageMaker domain.
- SageMaker accounts that host the following SageMaker images:
  - DLC framework images
  - Sklearn, Spark, XGBoost processing images

To configure a list of additional trusted AWS accounts, specify the VpcOnlyTrustedAccounts value as follows:

```
aws --region region \
  sagemaker update-domain --domain-id domain-id \
  --domain-settings-for-update '{"DockerSettings": {"EnableDockerAccess": "ENABLED", "VpcOnlyTrustedAccounts": ["account-list"]}}'
```
Docker support

Studio also supports limited Docker access capabilities with the following restrictions:

- Usage of Docker networks is not supported.
- Docker volume usage is not supported during container run. Only volume bind mount inputs are allowed during container orchestration. The volume bind mount inputs must be located on the Amazon Elastic File System (Amazon EFS) volume for Studio Classic. For JupyterLab and Code Editor applications, it must be located on the Amazon Elastic Block Store (Amazon EBS) volume.
- Container inspect operations are allowed.
- Container port to host mapping is not allowed. However, you can specify a port for hosting. The endpoint is then accessible from Studio using the following URL:
  
  `http://localhost:port`

Docker operations supported

The following table lists all of the Docker API endpoints that are supported in Studio, including any support limitations. If an API endpoint is missing from the table, Studio doesn't support it.

<table>
<thead>
<tr>
<th>API Documentation</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>SystemAuth</td>
<td></td>
</tr>
<tr>
<td>SystemEvents</td>
<td></td>
</tr>
<tr>
<td>SystemVersion</td>
<td></td>
</tr>
<tr>
<td>SystemPing</td>
<td></td>
</tr>
<tr>
<td>SystemPingHead</td>
<td></td>
</tr>
<tr>
<td>ContainerCreate</td>
<td>• Containers cannot be run in Docker default bridge or custom Docker networks. Containers are run in the same network as the Studio application container.</td>
</tr>
<tr>
<td>API Documentation</td>
<td>Limitations</td>
</tr>
<tr>
<td>--------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>• Users can only use the following value for the network name: <code>sagemaker</code>. For example:</td>
<td>• Users can only use the following value for the network name: <code>sagemaker</code>. For example:</td>
</tr>
<tr>
<td></td>
<td><code>docker run --net sagemaker parameter -values</code></td>
</tr>
<tr>
<td>• Only bind mounts are allowed for volume usage. The host directory should exist on Amazon EFS for KernelGateway applications or Amazon EBS for other applications.</td>
<td>• Only bind mounts are allowed for volume usage. The host directory should exist on Amazon EFS for KernelGateway applications or Amazon EBS for other applications.</td>
</tr>
<tr>
<td>• Containers cannot run in privileged mode or with elevated secure computing permissions.</td>
<td>• Containers cannot run in privileged mode or with elevated secure computing permissions.</td>
</tr>
</tbody>
</table>

- **ContainerStart**
- **ContainerStop**
- **ContainerKill**
- **ContainerDelete**
- **ContainerList**
- **ContainerLogs**
- **ContainerInspect**
- **ContainerWait**
- **ContainerAttach**
- **ContainerPrune**
- **ContainerResize**
- **ImageCreate**

VPC-only mode support is limited to Amazon ECR images in allowlisted accounts.
<table>
<thead>
<tr>
<th>API Documentation</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ImagePrune</strong></td>
<td></td>
</tr>
<tr>
<td><strong>ImagePush</strong></td>
<td>VPC-only mode support is limited to Amazon ECR images in allowlisted accounts.</td>
</tr>
<tr>
<td><strong>ImageList</strong></td>
<td></td>
</tr>
<tr>
<td><strong>ImageInspect</strong></td>
<td></td>
</tr>
<tr>
<td><strong>ImageGet</strong></td>
<td></td>
</tr>
<tr>
<td><strong>ImageDelete</strong></td>
<td></td>
</tr>
</tbody>
</table>
| **ImageBuild**    | • VPC-only mode support is limited to Amazon ECR images in allowlisted accounts.  
• Users can only use the following value for the network name: sagemaker. For example:  
  docker build --network sagemaker parameter-values |

**Docker installation**

To use Docker, you must manually install Docker from the terminal of your Studio application. The steps to install Docker are different if the domain has access to the internet or not.

**Internet access**

If the domain is created with public internet access or in VPC-only mode with limited internet access, use the following steps to install Docker.

1. (Optional) If your domain is created in VPC-only mode with limited internet access, create a public NAT gateway with access to the Docker website. For instructions, see [NAT gateways](#).
2. Navigate to the terminal of the Studio application that you want to install Docker in.
3. To return the operating system of the application, run the following command from the terminal:
4. Install Docker following the instructions for the operating system of the application in the Amazon SageMaker Local Mode Examples repository. For example, install Docker on Ubuntu following the script at https://github.com/aws-samples/amazon-sagemaker-local-mode/blob/main/sagemaker_studio_docker_cli_install/sagemaker-ubuntu-focal-docker-cli-install.sh with the following considerations:

- If chained commands fail, run commands one at a time.
- Studio only supports Docker version 20.10.X. and Docker Engine API version 1.41.
- The following packages aren't required to use the Docker CLI in Studio and their installation can be skipped:
  - containerd.io
  - docker-ce
  - docker-buildx-plugin

**Note**
You do not need to start the Docker service in your applications. The instance that hosts the Studio application runs Docker service by default. All Docker API calls are routed through the Docker service automatically.

5. Use the exposed Docker socket for Docker interactions within Studio applications. By default, the following socket is exposed:

```
unix:///docker/proxy.sock
```

The following Studio application environmental variable for the default USER uses this exposed socket:

```
DOCKER_HOST
```
No internet access

If the domain is created in VPC-only mode with no internet access, use the following steps to install Docker.

1. Navigate to the terminal of the Studio application that you want to install Docker in.
2. Run the following command from the terminal to return the operating system of the application:
   ```
   cat /etc/os-release
   ```
3. Download the required Docker .deb files to your local machine. For instructions about downloading the required files for the operating system of the Studio application, see Install Docker Engine.

   For example, install Docker from a package on Ubuntu following the steps 1–4 in Install from a package with the following considerations:
   
   - Install Docker from a package. Using other methods to install Docker will fail.
   - Install the latest packages corresponding to Docker version 20.10.X.
   - The following packages aren't required to use the Docker CLI in Studio. You don't need to install the following:
     - containerd.io
     - docker-ce
     - docker-buildx-plugin

   ![Note]

   You do not need to start the Docker service in your applications. The instance that hosts the Studio application runs Docker service by default. All Docker API calls are routed through the Docker service automatically.

4. Upload the .deb files to the Amazon EFS file system or to the Amazon EBS file system of the application.
5. Manually install the docker-ce-cli and docker-compose-plugin .deb packages from the Studio application terminal. For more information and instructions, see step 5 in Install from a package on the Docker docs website.
6. Use the exposed Docker socket for Docker interactions within Studio applications. By default, the following socket is exposed:

```
unix:///docker/proxy.sock
```

The following Studio application environmental variable for the default USER uses this exposed socket:

```
DOCKER_HOST
```

**View, stop, or delete your Studio running instances, applications, and spaces**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).

The following topics include information and instructions about how to view, stop, or delete your Studio running instances, applications, and spaces. For more information about Studio spaces, see [Amazon SageMaker Studio spaces](#).

We briefly give an overview of the differences between a space, application, and instance in the following bullet points:

- When you create a space, you are creating the necessary resources to run an application. This includes an Amazon Elastic Block Store (Amazon EBS) volume where your data is stored. When you delete a space, you are also deleting your data stored in the space.
- When you open an application, you will need to start an instance for the application to run on. When you close an application, you will not automatically stop and delete the instance. You can reopen the application while the instance is running.
When you use the `DeleteApp` API you also stop and delete the instance. You can restart the instance and application after using this API.

- For the instructions on this page, the action to stop an instance or delete an instance has the same effect. When you stop or delete an instance, you also stop the application.

Similarly, to stop an instance is the same as to stop or delete an application.

**Topics**

- [View your Studio running instances, applications, and spaces](#)
- [Delete or stop your Studio running instances, applications, and spaces](#)

**View your Studio running instances, applications, and spaces**

**View your Studio running instances and applications**

The **Running instances** page gives information about all running application instances that were created in Amazon SageMaker Studio by the user, or were shared with the user.

You can view and stop running instances for all of your applications and spaces. If an instance is stopped, it does not appear on this page. Stopped instances can be viewed from the landing page for their respective application types.

You can view a list of running applications and their details in Studio.

**To view running instances**

1. Launch Studio following the steps in [Launch Amazon SageMaker Studio](#).
2. On the left navigation pane, choose **Running instances**.
3. From the **Running instances** page, you can view a list of running applications and details about those applications.

   To view non-running instances, from the left navigation pane choose, the relevant application under **Applications**. The non-running applications will have the **Stopped** status under the **Status** column.
View your Studio spaces

The **Spaces** section within your **Domain details** page gives information about Studio spaces within your domain. You can view, create, and delete spaces on this page.

The spaces that you can view in the **Spaces** section are running spaces for the following:

- JupyterLab private space. For information about JupyterLab, see [SageMaker JupyterLab](#).
- Code Editor private space. For information about Code Editor, based on Code-OSS, Visual Studio Code - Open Source, see [Get started with Code Editor in Amazon SageMaker Studio](#).
- Studio Classic shared space. For information about Studio Classic shared space, see [Collaborate with shared spaces](#).

There are no spaces for SageMaker Canvas, Studio Classic (private), or RStudio.

To view Studio spaces in a domain

2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.
3. Choose the domain where you want to view the spaces.
4. On the **Domain details** page, choose the **Space management** tab to open the **Spaces** section.

Delete or stop your Studio running instances, applications, and spaces

To avoid additional charges from unused Studio running instances, applications, or spaces, you can stop or delete them. This page will provide some information about the differences between stopping or deleting your Studio running instances, applications, or spaces, followed by instructions.

For more information about the differences between Studio spaces, application, and instances, see [View, stop, or delete your Studio running instances, applications, and spaces](#).

Delete or stop your Amazon SageMaker Studio application or running instance

To avoid additional charges from unused running applications, you can stop and delete those applications and running instances. The following provides some information on stopping or deleting an application or instance:
• In the following instructions, to delete an application (uses the DeleteApp API) has the same effect as to stop the instance for the application. Following the instructions to delete an application or stop an instance, both stops and deletes the application and the instance for the application.

• After you delete an application or stop an instance, you can start up the instance and application again later.

• When you delete an application or stop an instance, the files in the space will persist. You can run the application again and expect to have access to the same files that are stored in the space, as you did before deleting the application.

• When you delete an application or stop an instance, the metadata for the application will be deleted within 24 hours. For more information, see the note in the CreationTime response element for the DescribeApp API.

The following tabs provide instructions to stop and delete an application from your domain using the Studio UI, the SageMaker console, or the AWS CLI.

**Note**

To view and stop all of your Studio running instances in one location, we recommend the Use the Studio UI to delete your domain applications workflow from the following options.

---

**Use the Studio UI to delete your domain applications**

To delete your Studio applications using the Studio UI, use the following instructions.

**To delete your domain applications (Studio UI)**

1. Launch Studio. This process may differ depending on your setup. For information about launching Studio, see Launch Amazon SageMaker Studio.

2. From the left navigation pane, choose Running instances.

   If the table on the page is empty, you don't have any running instances or applications in your spaces.

3. In the table under the Name and Application columns, find the space name and the application that you want to stop and delete.
4. Choose the corresponding **Stop** button to stop and delete the application.

**Delete domain applications using the SageMaker console**

To view or stop Studio running instances from a centralized location, see [Use the Studio UI to delete your domain applications](#). Otherwise, use the following instructions.

In the SageMaker console, you can only stop the running Studio applications for the spaces that you are able to view in the **Spaces** section of the console. For a list of the viewable spaces, see [View your Studio spaces](#).

These steps show how to delete your Studio applications by using the SageMaker console.

**To delete applications instructions (console)**

2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.
3. Choose the domain that you want to revert.
4. On the **Domain details** page, choose the **Space management** tab.
5. **Important**

   In the **Space management** tab, you have the option to delete the space. There is a difference between deleting the space and deleting an application. If you delete the space, you will lose access to the data within that space. Do not delete the space unless you're sure that you want to.

   To stop and delete the application, in the **Space management** tab and under the **Name** column, choose the space for the application.
6. In the **Apps** section and under the **App type** column, search for the app to stop and delete.
7. Under the **Action** column, choose the corresponding **Delete app** button.
8. In the pop-up box, choose **Yes, delete app**. After you do so the delete input field becomes available.
9. Enter **delete** in the delete input field to confirm deletion.
10. Choose **Delete**.
Delete your domain applications using the AWS CLI

To view or stop any of your Studio running instances from a centralized location, see Use the Studio UI to delete your domain applications. Otherwise, use the following instructions.

The following code examples use the DeleteApp API to delete an application in an example domain.

To stop your running JupyterLab or Code Editor instances, use the following code example:

```bash
aws sagemaker delete-app \
--domain-id example-domain-id \ 
--region AWS Region \ 
--app-name default \ 
--app-type example-app-type \ 
--space-name example-space-name
```

• To obtain your example-domain-id, use the following instructions:

  To get example-domain-id

  1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
  2. From the left navigation pane, expand Admin configurations and choose Domains.
  3. Choose the relevant domain.
  5. Copy the Domain ID.

• To obtain your AWS Region, use the following instructions to ensure you are using the correct AWS Region for your domain:

  To get AWS Region

  1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
  2. From the left navigation pane, expand Admin configurations and choose Domains.
  3. Choose the relevant domain.
  4. On the Domain details page, verify that this is the relevant domain.
  5. Expand the region dropdown list from the top right of the SageMaker console, and use the corresponding AWS Region ID to the right of your AWS Region name. For example, us-west-1.
For example-app-type, use the application type that's relevant to the application that you want to stop. For example, replace example-app-type with one of the following application types:

- JupyterLab application type: JupyterLab. For information about JupyterLab, see SageMaker JupyterLab.
- Code Editor application type: CodeEditor. For information about Code Editor, based on Code-OSS, Visual Studio Code - Open Source, see Get started with Code Editor in Amazon SageMaker Studio.

To obtain your example-space-name, use the following steps:

**To get example-space-name**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
4. On the Domain details page, choose the Space management tab.
5. Copy the relevant space name.

To stop running instances for SageMaker Canvas, Studio Classic, or RStudio, use the following code example:

```bash
aws sagemaker delete-app \
   --domain-id example-domain-id \
   --region AWS Region \
   --app-name default \
   --app-type example-app-type \
   --user-profile example-user-name
```

For example-app-type, use the application type relevant to the application that you want to stop. For example, replace example-app-type with one of the following application types:

- SageMaker Canvas application type: Canvas. For information about SageMaker Canvas, see Amazon SageMaker Canvas.
- Studio Classic application type: JupyterServer. For information about Studio Classic, see Amazon SageMaker Studio Classic.
- RStudio application type: RStudioServerPro. For information about RStudio, see RStudio on Amazon SageMaker.
To obtain your `example-user-name`, navigate to the **Domain details** page.

- Next, choose the **User profiles** tab, and copy the relevant space name.

For alternative instructions to delete your running Studio applications, see:

- **JupyterLab**: [Delete unused resources](#).
- **Code Editor**: [Log out and shut down resources](#).
- **SageMaker Canvas**: [Logging out of Amazon SageMaker Canvas](#).
- **Studio Classic**: [Shut Down and Update SageMaker Studio Classic and Studio Classic Apps](#).
- **RStudio**: [Shut down and restart RStudio](#).

## Delete a Studio space

![Important](#)

**Important**

After you delete your space, you will lose all of the data stored in the space. We recommend that you back up your data before deleting your space.

To delete a Studio space, you will need to have administrator permissions or at least have permissions to update domain, IAM and Amazon S3.

- Spaces are used to manage the storage and resource needs of the relevant application. When you delete a space, the storage volume also deletes. Therefore, you lose access to the files stored on that space. For more information about Studio spaces, see [Amazon SageMaker Studio spaces](#).

  We recommend that you back up your data if you choose to delete a space.

- After you delete a space, you can't access that space again.

You can delete the Studio spaces that are viewable in the **Spaces** section of the console. For a list of the viewable spaces, see [View your Studio spaces](#).

There are no spaces for SageMaker Canvas, Studio Classic (private), and RStudio. To stop and delete your SageMaker Canvas, Studio Classic (private), or RStudio applications, see [Delete or stop your Amazon SageMaker Studio application or running instance](#).
Delete a space using the SageMaker console

The **Spaces** section within your **Domain details** page gives information about Studio spaces within your domain. You can view, create, and delete spaces on this page.

**To view Studio spaces in a domain**

2. From the left navigation pane, expand **Admin configurations** and choose **Domains**.
3. Choose the domain where you want to view the spaces.
4. On the **Domain details**, choose **Space management** to open the **Spaces** section.
5. Select the space to delete.
6. Choose **Delete**.
7. In the pop-up box titled **Delete space**, you have two options:
   - If you already shut down all applications in the space, choose **Yes, delete space**.
   - If you still have applications running in the space, choose **Yes, shut down all apps and delete space**.
8. Enter **delete** in the delete input field to confirm deletion.
9. To delete the space, you have two options:
   - If you already shut down all applications in the space, choose **Delete space**.
   - If you still have applications running in the space, choose **Shut down all apps and delete space**.

**Delete a space using the AWS CLI**

Before you can delete a space using the AWS CLI, you must delete the application associated with it. For information about stopping your Studio applications, see [Delete or stop your Amazon SageMaker Studio application or running instance](#).

Use the following AWS CLI command to delete a space within a domain:

```
aws sagemaker delete-space
--domain-id example-domain-id
--region AWS Region
--space-name example-space-name
```
To obtain your example-domain-id, use the following instructions:

To get example-domain-id

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
5. Copy the Domain ID.

To obtain your AWS Region, use the following instructions to ensure you are using the correct AWS Region for your domain:

To get AWS Region

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
4. On the Domain details page, verify that this is the relevant domain.
5. Expand the region dropdown list from the top right of the SageMaker console, and use the corresponding AWS Region ID to the right of your AWS Region name. For example, us-west-1.

To obtain your example-space-name, use the following steps:

To get example-space-name

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation pane, expand Admin configurations and choose Domains.
3. Choose the relevant domain.
4. On the Domain details page, choose the Space management tab.
5. Copy the relevant space name.
Amazon SageMaker Studio pricing

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

There is no additional charge for using the Amazon SageMaker Studio UI.

The following do incur costs:

- Amazon Elastic Block Store or Amazon Elastic File System volumes that are mounted with your applications.
- Any jobs and resources that users launch from Studio applications.
- Launching a JupyterLab application, even if no resources or jobs launched in the application.

For information about how Amazon SageMaker Studio Classic is billed, see Amazon SageMaker Studio Classic Pricing.

For more information about billing along with pricing examples, see Amazon SageMaker Pricing.

Troubleshooting

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

This section shows how to troubleshoot common problems in Amazon SageMaker Studio.

Cannot delete Code Editor, based on Code-OSS, Visual Studio Code - Open Source or JupyterLab application
This issue occurs when a user creates an application from Amazon SageMaker Studio that is only available in Studio, then reverts to the Studio Classic experience as their default. As a result, the user cannot delete an application for Code Editor, based on Code-OSS, Visual Studio Code - Open Source or JupyterLab because they can't access the Studio UI.

To resolve this issue, notify your administrator so that they can delete the application manually using the AWS Command Line Interface (AWS CLI).

Amazon SageMaker Studio Classic

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker Studio Classic is a web-based integrated development environment (IDE) for machine learning that lets you build, train, debug, deploy, and monitor your machine learning models. Studio Classic provides all the tools you need to take your models from data preparation to experimentation to production while boosting your productivity. In a single unified visual interface, customers can perform the following tasks:

- Write and execute code in Jupyter notebooks
- Prepare data for machine learning
- Build and train machine learning models
- Deploy the models and monitor the performance of their predictions
- Track and debug the machine learning experiments

For information on the onboarding steps to sign in to Studio Classic, see Amazon SageMaker domain overview.

For the AWS Regions supported by Studio Classic, see Supported Regions and Quotas.

Topics
Studio Classic Features

Studio Classic includes the following features:

- *SageMaker Autopilot*
- *SageMaker Clarify*
- *SageMaker Data Wrangler*
- *SageMaker Debugger*
- *SageMaker Experiments*
- *SageMaker Feature Store*
- *SageMaker JumpStart*
- *Amazon SageMaker Model Building Pipelines*
- *SageMaker Model Registry*
- *SageMaker Projects*
- *SageMaker Studio Classic Notebooks*
- *SageMaker Studio Universal Notebook*
Amazon SageMaker Studio Classic UI Overview

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker Studio Classic extends the capabilities of JupyterLab with custom resources that can speed up your Machine Learning (ML) process by harnessing the power of AWS compute. Previous users of JupyterLab will notice the similarity of the user interface. The most prominent additions are detailed in the following sections. For an overview of the original JupyterLab interface, see The JupyterLab Interface.

The following image shows the default view upon launching Amazon SageMaker Studio Classic. The left navigation panel displays all top-level categories of features, and a Studio Classic Home page is open in the main working area. Come back to this central point of orientation by choosing the Home icon at any time, then selecting the Home node in the navigation menu.

Try the Getting started notebook for an in-product hands-on guide on how to set up and get familiar with Amazon SageMaker Studio Classic features. On the Quick actions section of the Studio Classic Home page, choose Open the Getting started notebook.
Note

This chapter is based on Studio Classic's updated user interface (UI) available on version v5.38.x and above on JupyterLab3.

- To retrieve your version of Studio Classic UI, from the Studio Classic Launcher, open a System Terminal, then

  1. Run `conda activate studio`
  2. Run `jupyter labextension list`
  3. Search for the version displayed after `@amzn/sagemaker-ui` version in the output.

- For information about updating Amazon SageMaker Studio Classic, see Shut down and Update SageMaker Studio Classic.

Topics

- Studio Classic Home page
Studio Classic Home page

The Home page provides access to common tasks and workflows. In particular, it includes a list of Quick actions for common tasks such as Open Launcher to create notebooks and other resources and Import & prepare data visually to create a new flow in Data Wrangler. The Home page also offers tooltips on key controls in the UI.

The Prebuilt and automated solutions help you get started quickly with SageMaker's low-code solutions such as Amazon SageMaker JumpStart and Autopilot.

In Workflows and tasks, you can find a list of relevant tasks for each step of your ML workflow that takes you to the right tool for the job. For example, Transform, analyse, and export data takes you to Amazon SageMaker Data Wrangler and opens the workflow to create a new data flow, or View all experiments takes you to SageMaker Experiments and opens the experiments list view.

Upon Studio Classic launch, the Home page is open in the main working area. You can customize your SageMaker Home page by choosing Customize Layout at the top right of the Home tab.

Studio Classic layout

The Amazon SageMaker Studio Classic interface consists of a menu bar at the top, a collapsible left sidebar displaying a variety of icons such as the Home icon and the File Browser, a status bar at the bottom of the screen, and a central area divided horizontally into two panes. The left pane is a collapsible navigation panel. The right pane, or main working area, contains one or more tabs for resources such as launchers, notebooks, terminals, metrics, and graphs, and can be further divided.

Report a bug in Studio Classic or choose the notification icon to view notifications from Studio Classic, such as new Studio Classic versions and new SageMaker features, on the right corner of the menu bar. To update to a new version of Studio Classic, see Shut Down and Update SageMaker Studio Classic and Studio Classic Apps.

The following sections describe the Studio Classic main user interface areas.
**Left sidebar**

The *left sidebar* includes the following icons. When hovering over an icon, a tooltip displays the icon name. A single click on an icon opens up the left navigation panel with the described functionality. A double click minimizes the left navigation panel.

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Home Icon" /></td>
<td><strong>Home</strong></td>
</tr>
</tbody>
</table>

Choose the **Home** icon to open a top-level navigation menu in the *left navigation* panel.

Using the **Home** navigation menu, you can discover and navigate to the right tools for each step of your ML workflow. The menu also provides shortcuts to quick-start solutions and learning resources such as documentation and guided tutorials.

The menu categories group relevant features together. Choosing **Data**, for example, expands the relevant SageMaker capabilities for your data preparations tasks. From here, you can prepare your data with Data Wrangler, create and store ML features with Amazon SageMaker Feature Store, and manage Amazon EMR clusters for large-scale data processing. The categories are ordered following a typical ML workflow from preparing data, to building, training, and deploying ML models (data, pipelines, models, and deployments).

When you choose a specific node (such as Data Wrangler), a corresponding page opens in the main working area.

Choose **Home** in the navigation menu to open the [Studio Classic Home page](#).

| ![File Browser Icon](image) | **File Browser** |

The **File Browser** displays lists of your notebooks, experiments, trials, trial components, endpoints, and low-code solutions.

Whether you are in a personal or shared space determines who has access to your files. You can identify which type of space you are in by...
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>looking at the top right corner. If you are in a personal app, you see a user icon followed by <code>[user_name] / Personal Studio</code> and if you are in a collaborative space, you see a globe icon followed by <code>&quot;[user_name] / [space_name]. &quot;</code></td>
<td></td>
</tr>
</tbody>
</table>

- **Personal Studio Classic app**: A private Amazon EFS directory that only you can access.

- **Collaborative space**: A shared Amazon EFS directory with other members of your team for group access to notebooks and resources. Working in a shared space allows for real-time team collaboration on notebooks.

- **Studio Classic launcher**: Choose the plus (+) sign on the menu at the top of the file browser to open the [Amazon SageMaker Studio Classic Launcher](https://aws.amazon.com).

- **Upload files**: Choose the **Upload Files** icon (↑) to add files to Studio Classic or drag and drop them from your desktop.

- **Open files**: Double-click a file to open the file in a new tab or right-click and select **Open**.

- **Panel management**: To work in adjacent files, choose a tab that contains a notebook, Python, or text file, then choose **New View for File**.

  For hierarchical entries, a selectable breadcrumb at the top of the browser shows your location in the hierarchy.
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Icon" /> <strong>Property Inspector</strong>&lt;br&gt;The Property Inspector is a notebook cell tools inspector which displays contextual property settings when open.</td>
<td></td>
</tr>
<tr>
<td><img src="image" alt="Icon" /> <strong>Running Terminals and Kernels</strong>&lt;br&gt;You can check the list of all the <em>kernels</em> and <em>terminals</em> currently running across all notebooks, code consoles, and directories. You can shut down individual resources, including notebooks, terminals, kernels, apps, and instances. You can also shut down all resources in one of these categories at the same time.&lt;br&gt;&lt;br&gt;For more information, see <a href="#">Shut Down Resources</a>.</td>
<td></td>
</tr>
<tr>
<td><img src="image" alt="Icon" /> <strong>Git</strong>&lt;br&gt;You can connect to a Git repository and then access a full range of Git tools and operations.&lt;br&gt;&lt;br&gt;For more information, see <a href="#">Clone a Git Repository in SageMaker Studio Classic</a>.</td>
<td></td>
</tr>
<tr>
<td><img src="image" alt="Icon" /> <strong>Table of Contents</strong>&lt;br&gt;You can navigate the structure of a document when a notebook or Python files are open.&lt;br&gt;A table of contents is auto-generated in the left navigation panel when you have a notebook, Markdown files, or Python files opened. The entries are clickable and scroll the document to the heading in question.</td>
<td></td>
</tr>
<tr>
<td>Icon</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| ![Puzzle](image) | **Extensions**  
You can turn on and manage third-party JupyterLab extensions. You can check the already installed extensions and search for extensions by typing the name in the search bar. When you have found the extension you want to install, choose Install. After installing your new extensions, be sure to restart JupyterLab by refreshing your browser.  
For more information, see [JupyterLab Extensions documentation](#). |

**Left navigation panel**

The left navigation panel content varies with the Icon selected in the left sidebar.

For example, choosing the **Home** icon displays the navigation menu. Choosing **File browser** lists all the files and directories available in your workspace (notebooks, experiments, data flows, trials, trial components, endpoints, or low-code solutions).

In the navigation menu, choosing a node brings up the corresponding feature page in the main working area. For example, choosing **Data Wrangler** in the **Data** menu opens up the **Data Wrangler** tab listing all existing flows.

**Main working area**

The main working area consists of multiple tabs that contain your open notebooks, terminals, and detailed information about your experiments and endpoints. In the main working area, you can arrange documents (such as notebooks and text files) and other activities (such as terminals and code consoles) into panels of tabs that you can resize or subdivide. Drag a tab to the center of a tab panel to move the tab to the panel. Subdivide a tab panel by dragging a tab to the left, right, top, or bottom of the panel. The tab for the current activity is marked with a colored top border (blue by default).

---

**Note**

All feature pages provide in-product contextual help. To access help, choose **Show information**. The help interface provides a brief introduction to the tool and links to additional resources, such as videos, tutorials, or blogs.
Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

After you have onboarded to an Amazon SageMaker domain, you can launch an Amazon SageMaker Studio Classic application from either the SageMaker console or the AWS CLI. For more information about onboarding to a domain, see Amazon SageMaker domain overview.

Topics

- Launch Studio Classic Using the Amazon SageMaker Console
- Launch Studio Classic Using the AWS CLI

Launch Studio Classic Using the Amazon SageMaker Console

The process to navigate to Studio Classic from the Amazon SageMaker Console differs depending on if Studio Classic or Amazon SageMaker Studio are set as the default experience for your domain. For more information about setting the default experience for your domain, see Migrating from Amazon SageMaker Studio Classic.

Topics

- Prerequisite

Prerequisite

To complete this procedure, you must onboard to a domain by following the steps in Onboard to Amazon SageMaker domain.

Launch Studio Classic if Studio is your default experience

1. Navigate to Studio following the steps in Launch Amazon SageMaker Studio.
2. From the Studio UI, find the applications pane on the left side.
3. From the applications pane, select **Studio Classic**.
4. From the Studio Classic landing page, select the Studio Classic instance to open.
5. Choose “Open”.

**Launch Studio Classic if Studio Classic is your default experience**

When Studio Classic is your default experience, you can launch a Amazon SageMaker Studio Classic application from the SageMaker console using the Studio Classic landing page or the Amazon SageMaker domain details page. The following sections demonstrate how to launch the Studio Classic application from the SageMaker console.

**Launch Studio Classic from the domain details page**

The following sections describe how to launch a Studio Classic application from the domain details page. The steps to launch the Studio Classic application after you have navigated to the domain details page differ depending on if you’re launching a personal application or a shared space.

**Navigate to the domain details page**

The following procedure shows how to navigate to the domain details page.

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domain, select the domain that you want to launch the Studio Classic application in.

**Launch a user profile app**

The following procedure shows how to launch a Studio Classic application that is scoped to a user profile.

1. On the domain details page, choose the **User profiles** tab.
2. Identify the user profile that you want to launch the Studio Classic application for.
3. Choose **Launch** for your selected user profile, then choose **Studio Classic**.

**Launch a shared space app**
The following procedure shows how to launch a Studio Classic application that is scoped to a shared space.

1. On the domain details page, choose the **Space management** tab.
2. Identify the shared space that you want to launch the Studio Classic application for.
3. Choose **Launch Studio Classic** for your selected shared space.

**Launch Studio Classic from the Studio Classic landing page**

The following procedure describes how to launch a Studio Classic application from the Studio Classic landing page.

**Launch Studio Classic**

2. On the left navigation pane, choose Studio Classic.
3. Under **Get started**, select the domain that you want to launch the Studio Classic application in. If your user profile only belongs to one domain, you do not see the option for selecting a domain.
4. Select the user profile that you want to launch the Studio Classic application for. If there is no user profile in the domain, choose **Create user profile**. For more information, see Add and Remove User Profiles.
5. Choose **Launch Studio Classic**. If the user profile belongs to a shared space, choose Open Spaces.
6. To launch a Studio Classic application scoped to a user profile, choose **Launch personal Studio Classic**.
7. To launch a shared Studio Classic application, choose the **Launch shared Studio Classic** button next to the shared space that you want to launch into.

**Launch Studio Classic Using the AWS CLI**

You can use the AWS Command Line Interface (AWS CLI) to launch Amazon SageMaker Studio Classic by creating a presigned domain URL.

**Prerequisites**

Before you begin, complete the following prerequisites:
• Onboard to Amazon SageMaker domain. For more information, see Onboard to Amazon SageMaker domain.

• Update the AWS CLI by following the steps in Installing the current AWS CLI Version.

• From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.

The following code snippet demonstrates how to launch Amazon SageMaker Studio Classic from the AWS CLI using a presigned domain URL. For more information, see create-presigned-domain-url.

```
aws sagemaker create-presigned-domain-url \
  --region region \n  --domain-id domain-id \n  --space-name space-name \n  --user-profile-name user-profile-name \n  --session-expiration-duration-in-seconds 43200
```

## JupyterLab Versioning

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The Amazon SageMaker Studio Classic interface is based on JupyterLab, which is a web-based interactive development environment for notebooks, code, and data. Studio Classic now supports using both JupyterLab 1 and JupyterLab 3. The default version of JupyterLab in Studio Classic is JupyterLab 3. If you created your Amazon SageMaker domain and user profile using the AWS Management Console before 08/31/2022 or using the AWS Command Line Interface before 02/22/23, then your Studio Classic instance defaults to JupyterLab 1. After 08/31/2022, JupyterLab version 1 on Amazon SageMaker Studio Classic only receives security fixes. You can choose the version that you want to run. However, you can run only a single instance of JupyterLab at one time per user profile. You can’t run multiple versions of JupyterLab simultaneously.
After 03/31/23, Studio Classic only supports the creation of JupyterLab 3 applications. After that date, Studio Classic stops supporting JupyterLab 1 application creation. On 04/30/2023, Studio Classic removes all existing applications that run JupyterLab 1. Update your existing JupyterLab1 applications to JupyterLab 3 before 04/30/2023 following the steps in View and update the JupyterLab version of an application from the console.

Topics
- JupyterLab 3
- Restricting default JupyterLab version using an IAM policy condition key
- Setting a default JupyterLab version
- View and update the JupyterLab version of an application from the console
- Installing JupyterLab and Jupyter Server extensions

JupyterLab 3

JupyterLab 3 includes the following features that are not available in previous versions. For more information about these features, see JupyterLab 3.0 is released!

- Visual debugger when using the Base Python 2.0 and Data Science 2.0 kernels.
- File browser filter
- Table of Contents (TOC)
- Multi-language support
- Simple mode
- Single interface mode

Important changes to JupyterLab 3

Consider the following when using JupyterLab 3:

- When setting the JupyterLab version using the AWS CLI, select the corresponding image for your Region and JupyterLab version from the image list in From the AWS CLI.
- In JupyterLab 3, you must activate the studio conda environment before installing extensions. For more information, see Installing JupyterLab and Jupyter Server extensions.
- Debugger is only supported when using the following images:
Restricting default JupyterLab version using an IAM policy condition key

You can use IAM policy condition keys to restrict the version of JupyterLab that your users can launch.

The following policy shows how to limit the JupyterLab version at the domain level.

```
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "Block users from creating JupyterLab 3 apps at the domain level",
         "Effect": "Deny",
         "Action": [
            "sagemaker:CreateDomain",
            "sagemaker:UpdateDomain"
         ],
         "Resource": "*",
         "Condition": {
            "ForAnyValue:StringLike": {
               "sagemaker:ImageArns": "*image/jupyter-server-3"
            }
         }
      }
   ]
}
```

The following policy shows how to limit the JupyterLab version at the user profile level.

```
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "Block users from creating JupyterLab 3 apps at the user profile level",
         "Effect": "Deny",
         "Action": [
            "sagemaker:CreateDomain",
            "sagemaker:UpdateDomain"
         ],
         "Resource": "*",
         "Condition": {
            "ForAnyValue:StringLike": {
               "sagemaker:ImageArns": "*image/jupyter-server-3"
            }
         }
      }
   ]
}
```
The following policy shows how to limit the JupyterLab version at the application level. The `CreateApp` request must include the image ARN for this policy to apply.

```
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "Block users from creating JupyterLab 3 apps at the application level",
         "Effect": "Deny",
         "Action": "sagemaker:CreateApp",
         "Resource": "*",
         "Condition": {
            "ForAnyValue:StringLike": {
               "sagemaker:ImageArns": "*image/jupyter-server-3"
            }
         }
      }
   ]
}
```

### Setting a default JupyterLab version

The following sections show how to set a default JupyterLab version for Studio Classic using either the console or the AWS CLI.
From the console

You can select the default JupyterLab version to use on either the domain or user profile level during resource creation. To set the default JupyterLab version using the console, see Amazon SageMaker domain overview.

From the AWS CLI

You can select the default JupyterLab version to use on either the domain or user profile level using the AWS CLI.

To set the default JupyterLab version using the AWS CLI, you must include the ARN of the desired default JupyterLab version as part of an AWS CLI command. This ARN differs based on the version and the Region of the SageMaker domain.

The following table lists the ARNs of the available JupyterLab versions for each Region:

<table>
<thead>
<tr>
<th>Region</th>
<th>JL1</th>
<th>JL3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region</td>
<td>JupyterLab Versioning</td>
<td></td>
</tr>
<tr>
<td>----------------</td>
<td>--------------------------------------------------------------------------------------</td>
<td></td>
</tr>
</tbody>
</table>
Create or update domain


The following shows how to create a domain with JupyterLab 3 as the default, using the AWS CLI:

```
aws --region <REGION>
  sagemaker create-domain \
  --domain-name <NEW_DOMAIN_NAME>
```
The following shows how to update a domain to use JupyterLab 3 as the default, using the AWS CLI:

```bash
aws --region <REGION> \
  sagemaker update-domain \
  --domain-id <YOUR_DOMAIN_ID> \
  --default-user-settings '{
    "JupyterServerAppSettings": {
      "DefaultResourceSpec": {
        "InstanceType": "system"
      }
    }
  }'
```

Create or update user profile

You can set a default JupyterServer version at the user profile level by invoking `CreateUserProfile` or `UpdateUserProfile` and passing the `UserSettings.JupyterServerAppSettings.DefaultResourceSpec.SageMakerImageArn` field.

The following shows how to create a user profile with JupyterLab 3 as the default on an existing domain, using the AWS CLI:

```bash
aws --region <REGION> \
  sagemaker create-user-profile \
```

JupyterLab Versioning
The following shows how to update a user profile to use JupyterLab 3 as the default, using the AWS CLI:

```bash
aws --region <REGION> \
  sagemaker update-user-profile \
    --domain-id <YOUR_DOMAIN_ID> \
    --user-profile-name <EXISTING_USERPROFILE_NAME> \
    --user-settings '{
      "JupyterServerAppSettings": {
        "DefaultResourceSpec": {
          "InstanceType": "system"
        }
      }
    }'
```

View and update the JupyterLab version of an application from the console

The following shows how to view and update the JupyterLab version of an application.

1. Navigate to the SageMaker domains page.
2. Select a domain to view its user profiles.
3. Select a user to view their applications.
4. To view the JupyterLab version of an application, select the application's name.
5. To update the JupyterLab version, select Action.
6. From the dropdown menu, select Change JupyterLab version.
7. From the **Studio Classic settings** page, select the JupyterLab version from the dropdown menu.

8. After the JupyterLab version for the user profile has been successfully updated, restart the JupyterServer application to make the version changes effective. For more information about restarting a JupyterServer application, see [Shut down and Update SageMaker Studio Classic](#).

**Installing JupyterLab and Jupyter Server extensions**

The process for installing JupyterLab and Jupyter Server extensions differs depending on the JupyterLab version of your Studio Classic instance. In JupyterLab 1, you can open the terminal and install extensions without activating any conda environment. In JupyterLab 3, you must activate the studio conda environment before installing extensions. The method for this differs if you're installing the extensions from within Studio Classic or using a lifecycle configuration script.

**Installing Extension from within Studio Classic**

To install extensions from within Studio Classic, you must activate the studio environment before you install extensions.

```bash
# Before installing extensions
conda activate studio

# Install your extensions
pip install <JUPYTER_EXTENSION>

# After installing extensions
conda deactivate
```

**Installing Extensions using a lifecycle configuration script**

If you're installing JupyterLab and Jupyter Server extensions in your lifecycle configuration script, you must modify your script so that it works with JupyterLab 3. The following sections show the code needed for existing and new lifecycle configuration scripts.

**Existing lifecycle configuration script**

If you're reusing an existing lifecycle configuration script that must work with both versions of JupyterLab, use the following code in your script:

```bash
# Before installing extension
```
export
AWS_SAGEMAKER_JUPYTERSERVER_IMAGE="${AWS_SAGEMAKER_JUPYTERSERVER_IMAGE:-'jupyter-server'}"
if [ "$AWS_SAGEMAKER_JUPYTERSERVER_IMAGE" = "jupyter-server-3" ]; then
  eval "$(conda shell.bash hook)"
  conda activate studio
fi;

# Install your extensions
pip install <JUPYTER_EXTENSION>

# After installing extension
if [ "$AWS_SAGEMAKER_JUPYTERSERVER_IMAGE" = "jupyter-server-3" ]; then
  conda deactivate
fi;

New lifecycle configuration script

If you're writing a new lifecycle configuration script that only uses JupyterLab 3, you can use the following code in your script:

# Before installing extension
eval "$(conda shell.bash hook)"
conda activate studio

# Install your extensions
pip install <JUPYTER_EXTENSION>
conda deactivate

Use the Amazon SageMaker Studio Classic Launcher

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
You can use the Amazon SageMaker Studio Classic Launcher to create notebooks and text files, and to launch terminals and interactive Python shells.

You can open Studio Classic Launcher in any of the following ways:

- Choose **Amazon SageMaker Studio Classic** at the top left of the Studio Classic interface.
- Use the keyboard shortcut **Ctrl + Shift + L**.
- From the Studio Classic menu, choose **File** and then choose **New Launcher**.
- If the SageMaker file browser is open, choose the plus (+) sign in the Studio Classic file browser menu.
- In the **Quick actions** section of the **Home** tab, choose **Open Launcher**. The Launcher opens in a new tab. The **Quick actions** section is visible by default but can be toggled off. Choose **Customize Layout** to turn this section back on.

The Launcher consists of the following two sections:

**Topics**

- **Notebooks and compute resources**
Utilities and files

Notebooks and compute resources

In this section, you can create a notebook, open an image terminal, or open a Python console.

To create or launch one of those items:

1. Choose **Change environment** to select a SageMaker image, a kernel, an instance type, and, optionally, add a lifecycle configuration script that runs on image start-up. For more information on lifecycle configuration scripts, see Use lifecycle configurations with Amazon SageMaker Studio Classic. For more information about kernel updates, see Change an Image or a Kernel.
2. Select an item.

---

Note

When you choose an item from this section, you might incur additional usage charges. For more information, see Usage Metering.

---

The following items are available:

- **Notebook**
  
  Launches the notebook in a kernel session on the chosen SageMaker image.

  Creates the notebook in the folder that you have currently selected in the file browser. To view the file browser, in the left sidebar of Studio Classic, choose the File Browser icon.

- **Console**

  Launches the shell in a kernel session on the chosen SageMaker image.

  Opens the shell in the folder that you have currently selected in the file browser.

- **Image terminal**

  Launches the terminal in a terminal session on the chosen SageMaker image.

  Opens the terminal in the root folder for the user (as shown by the Home folder in the file browser).
## Note

By default, CPU instances launch on a `ml.t3.medium` instance, while GPU instances launch on a `ml.g4dn.xlarge` instance.

## Utilities and files

In this section, you can add contextual help in a notebook; create Python, Markdown and text files; and open a system terminal.

### Note

Items in this section run in the context of Amazon SageMaker Studio Classic and don't incur usage charges.

The following items are available:

- **Show Contextual Help**

  Opens a new tab that displays contextual help for functions in a Studio Classic notebook. To display the help, choose a function in an active notebook. To make it easier to see the help in context, drag the help tab so that it's adjacent to the notebook tab. To open the help tab from within a notebook, press Ctrl + I.

  The following screenshot shows the contextual help for the `Experiment.create` method.
• **System terminal**

Opens a bash shell in the root folder for the user (as shown by the **Home** folder in the file browser).

• **Text File and Markdown File**

Creates a file of the associated type in the folder that you have currently selected in the file browser. To view the file browser, in the left sidebar, choose the **File Browser** icon (folder icon).
Collaborate with shared spaces

Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

An Amazon SageMaker Studio Classic shared space consists of a shared JupyterServer application and a shared directory. All user profiles in a domain have access to all shared spaces in the domain. Amazon SageMaker automatically scopes resources in a shared space within the context of the Amazon SageMaker Studio Classic application that you launch in that shared space. Resources in a shared space include notebooks, files, experiments, and models.

A Studio Classic shared space only supports Studio Classic and KernelGateway applications. A shared space only supports the use of a JupyterLab 3 image Amazon Resource Name (ARN). For more information, see JupyterLab Versioning.

Amazon SageMaker automatically tags all SageMaker resources that you create within the scope of a shared space. You can use these tags to monitor costs and plan budgets using tools, such as AWS Budgets.

A shared space uses the same VPC settings as the domain that it's created in.

Note

Shared spaces do not support the use of Amazon SageMaker Data Wrangler or Amazon EMR cross-account clusters.

Automatic tagging

All resources created in a shared space are automatically tagged with a domain ARN tag and shared space ARN tag. The domain ARN tag is based on the domain ID, while the shared space ARN tag is based on the shared space name.

You can use these tags to monitor AWS CloudTrail usage. For more information, see Log Amazon SageMaker API Calls with AWS CloudTrail.
You can also use these tags to monitor costs with AWS Billing and Cost Management. For more information, see Using AWS cost allocation tags.

**Real time co-editing of notebooks**

A key benefit of a shared space is that it facilitates collaboration between members of the shared space in real time. Users collaborating in a workspace get access to a shared Studio Classic application where they can access, read, and edit their notebooks in real time. Real time collaboration is only supported for JupyterServer applications within a shared space.

Users with access to a shared space can simultaneously open, view, edit, and execute Jupyter notebooks in the shared Studio Classic application in that space.

The notebook indicates each co-editing user with a different cursor that shows the user profile name. While multiple users can view the same notebook, co-editing is best suited for small groups of two to five users.

To track changes being made by multiple users, we strongly recommended using Studio Classic's built-in Git-based version control.

**JupyterServer 2**

To use shared spaces, Jupyter Server version 2 is required. Certain JupyterLab extensions and packages can forcefully downgrade Jupyter Server to version 1. This prevents the use of shared space. Run the following from the command prompt to change the version number and continue using shared spaces.

```
conda activate studio
pip install jupyter-server==2.0.0rc3
```

**Customize a shared space**

To attach a lifecycle configuration or custom image to a shared space, you must use the AWS CLI. For more information about creating and attaching lifecycle configurations, see Create and associate a lifecycle configuration. For more information about creating and attaching custom images, see Bring your own SageMaker image.
Create a shared space

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following topic demonstrates how to create an Amazon SageMaker Studio Classic shared space in an existing Amazon SageMaker domain. If you created your domain without support for shared spaces, you must add support for shared spaces to your existing domain before you can create a shared space.

Topics

- Add shared space support to an existing domain
- Create a shared space

Add shared space support to an existing domain

You can use the SageMaker console or the AWS CLI to add support for shared spaces to an existing domain. If the domain is using VPC only network access, then you can only add shared space support using the AWS CLI.

Console

Complete the following procedure to add support for shared spaces to an existing domain from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to open the domain settings page for.
5. On the domain details page, choose the domain settings tab.
6. Choose Edit.
7. For **Space default execution role**, set an IAM role that is used by default for all shared spaces created in the domain.

8. Choose **Next**.

9. Choose **Next**.

10. Choose **Next**.

11. Choose **Submit**.

**AWS CLI**

Run the following command from the terminal of your local machine to add default shared space settings to a domain from the AWS CLI. If you are adding default shared space settings to a domain within an Amazon VPC, you must also include a list of security groups. Shared spaces only support the use of JupyterLab 3 image ARNs. For more information, see [JupyterLab Versioning](#).

```bash
# Public Internet domain
aws --region region \n sagemaker update-domain \n   --domain-id domain-id \n   --default-space-settings "ExecutionRole=execution-role-arn, JupyterServerAppSettings={DefaultResourceSpec={InstanceType=system, SageMakerImageArn=sagemaker-image-arn}}"

# VPCOnly domain
aws --region region \n sagemaker update-domain \n   --domain-id domain-id \n   --default-space-settings "ExecutionRole=execution-role-arn, JupyterServerAppSettings={DefaultResourceSpec={InstanceType=system, SageMakerImageArn=sagemaker-image-arn}}, SecurityGroups=[security-groups]"
```

Verify that the default shared space settings have been updated.

```bash
aws --region region \n sagemaker describe-domain \n   --domain-id domain-id
```
Create a shared space

The following sections demonstrate how to create a shared space from the Amazon SageMaker console, Amazon SageMaker Studio, or the AWS CLI.

Create from Studio

Complete the following procedure to create a shared space in a domain from Studio.

1. Navigate to Studio following the steps in Launch Amazon SageMaker Studio.
2. From the Studio UI, find the applications pane on the left side.
3. From the applications pane, select Studio Classic.
4. Choose Create Studio Classic space
5. In the pop up window, enter a name for the space.
6. Choose Create space.

Create from the console

Complete the following procedure to create a shared space in a domain from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to create a shared space for.
5. On the domain details page, choose the Space management tab.
6. Choose Create.
7. Enter a name for your shared space. shared space names within a domain must be unique. The execution role for the shared space is set to the domain IAM execution role.

Create from AWS CLI

This section shows how to create a shared space from the AWS CLI.

You cannot set the execution role of a shared space when creating or updating it. The DefaultDomainExecRole can only be set when creating or updating the domain.
spaces only support the use of JupyterLab 3 image ARNs. For more information, see JupyterLab Versioning.

To create a shared space from the AWS CLI, run the following command from the terminal of your local machine.

```bash
aws --region region \
sagemaker create-space \
--domain-id domain-id \
--space-name space-name \
--space-settings '{
  "JupyterServerAppSettings": {
    "DefaultResourceSpec": {
      "SageMakerImageArn": "sagemaker-image-arn",
      "InstanceType": "system"
    }
  }
}'
```

List and Describe shared spaces

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

This guide shows how to access a list of Amazon SageMaker Studio Classic shared spaces in an Amazon SageMaker domain with the Amazon SageMaker console, Amazon SageMaker Studio, or the AWS CLI. It also shows how to view details of a shared space from the AWS CLI.

Topics

- List shared spaces
- View shared space details
List shared spaces

The following topic describes how to view a list of shared spaces within a domain from the SageMaker console or the AWS CLI.

List shared spaces from Studio

Complete the following procedure to view a list of the shared spaces in a domain from Studio.

1. Navigate to Studio following the steps in Launch Amazon SageMaker Studio.
2. From the Studio UI, find the applications pane on the left side.
3. From the applications pane, select Studio Classic. This page lists all of the Studio Classic spaces in the domain that you have access to.

List shared spaces from the console

Complete the following procedure to view a list of the shared spaces in a domain from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to view the list of shared spaces for.
5. On the domain details page, choose the Space management tab.

List shared spaces from the AWS CLI

To list the shared spaces in a domain from the AWS CLI, run the following command from the terminal of your local machine.

```
aws --region region \
  sagemaker list-spaces \
  --domain-id domain-id
```

View shared space details

The following section describes how to view shared space details from the SageMaker console, Studio, or the AWS CLI.
View shared spaces details from Studio

Complete the following procedure to view the details of a shared spaces in a domain from Studio.

1. Navigate to Studio following the steps in Launch Amazon SageMaker Studio.
2. From the Studio UI, find the applications pane on the left side.
3. From the applications pane, select Studio Classic. This page lists all of the Studio Classic spaces in the domain that you have access to.
4. Select the name of the space that you want to view more details for.

View shared space details from the console

You can view the details of a shared space from the SageMaker console using the following procedure.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to view the list of shared spaces for.
5. On the domain details page, choose the Space management tab.
6. Select the name of the space to open a new page that lists details about the shared space.

View shared space details from the AWS CLI

To view the details of a shared space from the AWS CLI, run the following command from the terminal of your local machine.

```
aws --region region \n    sagemaker describe-space \n    --domain-id domain-id \n    --space-name space-name
```
Edit a shared space

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You can only edit the details for an Amazon SageMaker Studio Classic shared space using the AWS CLI. This is not currently supported from the Amazon SageMaker console. You can only update workspace attributes when there are no running applications in the shared space.

To edit the details of a shared space from the AWS CLI, run the following command from the terminal of your local machine. Shared spaces only support the use of JupyterLab 3 image ARNs. For more information, see JupyterLab Versioning.

```bash
aws --region region \ 
sagemaker update-space \ 
--domain-id domain-id \ 
--space-name space-name \ 
--query SpaceArn --output text \ 
--space-settings '{
    "JupyterServerAppSettings": {
        "DefaultResourceSpec": {
            "SageMakerImageArn": "sagemaker-image-arn",
            "InstanceType": "system"
        }
    }
}'
```

Delete a shared space

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the
The following topic shows how to delete an Amazon SageMaker Studio Classic shared space from the Amazon SageMaker console or AWS CLI. A shared space can only be deleted if it has no running applications.

**Topics**

- [Console](#)
- [AWS CLI](#)

**Console**

Complete the following procedure to delete a shared space in the Amazon SageMaker domain from the SageMaker console.

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the domain that you want to create a shared space for.
5. On the **domain details** page, choose the **Space management** tab.
6. Select the shared space that you want to delete. The shared space must not contain any non-failed apps.
7. Choose **Delete**. This opens a new window.
8. Choose **Yes, delete space**.
9. Enter **delete** in the field.
10. Choose **Delete space**.

**AWS CLI**

To delete a shared space from the AWS CLI, run the following command from the terminal of your local machine.

```bash
aws --region region
```
Use Amazon SageMaker Studio Classic Notebooks

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker Studio Classic notebooks are collaborative notebooks that you can launch quickly because you don't need to set up compute instances and file storage beforehand. A set of instance types, known as Fast launch types are designed to launch in under two minutes. Studio Classic notebooks provide persistent storage, which enables you to view and share notebooks even if the instances that the notebooks run on are shut down.

You can share your notebooks with others, so that they can easily reproduce your results and collaborate while building models and exploring your data. You provide access to a read-only copy of the notebook through a secure URL. Dependencies for your notebook are included in the notebook's metadata. When your colleagues copy the notebook, it opens in the same environment as the original notebook.

A Studio Classic notebook runs in an environment defined by the following:

- Amazon EC2 instance type – The hardware configuration the notebook runs on. The configuration includes the number and type of processors (vCPU and GPU), and the amount and type of memory. The instance type determines the pricing rate.
- SageMaker image – A container image that is compatible with SageMaker Studio Classic. The image consists of the kernels, language packages, and other files required to run a notebook in Studio Classic. There can be multiple images in an instance. For more information, see Bring your own SageMaker image.
- KernelGateway app – A SageMaker image runs as a KernelGateway app. The app provides access to the kernels in the image. There is a one-to-one correspondence between a SageMaker image and a KernelGateway app.
• Kernel – The process that inspects and runs the code contained in the notebook. A kernel is defined by a kernel spec in the image. There can be multiple kernels in an image.

You can change any of these resources from within the notebook.

The following diagram outlines how a notebook kernel runs in relation to the KernelGateway App, User, and domain.

Sample SageMaker Studio Classic notebooks are available in the aws_sagemaker_studio folder of the Amazon SageMaker example GitHub repository. Each notebook comes with the necessary SageMaker image that opens the notebook with the appropriate kernel.

We recommend that you familiarize yourself with the SageMaker Studio Classic interface and the Studio Classic notebook toolbar before creating or using a Studio Classic notebook. For more information, see Amazon SageMaker Studio Classic UI Overview and Use the Studio Classic Notebook Toolbar.
How Are Amazon SageMaker Studio Classic Notebooks Different from Notebook Instances?

Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

When you're starting a new notebook, we recommend that you create the notebook in Amazon SageMaker Studio Classic instead of launching a notebook instance from the Amazon SageMaker console. There are many benefits to using a Studio Classic notebook, including the following:

- **Faster**: Starting a Studio Classic notebook is faster than launching an instance-based notebook. Typically, it is 5-10 times faster than instance-based notebooks.
- **Easy notebook sharing**: Notebook sharing is an integrated feature in Studio Classic. Users can generate a shareable link that reproduces the notebook code and also the SageMaker image required to execute it, in just a few clicks.
• **Latest Python SDK:** Studio Classic notebooks come pre-installed with the latest Amazon SageMaker Python SDK.

• **Access all Studio Classic features:** Studio Classic notebooks are accessed from within Studio Classic. This enables you to build, train, debug, track, and monitor your models without leaving Studio Classic.

• **Persistent user directories:** Each member of a Studio team gets their own home directory to store their notebooks and other files. The directory is automatically mounted onto all instances and kernels as they're started, so their notebooks and other files are always available. The home directories are stored in Amazon Elastic File System (Amazon EFS) so that you can access them from other services.

• **Direct access:** When using IAM Identity Center, you use your IAM Identity Center credentials through a unique URL to directly access Studio Classic. You don't have to interact with the AWS Management Console to run your notebooks.

• **Optimized images:** Studio Classic notebooks are equipped with a set of predefined SageMaker image settings to get you started faster.

---

**Note**

Studio Classic notebooks don't support *local mode*. However, you can use a notebook instance to train a sample of your dataset locally, and then use the same code in a Studio Classic notebook to train on the full dataset.

---

When you open a notebook in SageMaker Studio Classic, the view is an extension of the JupyterLab interface. The primary features are the same, so you'll find the typical features of a Jupyter notebook and JupyterLab. For more information about the Studio Classic interface, see Amazon SageMaker Studio Classic UI Overview.

---

**Get Started**

---

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
To get started, you or your organization's administrator need to complete the SageMaker domain onboarding process. For more information, see Amazon SageMaker domain overview.

You can access a Studio Classic notebook in any of the following ways:

- You receive an email invitation to access Studio Classic through your organization's IAM Identity Center, which includes a direct link to login to Studio Classic without having to use the Amazon SageMaker console. You can proceed to the the section called “Next Steps”.
- You receive a link to a shared Studio Classic notebook, which includes a direct link to log in to Studio Classic without having to use the SageMaker console. You can proceed to the the section called “Next Steps”.
- You onboard to a domain and then log in to the SageMaker console. For more information, see Amazon SageMaker domain overview.

Launch Amazon SageMaker

Complete the steps in Launch Amazon SageMaker Studio Classic to launch Studio Classic.

Next Steps

Now that you're in Studio Classic, you can try any of the following options:

- To create a Studio Classic notebook or explore Studio Classic end-to-end tutorial notebooks – See Amazon SageMaker Studio Classic Tour in the next section.
- To familiarize yourself with the Studio Classic interface – See Amazon SageMaker Studio Classic UI Overview or try the Getting started notebook by selecting Open the Getting started notebook in the Quick actions section of the Studio Classic Home page.

Amazon SageMaker Studio Classic Tour

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
For a walkthrough that takes you on a tour of the main features of Amazon SageMaker Studio Classic, see the `xgboost_customer_churn_studio.ipynb` sample notebook from the `aws/amazon-sagemaker-examples` GitHub repository. The code in the notebook trains multiple models and sets up the SageMaker Debugger and SageMaker Model Monitor. The walkthrough shows you how to view the trials, compare the resulting models, show the debugger results, and deploy the best model using the Studio Classic UI. You don't need to understand the code to follow this walkthrough.

**Prerequisites**

To run the notebook for this tour, you need:

- An IAM account to sign in to Studio. For information, see [Amazon SageMaker domain overview](#).
- Basic familiarity with the Studio user interface and Jupyter notebooks. For information, see [Amazon SageMaker Studio Classic UI Overview](#).
- A copy of the `aws/amazon-sagemaker-examples` repository in your Studio environment.

**To clone the repository**

1. Launch Studio Classic following the steps in [Launch Amazon SageMaker Studio Classic](#). For users in IAM Identity Center, sign in using the URL from your invitation email.
2. On the top menu, choose **File**, then **New**, then **Terminal**.
3. At the command prompt, run the following command to clone the `aws/amazon-sagemaker-examples` GitHub repository.

   ```bash
   $ git clone https://github.com/aws/amazon-sagemaker-examples.git
   ```

**To navigate to the sample notebook**

1. From the **File Browser** on the left menu, select `amazon-sagemaker-examples`.
2. Navigate to the example notebook with the following path.

   ```bash
   ~/amazon-sagemaker-examples/aws_sagemaker_studio/getting_started/xgboost_customer_churn_studio.ipynb
   ```
3. Follow the notebook to learn about Studio Classic's main features.
Create or Open an Amazon SageMaker Studio Classic Notebook

Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

When you Create a Notebook from the File Menu in Amazon SageMaker Studio Classic or Open a notebook in Studio Classic for the first time, you are prompted to set up your environment by choosing a SageMaker image, a kernel, an instance type, and, optionally, a lifecycle configuration script that runs on image start-up. SageMaker launches the notebook on an instance of the chosen type. By default, the instance type is set to ml.t3.medium (available as part of the AWS Free Tier) for CPU-based images. For GPU-based images, the default instance type is ml.g4dn.xlarge.

If you create or open additional notebooks that use the same instance type, whether or not the notebooks use the same kernel, the notebooks run on the same instance of that instance type.

After you launch a notebook, you can change its instance type, SageMaker image, and kernel from within the notebook. For more information, see Change an Instance Type and Change an Image or a Kernel.

Note
You can have only one instance of each instance type. Each instance can have multiple SageMaker images running on it. Each SageMaker image can run multiple kernels or terminal instances.
Billing occurs per instance and starts when the first instance of a given instance type is launched. If you want to create or open a notebook without the risk of incurring charges, open the notebook from the File menu and choose No Kernel from the Select Kernel dialog box. You can read and edit a notebook without a running kernel but you can't run cells.

Billing ends when the SageMaker image for the instance is shut down. For more information, see Usage Metering.

For information about shutting down the notebook, see Shut Down Resources.

Topics

- Open a notebook in Studio Classic
- Create a Notebook from the File Menu
- Create a Notebook from the Launcher
- List of the available instance types, images, and kernels

Open a notebook in Studio Classic

Amazon SageMaker Studio Classic can only open notebooks listed in the Studio Classic file browser. For instructions on uploading a notebook to the file browser, see Upload Files to SageMaker Studio Classic or Clone a Git Repository in SageMaker Studio Classic.

To open a notebook

1. In the left sidebar, choose the File Browser icon ( ) to display the file browser.
2. Browse to a notebook file and double-click it to open the notebook in a new tab.

Create a Notebook from the File Menu

To create a notebook from the File menu

1. From the Studio Classic menu, choose File, choose New, and then choose Notebook.
2. In the Change environment dialog box, use the dropdown menus to select your Image, Kernel, Instance type, and Start-up script, then choose Select. Your notebook launches and opens in a new Studio Classic tab.
Create a Notebook from the Launcher

To create a notebook from the Launcher

1. To open the Launcher, choose Amazon SageMaker Studio Classic at the top left of the Studio Classic interface or use the keyboard shortcut Ctrl + Shift + L.

   To learn about all the available ways to open the Launcher, see Use the Amazon SageMaker Studio Classic Launcher

2. In the Launcher, in the Notebooks and compute resources section, choose Change environment.
3. In the **Change environment** dialog box, use the dropdown menus to select your **Image**, **Kernel**, **Instance type**, and **Start-up script**, then choose **Select**.

4. In the Launcher, choose **Create notebook**. Your notebook launches and opens in a new Studio Classic tab.

To view the notebook's kernel session, in the left sidebar, choose the **Running Terminals and Kernels** icon ()

You can stop the notebook's kernel session from this view.

**List of the available instance types, images, and kernels**

For a list of all available resources, see:

- [Available Studio Classic Instance Types](#)
- [Available Amazon SageMaker Images](#)

**Use the Studio Classic Notebook Toolbar**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the
Amazon SageMaker Studio Classic notebooks extend the JupyterLab interface. For an overview of the original JupyterLab interface, see The JupyterLab Interface.

The following image shows the toolbar and an empty cell from a Studio Classic notebook.

When you pause on a toolbar icon, a tooltip displays the icon function. Additional notebook commands are found in the Studio Classic main menu. The toolbar includes the following icons:

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="icon" alt="Save and checkpoint" /></td>
<td><strong>Save and checkpoint</strong>&lt;br&gt;Saves the notebook and updates the checkpoint file. For more information, see <a href="#">Get the Difference Between the Last Checkpoint</a>.</td>
</tr>
<tr>
<td><img src="icon" alt="Insert cell" /></td>
<td><strong>Insert cell</strong>&lt;br&gt;Inserts a code cell below the current cell. The current cell is noted by the blue vertical marker in the left margin.</td>
</tr>
<tr>
<td><img src="icon" alt="Cut, copy, and paste cells" /></td>
<td><strong>Cut, copy, and paste cells</strong>&lt;br&gt;Cuts, copies, and pastes the selected cells.</td>
</tr>
<tr>
<td><img src="icon" alt="Run cells" /></td>
<td><strong>Run cells</strong>&lt;br&gt;Runs the selected cells and then makes the cell that follows the last selected cell the new selected cell.</td>
</tr>
<tr>
<td><img src="icon" alt="Interrupt kernel" /></td>
<td><strong>Interrupt kernel</strong></td>
</tr>
<tr>
<td>Icon</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
</tr>
<tr>
<td><img src="image.png" alt="Interrupt" /></td>
<td>Interrupts the kernel, which cancels the currently running operation. The kernel remains active.</td>
</tr>
</tbody>
</table>
| ![Restart kernel](image.png) | **Restart kernel**
Restarts the kernel. Variables are reset. Unsaved information is not affected. |
| ![Restart kernel and run all cells](image.png) | **Restart kernel and run all cells**
Restarts the kernel, then run all the cells of the notebook. |
| ![Cell type](image.png) | **Cell type**
Displays or changes the current cell type. The cell types are:
- Code – Code that the kernel runs.
- Markdown – Text rendered as markdown.
- Raw – Content, including Markdown markup, that's displayed as text. |
| ![Launch terminal](image.png) | **Launch terminal**
Launches a terminal in the SageMaker image hosting the notebook. For an example, see [Get App Metadata](#).

| ![Checkpoint diff](image.png) | **Checkpoint diff**
Opens a new tab that displays the difference between the notebook and the checkpoint file. For more information, see [Get the Difference Between the Last Checkpoint](#).

| ![Git diff](image.png) | **Git diff**
Only enabled if the notebook is opened from a Git repository. Opens a new tab that displays the difference between the notebook and the last Git commit. For more information, see [Get the Difference Between the Last Commit](#).
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2 vCPU + 4 GiB</strong></td>
<td><strong>Instance type</strong>&lt;br&gt;Displays or changes the instance type the notebook runs in. The format is as follows:&lt;br&gt;&lt;br&gt;number of vCPUs + amount of memory + number of GPUs&lt;br&gt;&lt;br&gt;Unknown indicates the notebook was opened without specifying a kernel. The notebook runs on the SageMaker Studio instance and doesn't accrue runtime charges. You can't assign the notebook to an instance type. You must specify a kernel and then Studio assigns the notebook to a default type. &lt;br&gt;&lt;br&gt;For more information, see <a href="#">Create or Open an Amazon SageMaker Studio Classic Notebook</a> and <a href="#">Change an Instance Type</a>.</td>
</tr>
<tr>
<td><strong>Cluster</strong></td>
<td><strong>Cluster</strong>&lt;br&gt;Connect your notebook to an Amazon EMR cluster and scale your ETL jobs or run large-scale model training using Apache Spark, Hive, or Presto. &lt;br&gt;&lt;br&gt;For more information, see <a href="#">Prepare data using Amazon EMR</a>.</td>
</tr>
<tr>
<td><strong>Python 3 (Data Science)</strong></td>
<td><strong>Kernel and SageMaker Image</strong>&lt;br&gt;Displays or changes the kernel that processes the cells in the notebook. The format is as follows:&lt;br&gt;&lt;br&gt;Kernel (SageMaker Image)&lt;br&gt;&lt;br&gt;No Kernel indicates the notebook was opened without specifying a kernel. You can edit the notebook but you can't run any cells. &lt;br&gt;&lt;br&gt;For more information, see <a href="#">Change an Image or a Kernel</a>.</td>
</tr>
<tr>
<td>Icon</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
</tr>
<tr>
<td>🔄</td>
<td>Kernel busy status</td>
</tr>
</tbody>
</table>

Displays the busy status of the kernel. When the edge of the circle and its interior are the same color, the kernel is busy. The kernel is busy when it is starting and when it is processing cells. Additional kernel states are displayed in the status bar at the bottom-left corner of SageMaker Studio.

<table>
<thead>
<tr>
<th>Share</th>
<th>Share notebook</th>
</tr>
</thead>
</table>

Shares the notebook. For more information, see [Share and Use an Amazon SageMaker Studio Classic Notebook](#).

To select multiple cells, click in the left margin outside of a cell. Hold down the Shift key and use K or the Up key to select previous cells, or use J or the Down key to select following cells.

## Install External Libraries and Kernels in Amazon SageMaker Studio Classic

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

Amazon SageMaker Studio Classic notebooks come with multiple images already installed. These images contain kernels and Python packages including scikit-learn, Pandas, NumPy, TensorFlow, PyTorch, and MXNet. You can also install your own images that contain your choice of packages and kernels. For more information on installing your own image, see [Bring your own SageMaker image](#).

The different Jupyter kernels in Amazon SageMaker Studio Classic notebooks are separate conda environments. For information about conda environments, see [Managing environments](#).
Package installation tools

The method that you use to install Python packages from the terminal differs depending on the image. Studio Classic supports the following package installation tools:

- **Notebooks** – The following commands are supported. If one of the following does not work on your image, try the other one.
  - `%conda install`
  - `%pip install`
- **The Jupyter terminal** – You can install packages using pip and conda directly. You can also use `apt-get install` to install system packages from the terminal.

Note

We do not recommend using `pip install -u` or `pip install --user`, because those commands install packages on the user's Amazon EFS volume and can potentially block JupyterServer app restarts. Instead, use a lifecycle configuration to reinstall the required packages on app restarts as shown in [Install packages using lifecycle configurations](#).

We recommend using `%pip` and `%conda` to install packages from within a notebook because they correctly take into account the active environment or interpreter being used. For more information, see [Add `%pip` and `%conda` magic functions](#). You can also use the system command syntax (lines starting with `!`) to install packages. For example, `!pip install` and `!conda install`.

Conda

Conda is an open source package management system and environment management system that can install packages and their dependencies. SageMaker supports using conda with either of these two main channels: the default channel or the conda-forge channel. For more information, see [Conda channels](#). The conda-forge channel is a community channel where contributors can upload packages.

Note

Installing packages from conda-forge can take up to 10 minutes. Timing relates to how conda resolves the dependency graph.
All of the SageMaker provided environments are functional. User installed packages may not function correctly.

Conda has two methods for activating environments: `conda activate` and `source activate`. For more information, see Managing environment.

**Supported conda operations**

- `conda install` of a package in a single environment
- `conda install` of a package in all environments
- Installing a package from the main conda repository
- Installing a package from conda-forge
- Changing the conda install location to use Amazon EBS
- Supporting both `conda activate` and `source activate`

**Pip**

Pip is the tool for installing and managing Python packages. Pip searches for packages on the Python Package Index (PyPI) by default. Unlike conda, pip doesn't have built in environment support. Therefore, pip isn't as thorough as conda when it comes to packages with native or system library dependencies. Pip can be used to install packages in conda environments. You can use alternative package repositories with pip instead of the PyPI.

**Supported pip operations**

- Using pip to install a package without an active conda environment
- Using pip to install a package in a conda environment
- Using pip to install a package in all conda environments
- Changing the pip install location to use Amazon EBS
- Using an alternative repository to install packages with pip

**Unsupported**

SageMaker aims to support as many package installation operations as possible. However, if the packages were installed by SageMaker and you use the following operations on these packages, it might make your environment unstable:
• Uninstalling
• Downgrading
• Upgrading

Due to potential issues with network conditions or configurations, or the availability of conda or PyPi, packages may not install in a fixed or deterministic amount of time.

Note

Attempting to install a package in an environment with incompatible dependencies can result in a failure. If issues occur, you can contact the library maintainer about updating the package dependencies. When you modify the environment, such as removing or updating existing packages, this may result in instability of that environment.

Install packages using lifecycle configurations

Install custom images and kernels on the Studio Classic instance's Amazon EBS volume so that they persist when you stop and restart the notebook, and that any external libraries you install are not updated by SageMaker. To do that, use a lifecycle configuration that includes both a script that runs when you create the notebook (on-create) and a script that runs each time you restart the notebook (on-start). For more information about using lifecycle configurations with Studio Classic, see Use lifecycle configurations with Amazon SageMaker Studio Classic. For sample lifecycle configuration scripts, see SageMaker Studio Classic Lifecycle Configuration Samples.

Share and Use an Amazon SageMaker Studio Classic Notebook

Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You can share your Amazon SageMaker Studio Classic notebooks with your colleagues. The shared notebook is a copy. After you share your notebook, any changes you make to your original
notebook aren't reflected in the shared notebook and any changes your colleague's make in their shared copies of the notebook aren't reflected in your original notebook. If you want to share your latest version, you must create a new snapshot and then share it.

Topics
- Share a Notebook
- Use a Shared Notebook
- Shared spaces and realtime collaboration

Share a Notebook

The following screenshot shows the menu from a Studio Classic notebook.

To share a notebook

1. In the upper-right corner of the notebook, choose Share.
2. (Optional) In Create shareable snapshot, choose any of the following items:
   - Include Git repo information – Includes a link to the Git repository that contains the notebook. This enables you and your colleague to collaborate and contribute to the same Git repository.
   - Include output – Includes all notebook output that has been saved.

   ![Share](image)

   **Note**

   If you're an user in IAM Identity Center and you don't see these options, your IAM Identity Center administrator probably disabled the feature. Contact your administrator.

3. Choose Create.
4. After the snapshot is created, choose Copy link and then choose Close.
5. Share the link with your colleague.
After selecting your sharing options, you are provided with a URL. You can share this link with users that have access to Amazon SageMaker Studio Classic. When the user opens the URL, they're prompted to log in using IAM Identity Center or IAM authentication. This shared notebook becomes a copy, so changes made by the recipient will not be reproduced in your original notebook.

**Use a Shared Notebook**

You use a shared notebook in the same way you would with a notebook that you created yourself. You must first login to your account, then open the shared link. If you don't have an active session, you receive an error.

When you choose a link to a shared notebook for the first time, a read-only version of the notebook opens. To edit the shared notebook, choose Create a Copy. This copies the shared notebook to your personal storage.

The copied notebook launches on an instance of the instance type and SageMaker image that the notebook was using when the sender shared it. If you aren't currently running an instance of the instance type, a new instance is started. Customization to the SageMaker image isn't shared. You can also inspect the notebook snapshot by choosing Snapshot Details.

The following are some important considerations about sharing and authentication:

- If you have an active session, you see a read-only view of the notebook until you choose Create a Copy.
- If you don't have an active session, you need to log in.
- If you use IAM to login, after you login, select your user profile then choose Open Studio Classic. Then you need to choose the link you were sent.
- If you use IAM Identity Center to login, after you login the shared notebook is opened automatically in Studio.

**Shared spaces and realtime collaboration**

A shared space consists of a shared JupyterServer application and a shared directory. A key benefit of a shared space is that it facilitates collaboration between members of the shared space in real time. Users collaborating in a workspace get access to a shared Studio Classic application where they can access, read, and edit their notebooks in real time. Real time collaboration is only supported for JupyterServer applications within a shared space. Users with access to a shared space can simultaneously open, view, edit, and execute Jupyter notebooks in the shared Studio Classic.
application in that space. For more information about shared spaced and real time collaboration, see [Collaborate with shared spaces](#).

### Get Studio Classic Notebook and App Metadata

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

You can access notebook metadata and App metadata using the Amazon SageMaker Studio Classic UI.

#### Topics

- [Get Studio Classic Notebook Metadata](#)
- [Get App Metadata](#)

### Get Studio Classic Notebook Metadata

Jupyter notebooks contain optional metadata that you can access through the Amazon SageMaker Studio Classic UI.

**To view the notebook metadata:**

1. In the right sidebar, choose the **Property Inspector** icon (].[231x231])
2. Open the **Advanced Tools** section.

The metadata should look similar to the following.

```
{
    "instance_type": "ml.t3.medium",
    "kernelspec": {
        "display_name": "Python 3 (Data Science)"
    },
```
Get App Metadata

When you create a notebook in Amazon SageMaker Studio Classic, the App metadata is written to a file named `resource-metadata.json` in the folder `/opt/ml/metadata/`. You can get the App metadata by opening an Image terminal from within the notebook. The metadata gives you the following information, which includes the SageMaker image and instance type the notebook runs in:

- **AppType** – KernelGateway
- **DomainId** – Same as the Studio ClassicID
- **UserProfileName** – The profile name of the current user
- **ResourceArn** – The Amazon Resource Name (ARN) of the App, which includes the instance type
- **ResourceName** – The name of the SageMaker image

Additional metadata might be included for internal use by Studio Classic and is subject to change.

**To get the App metadata**

1. In the center of the notebook menu, choose the **Launch Terminal** icon ( ).

   This opens a terminal in the SageMaker image that the notebook runs in.
2. Run the following commands to display the contents of the `resource-metadata.json` file.

```
$ cd /opt/ml/metadata/
cat resource-metadata.json
```

The file should look similar to the following.

```
{
    "AppType": "KernelGateway",
    "DomainId": "d-xxxxxxxxxxxx",
    "UserProfileName": "profile-name",
    "ResourceName": "datascience--1-0-ml",
    "AppImageVersion": ""
}
```

### Get Notebook Differences

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

You can display the difference between the current notebook and the last checkpoint or the last Git commit using the Amazon SageMaker UI.

The following screenshot shows the menu from a Studio Classic notebook.

### Topics

- [Get the Difference Between the Last Checkpoint](#)
• **Get the Difference Between the Last Commit**

**Get the Difference Between the Last Checkpoint**

When you create a notebook, a hidden checkpoint file that matches the notebook is created. You can view changes between the notebook and the checkpoint file or revert the notebook to match the checkpoint file.

By default, a notebook is auto-saved every 120 seconds and also when you close the notebook. However, the checkpoint file isn't updated to match the notebook. To save the notebook and update the checkpoint file to match, you must choose the **Save notebook and create checkpoint** icon ( ) on the left of the notebook menu or use the Ctrl + S keyboard shortcut.

To view the changes between the notebook and the checkpoint file, choose the **Checkpoint diff** icon ( ) in the center of the notebook menu.

To revert the notebook to the checkpoint file, from the main Studio Classic menu, choose **File** then **Revert Notebook to Checkpoint**.

**Get the Difference Between the Last Commit**

If a notebook is opened from a Git repository, you can view the difference between the notebook and the last Git commit.

To view the changes in the notebook from the last Git commit, choose the **Git diff** icon ( ) in the center of the notebook menu.

**Manage Resources**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the
You can change the instance type, and SageMaker image and kernel from within an Amazon SageMaker Studio Classic notebook. To create a custom kernel to use with your notebooks, see Bring your own SageMaker image.

**Topics**

- Change an Instance Type
- Change an Image or a Kernel
- Shut Down Resources

**Change an Instance Type**

When you open a new Studio Classic notebook for the first time, you are assigned a default Amazon Elastic Compute Cloud (Amazon EC2) instance type to run the notebook. When you open additional notebooks on the same instance type, the notebooks run on the same instance as the first notebook, even if the notebooks use different kernels.

You can change the instance type that your Studio Classic notebook runs on from within the notebook.

The following information only applies to Studio Classic notebooks. For information about how to change the instance type of a Amazon SageMaker notebook instance, see Update a Notebook Instance.

⚠️ **Important**

If you change the instance type, unsaved information and existing settings for the notebook are lost, and installed packages must be re-installed.

The previous instance type continues to run even if no kernel sessions or apps are active. You must explicitly stop the instance to stop accruing charges. To stop the instance, see Shut Down Resources.

The following screenshot shows the menu from a Studio Classic notebook. The processor and memory of the instance type powering the notebook are displayed as **2 vCPU + 4 GiB**.
To change the instance type

1. Choose the processor and memory of the instance type powering the notebook. This opens a pop up window.
2. From the Set up notebook environment pop up window, select the Instance type dropdown menu.
3. From the Instance type dropdown, choose one of the instance types that are listed.
4. After choosing a type, choose Select.
5. Wait for the new instance to become enabled, and then the new instance type information is displayed.

For a list of the available instance types, see Available Studio Classic Instance Types.

Change an Image or a Kernel

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

With Amazon SageMaker Studio Classic notebooks, you can change the notebook's image or kernel from within the notebook.

The following screenshot shows the menu from a Studio Classic notebook. The current SageMaker kernel and image are displayed as Python 3 (Data Science), where Python 3 denotes the kernel and Data Science denotes the SageMaker image that contains the kernel. The color of the circle to the right indicates the kernel is idle or busy. The kernel is busy when the center and the edge of the circle are the same color.
To change a notebook's image or kernel

1. Choose the image/kernel name in the notebook menu.
2. From the Set up notebook environment pop up window, select the Image or Kernel dropdown menu.
3. From the dropdown menu, choose one of the images or kernels that are listed.
4. After choosing an image or kernel, choose Select.
5. Wait for the kernel's status to show as idle, which indicates the kernel has started.

For a list of available SageMaker images and kernels, see Available Amazon SageMaker Images.

Shut Down Resources

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You can shut down individual Amazon SageMaker resources, including notebooks, terminals, kernels, apps, and instances from Studio Classic. You can also shut down all of the resources in one of these categories at the same time. Amazon SageMaker Studio Classic does not support shutting down resources from within a notebook.

🔍 Note
When you shut down a Studio Classic notebook instance, additional resources that you created in Studio Classic are not deleted. For example, additional resources can include SageMaker endpoints, Amazon EMR clusters, and Amazon S3 buckets. To stop the accrual
of charges, you must manually delete these resources. For information about finding resources that are accruing charges, see Analyzing your costs with AWS Cost Explorer.

The following topics demonstrate how to delete these SageMaker resources.

Topics

- Shut Down an Open Notebook
- Shut Down Resources

Shut Down an Open Notebook

When you shut down a Studio Classic notebook, the notebook is not deleted. The kernel that the notebook is running on is shut down and any unsaved information in the notebook is lost. You can shut down an open notebook from the Studio Classic File menu or from the Running Terminal and Kernels pane. The following procedure shows how to shut down an open notebook from the Studio Classic File menu.

To shut down an open notebook from the File menu

1. Launch Studio Classic by following the steps in Launch Amazon SageMaker Studio Classic.
2. (Optional) Save the notebook contents by choosing File, then Save Notebook.
3. Choose File.
4. Choose Close and Shutdown Notebook. This opens a pop-up window.
5. From the pop-up window, choose OK.

Shut Down Resources

You can reach the Running Terminals and Kernels pane of Amazon SageMaker Studio Classic by selecting the icon. The Running Terminals and Kernels pane consists of four sections. Each section lists all the resources of that type. You can shut down each resource individually or shut down all the resources in a section at the same time.
When you choose to shut down all resources in a section, the following occurs:

- **RUNNING INSTANCES/RUNNING APPS** – All instances, apps, notebooks, kernel sessions, consoles/shells, and image terminals are shut down. System terminals aren't shut down.
- **KERNEL SESSIONS** – All kernels, notebooks and consoles/shells are shut down.
- **TERMINAL SESSIONS** – All image terminals and system terminals are shut down.

**To shut down resources**

1. Launch Studio Classic by following the steps in [Launch Amazon SageMaker Studio Classic](#).
2. Choose the **Running Terminals and Kernels** icon ( 
   ![icon]
).
3. Do either of the following:
   - To shut down a specific resource, choose the **Shut Down** icon ( 
     ![icon]
   ) on the same row as the resource.
For running instances, a confirmation dialog box lists all of the resources that SageMaker will shut down. A confirmation dialog box displays all running apps. To proceed, choose **Shut down all**.

**Note**

A confirmation dialog box isn't displayed for kernel sessions or terminal sessions.

- To shut down all resources in a section, choose the X to the right of the section label. A confirmation dialog box is displayed. Choose **Shut down all** to proceed.

**Note**

When you shut down these Studio Classic resources, any additional resources created from Studio Classic, such as SageMaker endpoints, Amazon EMR clusters, and Amazon S3 buckets are not deleted. You must manually delete these resources to stop the accrual of charges. For information about finding resources that are accruing charges, see [Analyzing your costs with AWS Cost Explorer](https://aws.amazon.com/costs-explorer/).

**Usage Metering**

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](https://aws.amazon.com/sagemaker/).

There is no additional charge for using Amazon SageMaker Studio Classic. The costs incurred for running Amazon SageMaker Studio Classic notebooks, interactive shells, consoles, and terminals are based on Amazon Elastic Compute Cloud (Amazon EC2) instance usage.

When you run the following resources, you must choose a SageMaker image and kernel:
From the Studio Classic Launcher

- Notebook
- Interactive Shell
- Image Terminal

From the File menu

- Notebook
- Console

When launched, the resource is run on an Amazon EC2 instance of the chosen instance type. If an instance of that type was previously launched and is available, the resource is run on that instance.

For CPU based images, the default suggested instance type is ml.t3.medium. For GPU based images, the default suggested instance type is ml.g4dn.xlarge.

The costs incurred are based on the instance type. You are billed separately for each instance.

Metering starts when an instance is created. Metering ends when all the apps on the instance are shut down, or the instance is shut down. For information about how to shut down an instance, see Shut Down Resources.

⚠️ Important

You must shut down the instance to stop incurring charges. If you shut down the notebook running on the instance but don't shut down the instance, you will still incur charges. When you shut down the Studio Classic notebook instances, any additional resources, such as SageMaker endpoints, Amazon EMR clusters, and Amazon S3 buckets created from Studio Classic are not deleted. Delete those resources to stop accrual of charges.

When you open multiple notebooks on the same instance type, the notebooks run on the same instance even if they are using different kernels. You are billed only for the time that one instance is running.

You can change the instance type from within the notebook after you open it. For more information, see Change an Instance Type.
Available Resources

Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following sections list the available resources for Amazon SageMaker Studio Classic notebooks.

Topics

- Available Studio Classic Instance Types
- Available Amazon SageMaker Images

Available Studio Classic Instance Types

Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker Studio Classic notebooks run on Amazon Elastic Compute Cloud (Amazon EC2) instances. The following Amazon EC2 instance types are available for use with Studio Classic notebooks. For detailed information on which instance types fit your use case, and their performance capabilities, see Amazon Elastic Compute Cloud Instance types. For information about pricing for these instance types, see Amazon EC2 Pricing.

For information about available Amazon SageMaker Notebook Instance types, see CreateNotebookInstance.
### Note

For most use cases, you should use a `ml.t3.medium`. This is the default instance type for CPU-based SageMaker images, and is available as part of the [AWS Free Tier](https://aws.amazon.com/free/).

### Topics

- [CPU instances](#)
- [Instances with 1 or more GPUs](#)

### CPU instances

The following table lists the Amazon EC2 CPU instance types with no GPU attached that are available for use with Studio Classic notebooks. It also lists information about the specifications of each instance type. The default instance type for CPU-based images is `ml.t3.medium`.

For detailed information on which instance types fit your use case, and their performance capabilities, see [Amazon Elastic Compute Cloud Instance types](https://aws.amazon.com/ec2/instance-types/). For information about pricing for these instance types, see [Amazon EC2 Pricing](https://aws.amazon.com/ec2/pricing).

#### Note

Fast launch instances types are optimized to start in under two minutes.

### CPU instances

<table>
<thead>
<tr>
<th>Instance</th>
<th>Use case</th>
<th>Fast launch</th>
<th>vCPU</th>
<th>Memory (GiB)</th>
<th>Instance Storage (GB)</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>ml.t3.medium</code></td>
<td>General purpose</td>
<td>Yes</td>
<td>2</td>
<td>4</td>
<td>Amazon EBS Only</td>
</tr>
<tr>
<td><code>ml.t3.large</code></td>
<td>General purpose</td>
<td>No</td>
<td>2</td>
<td>8</td>
<td>Amazon EBS Only</td>
</tr>
</tbody>
</table>

Use Studio Classic Notebooks
<table>
<thead>
<tr>
<th>Instance</th>
<th>Use case</th>
<th>Fast launch</th>
<th>vCPU</th>
<th>Memory (GiB)</th>
<th>Instance Storage (GB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.t3.xlarge</td>
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<td>4</td>
<td>16</td>
<td>Amazon EBS Only</td>
</tr>
<tr>
<td>ml.t3.2xlarge</td>
<td>General purpose</td>
<td>No</td>
<td>8</td>
<td>32</td>
<td>Amazon EBS Only</td>
</tr>
<tr>
<td>ml.m5.large</td>
<td>General purpose</td>
<td>Yes</td>
<td>2</td>
<td>8</td>
<td>Amazon EBS Only</td>
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<td>ml.m5.xlarge</td>
<td>General purpose</td>
<td>No</td>
<td>4</td>
<td>16</td>
<td>Amazon EBS Only</td>
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<tr>
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<td>256</td>
<td>Amazon EBS Only</td>
</tr>
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<td>Use case</td>
<td>Fast launch</td>
<td>vCPU</td>
<td>Memory (GiB)</td>
<td>Instance Storage (GB)</td>
</tr>
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<td>Instance</td>
<td>Use case</td>
<td>Fast launch</td>
<td>vCPU</td>
<td>Memory (GiB)</td>
<td>Instance Storage (GB)</td>
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<td>16</td>
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<td>Instance</td>
<td>Use case</td>
<td>Fast launch</td>
<td>vCPU</td>
<td>Memory (GiB)</td>
<td>Instance Storage (GB)</td>
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<td>ml.r5.xlarge</td>
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<td>32</td>
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<td>768</td>
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</table>

**Instances with 1 or more GPUs**

The following table lists the Amazon EC2 instance types with 1 or more GPUs attached that are available for use with Studio Classic notebooks. It also lists information about the specifications of each instance type. The default instance type for GPU-based images is `ml.g4dn.xlarge`.
For detailed information on which instance types fit your use case, and their performance capabilities, see Amazon Elastic Compute Cloud Instance types. For information about pricing for these instance types, see Amazon EC2 Pricing.

**Note**

Fast launch instances types are optimized to start in under two minutes.

### Instances with 1 or more GPUs

<table>
<thead>
<tr>
<th>Instance</th>
<th>Use case</th>
<th>Fast launch</th>
<th>GPUs</th>
<th>vCPU</th>
<th>Memor (GiB)</th>
<th>GPU Memor (GiB)</th>
<th>Instance Storage (GB)</th>
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<td>64</td>
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<td>64</td>
<td>488</td>
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<td>8</td>
<td>96</td>
<td>768</td>
<td>256</td>
<td>2 x 900 NVMe SSD</td>
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<tr>
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<td>320 GB HBM2</td>
<td>8 x 1000 NVMe SSD</td>
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<td>Use case</td>
<td>Fast launch</td>
<td>GPUs</td>
<td>vCPU</td>
<td>Memor (GiB)</td>
<td>GPU Memor (GiB)</td>
<td>Instance Storage (GB)</td>
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<td>1152</td>
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<td>64</td>
<td>16</td>
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<td>256</td>
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<td>1 x 900 NVMe SSD</td>
</tr>
<tr>
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<td>Use case</td>
<td>Fast launch</td>
<td>GPUs</td>
<td>vCPU</td>
<td>Memory (GiB)</td>
<td>GPU Memory (GiB)</td>
<td>Instance Storage (GB)</td>
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</tr>
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<td>64</td>
<td>24</td>
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<td>128</td>
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<td>1 x 900 NVMe SSD</td>
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<tr>
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<td>48</td>
<td>192</td>
<td>96</td>
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<td>384</td>
<td>96</td>
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## Image ARN format

The following table lists the image ARN and URI format for each Region. To create the full ARN for an image, replace the `resource-identifier` placeholder with the corresponding resource identifier for the image from the SageMaker images and kernels table. To create the full URI for an

<table>
<thead>
<tr>
<th>Instance</th>
<th>Use case</th>
<th>Fast launch</th>
<th>GPUs</th>
<th>vCPU</th>
<th>Memor (GiB)</th>
<th>GPU Memor (GiB)</th>
<th>Instance Storage (GB)</th>
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<td>768</td>
<td>192</td>
<td>2 x 3800 NVMe SSD</td>
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</table>

### Available Amazon SageMaker Images

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

This page lists the SageMaker images and associated kernels that are available in Amazon SageMaker Studio Classic, as well as the format needed to create the ARN for each image. SageMaker images contain the latest Amazon SageMaker Python SDK and the latest version of the kernel. For more information, see Deep Learning Containers Images.

### Topics

- Image ARN format
- Supported URI tags
- Supported images
- Images slated for deprecation
image, replace the `tag` placeholder with the corresponding cpu or gpu tag. For the list of tags you can use, see Supported URI tags.

**Note**

SageMaker Distribution images use a distinct set of image ARNs, which are listed in the following table.

<table>
<thead>
<tr>
<th>Region</th>
<th>Image ARN Format</th>
<th>SageMaker Distribution Image ARN Format</th>
<th>SageMaker Distribution Image URI Format</th>
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<td>Region</td>
<td>ARN (Amazon Resource Name) with ARN Tagging &amp; Image</td>
<td>ARN (Amazon Resource Name) with ARN Tagging &amp; Image</td>
<td>ARN (Amazon Resource Name) with ARN Tagging &amp; Image</td>
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<td>ARN of Amazon SageMaker</td>
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<tr>
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<td>-------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------</td>
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</tbody>
</table>
Supported URI tags

The following list shows the tags you can include in your image URI.

- 1-cpu
- 1-gpu
- 0-cpu
- 0-gpu

The following examples show URIs with various tag formats:

- 542918446943.dkr.ecr.us-west-2.amazonaws.com/sagemaker-distribution-prod:1-cpu
- 542918446943.dkr.ecr.us-west-2.amazonaws.com/sagemaker-distribution-prod:0-gpu

Supported images

The following table gives information about the SageMaker images and associated kernels that are available in Amazon SageMaker Studio Classic, as well as the resource identifier and Python version included in the image.

SageMaker images and kernels

<table>
<thead>
<tr>
<th>SageMaker Image</th>
<th>Description</th>
<th>Resource Identifier</th>
<th>Kernels (and Identifier)</th>
<th>Python Version</th>
</tr>
</thead>
<tbody>
<tr>
<td>SageMaker Distribution v0 CPU</td>
<td>SageMaker Distribution v0 CPU is a Python 3.8 image that includes popular sagemaker-distribution-cpu-v0</td>
<td>sagemaker-distribution-cpu-v0</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
</tr>
<tr>
<td>SageMaker Image</td>
<td>Description</td>
<td>Resource Identifier</td>
<td>Kernels (and Identifier)</td>
<td>Python Version</td>
</tr>
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<td>--------------------------</td>
<td>----------------</td>
</tr>
<tr>
<td></td>
<td>frameworks for machine learning, data science and visualization on CPU. This includes deep learning frameworks like PyTorch, TensorFlow and Keras; popular Python packages like numpy, scikit-learn and pandas; and IDEs like Jupyter Lab. For more information, see the Amazon SageMaker Distribution repo.</td>
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<td>SageMaker Image</td>
<td>Description</td>
<td>Resource Identifier</td>
<td>Kernels (and Identifier)</td>
<td>Python Version</td>
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</tr>
<tr>
<td>SageMaker Distribution v0 GPU</td>
<td>SageMaker Distribution v0 GPU is a Python 3.8 image that includes popular frameworks for machine learning, data science and visualization on GPU. This includes deep learning frameworks like PyTorch, Tensorflow and Keras; popular Python packages like numpy, scikit-learn and pandas; and IDEs like Jupyter Lab. For more information, see the Amazon SageMaker Distribution repo.</td>
<td>sagemaker-distribution-gpu-v0</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
</tr>
<tr>
<td>SageMaker Image</td>
<td>Description</td>
<td>Resource Identifier</td>
<td>Kernels (and Identifier)</td>
<td>Python Version</td>
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</tr>
<tr>
<td>SageMaker Distribution v1 CPU</td>
<td>SageMaker Distribution v1 CPU is a Python 3.10 image that includes popular frameworks for machine learning, data science and data analytics on CPU. This includes deep learning frameworks like PyTorch, TensorFlow and Keras; popular Python packages like numpy, scikit-learn and pandas; and IDEs like Jupyter Lab. For more information, see the Amazon SageMaker Distribution repo.</td>
<td>sagemaker-distribution-cpu-v1</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
</tr>
<tr>
<td>SageMaker Image</td>
<td>Description</td>
<td>Resource Identifier</td>
<td>Kernels (and Identifier)</td>
<td>Python Version</td>
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<tr>
<td>SageMaker Distribution v1 GPU</td>
<td>SageMaker Distribution v1 GPU is a Python 3.10 image that includes popular frameworks for machine learning, data science and data analytics on GPU. This includes deep learning frameworks like PyTorch, TensorFlow and Keras; popular Python packages like numpy, scikit-learn and pandas; and IDEs like Jupyter Lab. For more information, see the Amazon SageMaker Distribution repo.</td>
<td>sagemaker-distribution-gpu-v1</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
</tr>
<tr>
<td>Base Python 3.0</td>
<td>Official Python 3.10 image from DockerHub with boto3 and AWS CLI included.</td>
<td>sagemaker-base-python-310-v1</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
</tr>
<tr>
<td>SageMaker Image</td>
<td>Description</td>
<td>Resource Identifier</td>
<td>Kernels (and Identifier)</td>
<td>Python Version</td>
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</tr>
<tr>
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<td>Official Python 3.8 image from DockerHub with boto3 and AWS CLI included.</td>
<td>sagemaker-base-python-38</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
</tr>
<tr>
<td>Data Science 3.0</td>
<td>Data Science 3.0 is a Python 3.10 conda image based on anaconda version 2022.10 with the most commonly used Python packages and libraries, such as NumPy and SciKit Learn.</td>
<td>sagemaker-data-science-310-v1</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
</tr>
<tr>
<td>Data Science 2.0</td>
<td>Data Science 2.0 is a Python 3.8 conda image based on anaconda version 2021.11 with the most commonly used Python packages and libraries, such as NumPy and SciKit Learn.</td>
<td>sagemaker-data-science-38</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
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<td>SageMaker Image</td>
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<td>Geospatial 1.0</td>
<td>Amazon SageMaker geospatial is a Python image consisting of commonly used geospatial libraries such as GDAL, Fiona, GeoPandas, Shapely, and Rasterio, and allows you to visualize geospatial data within SageMaker. For more information, see <a href="https://docs.aws.amazon.com/sagemaker/latest/dg/geospatial.html">Amazon SageMaker geospatial Notebook SDK</a>.</td>
<td>sagemaker-geospatial-1.0</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
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<tr>
<td>PyTorch 2.0.0</td>
<td>The AWS Deep Learning Container for PyTorch 2.0.0 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="https://docs.aws.amazon.com/sagemaker/latest/dg/containers-release-notes.html">Release Notes for Deep Learning Containers</a>.</td>
<td>pytorch-2.0.0-cpu-py310</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
</tr>
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<td>SageMaker Image</td>
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<tr>
<td>PyTorch 2.0.0 Python 3.10 GPU Optimized</td>
<td>The AWS Deep Learning Container(s) for PyTorch 2.0.0 with CUDA 11.8 include container(s) for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>pytorch-2.0.0-gpu-py310</td>
<td>Python 3 (python3)</td>
<td>Python 3.10</td>
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<td>PyTorch 1.13 Python 3.9 CPU Optimized</td>
<td>The AWS Deep Learning Container(s) for PyTorch 1.13 with CUDA 11.3 include container(s) for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>pytorch-1.13-cpu-py39</td>
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<td>The AWS Deep Learning Container for PyTorch 1.13 with CUDA 11.7 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see Release Notes for Deep Learning Containers.</td>
<td>pytorch-1.13-gpu-py39</td>
<td>Python 3 (python3)</td>
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<td>PyTorch 1.12 Python 3.8 CPU Optimized</td>
<td>The AWS Deep Learning Container for PyTorch 1.12 with CUDA 11.3 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see AWS Deep Learning Containers for PyTorch 1.12.0.</td>
<td>pytorch-1.12-cpu-py38</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
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<tr>
<td>SageMaker Image</td>
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<tr>
<td>PyTorch 1.12</td>
<td>The AWS Deep Learning Containers for PyTorch 1.12 with CUDA 11.3 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see AWS Deep Learning Containers for PyTorch 1.12.0.</td>
<td>pytorch-1.12-gpu-py38</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
</tr>
<tr>
<td>PyTorch 1.10</td>
<td>The AWS Deep Learning Containers for PyTorch 1.10 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see AWS Deep Learning Containers for PyTorch 1.10.2 on SageMaker.</td>
<td>pytorch-1.10-cpu-py38</td>
<td>Python 3 (python3)</td>
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<td>PyTorch 1.10 Python 3.8 GPU Optimized</td>
<td>The AWS Deep Learning Containers for PyTorch 1.10 with CUDA 11.3 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="#">AWS Deep Learning Containers for PyTorch 1.10.2 on SageMaker</a>.</td>
<td>pytorch-1.10-gpu-py38</td>
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| SparkAnalytics 2.0  | Anaconda Individual Edition with PySpark and Spark kernels. For more information, see sparkmagic. | sagemaker-sparkanalytics-310-v1 | • SparkMagic Spark (conda-env-sm_sparkmagic-sparkkernel)  
• SparkMagic PySpark (conda-env-sm_sparkmagic-py.sparkkernel)  
• Glue Spark (conda-env-sm_glue_is-glue.spark)  
• Glue Python [PySpark and Ray] (conda-env-sm_glue_is-glue_pyspark) | Python 3.10        |
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<th>SageMaker Image</th>
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<td>SparkAnalytics 1.0</td>
<td>Anaconda Individual Edition with PySpark and Spark kernels. For more information, see sparkmagic.</td>
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<td>- SparkMagic Spark (conda-env-sm_sparkmagic-sparkkernel)</td>
<td>Python 3.8</td>
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<td>- SparkMagic PySpark (conda-env-sm_sparkmagic-py.sparkkernel)</td>
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<td>- Glue Spark (conda-env-sm_glue_is-glue.spark)</td>
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<td>- Glue Python [PySpark and Ray] (conda-env-sm_glue_is-glue_pyspark)</td>
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<tr>
<td>TensorFlow 2.12.0 Python 3.10 CPU Optimized</td>
<td>The AWS Deep Learning Container s for TensorFlow 2.12.0 with CUDA 11.2 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>tensorflow-2.12.0-cpu-py310-ubuntu20.04-sagemaker-v1.0</td>
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<td>TensorFlow 2.12.0 Python 3.10 GPU Optimized</td>
<td>The AWS Deep Learning Container s for TensorFlow 2.12.0 with CUDA 11.8 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>tensorflow-2.12.0-gpu-py310-cu118-ubuntu20.04-sagemaker-v1</td>
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<td>Python 3.10</td>
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<tr>
<td>TensorFlow 2.11.0 Python 3.9 CPU</td>
<td>The AWS Deep Learning Container s for TensorFlow 2.11.0 with CUDA 11.2 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>tensorflow-2.11.0-cpu-py39-ubuntu20.04-sagemaker-v1.1</td>
<td>Python 3 (python3)</td>
<td>Python 3.9</td>
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<tr>
<td>TensorFlow 2.11.0 Python 3.9 GPU</td>
<td>The AWS Deep Learning Container s for TensorFlow 2.11.0 with CUDA 11.2 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>tensorflow-2.11.0-gpu-py39-cu112-ubuntu20.04-sagemaker-v1.1</td>
<td>Python 3 (python3)</td>
<td>Python 3.9</td>
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<tr>
<td>TensorFlow 2.10 Python 3.9 CPU Optimized</td>
<td>The AWS Deep Learning Containers for TensorFlow 2.10 with CUDA 11.2 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>tensorflow-2.10.1-cpu-py39-ubuntu20.04-sagemaker-v1.2</td>
<td>Python 3 (python3)</td>
<td>Python 3.9</td>
</tr>
<tr>
<td>TensorFlow 2.10 Python 3.9 GPU Optimized</td>
<td>The AWS Deep Learning Containers for TensorFlow 2.10 with CUDA 11.2 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="#">Release Notes for Deep Learning Containers</a>.</td>
<td>tensorflow-2.10.1-gpu-py39-ubuntu20.04-sagemaker-v1.2</td>
<td>Python 3 (python3)</td>
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<td>SageMaker Image</td>
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<tr>
<td>TensorFlow 2.6 Python 3.8 CPU</td>
<td>The AWS Deep Learning Containers for TensorFlow 2.6 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="https://aws.amazon.com">AWS Deep Learning Containers for TensorFlow 2.6</a>.</td>
<td>tensorflow-2.6-cpu-py38-ubuntu20.04-v1</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
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<td>Optimized</td>
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<tr>
<td>TensorFlow 2.6 Python 3.8 GPU</td>
<td>The AWS Deep Learning Containers for TensorFlow 2.6 with CUDA 11.2 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="https://aws.amazon.com">AWS Deep Learning Containers for TensorFlow 2.6</a>.</td>
<td>tensorflow-2.6-gpu-py38-cu112-ubuntu20.04-v1</td>
<td>Python 3 (python3)</td>
<td>Python 3.8</td>
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<tr>
<td>Optimized</td>
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</tbody>
</table>
Images slated for deprecation

SageMaker ends support for images the day after any of the packages in the image reach end-of-life by their publisher.

The following SageMaker images are slated for deprecation. These images are based on Python 3.7, which reached end-of-life on June 27, 2023. Starting on October 30, 2023, SageMaker will discontinue support for these images and they will not be selectable from the Studio Classic UI. To avoid non-compliance issues, if you're using any of these images, we recommend that you move to an image with a later version.

SageMaker images slated for deprecation

<table>
<thead>
<tr>
<th>SageMaker Image</th>
<th>Deprecation date</th>
<th>Description</th>
<th>Resource Identifier</th>
<th>Kernels</th>
<th>Python Version</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data Science</td>
<td>October 30, 2023</td>
<td>Data Science is a Python 3.7 conda image with the most commonly used Python packages and libraries, such as NumPy and SciKit Learn.</td>
<td>datascience-1.0</td>
<td>Python 3</td>
<td>Python 3.7</td>
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<tr>
<td>SageMaker JumpStart Data Science 1.0</td>
<td>October 30, 2023</td>
<td>SageMaker JumpStart Data Science 1.0 is a SageMaker JumpStart image that includes commonly used packages and libraries.</td>
<td>sagemaker-jumpstart-data-science-1.0</td>
<td>Python 3</td>
<td>Python 3.7</td>
</tr>
<tr>
<td>SageMaker JumpStart Image</td>
<td>Deprecation date</td>
<td>Description</td>
<td>Resource Identifier</td>
<td>Kernels</td>
<td>Python Version</td>
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<tr>
<td>SageMaker JumpStart MXNet 1.0</td>
<td>October 30, 2023</td>
<td>SageMaker JumpStart MXNet 1.0 is a SageMaker JumpStart image that includes MXNet.</td>
<td>sagemaker-jumpstart-mxnet-1.0</td>
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<tr>
<td>SageMaker JumpStart PyTorch 1.0</td>
<td>October 30, 2023</td>
<td>SageMaker JumpStart PyTorch 1.0 is a SageMaker JumpStart image that includes PyTorch.</td>
<td>sagemaker-jumpstart-pytorch-1.0</td>
<td>Python 3</td>
<td>Python 3.7</td>
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<tr>
<td>SageMaker JumpStart TensorFlow 1.0</td>
<td>October 30, 2023</td>
<td>SageMaker JumpStart TensorFlow 1.0 is a SageMaker JumpStart image that includes TensorFlow.</td>
<td>sagemaker-jumpstart-tensorflow-1.0</td>
<td>Python 3</td>
<td>Python 3.7</td>
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<tr>
<td>SageMaker Image</td>
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<td>Description</td>
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<tr>
<td>SparkMagic</td>
<td>October 30, 2023</td>
<td>Anaconda Individual Edition with PySpark and Spark kernels. For more information, see <a href="https://sparkmagic.com">sparkmagic</a>.</td>
<td>sagemaker-sparkmagic</td>
<td>• PySpark</td>
<td>Python 3.7</td>
</tr>
<tr>
<td>TensorFlow 2.3 CPU Optimized</td>
<td>October 30, 2023</td>
<td>The AWS Deep Learning Containers for TensorFlow 2.3 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="https://aws.amazon.com/deeplearning/containers/tensorflow-2.3-cpu-py37-ubuntu18.04-v1">AWS Deep Learning Containers with TensorFlow 2.3.0</a>.</td>
<td>tensorflow-2.3-cpu-py37-ubuntu18.04-v1</td>
<td>Python 3</td>
<td>Python 3.7</td>
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<td>SageMaker Image</td>
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<tr>
<td>TensorFlow 2.3 Python 3.7 GPU Optimized</td>
<td>October 30, 2023</td>
<td>The AWS Deep Learning Containers for TensorFlow 2.3 with CUDA 11.0 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see <a href="https://aws.amazon.com/marketplace/pp/pt2-lambda-ec2-sdk">AWS Deep Learning Containers for TensorFlow 2.3.1 with CUDA 11.0</a>.</td>
<td>tensorflow-2.3-gpu-py37-cu10-ubuntu18.04-v3</td>
<td>Python 3</td>
<td>Python 3.7</td>
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<td>SageMaker Image</td>
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<tr>
<td>TensorFlow 1.15 Python 3.7 CPU Optimized</td>
<td>October 30, 2023</td>
<td>The AWS Deep Learning Containers for TensorFlow 1.15 include containers for training on CPU, optimized for performance and scale on AWS. For more information, see <a href="https://aws.amazon.com/deeplearningcontainers/tensorflow-1.15-cpu-py37-ubuntu18.04-v7">AWS Deep Learning Containers v7.0 for TensorFlow</a>.</td>
<td>tensorflow-1.15-cp u-py37-ubuntu18.04-v7</td>
<td>Python 3</td>
<td>Python 3.7</td>
</tr>
</tbody>
</table>
## Customize Amazon SageMaker Studio Classic

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

There are four options for customizing your Amazon SageMaker Studio Classic environment. You bring your own SageMaker image, use a lifecycle configuration script, attach suggested Git repos to Studio Classic, or create kernels using persistent Conda environments in Amazon EFS. Use each option individually, or together.

<table>
<thead>
<tr>
<th>SageMaker Image</th>
<th>Deprecation date</th>
<th>Description</th>
<th>Resource Identifier</th>
<th>Kernels</th>
<th>Python Version</th>
</tr>
</thead>
<tbody>
<tr>
<td>TensorFlow 1.15 Python 3.7 GPU Optimized</td>
<td>October 30, 2023</td>
<td>The AWS Deep Learning Containers for TensorFlow 1.15 with CUDA 11.0 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see AWS Deep Learning Containers v7.0 for TensorFlow.</td>
<td>tensorflow-1.15-gpu-py37-cu110-ubuntu18.04-v8</td>
<td>Python 3</td>
<td>Python 3.7</td>
</tr>
</tbody>
</table>

### TensorFlow 1.15 Python 3.7 GPU Optimized

The AWS Deep Learning Containers for TensorFlow 1.15 with CUDA 11.0 include containers for training on GPU, optimized for performance and scale on AWS. For more information, see AWS Deep Learning Containers v7.0 for TensorFlow.

- **Resource Identifier**: `tensorflow-1.15-gpu-py37-cu110-ubuntu18.04-v8`
- **Kernels**: Python 3
- **Python Version**: Python 3.7
• **Bring your own SageMaker image:** A SageMaker image is a file that identifies the kernels, language packages, and other dependencies required to run a Jupyter notebook in Amazon SageMaker Studio Classic. Amazon SageMaker provides many built-in images for you to use. If you need different functionality, you can bring your own custom images to Studio Classic.

• **Use lifecycle configurations with Amazon SageMaker Studio Classic:** Lifecycle configurations are shell scripts triggered by Amazon SageMaker Studio Classic lifecycle events, such as starting a new Studio Classic notebook. You can use lifecycle configurations to automate customization for your Studio Classic environment. For example, you can install custom packages, configure notebook extensions, preload datasets, and set up source code repositories.

• **Attach suggested Git repos to Studio Classic:** You can attach suggested Git repository URLs at the Amazon SageMaker domain or user profile level. Then, you can select the repo URL from the list of suggestions and clone that into your environment using the Git extension in Studio Classic.

• **Persist Conda environments to the Studio Classic Amazon EFS volume:** Studio Classic uses an Amazon EFS volume as a persistent storage layer. You can save your Conda environment on this Amazon EFS volume, then use the saved environment to create kernels. Studio Classic automatically picks up all valid environments saved in Amazon EFS as KernelGateway kernels. These kernels persist through restart of the kernel, app, and Studio Classic. For more information, see the **Persist Conda environments to the Studio Classic EFS volume** section in Four approaches to manage Python packages in Amazon SageMaker Studio Classic notebooks.

The following topics show how to use these three options to customize your Amazon SageMaker Studio Classic environment.

**Topics**

• [Bring your own SageMaker image](#)

• [Use lifecycle configurations with Amazon SageMaker Studio Classic](#)

• [Attach Suggested Git Repos to Studio Classic](#)

**Bring your own SageMaker image**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the...
A SageMaker image is a file that identifies the kernels, language packages, and other dependencies required to run a Jupyter notebook in Amazon SageMaker Studio Classic. These images are used to create an environment that you then run Jupyter notebooks from. Amazon SageMaker provides many built-in images for you to use. For the list of built-in images, see Available Amazon SageMaker Images.

If you need different functionality, you can bring your own custom images to Studio Classic. You can create images and image versions, and attach image versions to your domain or shared space, using the SageMaker control panel, the AWS SDK for Python (Boto3), and the AWS Command Line Interface (AWS CLI). You can also create images and image versions using the SageMaker console, even if you haven’t onboarded to a SageMaker domain. SageMaker provides sample Dockerfiles to use as a starting point for your custom SageMaker images in the SageMaker Studio Classic Custom Image Samples repository.

The following topics explain how to bring your own image using the SageMaker console or AWS CLI, then launch the image in Studio Classic. For a similar blog article, see Bringing your own R environment to Amazon SageMaker Studio Classic. For notebooks that show how to bring your own image for use in training and inference, see Amazon SageMaker Studio Classic Container Build CLI.

Key terminology

The following section defines key terms for bringing your own image to use with Studio Classic.

- **Dockerfile**: A Dockerfile is a file that identifies the language packages and other dependencies for your Docker image.

- **Docker image**: The Docker image is a built Dockerfile. This image is checked into Amazon ECR and serves as the basis of the SageMaker image.

- **SageMaker image**: A SageMaker image is a holder for a set of SageMaker image versions based on Docker images. Each image version is immutable.

- **Image version**: An image version of a SageMaker image represents a Docker image and is stored in an Amazon ECR repository. Each image version is immutable. These image versions can be attached to a domain or shared space and used with Studio Classic.
Topics

- Custom SageMaker image specifications
- Prerequisites
- Add a Docker image compatible with Studio Classic to Amazon ECR
- Create a custom SageMaker image
- Attach a custom SageMaker image
- Launch a custom SageMaker image in Amazon SageMaker Studio Classic
- Clean up resources

Custom SageMaker image specifications

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following specifications apply to the container image that is represented by a SageMaker image version.

Running the image

ENTRYPOINT and CMD instructions are overridden to enable the image to run as a KernelGateway app.

Port 8888 in the image is reserved for running the KernelGateway web server.

Stopping the image

The DeleteApp API issues the equivalent of a docker stop command. Other processes in the container won’t get the SIGKILL/SIGTERM signals.

Kernel discovery

SageMaker recognizes kernels as defined by Jupyter kernel specs.

You can specify a list of kernels to display before running the image. If not specified, python3 is displayed. Use the DescribeAppImageConfig API to view the list of kernels.
Conda environments are recognized as kernel specs by default.

**File system**

The `/opt/.sagemakerinternal` and `/opt/ml` directories are reserved. Any data in these directories might not be visible at runtime.

**User data**

Each user in a domain gets a user directory on a shared Amazon Elastic File System volume in the image. The location of the current user's directory on the Amazon EFS volume is configurable. By default, the location of the directory is `/home/sagemaker-user`.

SageMaker configures POSIX UID/GID mappings between the image and the host. This defaults to mapping the root user's UID/GID (0/0) to the UID/GID on the host.

You can specify these values using the [CreateAppImageConfig](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateAppImageConfig.html) API.

**GID/UID limits**

Amazon SageMaker Studio Classic only supports the following DefaultUID and DefaultGID combinations:

- DefaultUID: 1000 and DefaultGID: 100, which corresponds to a non-privileged user.
- DefaultUID: 0 and DefaultGID: 0, which corresponds to root access.

**Metadata**

A metadata file is located at `/opt/ml/metadata/resource-metadata.json`. No additional environment variables are added to the variables defined in the image. For more information, see [Get App Metadata](https://docs.aws.amazon.com/sagemaker/latest/dg/API_GetAppMetadata.html).

**GPU**

On a GPU instance, the image is run with the `--gpus` option. Only the CUDA toolkit should be included in the image not the NVIDIA drivers. For more information, see [NVIDIA User Guide](https://docs.nvidia.com/cuda/index.html).

**Metrics and logging**

Logs from the KernelGateway process are sent to Amazon CloudWatch in the customer's account. The name of the log group is `/aws/sagemaker/studio`. The name of the log stream is `$domainID/$userProfileName/KernelGateway/$appName`.

**Image size**

Limited to 25 GB. To view the size of your image, run `docker image ls`. 

---

**Customize Studio Classic**
Sample Dockerfile

The following sample Dockerfile creates an image based Amazon Linux 2, installs third party packages and the python3 kernel, and sets the scope to the non-privileged user.

```bash
FROM public.ecr.aws/amazonlinux/amazonlinux:2

ARG NB_USER="sagemaker-user"
ARG NB_UID="1000"
ARG NB_GID="100"

RUN \
    yum install --assumeyes python3 shadow-utils && \
    useradd --create-home --shell /bin/bash --gid "$NB_GID" --uid $NB_UID \\
    $NB_USER && \
    yum clean all && \
    python3 -m pip install ipykernel && \
    python3 -m ipykernel install

USER $NB_UID
```

Prerequisites

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You must satisfy the following prerequisites to bring your own container for use with Amazon SageMaker Studio Classic.

- The Docker application. For information about setting up Docker, see Orientation and setup.
- Install the AWS CLI by following the steps in Getting started with the AWS CLI.
- A local copy of any Dockerfile for creating a Studio Classic compatible image. For sample custom images, see the SageMaker Studio Classic custom image samples repository.
- Permissions to access the Amazon Elastic Container Registry (Amazon ECR) service. For more information, see Amazon ECR Managed Policies.
• An AWS Identity and Access Management execution role that has the AmazonSageMakerFullAccess policy attached. If you have onboarded to Amazon SageMaker domain, you can get the role from the Domain Summary section of the SageMaker control panel.

• Install the Studio Classic image build CLI by following the steps in SageMaker Docker Build. This CLI enables you to build a Dockerfile using AWS CodeBuild.

Add a Docker image compatible with Studio Classic to Amazon ECR

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You perform the following steps to add a container image to Amazon ECR:

• Create an Amazon ECR repository.
• Authenticate to Amazon ECR.
• Build a Docker image compatible with Studio Classic.
• Push the image to the Amazon ECR repository.

🔍 Note
The Amazon ECR repository must be in the same AWS Region as Studio Classic.

To build and add a container image to Amazon ECR

1. Create an Amazon ECR repository using the AWS CLI. To create the repository using the Amazon ECR console, see Creating a repository.

   ```bash
   aws ecr create-repository \
   --repository-name smstudio-custom \
   ```
2. Build the Dockerfile using the Studio Classic image build CLI. The period (.) specifies that the Dockerfile should be in the context of the build command. This command builds the image and uploads the built image to the ECR repo. It then outputs the image URI.

```
sm-docker build . --repository smstudio-custom:custom
```

The response should look similar to the following.

```
Image URI: <acct-id>.dkr.ecr.<region>.amazonaws.com/<image_name>
```

### Create a custom SageMaker image

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](https://aws.amazon.com/sagemaker).

This topic describes how you can create a custom SageMaker image using the SageMaker console or AWS CLI.
When you create an image from the console, SageMaker also creates an initial image version. The image version represents a container image in Amazon Elastic Container Registry (ECR). The container image must satisfy the requirements to be used in Amazon SageMaker Studio Classic. For more information, see Custom SageMaker image specifications. For information on testing your image locally and resolving common issues, see the SageMaker Studio Classic Custom Image Samples repo.

After you have created your custom SageMaker image, you must attach it to your domain or shared space to use it with Studio Classic. For more information, see Attach a custom SageMaker image.

Create a SageMaker image from the console

The following section demonstrates how to create a custom SageMaker image from the SageMaker console.

To create an image

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
4. On the Custom images page, choose Create image.
5. For Image source, enter the registry path to the container image in Amazon ECR. The path is in the following format:

   acct-id.dkr.ecr.region.amazonaws.com/repo-name[:tag] or [@digest]

6. Choose Next.
7. Under Image properties, enter the following:

   - Image name – The name must be unique to your account in the current AWS Region.
   - (Optional) Display name – The name displayed in the Studio Classic user interface. When not provided, Image name is displayed.
   - (Optional) Description – A description of the image.
   - IAM role – The role must have the AmazonSageMakerFullAccess policy attached. Use the dropdown menu to choose one of the following options:
     - Create a new role – Specify any additional Amazon Simple Storage Service (Amazon S3) buckets that you want users of your notebooks to have access to. If you don’t want to allow access to additional buckets, choose None.
SageMaker attaches the AmazonSageMakerFullAccess policy to the role. The role allows users of your notebooks access to the S3 buckets listed next to the checkmarks.

- Enter a custom IAM role ARN – Enter the Amazon Resource Name (ARN) of your IAM role.
- Use existing role – Choose one of your existing roles from the list.
- (Optional) Image tags – Choose Add new tag. You can add up to 50 tags. Tags are searchable using the Studio Classic user interface, the SageMaker console, or the SageMaker Search API.

8. Choose Submit.

The new image is displayed in the Custom images list and briefly highlighted. After the image has been successfully created, you can choose the image name to view its properties or choose Create version to create another version.

To create another image version

1. Choose Create version on the same row as the image.
2. For Image source, enter the registry path to the Amazon ECR container image. The container image shouldn't be the same image as used in a previous version of the SageMaker image.

Create a SageMaker image from the AWS CLI

You perform the following steps to create a SageMaker image from the container image using the AWS CLI.

- Create an Image.
- Create an ImageVersion.
- Create a configuration file.
- Create an AppImageConfig.

To create the SageMaker image entities

1. Create a SageMaker image.

```bash
aws sagemaker create-image \
    --image-name custom-image \
```
The response should look similar to the following.

```json
{
    "ImageArn": "arn:aws:sagemaker:us-east-2:acct-id:image/custom-image"
}
```

2. Create a SageMaker image version from the container image.

```bash
aws sagemaker create-image-version \\
    --image-name custom-image \\
    --base-image <acct-id>.dkr.ecr.<region>.amazonaws.com/smstudio-custom:custom-image
```

The response should look similar to the following.

```json
{
    "ImageVersionArn": "arn:aws:sagemaker:us-east-2:acct-id:image-version/custom-image/1"
}
```

3. Check that the image version was successfully created.

```bash
aws sagemaker describe-image-version \\
    --image-name custom-image \\
    --version-number 1
```

The response should look similar to the following.

```json
{
    "ImageVersionStatus": "CREATED"
}
```

⚠️ Note

If the response is "ImageVersionStatus": "CREATED_FAILED", the response also includes the failure reason. A permissions issue is a common cause of failure.
You also can check your Amazon CloudWatch logs if you experience a failure when starting or running the KernelGateway app for a custom image. The name of the log group is /aws/sagemaker/studio. The name of the log stream is $domainID/$userProfileName/KernelGateway/$appName.

4. Create a configuration file, named app-image-config-input.json. The Name value of KernelSpecs must match the name of the kernelSpec available in the Image associated with this AppImageConfig. This value is case sensitive. You can find the available kernelSpecs in an image by running jupyter-kernelspec list from a shell inside the container. MountPath is the path within the image to mount your Amazon Elastic File System (Amazon EFS) home directory. It needs to be different from the path you use inside the container because that path will be overridden when your Amazon EFS home directory is mounted.

**Note**

The following DefaultUID and DefaultGID combinations are the only accepted values:

- DefaultUID: 1000 and DefaultGID: 100
- DefaultUID: 0 and DefaultGID: 0

```json
{
    "AppImageConfigName": "custom-image-config",
    "KernelGatewayImageConfig": {
        "KernelSpecs": [
            {
                "Name": "python3",
                "DisplayName": "Python 3 (ipykernel)"
            }
        ],
        "FileSystemConfig": {
            "MountPath": "/home/sagemaker-user",
            "DefaultUid": 1000,
            "DefaultGid": 100
        }
    }
}
```
5. Create the AppImageConfig using the file created in the previous step.

```bash
aws sagemaker create-app-image-config \
    --cli-input-json file://app-image-config-input.json
```

The response should look similar to the following.

```json
{
}
```

## Attach a custom SageMaker image

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

To use a custom SageMaker image, you must attach a version of the image to your domain or shared space. When you attach an image version, it appears in the SageMaker Studio Classic Launcher and is available in the **Select image** dropdown list, which users use to launch an activity or change the image used by a notebook.

To make a custom SageMaker image available to all users within a domain, you attach the image to the domain. To make an image available to all users within a shared space, you can attach the image to the shared space. To make an image available to a single user, you attach the image to the user's profile. When you attach an image, SageMaker uses the latest image version by default. You can also attach a specific image version. After you attach the version, you can choose the version from the SageMaker Launcher or the image selector when you launch a notebook.

There is a limit to the number of image versions that can be attached at any given time. After you reach the limit, you must detach a version in order to attach another version of the image.
The following sections demonstrate how to attach a custom SageMaker image to your domain using either the SageMaker console or the AWS CLI. You can only attach a custom image to a share space using the AWS CLI.

**Attach the SageMaker image to a domain**

**Attach the SageMaker image using the Console**

This topic describes how you can attach an existing custom SageMaker image version to your domain using the SageMaker control panel. You can also create a custom SageMaker image and image version, and then attach that version to your domain. For the procedure to create an image and image version, see [Create a custom SageMaker image](#).

To attach an existing image

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the **Domains** page, select the domain to attach the image to.
5. From the **Domain details** page, select the **Environment** tab.
6. On the **Environment** tab, under **Custom SageMaker Studio Classic images attached to domain**, choose **Attach image**.
7. For **Image source**, choose **Existing image**.
8. Choose an existing image from the list.
9. Choose a version of the image from the list.
10. Choose **Next**.
11. Verify the values for **Image name**, **Image display name**, and **Description**.
12. Choose the IAM role. For more information, see [Create a custom SageMaker image](#).
13. (Optional) Add tags for the image.
14. Specify the EFS mount path. This is the path within the image to mount the user's Amazon Elastic File System (EFS) home directory.
15. For **Image type**, select **SageMaker Studio image**
16. For **Kernel name**, enter the name of an existing kernel in the image. For information on how to get the kernel information from the image, see [DEVELOPMENT](#) in the SageMaker Studio Classic Custom Image Samples repository. For more information, see the **Kernel discovery** and **User data** sections of [Custom SageMaker image specifications](#).
17. (Optional) For **Kernel display name**, enter the display name for the kernel.
18. Choose **Add kernel**.
19. Choose **Submit**.

- Wait for the image version to be attached to the domain. When attached, the version is displayed in the **Custom images** list and briefly highlighted.

**Attach the SageMaker image using the AWS CLI**

The following sections demonstrate how to attach a custom SageMaker image when creating a new domain or updating your existing domain using the AWS CLI.

**Attach the SageMaker image to a new domain**

The following section demonstrates how to create a new domain with the version attached. These steps require that you specify the Amazon Virtual Private Cloud (VPC) information and execution role required to create the domain. You perform the following steps to create the domain and attach the custom SageMaker image:

- Get your default VPC ID and subnet IDs.
- Create the configuration file for the domain, which specifies the image.
- Create the domain with the configuration file.

**To add the custom SageMaker image to your domain**

1. Get your default VPC ID.

   ```bash
   aws ec2 describe-vpcs \
   --filters Name=isDefault,Values=true \
   --query "Vpcs[0].VpcId" --output text
   
   vpc-xxxxxxxx
   ```

   The response should look similar to the following.

2. Get your default subnet IDs using the VPC ID from the previous step.

   ```bash
   aws ec2 describe-subnets \
   --filters Name=vpc-id,Values=<vpc-id>
   ```
3. Create a configuration file named `create-domain-input.json`. Insert the VPC ID, subnet IDs, `ImageName`, and `AppImageConfigName` from the previous steps. Because `ImageVersionNumber` isn't specified, the latest version of the image is used, which is the only version in this case.

```json
{
    "DomainName": "domain-with-custom-image",
    "VpcId": "<vpc-id>",
    "SubnetIds": [
        "<subnet-ids>
    ],
    "DefaultUserSettings": {
        "ExecutionRole": "<execution-role>",
        "KernelGatewayAppSettings": {
            "CustomImages": [
                {
                    "ImageName": "custom-image",
                    "AppImageConfigName": "custom-image-config"
                }
            ]
        }
    },
    "AuthMode": "IAM"
}
```

4. Create the domain with the attached custom SageMaker image.

```
aws sagemaker create-domain \
    --cli-input-json file://create-domain-input.json
```

The response should look similar to the following.
Attach the SageMaker image to your current domain

If you have onboarded to a SageMaker domain, you can attach the custom image to your current domain. For more information about onboarding to a SageMaker domain, see Amazon SageMaker domain overview. You don't need to specify the VPC information and execution role when attaching a custom image to your current domain. After you attach the version, you must delete all the apps in your domain and reopen Studio Classic. For information about deleting the apps, see Delete an Amazon SageMaker domain.

You perform the following steps to add the SageMaker image to your current domain.

- Get your DomainID from SageMaker control panel.
- Use the DomainID to get the DefaultUserSettings for the domain.
- Add the imageName and AppImageConfig as a CustomImage to the DefaultUserSettings.
- Update your domain to include the custom image.

To add the custom SageMaker image to your domain

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the Domains page, select the domain to attach the image to.
5. From the Domain details page, select the Domain settings tab.
6. From the Domain settings tab, under General settings, find the DomainId. The ID is in the following format: d-xxxxxxxxxxxxxx.
7. Use the domain ID to get the description of the domain.

aws sagemaker describe-domain \ 
  --domain-id <d-xxxxxxxxxxxxxx>
The response should look similar to the following.

```
{
    "DomainId": "d-xxxxxxxxxxxx",
    "DefaultUserSettings": {
        "KernelGatewayAppSettings": {
            "CustomImages": [
                
            ],
            ...
        }
    }
}
```

8. Save the default user settings section of the response to a file named `default-user-settings.json`.

9. Insert the `ImageName` and `AppImageConfigName` from the previous steps as a custom image. Because `ImageVersionNumber` isn't specified, the latest version of the image is used, which is the only version in this case.

```
{
    "DefaultUserSettings": {
        "KernelGatewayAppSettings": {
            "CustomImages": [
                {
                    "ImageName": "string",
                    "AppImageConfigName": "string"
                },
                ...
            ]
        }
    }
}
```

10. Use the domain ID and default user settings file to update your domain.

```
aws sagemaker update-domain \
    --domain-id <d-xxxxxxxxxxxx> \
    --cli-input-json file://default-user-settings.json
```

The response should look similar to the following.
Attach the SageMaker image to a shared space

You can only attach the SageMaker image to a shared space using the AWS CLI. After you attach the version, you must delete all of the applications in your shared space and reopen Studio Classic. For information about deleting the apps, see [Delete an Amazon SageMaker domain](#).

You perform the following steps to add the SageMaker image to a shared space.

- Get your DomainID from SageMaker control panel.
- Use the DomainID to get the DefaultSpaceSettings for the domain.
- Add the ImageName and AppImageConfig as a CustomImage to the DefaultSpaceSettings.
- Update your domain to include the custom image for the shared space.

**To add the custom SageMaker image to your shared space**

2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the Domains page, select the domain to attach the image to.
5. From the Domain details page, select the Domain settings tab.
6. From the Domain settings tab, under General settings, find the DomainId. The ID is in the following format: d-xxxxxxxxxxxx.
7. Use the domain ID to get the description of the domain.

```
aws sagemaker describe-domain \
  --domain-id <d-xxxxxxxxxxxx>
```

The response should look similar to the following.

```
{
  "DomainArn": "arn:aws:sagemaker:us-east-2:acct-id:domain/d-xxxxxxxxxxxx"
}
```
8. Save the default space settings section of the response to a file named default-space-settings.json.

9. Insert the ImageName and AppImageConfigName from the previous steps as a custom image. Because ImageVersionNumber isn’t specified, the latest version of the image is used, which is the only version in this case.

```json
{
    "DefaultSpaceSettings": {
        "KernelGatewayAppSettings": {
            "CustomImages": [
                {
                    "ImageName": "string",
                    "AppImageConfigName": "string"
                }
            ],
            ...
        }
    }
}
```

10. Use the domain ID and default space settings file to update your domain.

```bash
aws sagemaker update-domain \
    --domain-id <d-xxxxxxxxxxxxx> \
    --cli-input-json file://default-space-settings.json
```

The response should look similar to the following.

```json
{
    "DomainArn": "arn:aws:sagemaker:us-east-2:acct-id:domain/d-xxxxxxxxxxxxx"
}
View the attached image in SageMaker

After you create the custom SageMaker image and attach it to your domain, the image appears in the **Environment** tab of the domain. You can only view the attached images for shared spaces using the AWS CLI by using the following command.

```
aws sagemaker describe-domain \
   --domain-id <d-xxxxxxxxxxxxx>
```

Launch a custom SageMaker image in Amazon SageMaker Studio Classic

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

After you create your custom SageMaker image and attach it to your domain or shared space, the custom image and kernel appear in selectors in the **Change environment** dialog box of the Studio Classic Launcher.

**To launch and select your custom image and kernel**

1. In Amazon SageMaker Studio Classic, open the Launcher. To open the Launcher, choose [Amazon SageMaker Studio Classic](#) at the top left of the Studio Classic interface or use the keyboard shortcut Ctrl + Shift + L.

   To learn about all the available ways to open the Launcher, see [Use the Amazon SageMaker Studio Classic Launcher](#).
2. In the Launcher, in the **Notebooks and compute resources** section, choose **Change environment**.

3. In the **Change environment** dialog, use the dropdown menus to select your **Image** from the **Custom Image** section, and your **Kernel**, then choose **Select**.

4. In the Launcher, choose **Create notebook** or **Open image terminal**. Your notebook or terminal launches in the selected custom image and kernel.

To change your image or kernel in an open notebook, see [Change an Image or a Kernel](#).

**Note**

If you encounter an error when launching the image, check your Amazon CloudWatch logs. The name of the log group is /aws/sagemaker/studio. The name of the log stream is $domainID/$userProfileName/KernelGateway/$appName.
Clean up resources

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following sections show how to clean up the resources you created in the previous sections from the SageMaker console or AWS CLI. You perform the following steps to clean up the resources:

- Detach the image and image versions from your domain.
- Delete the image, image version, and app image config.
- Delete the container image and repository from Amazon ECR. For more information, see Deleting a repository.

Clean up resources from the SageMaker console

The following section shows how to clean up resources from the SageMaker console.

When you detach an image from a domain, all versions of the image are detached. When an image is detached, all users of the domain lose access to the image versions. A running notebook that has a kernel session on an image version when the version is detached, continues to run. When the notebook is stopped or the kernel is shut down, the image version becomes unavailable.

To detach an image

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
4. Under Custom SageMaker Studio Classic images attached to domain, choose the image and then choose Detach.
5. (Optional) To delete the image and all versions from SageMaker, select Also delete the selected images .... This does not delete the associated container images from Amazon ECR.
6. Choose **Detach**.

**Clean up resources from the AWS CLI**

The following section shows how to clean up resources from the AWS CLI.

**To clean up resources**

1. Detach the image and image versions from your domain by passing an empty custom image list to the domain. Open the `default-user-settings.json` file you created in **Attach the SageMaker image to your current domain**. To detach the image and image version from a shared space, open the `default-space-settings.json` file.

2. Delete the custom images and then save the file.

   ```json
   "DefaultUserSettings": {
     "KernelGatewayAppSettings": {
       "CustomImages": [],
       ...
     },
     ...
   }
   ``

3. Use the domain ID and default user settings file to update your domain. To update your shared space, use the default space settings file.

   ```
   aws sagemaker update-domain \
   --domain-id <d-xxxxxxxxxxxx> \
   --cli-input-json file://default-user-settings.json
   ```

   The response should look similar to the following.

   ```
   {
     "DomainArn": "arn:aws:sagemaker:us-east-2:acct-id:domain/d-xxxxxxxxxxxx"
   }
   ```

4. Delete the app image config.

   ```
   aws sagemaker delete-app-image-config \
   --app-image-config-name custom-image-config
   ```
5. Delete the SageMaker image, which also deletes all image versions. The container images in ECR that are represented by the image versions are not deleted.

```bash
aws sagemaker delete-image \
--image-name custom-image
```

### Use lifecycle configurations with Amazon SageMaker Studio Classic

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Lifecycle configurations are shell scripts triggered by Amazon SageMaker Studio Classic lifecycle events, such as starting a new Studio Classic notebook. You can use lifecycle configurations to automate customization for your Studio Classic environment. This customization includes installing custom packages, configuring notebook extensions, preloading datasets, and setting up source code repositories.

Using lifecycle configurations gives you flexibility and control to configure Studio Classic to meet your specific needs. For example, you can create a minimal set of base container images with the most commonly used packages and libraries, then use lifecycle configurations to install additional packages for specific use cases across your data science and machine learning teams.

For example lifecycle configuration scripts, see the Studio Classic Lifecycle Configuration examples GitHub repository. For a blog on implementing lifecycle configuration, see Customize Amazon SageMaker Studio Classic using Lifecycle Configurations.

⚠️ **Note**

Each script has a limit of **16384 characters**.

### Topics

- Customize Studio Classic
Create and associate a lifecycle configuration

Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker provides interactive applications that enable Studio Classic's visual interface, code authoring, and run experience. This series shows how to create a lifecycle configuration and associate it with a SageMaker domain.

Application types can be either JupyterServer or KernelGateway.

- **JupyterServer applications**: This application type enables access to the visual interface for Studio Classic. Every user and shared space in Studio Classic gets its own JupyterServer application.
- **KernelGateway applications**: This application type enables access to the code run environment and kernels for your Studio Classic notebooks and terminals. For more information, see Jupyter Kernel Gateway.

For more information about Studio Classic's architecture and Studio Classic applications, see Use Amazon SageMaker Studio Classic Notebooks.

Topics
- Create a lifecycle configuration from the AWS CLI
- Create a lifecycle configuration from the SageMaker console
Create a lifecycle configuration from the AWS CLI

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following topic shows how to create a lifecycle configuration using the AWS CLI to automate customization for your Studio Classic environment.

Prerequisites

Before you begin, complete the following prerequisites:

- Update the AWS CLI by following the steps in Installing the current AWS CLI Version.
- From your local machine, run aws configure and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.
- Onboard to SageMaker domain by following the steps in Amazon SageMaker domain overview.

Step 1: Create a lifecycle configuration

The following procedure shows how to create a lifecycle configuration script that prints Hello World.

💡 Note

Each script can have up to 16,384 characters.

1. From your local machine, create a file named my-script.sh with the following content.

```bash
#!/bin/bash
set -eux
echo 'Hello World!'```
2. Convert your my-script.sh file into base64 format. This requirement prevents errors that occur from spacing and line break encoding.

   \[ \text{LCC CONTENT} = \text{openssl base64 -A -in my-script.sh} \]

3. Create a lifecycle configuration for use with Studio Classic. The following command creates a lifecycle configuration that runs when you launch an associated KernelGateway application.

   \[
   \text{aws sagemaker create-studio-lifecycle-config} \\
   \text{--region region} \\
   \text{--studio-lifecycle-config-name my-studio-lcc} \\
   \text{--studio-lifecycle-config-content } \text{LCC CONTENT} \\
   \text{--studio-lifecycle-config-app-type KernelGateway}
   \]

   Note the ARN of the newly created lifecycle configuration that is returned. This ARN is required to attach the lifecycle configuration to your application.

**Step 2: Attach the lifecycle configuration to your domain, user profile, or shared space**

To attach the lifecycle configuration, you must update the UserSettings for your domain or user profile, or the SpaceSettings for a shared space. Lifecycle configuration scripts that are associated at the domain level are inherited by all users. However, scripts that are associated at the user profile level are scoped to a specific user, while scripts that are associated at the shared space level are scoped to the shared space.

The following example shows how to create a new user profile with the lifecycle configuration attached. You can also create a new domain or space with a lifecycle configuration attached by using the create-domain and create-space commands, respectively.

Add the lifecycle configuration ARN from the previous step to the settings for the appropriate app type. For example, place it in the JupyterServerAppSettings of the user. You can add multiple lifecycle configurations at the same time by passing a list of lifecycle configurations. When a user launches a JupyterServer application with the AWS CLI, they can pass a lifecycle configuration to use instead of the default. The lifecycle configuration that the user passes must belong to the list of lifecycle configurations in JupyterServerAppSettings.

\[
\text{# Create a new UserProfile} \\
\text{aws sagemaker create-user-profile --domain-id domain-id} \\
\text{--user-profile-name user-profile-name}
\]
The following example shows how to update an existing shared space to attach the lifecycle configuration. You can also update an existing domain or user profile with a lifecycle configuration attached by using the `update-domain` or `update-user-profile` command. When you update the list of lifecycle configurations attached, you must pass all lifecycle configurations as part of the list. If a lifecycle configuration is not part of this list, it will not be attached to the application.

```bash
aws sagemaker update-space --domain-id domain-id \
--space-name space-name \
--region region \
--space-settings '{
"JupyterServerAppSettings": {
"LifecycleConfigArns":
    [lifecycle-configuration-arn-list]
}
}'
```

For information about setting a default lifecycle configuration for a resource, see [Set default lifecycle configurations](#).

**Step 3: Launch application with lifecycle configuration**

After you attach a lifecycle configuration to a domain, user profile, or space, the user can select it when launching an application with the AWS CLI. This section describes how to launch an application with an attached lifecycle configuration. For information about changing the default lifecycle configuration after launching a JupyterServer application, see [Set default lifecycle configurations](#).

Launch the desired application type using the `create-app` command and specify the lifecycle configuration ARN in the `resource-spec` argument.

- The following example shows how to create a JupyterServer application with an associated lifecycle configuration. When creating the JupyterServer, the app-name must be `default`. The lifecycle configuration ARN passed as part of the `resource-spec` parameter must be part
of the list of lifecycle configuration ARNs specified in UserSettings for your domain or user profile, or SpaceSettings for a shared space.

```bash
aws sagemaker create-app --domain-id domain-id \ 
  --region region \ 
  --user-profile-name user-profile-name \ 
  --app-type JupyterServer \ 
  --resource-spec LifecycleConfigArn=lifecycle-configuration-arn \ 
  --app-name default
```

- The following example shows how to create a KernelGateway application with an associated lifecycle configuration.

```bash
aws sagemaker create-app --domain-id domain-id \ 
  --region region \ 
  --user-profile-name user-profile-name \ 
  --app-type KernelGateway \ 
  --resource-spec LifecycleConfigArn=lifecycle-configuration-arn,SageMakerImageArn=sagemaker-image-arn,InstanceType=instance-type \ 
  --app-name app-name
```

Create a lifecycle configuration from the SageMaker console

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following topic shows how to create a lifecycle configuration from the Amazon SageMaker console to automate customization for your Studio Classic environment.

Prerequisites

Before you can begin this tutorial, complete the following prerequisite:

- Onboard to Amazon SageMaker Studio Classic. For more information, see Onboard to Amazon SageMaker Studio Classic.
Step 1: Create a new lifecycle configuration

You can create a lifecycle configuration by entering a script from the Amazon SageMaker console.

Note

Each script can have up to 16,384 characters.

The following procedure shows how to create a lifecycle configuration script that prints Hello World.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose Lifecycle configurations.
4. Choose the Studio tab.
5. Choose Create configuration.
6. Under Select configuration type, select the type of application that the lifecycle configuration should be attached to. For more information about selecting which application to attach the lifecycle configuration to, see Set default lifecycle configurations.
7. Choose Next.
8. In the section called Configuration settings, enter a name for your lifecycle configuration.
9. In the Scripts section, enter the following content.

```
#!/bin/bash
set -eux
echo 'Hello World!'
```

10. (Optional) Create a tag for your lifecycle configuration.
11. Choose Submit.

Step 2: Attach the lifecycle configuration to a domain or user profile

Lifecycle configuration scripts associated at the domain level are inherited by all users. However, scripts that are associated at the user profile level are scoped to a specific user.
You can attach multiple lifecycle configurations to a domain or user profile for both JupyterServer and KernelGateway applications.

⚠️ **Note**

To attach a lifecycle configuration to a shared space, you must use the AWS CLI. For more information, see [Create a lifecycle configuration from the AWS CLI](#).

The following sections show how to attach a lifecycle configuration to your domain or user profile.

**Attach to a domain**

The following shows how to attach a lifecycle configuration to your existing domain from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the domain to attach the lifecycle configuration to.
5. From the **Domain details**, choose the **Environment** tab.
6. Under **Lifecycle configurations for personal Studio apps**, choose **Attach**.
7. Under **Source**, choose **Existing configuration**.
8. Under **Studio lifecycle configurations**, select the lifecycle configuration that you created in the previous step.
9. Select **Attach to domain**.

**Attach to your user profile**

The following shows how to attach a lifecycle configuration to your existing user profile.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the domain that contains the user profile to attach the lifecycle configuration to.
5. Under **User profiles**, select the user profile.
6. From the **User Details** page, choose **Edit**.
7. On the left navigation, choose **Studio settings**.
8. Under **Lifecycle configurations attached to user**, choose **Attach**.
9. Under **Source**, choose **Existing configuration**.
10. Under **Studio lifecycle configurations**, select the lifecycle configuration that you created in the previous step.
11. Choose **Attach to user profile**.

### Step 3: Launch an application with the lifecycle configuration

After you attach a lifecycle configuration to a domain or user profile, you can launch an application with that attached lifecycle configuration. Choosing which lifecycle configuration to launch with depends on the application type.

- **JupyterServer**: When launching a JupyterServer application from the console, SageMaker always uses the default lifecycle configuration. You can't use a different lifecycle configuration when launching from the console. For information about changing the default lifecycle configuration after launching a JupyterServer application, see [Set default lifecycle configurations](#).

  To select a different attached lifecycle configuration, you must launch with the AWS CLI. For more information about launching a JupyterServer application with an attached lifecycle configuration from the AWS CLI, see [Create a lifecycle configuration from the AWS CLI](#).

- **KernelGateway**: You can select any of the attached lifecycle configurations when launching a KernelGateway application using the Studio Classic Launcher.

The following procedure describes how to launch a KernelGateway application with an attached lifecycle configuration from the SageMaker console.

2. Launch Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
3. In the Studio Classic UI, open the Studio Classic Launcher. For more information, see [Use the Amazon SageMaker Studio Classic Launcher](#).
4. In the Studio Classic Launcher, navigate to the **Notebooks and compute resources** section.
5. Click the **Change environment** button.
6. On the **Change environment** dialog, use the dropdown menus to select your **Image**, **Kernel**, **Instance type**, and a **Start-up script**. If there is no default lifecycle configuration, the **Start-up script** value defaults to **No script**. Otherwise, the **Start-up script** value is your default lifecycle configuration. After you select a lifecycle configuration, you can view the entire script.

7. Click **Select**.

8. Back to the Launcher, click the **Create notebook** to launch a new notebook kernel with your selected image and lifecycle configuration.

**Step 4: View logs for a lifecycle configuration**

You can view the logs for your lifecycle configuration after it has been attached to a domain or user profile.

1. First, provide access to CloudWatch for your AWS Identity and Access Management (IAM) role. Add read permissions for the following log group and log stream.
   - **Log group:** /aws/sagemaker/studio
   - **Log stream:** domain/user-profile/app-type/app-name/LifecycleConfigOnStart

   For information about adding permissions, see [Enabling logging from certain AWS services](#).

2. From within Studio Classic, navigate to the **Running Terminals and Kernels** icon to monitor your lifecycle configuration.

3. Select an application from the list of running applications. Applications with attached lifecycle configurations have an attached indicator icon.

4. Select the indicator icon for your application. This opens a new panel that lists the lifecycle configuration.

5. From the new panel, select **View logs**. This opens a new tab that displays the logs.
Set default lifecycle configurations

Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Although you can attach multiple lifecycle configuration scripts to a single resource, you can only set one default lifecycle configuration for each JupyterServer or KernelGateway application. The behavior of the default lifecycle configuration depends on whether it is set for JupyterServer or KernelGateway apps.

- **JupyterServer apps**: When set as the default lifecycle configuration script for JupyterServer apps, the lifecycle configuration script runs automatically when the user signs in to Studio Classic for the first time or restarts Studio Classic. Use this default lifecycle configuration to automate one-time setup actions for the Studio Classic developer environment, such as installing notebook extensions or setting up a GitHub repo. For an example of this, see Customize Amazon SageMaker Studio using Lifecycle Configurations.

- **KernelGateway apps**: When set as the default lifecycle configuration script for KernelGateway apps, the lifecycle configuration is selected by default in the Studio Classic launcher. Users can launch a notebook or terminal with the default script selected, or they can select a different one from the list of lifecycle configurations.

SageMaker supports setting a default lifecycle configuration for the following resources:

- Domains
- User profiles
- Shared spaces

While domains and user profiles support setting a default lifecycle configuration from both the Amazon SageMaker console and AWS Command Line Interface, shared spaces only support setting a default lifecycle configuration from the AWS CLI.
You can set a lifecycle configuration as the default when creating a new resource or updating an existing resource. The following topics demonstrate how to set a default lifecycle configuration using the SageMaker console and AWS CLI.

**Default lifecycle configuration inheritance**

Default lifecycle configurations set at the *domain* level are inherited by all users and shared spaces. Default lifecycle configurations set at the *user* and *shared space* level are scoped to only that user or shared space. User and space defaults override defaults set at the domain level.

A default KernelGateway lifecycle configuration set for a domain applies to all KernelGateway applications launched in the domain. Unless the user selects a different lifecycle configuration from the list presented in the Studio Classic launcher, the default lifecycle configuration is used. The default script also runs if *No Script* is selected by the user. For more information about selecting a script, see [Step 3: Launch an application with the lifecycle configuration](#).

**Topics**

- [Set defaults from the AWS CLI](#)
- [Set defaults from the SageMaker console](#)

**Set defaults from the AWS CLI**

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You can set default lifecycle configuration scripts from the AWS CLI for the following resources:

- Domains
- User profiles
- Shared spaces

The following sections outline how to set default lifecycle configuration scripts from the AWS CLI.
Prerequisites

Before you begin, complete the following prerequisites:

- Update the AWS CLI by following the steps in [Installing the current AWS CLI version](#).
- From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see [Understanding and getting your AWS credentials](#).
- Onboard to SageMaker domain by following the steps in [Amazon SageMaker domain overview](#).
- Create a lifecycle configuration following the steps in [Create and associate a lifecycle configuration](#).

Set a default lifecycle configuration when creating a new resource

To set a default lifecycle configuration when creating a new domain, user profile, or space, pass the ARN of your previously created lifecycle configuration as part of one of the following AWS CLI commands:

- `create-user-profile`
- `create-domain`
- `create-space`

You must pass the lifecycle configuration ARN for the following values in the KernelGateway or JupyterServer default settings:

- `DefaultResourceSpec:LifecycleConfigArn` - This specifies the default lifecycle configuration for the application type.

- `LifecycleConfigArns` - This is the list of all lifecycle configurations attached to the application type. The default lifecycle configuration must also be part of this list.
For example, the following API call creates a new user profile with a default lifecycle configuration.

```bash
aws sagemaker create-user-profile --domain-id domain-id \
  --user-profile-name user-profile-name \
  --region region \
  --user-settings '{
    "KernelGatewayAppSettings": {
      "DefaultResourceSpec": {
        "InstanceType": "ml.t3.medium",
        "LifecycleConfigArn": "lifecycle-configuration-arn"
      },
      "LifecycleConfigArns": ["lifecycle-configuration-arn-list"]
    }
  }'
```

### Set a default lifecycle configuration for an existing resource

To set or update the default lifecycle configuration for an existing resource, pass the ARN of your previously created lifecycle configuration as part of one of the following AWS CLI commands:

- `update-user-profile`
- `update-domain`
- `update-space`

You must pass the lifecycle configuration ARN for the following values in the KernelGateway or JupyterServer default settings:

- **DefaultResourceSpec.LifecycleConfigArn** - This specifies the default lifecycle configuration for the application type.

- **LifecycleConfigArns** - This is the list of all lifecycle configurations attached to the application type. The default lifecycle configuration must also be part of this list.

For example, the following API call updates a user profile with a default lifecycle configuration.

```bash
aws sagemaker update-user-profile --domain-id domain-id \
  --user-profile-name user-profile-name \
  --region region \
  --user-settings '{
```
"KernelGatewayAppSettings": {
    "DefaultResourceSpec": {
        "InstanceType": "ml.t3.medium",
        "LifecycleConfigArn": "lifecycle-configuration-arn"
    },
    "LifecycleConfigArns": ["lifecycle-configuration-arn-list"]
}
'}

The following API call updates a domain to set a new default lifecycle configuration.

```bash
aws sagemaker update-domain --domain-id domain-id \ 
    --region region \ 
    --default-user-settings '{
        "JupyterServerAppSettings": {
            "DefaultResourceSpec": {
                "InstanceType": "ml.t3.medium",
                "LifecycleConfigArn": "lifecycle-configuration-arn"
            },
            "LifecycleConfigArns": ["lifecycle-configuration-arn-list"]
        }
    }'
```

Set defaults from the SageMaker console

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

You can set default lifecycle configuration scripts from the SageMaker console for the following resources.

- Domains
- User profiles
You cannot set default lifecycle configuration scripts for shared spaces from the SageMaker console. For information about setting defaults for shared spaces, see Set defaults from the AWS CLI.

The following sections outline how to set default lifecycle configuration scripts from the SageMaker console.

Topics

• Prerequisites
• Set a default lifecycle configuration for a domain
• Set a default lifecycle configuration for a user profile

Prerequisites

Before you begin, complete the following prerequisites:

• Onboard to SageMaker domain by following the steps in Amazon SageMaker domain overview.
• Create a lifecycle configuration following the steps in Create and associate a lifecycle configuration.

Set a default lifecycle configuration for a domain

The following procedure shows how to set a default lifecycle configuration for a domain from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the list of domains, select the name of the domain to set the default lifecycle configuration for.
3. From the Domain details page, choose the Environment tab.
4. Under Lifecycle configurations for personal Studio apps, select the lifecycle configuration that you want to set as the default for the domain. You can set distinct defaults for JupyterServer and KernelGateway applications.
5. Choose Set as default. This opens a pop up window that lists the current defaults for JupyterServer and KernelGateway applications.
6. Choose Set as default to set the lifecycle configuration as the default for its respective application type.
Set a default lifecycle configuration for a user profile

The following procedure shows how to set a default lifecycle configuration for a user profile from the SageMaker console.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the list of domains, select the name of the domain that contains the user profile that you want to set the default lifecycle configuration for.
3. From the Domain details page, choose the User profiles tab.
4. Select the name of the user profile to set the default lifecycle configuration for. This opens a User Details page.
5. From the User Details page, choose Edit. This opens an Edit user profile page.
6. From the Edit user profile page, choose Step 2 Studio settings.
7. Under Lifecycle configurations attached to user, select the lifecycle configuration that you want to set as the default for the user profile. You can set distinct defaults for JupyterServer and KernelGateway applications.
8. Choose Set as default. This opens a pop up window that lists the current defaults for JupyterServer and KernelGateway applications.
9. Choose Set as default to set the lifecycle configuration as the default for its respective application type.

Debug lifecycle configurations

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following topics show how to get information about and debug your lifecycle configurations.

Topics

- Verify lifecycle configuration process from CloudWatch Logs
- JupyterServer app failure
- KernelGateway app failure
- Lifecycle configuration timeout

### Verify lifecycle configuration process from CloudWatch Logs

Lifecycle configurations only log STDOUT and STDERR.

STDOUT is the default output for bash scripts. You can write to STDERR by appending `>&2` to the end of a bash command. For example, `echo 'hello'>&2`.

Logs for your lifecycle configurations are published to your AWS account using Amazon CloudWatch. These logs can be found in the `/aws/sagemaker/studio` log stream in the CloudWatch console.

2. Choose Logs from the left side. From the dropdown menu, select Log groups.
4. Select the log group.
5. On the Log group details page, choose the Log streams tab.
6. To find the logs for a specific app, search the log streams using the following format:

   ```
domain-id/user-profile-name/app-type/app-name
```

   For example, to find the lifecycle configuration logs for domain `d-m85lcu8vbqmq`, user profile `i-sonic-js`, application type JupyterServer and application name `test-lcc-echo`, use the following search string:

   ```
d-m85lcu8vbqmq/i-sonic-js/JupyterServer/test-lcc-echo
```

7. Select the log stream appended with LifecycleConfigOnStart to view the script execution logs.

### JupyterServer app failure

If your JupyterServer app crashes because of an issue with the attached lifecycle configuration, Studio Classic displays the following error message on the Studio Classic startup screen.
Failed to create SageMaker Studio due to start-up script failure

Select the View script logs link to view the CloudWatch logs for your JupyterServer app.

In the case where the faulty lifecycle configuration is specified in the DefaultResourceSpec of your domain, user profile, or shared space, Studio Classic continues to use the lifecycle configuration even after restarting Studio Classic.

To resolve this error, follow the steps in Set default lifecycle configurations to remove the lifecycle configuration script from the DefaultResourceSpec or select another script as the default. Then launch a new JupyterServer app.

**KernelGateway app failure**

If your KernelGateway app crashes because of an issue with the attached lifecycle configuration, Studio Classic displays the error message in your Studio Classic Notebook.

Choose View script logs to view the CloudWatch logs for your KernelGateway app.

In this case, your lifecycle configuration is specified in the Studio Classic Launcher when launching a new Studio Classic Notebook.

To resolve this error, use the Studio Classic launcher to select a different lifecycle configuration or select No script.

**Note**

A default KernelGateway lifecycle configuration specified in DefaultResourceSpec applies to all KernelGateway images in the domain, user profile, or shared space unless the user selects a different script from the list presented in the Studio Classic launcher. The default script also runs if No Script is selected by the user. For more information on selecting a script, see Step 3: Launch an application with the lifecycle configuration.

**Lifecycle configuration timeout**

There is a lifecycle configuration timeout limitation of 5 minutes. If a lifecycle configuration script takes longer than 5 minutes to run, Studio Classic throws an error.

To resolve this error, ensure that your lifecycle configuration script completes in less than 5 minutes.
To help decrease the run time of scripts, try the following:

- Cut down on necessary steps. For example, limit which conda environments to install large packages in.
- Run tasks in parallel processes.
- Use the nohup command in your script to ensure that hangup signals are ignored and do not stop the execution of the script.

**Update and detach lifecycle configurations**

⚠️ **Important**
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

A lifecycle configuration script can't be changed after it's created. To update your script, you must create a new lifecycle configuration script and attach it to the respective domain, user profile, or shared space. For more information about creating and attaching the lifecycle configuration, see Create and associate a lifecycle configuration.

The following topic shows how to detach a lifecycle configuration using the AWS CLI and SageMaker console.

**Topics**

- **Prerequisites**
- **Detach using the AWS CLI**

**Prerequisites**

Before detaching a lifecycle configuration, you must complete the following prerequisite.

- To successfully detach a lifecycle configuration, no running application can be using the lifecycle configuration. You must first shut down the running applications as shown in Shut Down and Update SageMaker Studio Classic and Studio Classic Apps.
Detach using the AWS CLI

To detach a lifecycle configuration using the AWS CLI, remove the desired lifecycle configuration from the list of lifecycle configurations attached to the resource and pass the list as part of the respective command:

- `update-user-profile`
- `update-domain`
- `update-space`

For example, the following command removes all lifecycle configurations for KernelGateways attached to the domain.

```
aws sagemaker update-domain --domain-id domain-id \
    --region region \
    --default-user-settings '{
        "KernelGatewayAppSettings": {
            "LifecycleConfigArns": [
            ]
        }
    }'
```

Attach Suggested Git Repos to Studio Classic

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](http://example.com).

Amazon SageMaker Studio Classic offers a Git extension for you to enter the URL of a Git repository (repo), clone it into your environment, push changes, and view commit history. In addition to this Git extension, you can also attach suggested Git repository URLs at the Amazon SageMaker domain or user profile level. Then, you can select the repo URL from the list of suggestions and clone that into your environment using the Git extension in Studio Classic.
The following topics show how to attach Git repo URLs to a domain or user profile from the AWS CLI and SageMaker console. You'll also learn how to detach these repository URLs.

Topics

- **Attach a Git Repository from the AWS CLI**
- **Attach a Git Repository from the SageMaker Console**
- **Detach Git Repos**

### Attach a Git Repository from the AWS CLI

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](https://aws.amazon.com/sagemaker/).

The following topic shows how to attach a Git repository URL using the AWS CLI, so that Amazon SageMaker Studio Classic automatically suggests it for cloning. After you attach the Git repository URL, you can clone it by following the steps in [Clone a Git Repository in SageMaker Studio Classic](https://aws.amazon.com/sagemaker/).

**Prerequisites**

Before you begin, complete the following prerequisites:

- Update the AWS CLI by following the steps in [Installing the current AWS CLI Version](https://aws.amazon.com/sagemaker/).
- From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see [Understanding and getting your AWS credentials](https://aws.amazon.com/sagemaker/).
- Onboard to Amazon SageMaker domain. For more information, see [Amazon SageMaker domain overview](https://aws.amazon.com/sagemaker/).

**Attach the Git repo to a domain or user profile**

Git repo URLs associated at the domain level are inherited by all users. However, Git repo URLs that are associated at the user profile level are scoped to a specific user. You can attach multiple Git repo URLs to a domain or user profile by passing a list of repository URLs.
The following sections show how to attach a Git repo URL to your domain and user profile.

**Attach to a domain**

The following command attaches a Git repo URL to an existing domain.

```bash
aws sagemaker update-domain --region region --domain-id domain-id \
  --default-user-settings
JupyterServerAppSettings={CodeRepositories=[{RepositoryUrl="repository"]}
```

**Attach to a user profile**

The following shows how to attach a Git repo URL to an existing user profile.

```bash
aws sagemaker update-user-profile --domain-id domain-id --user-profile-name user-name\n  --user-settings
JupyterServerAppSettings={CodeRepositories=[{RepositoryUrl="repository"]}
```

**Attach a Git Repository from the SageMaker Console**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following topic shows how to associate a Git repository URL from the Amazon SageMaker console to clone it in your Studio Classic environment. After you associate the Git repository URL, you can clone it by following the steps in Clone a Git Repository in SageMaker Studio Classic.

**Prerequisites**

Before you can begin this tutorial, you must onboard to Amazon SageMaker domain. For more information, see Amazon SageMaker domain overview.

**Attach the Git repo to a domain or user profile**

Git repo URLs associated at the domain level are inherited by all users. However, Git repo URL that are associated at the user profile level are scoped to a specific user.
The following sections show how to attach a Git repo URL to a domain and user profile.

**Attach to a domain**

**To attach a Git repo URL to an existing domain**

2. On the left navigation pane, choose *Admin configurations*.
3. Under *Admin configurations*, choose *domains*.
4. Select the domain to attach the Git repo to.
5. On the *domain details* page, choose the *Environment* tab.
6. On the *Suggested code repositories for the domain* tab, choose *Attach*.
7. Under *Source*, enter the Git repository URL.
8. Select *Attach to domain*.

**Attach to a user profile**

The following shows how to attach a Git repository URL to an existing user profile.

**To attach a Git repository URL to a user profile**

2. On the left navigation pane, choose *Admin configurations*.
3. Under *Admin configurations*, choose *domains*.
4. Select the domain that includes the user profile to attach the Git repo to.
5. On the *domain details* page, choose the *User profiles* tab.
6. Select the user profile to attach the Git repo URL to.
7. On the *User details* page, choose *Edit*.
8. On the *Studio settings* page, choose *Attach* from the *Suggested code repositories for the user* section.
9. Under *Source*, enter the Git repository URL.
10. Choose *Attach to user*. 
This guide shows how to detach Git repository URLs from an Amazon SageMaker domain or user profile using the AWS CLI or Amazon SageMaker console.

Topics

- Detach a Git repo using the AWS CLI
- Detach the Git repo using the SageMaker console

Detach a Git repo using the AWS CLI

To detach all Git repo URLs from a domain or user profile, you must pass an empty list of code repositories. This list is passed as part of the JupyterServerAppSettings parameter in an update-domain or update-user-profile command. To detach only one Git repo URL, pass the code repositories list without the desired Git repo URL. This section shows how to detach all Git repo URLs from your domain or user profile using the AWS Command Line Interface (AWS CLI).

Detach from a domain

The following command detaches all Git repo URLs from a domain.

```bash
aws sagemaker update-domain --region region --domain-name domain-name \ --domain-settings JupyterServerAppSettings={CodeRepositories=[]}
```

Detach from a user profile

The following command detaches all Git repo URLs from a user profile.

```bash
aws sagemaker update-user-profile --domain-name domain-name --user-profile-name username
```
Detach the Git repo using the SageMaker console

The following sections show how to detach a Git repo URL from a domain or user profile using the SageMaker console.

Detach from a domain

Use the following steps to detach a Git repo URL from an existing domain.

To detach a Git repo URL from an existing domain

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Select the domain with the Git repo URL that you want to detach.
5. On the domain details page, choose the Environment tab.
6. On the Suggested code repositories for the domain tab, select the Git repository URL to detach.
7. Choose Detach.
8. From the new window, choose Detach.

Detach from a user profile

Use the following steps to detach a Git repo URL from a user profile.

To detach a Git repo URL from a user profile

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Select the domain that includes the user profile with the Git repo URL that you want to detach.
5. On the domain details page, choose the User profiles tab.
6. Select the user profile with the Git repo URL that you want to detach.
7. On the User details page, choose Edit.
8. On the **Studio settings** page, select the Git repo URL to detach from the **Suggested code repositories for the user** tab.

9. Choose **Detach**.

10. From the new window, choose **Detach**.

### Perform Common Tasks in Amazon SageMaker Studio Classic

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](https://aws.amazon.com/sagemakerstudio).

The following sections describe how to perform common tasks in Amazon SageMaker Studio Classic. For an overview of the Studio Classic interface, see [Amazon SageMaker Studio Classic UI Overview](https://aws.amazon.com/sagemakerstudioclassic/).

**Topics**

- [Upload Files to SageMaker Studio Classic](#)
- [Clone a Git Repository in SageMaker Studio Classic](#)
- [Stop a Training Job in SageMaker Studio Classic](#)
- [Use TensorBoard in Amazon SageMaker Studio Classic](#)
- [Using CodeWhisperer and CodeGuru extensions with SageMaker](#)
- [Manage Your Amazon EFS Storage Volume in SageMaker Studio Classic](#)
- [Provide Feedback on SageMaker Studio Classic](#)
- [Shut Down and Update SageMaker Studio Classic and Studio Classic Apps](#)

### Upload Files to SageMaker Studio Classic

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the
When you onboard to Amazon SageMaker Studio Classic, a home directory is created for you in the Amazon Elastic File System (Amazon EFS) volume that was created for your team. Studio Classic can only open files that have been uploaded to your directory. The Studio Classic file browser maps to your home directory.

**Note**

Studio Classic does not support uploading folders. While you can only upload individual files, you can upload multiple files at the same time.

### To upload files to your home directory

1. In the left sidebar, choose the **File Browser** icon ( 
   ![File Browser Icon](image)
2. In the file browser, choose the **Upload Files** icon ( 
   ![Upload Files Icon](image)
3. Select the files you want to upload and then choose **Open**.
4. Double-click a file to open the file in a new tab in Studio Classic.

### Clone a Git Repository in SageMaker Studio Classic

**Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
Classic offers a Git extension for you to enter the URL of a Git repo, clone it into your environment, push changes, and view commit history. If the repo is private and requires credentials to access, then you are prompted to enter your user credentials. This includes your username and personal access token. For more information about personal access tokens, see Managing your personal access tokens.

Admins can also attach suggested Git repository URLs at the Amazon SageMaker domain or user profile level. Users can then select the repo URL from the list of suggestions and clone that into Studio Classic. For more information about attaching suggested repos, see Attach Suggested Git Repos to Studio Classic.

The following procedure shows how to clone a GitHub repo from Studio Classic.

**To clone the repo**

1. In the left sidebar, choose the Git icon ( ).
2. Choose Clone a Repository. This opens a new window.
3. In the Clone Git Repository window, enter the URL in the following format for the Git repo that you want to clone or select a repository from the list of Suggested repositories.
   
   https://github.com/path-to-git-repo/repo.git

4. If you entered the URL of the Git repo manually, select Clone "git-url" from the dropdown menu.
5. Under Project directory to clone into, enter the path to the local directory that you want to clone the Git repo into. If this value is left empty, Studio Classic clones the repo into JupyterLab's root directory.
6. Choose Clone. This opens a new terminal window.
7. If the repo requires credentials, you are prompted to enter your username and personal access token. This prompt does not accept passwords, you must use a personal access token. For more information about personal access tokens, see Managing your personal access tokens.
8. Wait for the download to finish. After the repo has been cloned, the File Browser opens to display the cloned repo.
9. Double click the repo to open it.
10. Choose the Git icon to view the Git user interface which now tracks the repo.
11. To track a different repo, open the repo in the file browser and then choose the **Git** icon.

**Stop a Training Job in SageMaker Studio Classic**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

You can stop a training job with the Amazon SageMaker Studio Classic UI. When you stop a training job, its status changes to Stopping at which time billing ceases. An algorithm can delay termination in order to save model artifacts after which the job status changes to Stopped. For more information, see the `stop_training_job` method in the AWS SDK for Python (Boto3).

**To stop a training job**

1. Follow the [View, search, and compare experiment runs](#) procedure on this page until you open the **Describe Trial Component** tab.
2. At the upper-right side of the tab, choose **Stop training job**. The **Status** at the top left of the tab changes to **Stopped**.
3. To view the training time and billing time, choose **AWS Settings**.

**Use TensorBoard in Amazon SageMaker Studio Classic**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

The following doc outlines how to install and run TensorBoard in Amazon SageMaker Studio Classic.
Note

This guide shows how to open the TensorBoard application through a SageMaker Studio Classic notebook server of an individual SageMaker domain user profile. For a more comprehensive TensorBoard experience integrated with SageMaker Training and the access control functionalities of SageMaker domain, see Use TensorBoard to debug and analyze training jobs in Amazon SageMaker.

Prerequisites

This tutorial requires a SageMaker domain. For more information, see Amazon SageMaker domain overview

Set Up TensorBoardCallback

1. Launch Studio Classic, and open the Launcher. For more information, see Use the Amazon SageMaker Studio Classic Launcher
2. In the Amazon SageMaker Studio Classic Launcher, under Notebooks and compute resources, choose the Change environment button.
3. On the Change environment dialog, use the dropdown menus to select the TensorFlow 2.6 Python 3.8 CPU Optimized Studio Classic Image.
4. Back to the Launcher, click the Create notebook tile. Your notebook launches and opens in a new Studio Classic tab.
5. Run this code from within your notebook cells.
6. Import the required packages.

```python
import os
import datetime
import tensorflow as tf
```
7. Create a Keras model.

```python
mnist = tf.keras.datasets.mnist
(x_train, y_train),(x_test, y_test) = mnist.load_data()
x_train, x_test = x_train / 255.0, x_test / 255.0
def create_model():
```
8. Create a directory for your TensorBoard logs

```python
LOG_DIR = os.path.join(os.getcwd(), "logs/fit/" +
                         datetime.datetime.now().strftime("%Y%m%d-%H%M%S"))
```

9. Run training with TensorBoard.

```python
model = create_model()
model.compile(optimizer='adam',
              loss='sparse_categorical_crossentropy',
              metrics=['accuracy'])

tensorboard_callback = tf.keras.callbacks.TensorBoard(log_dir=LOG_DIR,
                                                        histogram_freq=1)

model.fit(x=x_train,
           y=y_train,
           epochs=5,
           validation_data=(x_test, y_test),
           callbacks=[tensorboard_callback])
```

10. Generate the EFS path for the TensorBoard logs. You use this path to set up your logs from the terminal.

```python
EFS_PATH_LOG_DIR = "".join(LOG_DIR.strip("/").split('/')[1:-1])
print (EFS_PATH_LOG_DIR)
```

Retrieve the EFS_PATH_LOG_DIR. You will need it in the TensorBoard installation section.

**Install TensorBoard**

1. Click on the Amazon SageMaker Studio Classic button on the top left corner of Studio Classic to open the Amazon SageMaker Studio Classic Launcher. This launcher must be opened
from your root directory. For more information, see Use the Amazon SageMaker Studio Classic Launcher.

2. In the Launcher, under Utilities and files, click System terminal.

3. From the terminal, run the following commands. Copy EFS_PATH_LOG_DIR from the Jupyter notebook. You must run this from the /home/sagemaker-user root directory.

```bash
pip install tensorboard
tensorboard --logdir <EFS_PATH_LOG_DIR>
```

Launch TensorBoard

1. To launch TensorBoard, copy your Studio Classic URL and replace lab? with proxy/6006/ as follows. You must include the trailing / character.

   ```text
   https://<YOUR_URL>.studio.region.sagemaker.aws/jupyter/default/proxy/6006/
   ```

2. Navigate to the URL to examine your results.

Using CodeWhisperer and CodeGuru extensions with SageMaker

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker Studio Classic is an integrated machine learning environment where you can build, train, deploy, and analyze your models all in the same application. This topic shows how to generate code recommendations and suggest improvements related to code issues by using Amazon CodeWhisperer and Amazon CodeGuru with Amazon SageMaker.

The following extensions support writing code by generating code recommendations and suggesting improvements related to code issues:

- Amazon CodeWhisperer
What is Amazon CodeWhisperer?

Amazon CodeWhisperer is a service powered by machine learning that helps improve developer productivity. CodeWhisperer achieves this by generating code recommendations based on developers' comments in natural language and their code in the IDE. During preview, Amazon CodeWhisperer is available for the Java, JavaScript, Python, C# and TypeScript programming languages. The service integrates with JupyterLab, Amazon SageMaker Studio Classic, Amazon SageMaker notebook instances, and other integrated development environments (IDEs).

For more information, see the Setting up CodeWhisperer with Amazon SageMaker Studio Classic.

What is Amazon CodeGuru?


CodeGuru Security improves the security of your code in the following ways:

- Proactively detects security policy violations and vulnerabilities.
- Provides recommendations for addressing security risks.
- Suggests improvements to inefficient methods.

From SageMaker, you can call CodeGuru Security by using the open-source Jupyter plugin. You can use CodeGuru Security to scan notebooks for a variety of issues that can affect the security, correctness, reproducibility, maintainability, and performance of your code. For more information, see Tutorial: Run scans with SageMaker Studio Classic and JupyterLab.

Manage Your Amazon EFS Storage Volume in SageMaker Studio Classic

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the
The first time a user on your team onboards to Amazon SageMaker Studio Classic, Amazon SageMaker creates an Amazon Elastic File System (Amazon EFS) volume for the team. A home directory is created in the volume for each user who onboards to Studio Classic as part of your team. Notebook files and data files are stored in these directories. Users don't have access to other team member's home directories. Amazon SageMaker domain does not support mounting custom or additional Amazon EFS volumes.

**Important**

Don't delete the Amazon EFS volume. If you delete it, the domain will no longer function and all of your users will lose their work.

---

**To find your Amazon EFS volume**

1. Open the [SageMaker console](https://console.aws.amazon.com/sagemaker/

2. On the left navigation pane, choose **Admin configurations**.

3. Under **Admin configurations**, choose **domains**.

4. From the **Domains** page, select the domain to find the ID for.

5. From the **Domain details** page, select the **Domain settings** tab.

6. Under **General settings**, find the **Domain ID**. The ID will be in the following format: d-xxxxxxxxxxxxx.

7. Pass the **Domain ID**, as DomainId, to the `describe_domain` method.

8. In the response from `describe_domain`, note the value for the HomeEfsFileSystemId key. This is the Amazon EFS file system ID.

9. Open the [Amazon EFS console](https://console.aws.amazon.com/efs/)

10. Make sure the AWS Region is the same Region that's used by Studio Classic.

11. Under **File systems**, choose the file system ID from the previous step.

12. To verify that you've chosen the correct file system, select the **Tags** heading. The value corresponding to the ManagedByAmazonSageMakerResource key should match the Studio Classic ID.
For information on how to access the Amazon EFS volume, see Using file systems in Amazon EFS.

To delete the Amazon EFS volume, see Deleting an Amazon EFS file system.

Provide Feedback on SageMaker Studio Classic

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Amazon SageMaker takes your feedback seriously. We encourage you to provide feedback.

To provide feedback

1. At the right of SageMaker Studio Classic, find the Feedback icon.
2. Choose a smiley emoji to let us know how satisfied you are with SageMaker Studio Classic and add any feedback you'd care to share with us.
3. Decide whether to share your identity with us, then choose Submit.

Shut Down and Update SageMaker Studio Classic and Studio Classic Apps

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

The following topics show how to shut down and update SageMaker Studio Classic and Studio Classic Apps.
Studio Classic provides a notification icon in the upper-right corner of the Studio Classic UI. This notification icon displays the number of unread notices. To read the notices, select the icon.

Studio Classic provides two types of notifications:

- Upgrade – Displayed when Studio Classic or one of the Studio Classic apps have released a new version. To update Studio Classic, see `Shut down and Update SageMaker Studio Classic`. To update Studio Classic apps, see `Shut down and Update Studio Classic Apps`.

- Information – Displayed for new features and other information.

To reset the notification icon, you must select the link in each notice. Read notifications may still display in the icon. This does not indicate that updates are still needed after you have updated Studio Classic and Studio Classic Apps.

To learn how to update `Amazon SageMaker Data Wrangler`, see `Shut down and Update Studio Classic Apps`.

To ensure that you have the most recent software updates, update Amazon SageMaker Studio Classic and your Studio Classic apps using the methods outlined in the following topics.

**Topics**

- `Shut down and Update SageMaker Studio Classic`
- `Shut down and Update Studio Classic Apps`

**Shut down and Update SageMaker Studio Classic**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see `Amazon SageMaker Studio`.

To update Amazon SageMaker Studio Classic to the latest release, you must shut down the JupyterServer app. You can shut down the JupyterServer app from the SageMaker console, from...
Amazon SageMaker Studio or from within Studio Classic. After the JupyterServer app is shut down, you must reopen Studio Classic through the SageMaker console or from Studio which creates a new version of the JupyterServer app.

You cannot delete the JupyterServer application while the Studio Classic UI is still open in the browser. If you delete the JupyterServer application while the Studio Classic UI is still open in the browser, SageMaker automatically re-creates the JupyterServer application.

Any unsaved notebook information is lost in the process. The user data in the Amazon EFS volume isn't impacted.

Some of the services within Studio Classic, like Data Wrangler, run on their own app. To update these services you must delete the app for that service. To learn more, see Shut down and Update Studio Classic Apps.

Note

A JupyterServer app is associated with a single Studio Classic user. When you update the app for one user it doesn't affect other users.

The following page shows how to update the JupyterServer App from the SageMaker console, from Studio, or from inside Studio Classic.

Shut down and update from the SageMaker console

2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Select the domain that includes the Studio Classic application that you want to update.
5. Under User profiles, select your user name.
6. Under Apps, in the row displaying JupyterServer, choose Action, then choose Delete.
7. Choose Yes, delete app.
8. Type delete in the confirmation box.
9. Choose Delete.
10. After the app has been deleted, launch a new Studio Classic app to get the latest version.
Shut down and update from Studio

1. Navigate to Studio following the steps in Launch Amazon SageMaker Studio.
2. From the Studio UI, find the applications pane on the left side.
3. From the applications pane, select Studio Classic.
4. From the Studio Classic landing page, select the Studio Classic instance to stop.
5. Choose Stop.
6. After the app has been stopped, select Run to use the latest version.

Shut down and update from inside Studio Classic

1. Launch Studio Classic.
2. On the top menu, choose File then Shut Down.
3. Choose one of the following options:
   - Shutdown Server – Shuts down the JupyterServer app. Terminal sessions, kernel sessions, SageMaker images, and instances aren't shut down. These resources continue to accrue charges.
   - Shutdown All – Shuts down all apps, terminal sessions, kernel sessions, SageMaker images, and instances. These resources no longer accrue charges.
4. Close the window.
5. After the app has been deleted, launch a new Studio Classic app to use the latest version.

Shut down and Update Studio Classic Apps

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

To update an Amazon SageMaker Studio Classic app to the latest release, you must first shut down the corresponding KernelGateway app from the SageMaker console. After the KernelGateway app
is shut down, you must reopen it through SageMaker Studio Classic by running a new kernel. The kernel automatically updates. Any unsaved notebook information is lost in the process. The user data in the Amazon EFS volume isn't impacted.

After an application has been shut down for 24 hours, SageMaker deletes all metadata for the application. To be considered an update and retain application metadata, applications must be restarted within 24 hours after the previous application has been shut down. After this time window, creation of an application is considered a new application rather than an update of the previous application.

**Note**

A KernelGateway app is associated with a single Studio Classic user. When you update the app for one user it doesn't effect other users.

**To update the KernelGateway app**

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. Select the domain that includes the application that you want to update.
5. Under **User profiles**, select your user name.
6. Under **Apps**, in the row displaying the **App name**, choose **Action**, then choose **Delete**.

   To update Data Wrangler, delete the app that starts with `sagemaker-data-wrang`.

7. Choose **Yes, delete app**.
8. Type **delete** in the confirmation box.
9. Choose **Delete**.
10. After the app has been deleted, launch a new kernel from within Studio Classic to use the latest version.
Amazon SageMaker Studio Classic Pricing

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

When the first member of your team onboards to Amazon SageMaker Studio Classic, Amazon SageMaker creates an Amazon Elastic File System (Amazon EFS) volume for the team. When this member, or any member of the team, opens Studio Classic, a home directory is created in the volume for the member. A storage charge is incurred for this directory. Subsequently, additional storage charges are incurred for the notebooks and data files stored in the member's home directory. For pricing information on Amazon EFS, see Amazon EFS Pricing.

Additional costs are incurred when other operations are run inside Studio Classic, for example, running a notebook, running training jobs, and hosting a model.

For information on the costs associated with using Studio Classic notebooks, see Usage Metering.

For information about billing along with pricing examples, see Amazon SageMaker Pricing.

If Amazon SageMaker Studio is your default experience, see Amazon SageMaker Studio pricing for more pricing information.

Troubleshooting Amazon SageMaker Studio Classic

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
This topic describes how to troubleshoot common Amazon SageMaker Studio Classic issues during setup and use. The following are common errors that might occur while using Amazon SageMaker Studio Classic. Each error is followed by its solution.

**Studio Classic application issues**

The following issues occur when launching and using the Studio Classic application.

- **Screen not loading: Clearing workspace and waiting doesn't help**

  When launching the Studio Classic application, a pop-up displays the following message. No matter which option is selected, Studio Classic does not load.

  ![Loading message](loading_message.png)

  The Studio Classic application can have a launch delay if multiple tabs are open in the Studio Classic workspace or several files are on Amazon EFS. This pop-up should disappear in a few seconds after the Studio Classic workspace is ready.

  If you continue to see a loading screen with a spinner after selecting either of the options, there could be connectivity issues with the Amazon Virtual Private Cloud used by Studio Classic.

  To resolve connectivity issues with the Amazon Virtual Private Cloud (Amazon VPC) used by Studio Classic, verify the following networking configurations:

  - If your domain is set up in Vpc0nly mode: Verify that there is an Amazon VPC endpoint for AWS STS, or a NAT Gateway for outbound traffic, including traffic over the internet. To do this, follow the steps in [Connect SageMaker Studio Notebooks in a VPC to External Resources](#).

  - If your Amazon VPC is set up with a custom DNS instead of the DNS provided by Amazon: Verify that the routes are configured using Dynamic Host Configuration Protocol (DHCP) for each Amazon VPC endpoint added to the Amazon VPC used by Studio Classic. For more information about setting default and custom DHCP option sets, see [DHCP option sets in Amazon VPC](#).

- **Internal Failure when launching Studio Classic**

  When launching Studio Classic, you are unable to view the Studio Classic UI. You also see an error similar to the following, with **Internal Failure** as the error detail.
The JupyterServer app default encountered a problem and was stopped.

This error can be caused by multiple factors. If completion of these steps does not resolve your issue, create an issue with https://aws.amazon.com/premiumsupport/.

- **Missing Amazon EFS mount target**: Studio Classic uses Amazon EFS for storage. The Amazon EFS volume needs a mount target for each subnet that the Amazon SageMaker domain is created in. If this Amazon EFS mount target is deleted accidentally, the Studio Classic application cannot load because it cannot mount the user's file directory. To resolve this issue, complete the following steps.

**To verify or create mount targets.**

1. Find the Amazon EFS volume that is associated with the domain by using the `DescribeDomain` API call.
2. Sign in to the AWS Management Console and open the Amazon EFS console at https://console.aws.amazon.com/efs/.
3. From the list of Amazon EFS volumes, select the Amazon EFS volume that is associated with the domain.
4. On the Amazon EFS details page, select the **Network** tab. Verify that there are mount targets for all of the subnets that the domain is set up in.
5. If mount targets are missing, add the missing Amazon EFS mount targets. For instructions, see [Creating and managing mount targets and security groups](#).
6. After the missing mount targets are created, launch the Studio Classic application.

- **Conflicting files in the user's `.local` folder**: If you're using JupyterLab version 1 on Studio Classic, conflicting libraries in your `.local` folder can cause issues when launching the Studio Classic application. To resolve this, update your user profile's default JupyterLab version to JupyterLab 3.0. For more information about viewing and updating the JupyterLab version, see [JupyterLab Versioning](#).

- **ConfigurationError: LifecycleConfig when launching Studio Classic**

You can't view the Studio Classic UI when launching Studio Classic. This is caused by issues with the default lifecycle configuration script attached to the domain.
To resolve lifecycle configuration issues

1. View the Amazon CloudWatch Logs for the lifecycle configuration to trace the command that caused the failure. To view the log, follow the steps in [Verify lifecycle configuration process from CloudWatch Logs](#).

2. Detach the default script from the user profile or domain. For more information, see [Update and detach lifecycle configurations](#).

3. Launch the Studio Classic application.

4. Debug your lifecycle configuration script. You can run the lifecycle configuration script from the system terminal to troubleshoot. When the script runs successfully from the terminal, you can attach the script to the user profile or the domain.

- **SageMaker Studio Classic core functionalities are not available.**

If you get this error message when opening Studio Classic, it may be due to Python package version conflicts. This occurs if you used the following commands in a notebook or terminal to install Python packages that have version conflicts with SageMaker package dependencies.

```
pip install
```

```
pip install --user
```

To resolve this issue, complete the following steps:

1. Uninstall recently installed Python packages. If you’re not sure which package to uninstall, create an issue with [https://aws.amazon.com/premiumsupport/](https://aws.amazon.com/premiumsupport/).

2. Restart Studio Classic:
   a. Shut down Studio Classic from the **File** menu.
   b. Wait for one minute.
   c. Reopen Studio Classic by refreshing the page or opening it from the AWS Management Console.

The problem should be resolved if you have uninstalled the package which caused the conflict. To install packages without causing this issue again, use `%pip install` without the `--user` flag.

If the issue persists, create a new user profile and set up your environment with that user profile.
If these solutions don't fix the issue, create an issue with https://aws.amazon.com/premiumsupport/.

- **Unable to open Studio Classic from the AWS Management Console.**

  If you are unable to open Studio Classic and cannot make a new running instance with all default settings, create an issue with https://aws.amazon.com/premiumsupport/.

**KernelGateway application issues**

The following issues are specific to KernelGateway applications that are launched in Studio Classic.

- **Cannot access the Kernel session**

  When the user launches a new notebook, they are unable to connect to the notebook session. If the KernelGateway application's status is *In Service*, you can verify the following to resolve the issue.

- **Check Security Group configurations**

  If the domain is set up in VPCOnly mode, the security group associated with the domain must allow traffic between the ports in the range 8192–65535 for connectivity between the JupyterServer and KernelGateway apps.

**To verify the security group rules**

1. Get the security groups associated with the domain using the `DescribeDomain` API call.
2. Sign in to the AWS Management Console and open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
3. From the left navigation, under **Security**, choose **Security Groups**.
4. Filter by the IDs of the security groups that are associated with the domain.
5. For each security group:
   a. Select the security group.
   b. From the security group details page, view the **Inbound rules**. Verify that traffic is allowed between ports in the range 8192–65535.
For more information about security group rules, see Control traffic to resources using security groups. For more information about requirements to use Studio Classic in VPCOnly mode, see Connect SageMaker Studio Notebooks in a VPC to External Resources.

- **Verify firewall and WebSocket connections**

  If the KernelGateway apps have an InService status and the user is unable to connect to the Studio Classic notebook session, verify the firewall and WebSocket settings.

  1. Launch the Studio Classic application. For more information, see Launch Amazon SageMaker Studio Classic.
  2. Open your web browser's developer tools.
  3. Choose the **Network** tab.
  4. Search for an entry that matches the following format.

     
     ```
     wss://<domain-id>.studio.<region>.sagemaker.aws/jupyter/default/api/kernels/<unique-code>/channels?session_id=<unique-code>
     ```

     If the status or response code for the entry is anything other than 101, then your network settings are preventing the connection between the Studio Classic application and the KernelGateway apps.

     To resolve this issue, contact the team that manages your networking settings to allow list the Studio Classic URL and enable WebSocket connections.

- **Unable to launch an app caused by exceeded resource quotas**

  When a user tries to launch a new notebook, the notebook creation fails with either of the following errors. This is caused by exceeding resource quotas.

  - Unable to start more Apps of AppType [KernelGateway] and ResourceSpec(instanceType=[]) for UserProfile []. Please delete an App with a matching AppType and ResourceSpec, then try again

  Studio Classic supports up to four running KernelGateway apps on the same instance. To resolve this issue, you can do either of the following:

  - Delete an existing KernelGateway application running on the instance, then restart the new notebook.
• Start the new notebook on a different instance type

For more information, see Change an Instance Type.

• An error occurred (ResourceLimitExceeded) when calling the CreateApp operation

In this case, the account does not have sufficient limits to create a Studio Classic application on the specified instance type. To resolve this, navigate to the Service Quotas console at https://console.aws.amazon.com/servicequotas/. In that console, request to increase the Studio KernelGateway Apps running on instance-type instance limit. For more information, see AWS service quotas.

### SageMaker JupyterLab

Create a JupyterLab space within Amazon SageMaker Studio to launch the JupyterLab application. A JupyterLab space is a private space within Studio that manages the storage and compute resources needed to run the JupyterLab application. The JupyterLab application is a web-based interactive development environment (IDE) for notebooks, code, and data. Use the JupyterLab application’s flexible and extensive interface to configure and arrange machine learning (ML) workflows.

By default, the JupyterLab application comes with the SageMaker Distribution image. The distribution image has popular packages, such as the following:

• PyTorch
• TensorFlow
• Keras
• NumPy
• Pandas
• Scikit-learn

Within the JupyterLab application, you can use Amazon CodeWhisperer, a generative AI powered code companion to generate, debug, and explain your code.

Build unified analytics and ML workflows in same Jupyter notebook. Run interactive Spark jobs on Amazon EMR and AWS Glue serverless infrastructure, right from your notebook. Monitor and
debug jobs faster using the inline Spark UI. In a few steps, you can automate your data prep by scheduling the notebook as a job.

The JupyterLab application helps you work collaboratively with your peers. Use the built-in Git integration within the JupyterLab IDE to share and version code. Bring your own file storage system if you have an Amazon EFS volume.

The JupyterLab application runs on a single Amazon Elastic Compute Cloud (Amazon EC2) instance and uses a single Amazon Elastic Block Store (Amazon EBS) volume for storage. You can switch faster instances or increase the Amazon EBS volume size for your needs.

The JupyterLab 4 application runs in a JupyterLab space within Studio. Studio Classic uses the JupyterLab 3 application. JupyterLab 4 provides the following benefits:

- A faster IDE than Amazon SageMaker Studio Classic, especially with large notebooks
- Improved document search
- A more performant and accessible text editor

For more information about JupyterLab, see JupyterLab Documentation.

**Topics**

- JupyterLab user guide
- JupyterLab administrator guide
- Migrating from SageMaker Studio Classic to SageMaker Studio

**JupyterLab user guide**

This guide shows JupyterLab users how to run analytics and machine learning workflows within SageMaker Studio. You can get fast storage and scale your compute up or down, depending on your needs.

JupyterLab only supports private spaces, where private spaces are scoped to a single user in a domain. For information about Studio spaces, see Amazon SageMaker Studio spaces.

To get started using JupyterLab, create a private space and launch your JupyterLab application. The private space running your JupyterLab application is a JupyterLab space. The JupyterLab space uses a single Amazon EC2 instance for your compute and a single Amazon EBS volume for your
storage. Everything in your space such as your code, git profile, and environment variables are stored on the same Amazon EBS volume. The volume has 3000 IOPS and a throughput of 125 megabytes per second (MBps). You can use the fast storage to open and run multiple Jupyter notebooks on the same instance. You can also switch kernels in a notebook very quickly.

Your administrator has configured the default Amazon EBS storage settings for your space. The default storage size is 5 GB, but you can increase the amount of space that you get. You can talk to your administrator to provide you with guidelines.

You can switch the Amazon EC2 instance type that you're using to run JupyterLab, scaling your compute up or down depending on your needs. The **Fast launch** instances start up much faster than the other instances.

Your administrator might provide you with a lifecycle configuration that customizes your environment. You can specify the lifecycle configuration when you create the space.

If your administrator gives you access to an Amazon EFS, you can configure your JupyterLab space to access it.

By default, the JupyterLab application uses the SageMaker distribution image. This includes support for many machine learning, analytics, and deep learning packages. However, if you need a custom image, your administrator can help provide access to the custom images.

The Amazon EBS volume persists independently from the life of an instance. You won't lose your data when you change instances. Use the conda and pip package management libraries to create reproducible custom environments that persist even when you switch instance types.

To get started using JupyterLab, create a space or choose the space that your administrator created for you and open JupyterLab.

Use the following procedure to create a space and open JupyterLab.

**To create a space and open JupyterLab**

1. Open Studio. For information about opening Studio, see [Launch Amazon SageMaker Studio](#).
2. Choose **JupyterLab**.
3. Choose **Create JupyterLab space**.
4. For **Name**, specify the name of the space.
5. Choose **Create space**.
6. (Optional) For **Instance**, specify the Amazon EC2 instance that runs the space.

7. (Optional) For **Image**, specify an image that your administrator provided to customize your environment.

8. (Optional) For **Space Settings**, specify the following:
   - **Storage (GB)** – Up to 100 GB or the amount that your administrator specifies.
   - **Lifecycle Configuration** – A lifecycle configuration that your administrator specifies.
   - **Attach custom EFS filesystem** – An Amazon EFS to which your administrator provides access.

9. Choose **Run space**.

10. Choose **Open JupyterLab**.

### Configure space

After you create a JupyterLab space, you can configure it to do the following:

- Change the instance type.
- Change the storage volume.
- (Admin set up required) Use a custom image.
- (Admin set up required) Use a lifecycle configuration.
- (Admin set up required) Attach a custom Amazon EFS.

⚠️ **Important**

You must stop the JupyterLab space every time you configure it. Use the following procedure to configure the space.

### To configure a space

1. Within Studio, navigate to the JupyterLab application page.
2. Choose the name of the space.
3. (Optional) For **Image**, specify an image that your administrator provided to customize your environment.
4. (Optional) For **Space Settings**, specify the following:

   - **Storage (GB)** – Up to 100 GB or the amount that your administrator configured for the space.
   - **Lifecycle Configuration** – A lifecycle configuration that your administrator provides.
   - **Attach custom EFS filesystem** – An Amazon EFS to which your administrator provides access.

5. Choose **Run space**.

When you open the JupyterLab application, your space has the updated configuration.

After you open JupyterLab, you can configure your environment using the terminal. To open the terminal, navigate to the **Launcher** and choose **Terminal**.

The following are examples of different ways that you can configure an environment in JupyterLab.

**Note**

Within Studio, you can use lifecycle configurations to customize your environment, but we recommend using a package manager instead. Using lifecycle configurations is a more error-prone method. It's easier to add or remove dependencies than it is to debug a lifecycle configuration script. It can also increase the JupyterLab startup time.

For information about lifecycle configurations, see **Using lifecycle configurations with JupyterLab**.

**Customize your environment using a package manager**

Use pip or conda to customize your environment. We recommend using package managers instead of lifecycle configuration scripts.

**Create and activate your custom environment**

This section provides examples of different ways that you can configure an environment in JupyterLab.

A basic conda environment has the minimum number of packages that are required for your workflows in SageMaker. Use the following template to create a basic conda environment:
# initialize conda for shell interaction
conda init

# create a new fresh environment
conda create --name test-env

# check if your new environment is created successfully
conda info --envs

# activate the new environment
conda activate test-env

# install packages in your new conda environment
conda install pip boto3 pandas ipykernel

# list all packages install in your new environment
conda list

# parse env name information from your new environment
export CURRENT_ENV_NAME=$(conda info | grep "active environment" | cut -d : -f 2 | tr -d ' ')

# register your new environment as Jupyter Kernel for execution
python3 -m ipykernel install --user --name $CURRENT_ENV_NAME --display-name "user-env: ($CURRENT_ENV_NAME)"

# to exit your new environment
conda deactivate

The following image shows the location of the environment that you've created.

To change your environment, choose it and select an option from the dropdown menu.
Choose **Select** to select a kernel for the environment.

**Clean up a conda environment**

Cleaning up conda environments that you’re not using can help free up disk space and improve performance. Use the following template to clean up a conda environment:

```bash
# list your environments to select an environment to clean
conda info --envs # or conda info -e

# once you've selected your environment to purge
conda remove --name test-env --all
```
# run conda environment list to ensure the target environment is purged
conda info --envs # or conda info -e

Create a conda environment with a specific Python version

Cleaning up conda environments that you’re not using can help free up disk space and improve performance. Use the following template to clean up a conda environment:

```bash
# create a conda environment with a specific python version
conda create --name py38-test-env python=3.8.10

# activate and test your new python version
conda activate py38-test-env & python3 --version

# Install ipykernel to facilitate env registration
conda install ipykernel

# parse env name information from your new environment
export CURRENT_ENV_NAME=$(conda info | grep "active environment" | cut -d : -f 2 | tr -d ' ')

# register your new environment as Jupyter Kernel for execution
python3 -m ipykernel install --user --name $CURRENT_ENV_NAME --display-name "user-env: ($CURRENT_ENV_NAME)"

# deactivate your py38 test environment
conda deactivate
```

Create a conda environment with a specific set of packages

Use the following template to create a conda environment with a specific version of Python and set of packages:

```bash
# prefill your conda environment with a set of packages,
conda create --name py38-test-env python=3.8.10 pandas matplotlib=3.7 scipy ipykernel

# activate your conda environment and ensure these packages exist
conda activate py38-test-env
```
Clone conda from an existing environment

Clone your conda environment to preserve its working state. You experiment in the cloned environment without having to worry about introducing breaking changes in your test environment.

Use the following command to clone an environment.

```bash
# create a fresh env from a base environment
conda create --name py310-base-ext --clone base # replace 'base' with another env

# activate your conda environment and ensure these packages exist
conda activate py310-base-ext

# install ipykernel to register your env
conda install ipykernel

# parse env name information from your new environment
export CURRENT_ENV_NAME=$(conda info | grep "active environment" | cut -d : -f 2 | tr -d ' ')

# register your new environment as Jupyter Kernel for execution
python3 -m ipykernel install --user --name $CURRENT_ENV_NAME --display-name "user-env: ($CURRENT_ENV_NAME)"

# deactivate your conda environment
conda deactivate
```
### Clone conda from a reference YAML file

Create a conda environment from a reference YAML file. The following is an example of a YAML file that you can use.

```yaml
# anatomy of a reference environment.yml
name: py311-new-env
channels:
  - conda-forge
dependencies:
  - python=3.11
  - numpy
  - pandas
  - scipy
  - matplotlib
  - pip
  - ipykernel
  - pip:
    - git+https://github.com/huggingface/transformers
```

Under `pip`, we recommend specifying only the dependencies that aren't available with conda.

Use the following commands to create a conda environment from a YAML file.

```bash
# create your conda environment
conda create -f environment.yml

# activate your env
conda activate py311-new-env
```

### Share environments between instance types

You can share conda environments by saving them to an Amazon EFS directory outside of your Amazon EBS volume. Another user can access the environment in the directory where you saved it.
Important

There are limitations with sharing your environments. For example, we don't recommend an environment meant to run on a GPU Amazon EC2 instance over an environment running on a CPU instance.

Use the following commands as a template to specify the target directory where you're creating a custom environment. You're creating a conda within a particular path. You create it within the Amazon EFS directory. You can spin up a new instance and do conda activate path and do it within the Amazon EFS.

```
# if you know your environment path for your conda environment
conda create --prefix /home/sagemaker-user/my-project/py39-test python=3.9

# activate the env with full path from prefix
conda activate home/sagemaker-user/my-project/py39-test

# parse env name information from your new environment
export CURRENT_ENV_NAME=$(conda info | grep "active environment" | awk -F' : ' '{print $2}' | awk -F'/' '{print $NF}')</n

# register your new environment as Jupyter Kernel for execution
python3 -m ipykernel install --user --name $CURRENT_ENV_NAME --display-name "user-env-prefix:($CURRENT_ENV_NAME)"

# deactivate your conda environment
conda deactivate
```

JupyterLab administrator guide

This guide for administrators describes SageMaker JupyterLab resources, such as those from Amazon Elastic Block Store (Amazon EBS) and Amazon Elastic Compute Cloud (Amazon EC2). The topics also show how to provide user access and change storage size.

A SageMaker JupyterLab space is composed of the following resources:

- A distinct Amazon EBS volume that stores all of the data, such as the code and the environment variables.
• The Amazon EC2 instance used to run the space.
• The image used to run JupyterLab.

Note

Applications do not have access to the EBS volume of other applications. For example, Code Editor, based on Code-OSS, Visual Studio Code - Open Source doesn't have access to the EBS volume for JupyterLab. For more information about EBS volumes, see Amazon Elastic Block Store (Amazon EBS).

You can use the Amazon SageMaker API to do the following:

• Change the default storage size of the EBS volume for your users.
• Change the maximum size of the EBS storage
• Specify the user settings for the application. For example, you can specify whether the user is using a custom image or a code repository.
• Specify the support application type.

The default size of the Amazon EBS volume is 5 GB. You can increase the volume size to a maximum of 16,384 GB. If you don't do anything, your users can increase their volume size to 100 GB. The volume size can be changed only once within a six hour period.

The kernels associated with the JupyterLab application run on the same Amazon EC2 instance that runs JupyterLab. When you create a space, the latest version of the SageMaker Distribution Image is used by default. For more information about SageMaker Distribution Images, see SageMaker Distribution Images.

Important

For information about updating the space to use the latest version of the SageMaker Distribution Image, see Updating the SageMaker Distribution Image.

The following sections walk you through the configurations that you need to perform as an administrator.
Give your users access to private spaces

To give users access to private spaces, you must attach a permissions policy to their IAM roles. You can also use the permissions policy to restrict private spaces and their associated applications to a specific user profile.

The following permission policy grants access to private spaces. This allows users to create their own private space and list other private spaces within their domain. A user with this policy can't access the private space of a different user. For information about Studio spaces, see Amazon SageMaker Studio spaces.

The policy provides users with permissions to the following:

- Private spaces.
- A user profile for accessing private spaces.

To provide permissions, you can attach the following policy to the IAM roles of your users. You can also use this policy to restrict private spaces, and their associated applications, to a specific user profile.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": "sagemaker:CreatePrivateSpace",
        }
    ]
}
```
"Action": [
    "sagemaker:CreateApp",
    "sagemaker:DeleteApp"
],
"Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:app/*",
"Condition": {
    "Null": {
        "sagemaker:OwnerUserProfileArn": "true"
    }
}
},
{
    "Sid": "SMStudioCreatePresignedDomainUrlForUserProfile",
    "Effect": "Allow",
    "Action": [
        "sagemaker:CreatePresignedDomainUrl"
    ],
    "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:user-profile/${sagemaker:DomainId}/${sagemaker:UserProfileName}"
},
{
    "Sid": "SMStudioAppPermissionsListAndDescribe",
    "Effect": "Allow",
    "Action": [
        "sagemaker:ListApps",
        "sagemaker:ListDomains",
        "sagemaker:ListUserProfiles",
        "sagemaker:ListSpaces",
        "sagemaker:DescribeApp",
        "sagemaker:DescribeDomain",
        "sagemaker:DescribeUserProfile",
        "sagemaker:DescribeSpace"
    ],
    "Resource": "*"
},
{
    "Sid": "SMStudioAppPermissionsTagOnCreate",
    "Effect": "Allow",
    "Action": [
        "sagemaker:AddTags"
    ],
    "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:*/**",
    "Condition": {
        "Null": {
            ...
"sagemaker:TaggingAction": "false"
],
"Sid": "SMStudioRestrictSharedSpacesWithoutOwners",
"Effect": "Allow",
"Action": [
  "sagemaker:CreateSpace",
  "sagemaker:UpdateSpace",
  "sagemaker:DeleteSpace"
],
"Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:space/${sagemaker:DomainId}/*",
"Condition": {
  "Null": {
    "sagemaker:OwnerUserProfileArn": "true"
  }
}
},
{
  "Sid": "SMStudioRestrictSpacesToOwnerUserProfile",
  "Effect": "Allow",
  "Action": [
    "sagemaker:CreateSpace",
    "sagemaker:UpdateSpace",
    "sagemaker:DeleteSpace"
  ],
  "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:space/${sagemaker:DomainId}/*",
  "Condition": {
    "ArnLike": {
      "sagemaker:OwnerUserProfileArn": "arn:aws:sagemaker:${AWS Region: $1112223333}:user-profile/${sagemaker:DomainId}/${sagemaker:UserProfileName}"
    },
    "StringEquals": {
      "sagemaker:SpaceSharingType": [
        "Private",
        "Shared"
      ]
    }
  }
}
Change the default storage size for your JupyterLab users

You can change the default storage settings for your users. You can also change the default storage settings based on your organizational requirements and the needs of your users.

To change the storage size, this section provides commands to do the following:

1. Update the Amazon EBS storage settings in the Amazon SageMaker domain (domain).
2. Create a user profile and specify the storage settings within it.

Use the following AWS Command Line Interface (AWS CLI) commands to change the default storage size.

Use the following AWS CLI command to update the domain:
Use the following AWS CLI command to create the user profile and specify the default storage settings:

```
aws --region $REGION sagemaker create-user-profile \
--domain-id $DOMAIN_ID \
--user-profile-name $USER_PROFILE_NAME \
--user-settings '{
    "SpaceStorageSettings": {
        "DefaultEbsStorageSettings":{
            "DefaultEbsVolumeSizeInGb":5,
            "MaximumEbsVolumeSizeInGb":100
        }
    }
}'}
```

Use the following AWS CLI commands to update the default storage settings in the user profile:

```
aws --region $REGION sagemaker update-user-profile \
--domain-id $DOMAIN_ID \
--user-profile-name $USER_PROFILE_NAME \
--user-settings '{
    "SpaceStorageSettings": {
        "DefaultEbsStorageSettings":{
            "DefaultEbsVolumeSizeInGb":25,
            "MaximumEbsVolumeSizeInGb":200
        }
    }
}'}
```

Use the following AWS CLI command to update the domain's default storage settings:

```
aws --region $REGION sagemaker update-domain \
--domain-id $DOMAIN_ID \
--default-user-settings '{
    "SpaceStorageSettings": {
        "DefaultEbsStorageSettings":{
            "DefaultEbsVolumeSizeInGb":5,
            "MaximumEbsVolumeSizeInGb":100
        }
    }
}'}
```
Using lifecycle configurations with JupyterLab

Lifecycle configurations are shell scripts that are triggered by JupyterLab lifecycle events, such as starting a new JupyterLab notebook. You can use lifecycle configurations to automate customization for your JupyterLab environment. This customization includes installing custom packages, configuring notebook extensions, preloading datasets, and setting up source code repositories.

Using lifecycle configurations gives you flexibility and control to configure JupyterLab to meet your specific needs. For example, you can create a minimal set of base container images with the most commonly used packages and libraries. Then you can use lifecycle configurations to install additional packages for specific use cases across your data science and machine learning teams.

Note

Each script has a limit of 16,384 characters.

Topics

- Create and associate a lifecycle configuration
- Debug lifecycle configurations
- Detach lifecycle configurations

Create and associate a lifecycle configuration

This topic includes instructions for creating and associating a lifecycle configuration with JupyterLab. You use the AWS Command Line Interface (AWS CLI) or the AWS Management Console to automate customization for your JupyterLab environment.

Lifecycle configurations are shell scripts triggered by JupyterLab lifecycle events, such as starting a new JupyterLab notebook. For more information about lifecycle configurations, see Using lifecycle configurations with JupyterLab.

Create a lifecycle configuration (AWS CLI)

Learn how to create a lifecycle configuration using the AWS Command Line Interface (AWS CLI) to automate customization for your Studio environment.
Prerequisites

Before you begin, complete the following prerequisites:

• Update the AWS CLI by following the steps in Installing the current AWS CLI Version.
• From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.
• Onboard to Amazon SageMaker domain. For conceptual information, see Amazon SageMaker domain overview. For a quickstart guide, see Quick onboard to Amazon SageMaker domain.

Step 1: Create a lifecycle configuration

The following procedure shows how to create a lifecycle configuration script that prints Hello World.

⚠️ Note

Each script can have up to 16,384 characters.

1. From your local machine, create a file named `my-script.sh` with the following content:

   ```bash
   #!/bin/bash
   set -eux
   echo 'Hello World!'
   ```

2. Use the following to convert your `my-script.sh` file into base64 format. This requirement prevents errors that occur from spacing and line break encoding.

   ```bash
   LCC_CONTENT=`openssl base64 -A -in my-script.sh`
   ```

3. Create a lifecycle configuration for use with Studio. The following command creates a lifecycle configuration that runs when you launch an associated JupyterLab application:

   ```bash
   aws sagemaker create-studio-lifecycle-config \
   --region region \ 
   --studio-lifecycle-config-name my-jl-lcc \ 
   --studio-lifecycle-config-content $LCC_CONTENT \ 
   --studio-lifecycle-config-app-type JupyterLab
   ```
Note the ARN of the newly created lifecycle configuration that is returned. This ARN is required to attach the lifecycle configuration to your application.

**Step 2: Attach the lifecycle configuration to your Amazon SageMaker domain (domain) and user profile**

To attach the lifecycle configuration, you must update the UserSettings for your domain or user profile. Lifecycle configuration scripts that are associated at the domain level are inherited by all users. However, scripts that are associated at the user profile level are scoped to a specific user.

You can create a new user profile, domain, or space with a lifecycle configuration attached by using the following commands:

- `create-user-profile`
- `create-domain`
- `create-space`

The following command creates a user profile with a lifecycle configuration. Add the lifecycle configuration ARN from the preceding step to the JupyterLabAppSettings of the user. You can add multiple lifecycle configurations at the same time by passing a list of them. When a user launches a JupyterLab application with the AWS CLI, they can specify a lifecycle configuration instead of using the default one. The lifecycle configuration that the user passes must belong to the list of lifecycle configurations in JupyterLabAppSettings.

```bash
# Create a new UserProfile
aws sagemaker create-user-profile --domain-id domain-id \   --user-profile-name user-profile-name \   --region region \   --user-settings '{"JupyterLabAppSettings": { "LifecycleConfigArns": [lifecycle-configuration-arn-list] }}'
```
Create a lifecycle configuration (Console)

Learn how to create a lifecycle configuration using the AWS Management Console to automate customization for your Studio environment.

Step 1: Create a lifecycle configuration

Use the following procedure to create a lifecycle configuration script that prints Hello World.

To create a lifecycle configuration

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose Lifecycle configurations.
4. Choose the JupyterLab tab.
5. Choose Create configuration.
6. For Name, specify the name of the lifecycle configuration.
7. For the text box under Scripts, specify the following lifecycle configuration:

```bash
#!/bin/bash
set -eux
echo 'Hello World!
```
8. Choose Create configuration.

Step 2: Attach the lifecycle configuration to your Amazon SageMaker domain (domain) and user profile

Lifecycle configuration scripts associated at the domain level are inherited by all users. However, scripts that are associated at the user profile level are scoped to a specific user.

You can attach multiple lifecycle configurations to a domain or user profile for JupyterLab.

Use the following procedure to attach a lifecycle configuration to a domain.

To attach a lifecycle configuration to a domain

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the domain to attach the lifecycle configuration to.
5. From the **Domain details**, choose the **Environment** tab.
6. Under **Lifecycle configurations for personal Studio apps**, choose **Attach**.
7. Under **Source**, choose **Existing configuration**.
8. Under **Studio lifecycle configurations**, select the lifecycle configuration that you created in the previous step.
9. Select **Attach to domain**.

Use the following procedure to attach a lifecycle configuration to a user profile.

**To attach a lifecycle configuration to a user profile**

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the domain that contains the user profile to attach the lifecycle configuration to.
5. Under **User profiles**, select the user profile.
6. From the **User Details** page, choose **Edit**.
7. On the left navigation, choose **Studio settings**.
8. Under **Lifecycle configurations attached to user**, choose **Attach**.
9. Under **Source**, choose **Existing configuration**.
10. Under **Studio lifecycle configurations**, select the lifecycle configuration that you created in the previous step.
11. Choose **Attach to user profile**.

**Debug lifecycle configurations**

The following topics show how to get information about and debug your lifecycle configurations.

**Topics**

- [Verify lifecycle configuration process from CloudWatch Logs](#)
Verify lifecycle configuration process from CloudWatch Logs

Lifecycle configurations only log STDOUT and STDERR.

STDOUT is the default output for bash scripts. You can write to STDERR by appending `>&2` to the end of a bash command. For example, `echo 'hello'>&2`.

Logs for your lifecycle configurations are published to your AWS account using Amazon CloudWatch. These logs can be found in the `/aws/sagemaker/studio` log stream in the CloudWatch console.

2. Choose Logs from the left navigation pane. From the dropdown menu, select Log groups.
4. Select the log group.
5. On the Log group details page, choose the Log streams tab.
6. To find the logs for a specific app, search the log streams using the following format:

   `domain-id/user-profile-name/app-type/app-name`

   The following search string finds the lifecycle configuration logs for the domain `d-m85lcu8vbqzm`, user profile `i-sonic-js`, application type `JupyterLab`, and application name `test-lcc-echo`:

   `d-m85lcu8vbqzm/i-sonic-js/JupyterLab/test-lcc-echo`

7. To view the script execution logs, select the log stream appended with LifecycleConfigOnStart.

Lifecycle configuration timeout

There is a lifecycle configuration timeout limitation of 5 minutes. If a lifecycle configuration script takes longer than 5 minutes to run, you get an error.

To resolve this error, make sure that your lifecycle configuration script completes in less than 5 minutes.
To help decrease the runtime of scripts, try the following:

- Reduce unnecessary steps. For example, limit which conda environments to install large packages in.
- Run tasks in parallel processes.
- Use the nohup command in your script to make sure that hangup signals are ignored so that the script runs without stopping.

**Detach lifecycle configurations**

To update your script, you must create a new lifecycle configuration script and attach it to the respective Amazon SageMaker domain (domain), user profile, or shared space. A lifecycle configuration script can't be changed after it's created. For more information about creating and attaching the lifecycle configuration, see [Create and associate a lifecycle configuration](#).

The following section shows how to detach a lifecycle configuration using the AWS Command Line Interface (AWS CLI).

**Detach using the AWS CLI**

To detach a lifecycle configuration using the (AWS CLI), remove the desired lifecycle configuration from the list of lifecycle configurations attached to the resource. You then pass the list as part of the respective command:

- `update-user-profile`
- `update-domain`
- `update-space`

For example, the following command removes all lifecycle configurations for the JupyterLab application that's attached to the domain.

```
aws sagemaker update-domain --domain-id domain-id \ 
--region region \ 
--default-user-settings '{
 "JupyterLabAppSettings": {
  "LifecycleConfigArns": [],
  }
}
```
**Attach Git repos**

JupyterLab offers a Git extension to enter the URL of a Git repository (repo), clone it into an environment, push changes, and view the commit history. You can also attach suggested Git repo URLs to a Amazon SageMaker domain (domain) or user profile.

The following sections show how to attach Git repo URLs to a domain or user profile from the AWS Command Line Interface (AWS CLI) and the SageMaker console. A section also provides AWS CLI commands to detach these repository URLs.

**Attach a Git repository (AWS CLI)**

This section shows how to attach a Git repository (repo) URL using the AWS CLI. After you attach the Git repo URL, you can clone it by following the steps in [Clone a Git repo in Amazon SageMaker Studio](#).

**Prerequisites**

Before you begin, complete the following prerequisites:

- Update the AWS CLI by following the steps in [Installing the current AWS Command Line Interface Version](#).
- From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see [Understanding and getting your AWS credentials](#).
- Onboard to Amazon SageMaker domain. For more information, see [Amazon SageMaker domain overview](#).

**Attach the Git repo to a Amazon SageMaker domain (domain) or user profile**

Git repo URLs that are associated at the domain level are inherited by all users. However, Git repo URLs that are associated at the user profile level are scoped to a specific user. You can attach multiple Git repo URLs to a Amazon SageMaker domain or to a user profile by passing a list of repository URLs.

The following sections show how to attach a Git repo URL to your domain and your user profile.

**Attach to a Amazon SageMaker domain**

The following command attaches a Git repo URL to an existing domain:

```bash
jupyterlab attach-repo --domain-domain-name repo-url
```
Attach a user profile

The following command attaches a Git repo URL to an existing user profile:

```bash
aws sagemaker update-user-profile --domain-id domain-id --user-profile-name user-name
   --user-settings
   JupyterLabAppSettings={CodeRepositories=[{RepositoryUrl="repository"}]}
```

Clone a Git repo in Amazon SageMaker Studio

Amazon SageMaker Studio connects to a local Git repo only. To access the files in the repo, clone the Git repo from within Studio. To do so, Studio offers a Git extension for you to enter the URL of a Git repo, clone it into your environment, push changes, and view commit history.

If the repo is private and requires credentials to access, you receive a prompt to enter your user credentials. Your credentials include your username and personal access token. For more information about personal access tokens, see Managing your personal access tokens.

Admins can also attach suggested Git repository URLs at the Amazon SageMaker domain or user profile level. Users can then select the repo URL from the list of suggestions and clone that into Studio. For more information about attaching suggested repos, see Attach Suggested Git Repos to Studio Classic.

Detach Git repo URLs

This section shows how to detach Git repository URLs from an Amazon SageMaker domain (domain) or a user profile. You can detach repo URLs by using the AWS Command Line Interface (AWS CLI) or the Amazon SageMaker console.

Detach a Git repo using the AWS CLI

To detach all Git repo URLs from a domain or user profile, you must pass an empty list of code repositories. This list is passed as part of the JupyterLabAppSettings parameter in an update-domain or update-user-profile command. To detach only one Git repo URL, pass the code repositories list without the desired Git repo URL.
**Detach from an Amazon SageMaker domain**

The following command detaches all Git repo URLs from a domain:

```bash
aws sagemaker update-domain --region region --domain-name domain-name \ --domain-settings JupyterLabAppSettings={CodeRepositories=[]}
```

**Detach from a user profile**

The following command detaches all Git repo URLs from a user profile:

```bash
aws sagemaker update-user-profile --domain-name domain-name --user-profile-name user-name\ --user-settings JupyterLabAppSettings={CodeRepositories=[]}
```

**Customize environments using custom images**

If you need functionality that is different than what's provided by SageMaker distribution, you can bring your own image with your custom extensions and packages. You can also use it to personalize the JupyterLab UI for your own branding or compliance needs.

For a tutorial that helps you create an image that your users can run in their JupyterLab environment, see [Provide users with access to custom images](#).

For requirements for your image, see [Dockerfile specifications](#).

**Topics**

- [Provide users with access to custom images](#)
- [Dockerfile specifications](#)

**Provide users with access to custom images**

This documentation provides step-by-step instructions to provide your users with access to custom images within their JupyterLab environments. You can use the information on this page to create custom environments for your user's workflows. The process involves utilizing:

- Docker
- AWS Command Line Interface
- Amazon Elastic Container Registry
After following the guidance on this page, JupyterLab users on the Amazon SageMaker domain will have access to the custom image and environment from their Jupyter spaces to empower their machine learning workflows.

⚠️ Important

This page assumes that you have the AWS Command Line Interface and Docker installed on your local machine.

To have your users successfully run their image within JupyterLab, you must do the following:

To have your users successfully run the image

1. Create the Dockerfile
2. Build the image from the Dockerfile
3. Upload the image to Amazon Elastic Container Registry
4. Attach the image to your Amazon SageMaker domain
5. Have your users access the image from your JupyterLab space

Step 1: Create the Dockerfile

Create a Dockerfile to define the steps needed to create the environment needed to run the application in your user's container.

⚠️ Important

Your Dockerfile must meet the specifications provided in Dockerfile specifications.

Use the following Dockerfile template to create an Amazon Linux 2 image:

```bash
FROM public.ecr.aws/amazonlinux/amazonlinux:2
ARG NB_USER="sagemaker-user"
```
ARG NB_UID="1000"
ARG NB_GID="100"
RUN yum install --assumeyes python3 shadow-utils && \
    useradd --create-home --shell /bin/bash --gid "${NB_GID}" --uid ${NB_UID}
${NB_USER} && \
    yum clean all && \
    python3 -m pip install jupyterlab

RUN python3 -m pip install --upgrade pip

RUN python3 -m pip install --upgrade urllib3==1.26.6

USER ${NB_UID}
CMD jupyter lab --ip 0.0.0.0 --port 8888 \
   --ServerApp.base_url="/jupyterlab/default" \
   --ServerApp.token='' \
   --ServerApp.allow_origin='*' 

Use the following Dockerfile template to create an Amazon SageMaker Distribution Image:

FROM public.ecr.aws/sagemaker/sagemaker-distribution:latest-cpu
ARG NB_USER="sagemaker-user"
ARG NB_UID=1000
ARG NB_GID=100

ENV MAMBA_USER=$NB_USER

USER root

RUN apt-get update
RUN micromamba install sagemaker-inference --freeze-installed --yes --channel conda-forge --name base

USER $MAMBA_USER

ENTRYPOINT ["jupyter-lab"]
CMD ["--ServerApp.ip=0.0.0.0", "--ServerApp.port=8888", "--ServerApp.allow_origin=''", "--ServerApp.token=''", "--ServerApp.base_url=/jupyterlab/default"]
Step 2: Build the Dockerfile

In the same directory as your Dockerfile, build your image using the following command:

```
docker build -t username/imagename:tag your-account-id.dkr.ecr.AWS Region.amazonaws.com/your-repository-name:tag
```

⚠️ Important
Your image must be tagged in the following format: `123456789012.dkr.ecr.your-region.amazonaws.com/your-repository-name:tag`
You won’t be able to push it to an Amazon Elastic Container Registry repository otherwise.

Step 3: Push the image to the Amazon Elastic Container Registry repository

After you’ve built your image, log in to your Amazon ECR repository using the following command:

```
aws ecr get-login-password --region AWS Region | docker login --username AWS --password-stdin 123456789012.dkr.ecr.AWS Region.amazonaws.com
```

After you’ve logged in, push your Dockerfile using the following command:

```
docker push 123456789012.dkr.ecr.AWS Region.amazonaws.com/your-repository-name:tag
```

Step 4: Attach image to the Amazon SageMaker domain of your users

After you’ve pushed the image, you must access it from your Amazon SageMaker domain. Use the following procedure to attach the image to a SageMaker domain:

1. Open the SageMaker console.
2. Under Admin configurations, choose domains.
3. From the list of domains, select a domain.
4. Open the **Environment** tab.

5. For **Custom images for personal Studio apps**, choose **Attach image**.

6. Specify the image source.

7. Choose **Next**.

8. Choose **Submit**.

Your users can now select the image that you've attached to their Domain from their JupyterLab space.

**Dockerfile specifications**

The image that you specify in your Dockerfile must match the specifications in the following sections to create the image successfully.

**Running the image**

- **Entrypoint** – We recommend embedding the entry point into the image using the Docker `CMD` or `Entrypoint` instructions. You can also configure `ContainerEntrypoint` and `ContainerArguments` that are passed to the container at runtime.

- **EnvVariables** – With Studio, you can configure `ContainerEnvironment` variables that are made available to a container. The environment variable is overwritten with the environment variables from SageMaker. To provide you with a better experience, the environment variables are usually `AWS_` and `SageMaker_` namespaced to give priority to platform environments.

The following are the environment variables:

- `AWS_REGION`
- `AWS_DEFAULT_REGION`
- `AWS_CONTAINER_CREDENTIALS_RELATIVE_URI`
- `SageMaker_SPACE_NAME`

**Specifications for the user and file system**

- **WorkingDirectory** – The Amazon EBS volume used as storage. It's attached to `/home/sagemaker-user`, the working directory of the container. Use the `WORKDIR` command to define the working directory.
Amazon SageMaker

Developer Guide

• UID – The user ID of the Docker container. UID=1000 is a supported value. You can add sudo
access to your users. The IDs are remapped to prevent a process running in the container from
having more privileges than necessary.
• GID – The group ID of the Docker container. GID=100 is a supported value. You can add sudo
access to your users. The IDs are remapped to prevent a process running in the container from
having more privileges than necessary.
• Metadata directories – The /opt/.sagemakerintenral and /opt/ml directories that are used
by AWS. The metadata ﬁle in /opt/ml contains metadata about resources such as DomainId.
Use the following command to show the ﬁle system contents:

cat /opt/ml/metadata/resource-metadata.json
{"AppType":"JupyterLab","DomainId":"example-domain-id","UserProfileName":"exampleuser-profile-name,"ResourceArn":"arn:aws:sagemaker:AWS
Region:111122223333;:app/domain-ID/user-ID/Jupyte
rLab/default","ResourceName":"default","AppImageVersion":"current"}

• Logging directories – /var/logs/studio are reserved for the logging directories of JupyterLab
and the extensions associated with it. We recommend that you don't use the folders in creating
your image.
Health check and URL for applications
• Base URL – The base URL for the BYOI application must be jupyterlab/default. You can
only have one application and it must always be named default.
• HealthCheck API – The HostAgent uses the HealthCheckAPI at port 8888 to check the
health of the JupyterLab application. jupyterlab/default/api/status is the endpoint for
the health check.
• Home/Default URL – The /opt/.sagemakerinternal and /opt/ml directories that
are used by AWS. The metadata ﬁle in /opt/ml contains metadata about resources such as
DomainId.
• Authentication – To enable authentication for your users, turn oﬀ the Jupyter notebooks token
or password based authentication and allow all origins.

JupyterLab administrator guide

618


The following is a sample Amazon Linux 2 Dockerfile that meets the preceding specifications:

```
FROM public.ecr.aws/amazonlinux/amazonlinux:2

ARG NB_USER="sagemaker-user"
ARG NB_UID="1000"
ARG NB_GID="100"
RUN yum install --assumeyes python3 shadow-utils && \
    useradd --create-home --shell /bin/bash --gid "${NB_GID}" --uid ${NB_UID} \
    ${NB_USER} && \
    yum clean all && \
    python3 -m pip install jupyterlab

RUN python3 -m pip install --upgrade pip
RUN python3 -m pip install --upgrade urllib3==1.26.6

USER ${NB_UID}
CMD jupyter lab --ip 0.0.0.0 --port 8888 \
    --ServerApp.base_url="/jupyterlab/default" \
    --ServerApp.token='' \
    --ServerApp.allow_origin='*' 
```

The following is a sample Amazon SageMaker Distribution Dockerfile that meets the preceding specifications:

```
FROM public.ecr.aws/sagemaker/sagemaker-distribution:latest-cpu
ARG NB_USER="sagemaker-user"
ARG NB_UID=1000
ARG NB_GID=100

ENV MAMBA_USER=$NB_USER

USER root

RUN apt-get update
RUN micromamba install sagemaker-inference --freeze-installed --yes --channel conda-forge --name base
```
Update the SageMaker Distribution Image

⚠️ Important
This topic assumes that you've created a space and given the user access to it. For more information, see Give your users access to private spaces.

Update the JupyterLab spaces that you've already created to use the latest version of the SageMaker Distribution Image. You can use either the Studio UI or the AWS Command Line Interface (AWS CLI) to update the image.

The following sections provide information about updating an image.

Update the image (UI)

Updating the image involves restarting the JupyterLab space of your user. Use the following procedure to update your user's JupyterLab space with the latest image.

To update the image (UI)

1. Open Studio. For information about opening Studio, see Launch Amazon SageMaker Studio.
2. Choose JupyterLab.
3. Select the JupyterLab space of your user.
4. Choose Stop space.
5. For Image, select an updated version of the SageMaker Distribution Image. For the latest image, choose Latest.
6. Choose Run space.
Update the image (AWS CLI)

This section assumes that you have the AWS Command Line Interface (AWS CLI) installed. For information about installing the AWS CLI, see Install or update to the latest version of the AWS CLI.

To update the image, you must do the following for your user's space:

1. Delete the JupyterLab application
2. Update the space
3. Create the application

⚠️ **Important**

You must have the following information ready before you start updating the image:

- **domain ID** – The ID of your user's Amazon SageMaker domain.
- **Application type** – JupyterLab.
- **Application name** – default.
- **Space name** – The name specified for the space.
- **Instance type** – The Amazon EC2 instance type that you're using to run the application. For example, `ml.t3.medium`.
- **SageMaker Image ARN** – The Amazon Resource Name (ARN) of the SageMaker Distribution Image. You can provide the latest version of the SageMaker Distribution Image by specifying either `sagemaker-distribution-cpu` or `sagemaker-distribution-gpu` as the resource identifier.

To delete the JupyterLab application, run the following command:

```bash
aws sagemaker delete-app \
--domain-id your-user's-domain-id \
--app-type JupyterLab \
--app-name default \
--space-name name-of-your-user's-space
```

To update your user's space, run the following command:
aws sagemaker update-space \
--space-name name-of-your-user's-space \
--domain-id your-user's-domain-id

If you've updated the space successfully, you'll see the space ARN in the response:

```
{
}
```

To create the application, run the following command:

```
aws sagemaker create-app \
--domain-id your-user's-domain-id \
--app-type JupyterLab \
--app-name default \
--space-name name-of-your-user's-space \
--resource-spec "InstanceType=instance-type,SageMakerImageArn=arn:aws:sagemaker:AWS Region:555555555555:image/sagemaker-distribution-resource-identifier"
```

Delete unused resources

To avoid incurring additional costs running JupyterLab, we recommend deleting unused resources in the following order:

1. JupyterLab applications
2. Spaces
3. User profiles
4. domains
Use the following AWS Command Line Interface (AWS CLI) commands to delete resources within a domain:

Delete a JupyterLab application

```
aws --region AWS Region sagemaker delete-app --domain-id example-domain-id --app-name default --app-type JupyterLab --space-name example-space-name
```

Delete a space

⚠️ Important

If you delete a space, you delete the Amazon EBS volume associated with it. We recommend backing up any valuable data before you delete your space.

```
aws --region AWS Region sagemaker delete-space --domain-id example-domain-id --space-name example-space-name
```

Delete a user profile

```
aws --region AWS Region sagemaker delete-user-profile --domain-id example-domain-id --user-profile example-user-profile
```

Quotas

JupyterLab, has quotas for the following:

- The sum of all Amazon EBS volumes within an AWS account.
- The instance types that are available for your users.
- The number of instances for a particular that your users can launch.
To get more storage and compute for your users, request an increase to your AWS quotas. For more information about requesting a quota increase, see Amazon SageMaker endpoints and quotas.

Migrating from SageMaker Studio Classic to SageMaker Studio

⚠️ Important

Before you can migrate users from Amazon SageMaker Studio Classic to Amazon SageMaker Studio, you must give them permissions in Studio. For more information, see Set Studio as the default experience for your domain.

Migrating your users from Amazon SageMaker Studio Classic to Amazon SageMaker Studio lets your users leverage the features and improved performance of Studio. This guide provides an overview of the migration process. It provides guidance on transferring user data, migrating lifecycle configurations, and ensuring the compatibility of JupyterLab extensions.

Migrating your users from Studio Classic to Studio involves the following:

- Migrating user data
- Bringing user lifecycle configurations
- Migrating user JupyterLab extensions

Migrate user data from Amazon SageMaker Studio Classic to Amazon SageMaker Studio

Studio Classic stores files in Amazon Elastic File System (Amazon EFS) volumes. Studio stores files in Amazon Elastic Block Store (Amazon EBS).

We recommend that you transfer user home directories from Amazon EFS to Amazon EBS. Transferring the home directories helps your users maintain their working environments after migration.

You can migrate home directories in various ways. The following is an example of a method that you can use:

1. Identify the EFS volume that you use for Studio Classic user storage.
2. For each user, copy their home directory from Amazon EFS to the corresponding EBS volume allocated for JupyterLab.

3. Update the new EBS volume permissions so users can access their migrated home directories in JupyterLab.

4. Communicate steps to users so they know what to expect when launching JupyterLab for the first time after migration.

You can use AWS DataSync to copy the contents of a Studio Classic EFS file system to an Amazon S3 bucket.

After the DataSync task finishes, the contents of the Studio Classic user home directories are copied to the target Amazon S3 bucket. The S3 object keys have a prefix matching the HomeEfsFileSystemUid value for each user profile. You can use the DescribeUserProfile operation to get the value.

You can implement a shell script to copy the home directory from Amazon S3 to the `/home/sagemaker-user` directory on the Amazon EBS volume. The script should do all of the following:

- Get the current user's profile and HomeEfsFileSystemUid value.
- Use the HomeEfsFileSystemUid to identify the S3 prefix for the contents of the user's home directory.
- Copy the objects under the S3 prefix to `/home/sagemaker-user`.

⚠️ **Important**

Before you run the script, check for the following:

- The Amazon EBS volume is large enough to store the objects that you're exporting.
- You aren't migrating hidden files and folders, such as `.bashrc` and `.condarc` if you aren't intending to do so.
- The AWS Identity and Access Management (IAM) execution role that's associated with Studio Classic user profiles has the policies configured to access only the respective home directory in Amazon S3.

You can run the script manually in the JupyterLab application or set it as a lifecycle configuration.
When you implement the script, do the following:

- Only copy the specific user's directory, not all Amazon S3 data.
- Resolve any errors or incomplete transfers.
- Preserve permissions, ownership, and metadata of files and directories.
- To avoid duplicates, delete previous copies of files in /home/sagemaker-user.
- Verify that the script can run multiple times without issue.

After you sync the home directories for all users, delete the exported home directories from Amazon S3. This will help reduce storage costs and detach the lifecycle configuration.

**Bring lifecycle configurations from Studio to JupyterLab**

Amazon SageMaker Studio Classic notebooks are based on kernel gateway architecture. The Jupyter Server kernel and the other kernels run on separate compute resources. Within Studio Classic, you frequently had to create separate lifecycle configurations for both kernel gateway and Jupyter Server applications. For more information about Studio Classic lifecycle configurations, see [SageMaker JupyterLab](https://docs.aws.amazon.com/sagemaker/latest/dg/jupyterlab.html).

In Amazon SageMaker Studio, both Jupyter Server and the kernels run on the same Amazon EC2 instance. The runtime environments of Studio JupyterLab applications are based on images from the SageMaker Distribution. The runtimes in Studio Classic run on different images, unless an image from SageMaker Distribution has been specified.

To migrate user environments successfully, we recommend changing your LCC scripts and testing them.

**Migrate JupyterLab extensions**

Studio Classic uses JupyterLab 3. Studio uses JupyterLab 4. Custom JupyterLab extensions in Studio Classic might not work in Studio, so we recommend checking extensions compatibility. You might have to upgrade them.

For more information about extensions, see [Extension Compatibility with JupyterLab 4.0](https://docs.aws.amazon.com/sagemaker/latest/dg/jupyterlab.html).

**Amazon SageMaker Notebook Instances**

An Amazon SageMaker notebook instance is a machine learning (ML) compute instance running the Jupyter Notebook App. SageMaker manages creating the instance and related resources. Use
Jupyter notebooks in your notebook instance to prepare and process data, write code to train models, deploy models to SageMaker hosting, and test or validate your models.

SageMaker also provides sample notebooks that contain complete code walkthroughs. These walkthroughs show how to use SageMaker to perform common machine learning tasks. For more information, see [Example Notebooks](#).

For information about pricing with Amazon SageMaker notebook instance, see [Amazon SageMaker Pricing](#).

**Maintenance**

SageMaker updates the underlying software for Amazon SageMaker Notebook Instances at least once every 90 days. Some maintenance updates, such as operating system upgrades, may require your application to be taken offline for a short period of time. It is not possible to perform any operations during this period while the underlying software is being updated. We recommend that you restart your notebooks at least once every 30 days to automatically consume patches.

For more information, contact [https://aws.amazon.com/premiumsupport/](https://aws.amazon.com/premiumsupport/).

**Topics**

- Use Notebook Instances to build models
- Amazon Linux 2 notebook instances
- JupyterLab versioning
- Create a Notebook Instance
- Access Notebook Instances
- Update a Notebook Instance
- Customize a Notebook Instance Using a Lifecycle Configuration Script
- Example Notebooks
- Set the Notebook Kernel
- Associate Git Repositories with SageMaker Notebook Instances
- Notebook Instance Metadata
- Monitor Jupyter Logs in Amazon CloudWatch Logs
Use Notebook Instances to build models

One of the best ways for machine learning (ML) practitioners to use Amazon SageMaker is to train and deploy ML models using SageMaker notebook instances. The SageMaker notebook instances help create the environment by initiating Jupyter servers on Amazon Elastic Compute Cloud (Amazon EC2) and providing preconfigured kernels with the following packages: the Amazon SageMaker Python SDK, AWS SDK for Python (Boto3), AWS Command Line Interface (AWS CLI), Conda, Pandas, deep learning framework libraries, and other libraries for data science and machine learning.

Machine Learning with the SageMaker Python SDK

To train, validate, deploy, and evaluate an ML model in a SageMaker notebook instance, use the SageMaker Python SDK. The SageMaker Python SDK abstracts AWS SDK for Python (Boto3) and SageMaker API operations. It enables you to integrate with and orchestrate other AWS services, such as Amazon Simple Storage Service (Amazon S3) for saving data and model artifacts, Amazon Elastic Container Registry (ECR) for importing and servicing the ML models, Amazon Elastic Compute Cloud (Amazon EC2) for training and inference.

You can also take advantage of SageMaker features that help you deal with every stage of a complete ML cycle: data labeling, data preprocessing, model training, model deployment, evaluation on prediction performance, and monitoring the quality of model in production.

If you're a first-time SageMaker user, we recommend you to use the SageMaker Python SDK, following the end-to-end ML tutorial. To find the open source documentation, see the Amazon SageMaker Python SDK.

Tutorial Overview

This Get Started tutorial walks you through how to create a SageMaker notebook instance, open a Jupyter notebook with a preconfigured kernel with the Conda environment for machine learning, and start a SageMaker session to run an end-to-end ML cycle. You'll learn how to save a dataset to a default Amazon S3 bucket automatically paired with the SageMaker session, submit a training job of an ML model to Amazon EC2, and deploy the trained model for prediction by hosting or batch inferencing through Amazon EC2.

This tutorial explicitly shows a complete ML flow of training the XGBoost model from the SageMaker built-in model pool. You use the US Adult Census dataset, and you evaluate the performance of the trained SageMaker XGBoost model on predicting individuals' income.
• **SageMaker XGBoost** – The XGBoost model is adapted to the SageMaker environment and preconfigured as Docker containers. SageMaker provides a suite of built-in algorithms that are prepared for using SageMaker features. To learn more about what ML algorithms are adapted to SageMaker, see Choose an Algorithm and Use Amazon SageMaker Built-in Algorithms. For the SageMaker built-in algorithm API operations, see First-Party Algorithms in the Amazon SageMaker Python SDK.

• **Adult Census dataset** – The dataset from the 1994 Census bureau database by Ronny Kohavi and Barry Becker (Data Mining and Visualization, Silicon Graphics). The SageMaker XGBoost model is trained using this dataset to predict if an individual makes over $50,000 a year or less.

**Topics**

• **Step 1: Create an Amazon SageMaker Notebook Instance**

• **Step 2: Create a Jupyter Notebook**

• **Step 3: Download, Explore, and Transform a Dataset**

• **Step 4: Train a Model**

• **Step 5: Deploy the Model to Amazon EC2**

• **Step 6: Evaluate the Model**

• **Step 7: Clean Up**

**Step 1: Create an Amazon SageMaker Notebook Instance**

An Amazon SageMaker notebook instance is a fully managed machine learning (ML) Amazon Elastic Compute Cloud (Amazon EC2) compute instance that runs the Jupyter Notebook App. You use the notebook instance to create and manage Jupyter notebooks for preprocessing data and to train and deploy machine learning models.

**To create a SageMaker notebook instance**

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Notebook instances, and then choose Create notebook instance.
3. On the Create notebook instance page, provide the following information (if a field is not mentioned, leave the default values):
   a. For Notebook instance name, type a name for your notebook instance.
b. For **Notebook Instance type**, choose `ml.t2.medium`. This is the least expensive instance type that notebook instances support, and it suffices for this exercise. If a `ml.t2.medium` instance type isn't available in your current AWS Region, choose `ml.t3.medium`.

c. For **Platform Identifier**, choose a platform type to create the notebook instance on. This platform type dictates the Operating System and the JupyterLab version that your notebook instance is created with. For information about platform identifier type, see [Amazon Linux 2 notebook instances](#). For information about JupyterLab versions, see [JupyterLab versioning](#).

d. For **IAM role**, choose **Create a new role**, and then choose **Create role**. This IAM role automatically gets permissions to access any S3 bucket that has `sagemaker` in the name. It gets these permissions through the `AmazonSageMakerFullAccess` policy, which SageMaker attaches to the role.

   > **Note**
   
   If you want to grant the IAM role permission to access S3 buckets without `sagemaker` in the name, you need to attach the `S3FullAccess` policy or limit the permissions to specific S3 buckets to the IAM role. For more information and examples of adding bucket policies to the IAM role, see [Bucket Policy Examples](#).

e. Choose **Create notebook instance**.

   In a few minutes, SageMaker launches an ML compute instance—in this case, a notebook instance—and attaches a 5 GB of Amazon EBS storage volume to it. The notebook instance has a preconfigured Jupyter notebook server, SageMaker and AWS SDK libraries, and a set of Anaconda libraries.

   For more information about creating a SageMaker notebook instance, see [Create a Notebook Instance](#).

**Optional** Change SageMaker Notebook Instance Settings

If you want to change the ML compute instance type or the size of the Amazon EBS storage of a SageMaker notebook instance that's already created, you can edit the notebook instance settings.

**To change and update the SageMaker Notebook instance type and the EBS volume**

1. On the **Notebook instances** page in the SageMaker console, choose your notebook instance.
2. Choose Actions, choose Stop, and then wait until the notebook instance fully stops.

3. After the notebook instance status changes to Stopped, choose Actions, and then choose Update settings.
   
   a. For Notebook instance type, choose a different ML instance type.
   
   b. For Volume size in GB, type a different integer to specify a new EBS volume size.

   **Note**
   
   EBS storage volumes are encrypted, so SageMaker can't determine the amount of available free space on the volume. Because of this, you can increase the volume size when you update a notebook instance, but you can't decrease the volume size. If you want to decrease the size of the ML storage volume in use, create a new notebook instance with the desired size.

4. At the bottom of the page, choose Update notebook instance.

5. When the update is complete, Start the notebook instance with the new settings.

For more information about updating SageMaker notebook instance settings, see Update a Notebook Instance.

(Optional) Advanced Settings for SageMaker Notebook Instances

The following tutorial video shows how to set up and use SageMaker notebook instances through the SageMaker console with advanced options, such as SageMaker lifecycle configuration and importing GitHub repositories. (Length: 26:04)

For complete documentation about SageMaker notebook instance, see Use Amazon SageMaker notebook Instances.

**Step 2: Create a Jupyter Notebook**

To start scripting for training and deploying your model, create a Jupyter notebook in the SageMaker notebook instance. Using the Jupyter notebook, you can conduct machine learning (ML) experiments for training and inference while accessing the SageMaker features and the AWS infrastructure.
To create a Jupyter notebook

1. Open the notebook instance as follows:
   b. On the Notebook instances page, open your notebook instance by choosing either Open JupyterLab for the JupyterLab interface or Open Jupyter for the classic Jupyter view.

   ! Note
   If the notebook instance status shows Pending in the Status column, your notebook instance is still being created. The status will change to InService when the notebook instance is ready for use.

2. Create a notebook as follows:

   - If you opened the notebook in the JupyterLab view, on the File menu, choose New, and then choose Notebook. For Select Kernel, choose conda_python3. This preinstalled environment includes the default Anaconda installation and Python 3.

   - If you opened the notebook in the classic Jupyter view, on the Files tab, choose New, and then choose conda_python3. This preinstalled environment includes the default Anaconda installation and Python 3.

3. Save the notebooks as follows:

   - In the JupyterLab view, choose File, choose Save Notebook As..., and then rename the notebook.

   - In the Jupyter classic view, choose File, choose Save as..., and then rename the notebook.

Step 3: Download, Explore, and Transform a Dataset

In this step, you load the Adult Census dataset to your notebook instance using the SHAP (SHapley Additive exPlanations) Library, review the dataset, transform it, and upload it to Amazon S3. SHAP is a game theoretic approach to explain the output of any machine learning model. For more information about SHAP, see Welcome to the SHAP documentation.

To run the following example, paste the sample code into a cell in your notebook instance.
Load Adult Census Dataset Using SHAP

Using the SHAP library, import the Adult Census dataset as shown following:

```python
import shap
X, y = shap.datasets.adult()
X_display, y_display = shap.datasets.adult(display=True)
feature_names = list(X.columns)
feature_names
```

**Note**
If the current Jupyter kernel does not have the SHAP library, install it by running the following `conda` command:

```bash
%conda install -c conda-forge shap
```

If you're using JupyterLab, you must manually refresh the kernel after the installation and updates have completed. Run the following IPython script to shut down the kernel (the kernel will restart automatically):

```python
import IPython
IPython.Application.instance().kernel.do_shutdown(True)
```

The `feature_names` list object should return the following list of features:

```python
['Age',
 'Workclass',
 'Education-Num',
 'Marital Status',
 'Occupation',
 'Relationship',
 'Race',
 'Sex',
 'Capital Gain',
 'Capital Loss',
 'Hours per week',
 'Country']
```
Tip

If you're starting with unlabeled data, you can use Amazon SageMaker Ground Truth to create a data labeling workflow in minutes. To learn more, see Label Data.

Overview the Dataset

Run the following script to display the statistical overview of the dataset and histograms of the numeric features.

```python
display(X.describe())
hist = X.hist(bins=30, sharey=True, figsize=(20, 10))
```
Tip
If you want to use a dataset that needs to be cleaned and transformed, you can simplify and streamline data preprocessing and feature engineering using Amazon SageMaker Data Wrangler. To learn more, see [Prepare ML Data with Amazon SageMaker Data Wrangler](#).

Split the Dataset into Train, Validation, and Test Datasets

Using Sklearn, split the dataset into a training set and a test set. The training set is used to train the model, while the test set is used to evaluate the performance of the final trained model. The dataset is randomly sorted with the fixed random seed: 80 percent of the dataset for training set and 20 percent of it for a test set.

```python
from sklearn.model_selection import train_test_split
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=1)
X_train_display = X_display.loc[X_train.index]
```

Split the training set to separate out a validation set. The validation set is used to evaluate the performance of the trained model while tuning the model's hyperparameters. 75 percent of the training set becomes the final training set, and the rest is the validation set.

```python
X_train, X_val, y_train, y_val = train_test_split(X_train, y_train, test_size=0.25, random_state=1)
X_train_display = X_display.loc[X_train.index]
X_val_display = X_display.loc[X_val.index]
```

Using the pandas package, explicitly align each dataset by concatenating the numeric features with the true labels.

```python
import pandas as pd
train = pd.concat([pd.Series(y_train, index=X_train.index, name='Income>50K', dtype=int), X_train], axis=1)
validation = pd.concat([pd.Series(y_val, index=X_val.index, name='Income>50K', dtype=int), X_val], axis=1)
test = pd.concat([pd.Series(y_test, index=X_test.index, name='Income>50K', dtype=int), X_test], axis=1)
```

Check if the dataset is split and structured as expected:
### train

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<th>Occupation</th>
<th>Relationship</th>
<th>Race</th>
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19536 rows × 13 columns

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6512 rows × 13 columns

### test
Convert the Train and Validation Datasets to CSV Files

Convert the `train` and `validation` dataframe objects to CSV files to match the input file format for the XGBoost algorithm.

```python
# Use 'csv' format to store the data
# The first column is expected to be the output column
train.to_csv('train.csv', index=False, header=False)
validation.to_csv('validation.csv', index=False, header=False)
```

Upload the Datasets to Amazon S3

Using the SageMaker and Boto3, upload the training and validation datasets to the default Amazon S3 bucket. The datasets in the S3 bucket will be used by a compute-optimized SageMaker instance on Amazon EC2 for training.

The following code sets up the default S3 bucket URI for your current SageMaker session, creates a new `demo-sagemaker-xgboost-adult-income-prediction` folder, and uploads the training and validation datasets to the data subfolder.

```python
import sagemaker, boto3, os
bucket = sagemaker.Session().default_bucket()
prefix = "demo-sagemaker-xgboost-adult-income-prediction"

boto3.Session().resource('s3').Bucket(bucket).Object(
    os.path.join(prefix, 'data/train.csv')).upload_file('train.csv')

```

---

**Table: Income vs Age vs Workclass vs Education-Num vs Marital Status vs Occupation vs Relationship vs Race vs Sex vs Capital Gain vs Capital Loss vs Hours per week vs Country**

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<th>Workclass</th>
<th>Education-Num</th>
<th>Marital Status</th>
<th>Occupation</th>
<th>Relationship</th>
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<td>12</td>
<td>3</td>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>20.0</td>
</tr>
<tr>
<td>29514</td>
<td>0</td>
<td>35.0</td>
<td>4</td>
<td>9</td>
<td>4</td>
<td>14</td>
<td>3</td>
<td>4</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>40.0</td>
</tr>
<tr>
<td>1600</td>
<td>0</td>
<td>30.0</td>
<td>4</td>
<td>7</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>45.0</td>
</tr>
<tr>
<td>639</td>
<td>1</td>
<td>52.0</td>
<td>6</td>
<td>16.0</td>
<td>2</td>
<td>10</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>60.0</td>
</tr>
</tbody>
</table>

6513 rows x 13 columns
Run the following AWS CLI to check if the CSV files are successfully uploaded to the S3 bucket.

```
! aws s3 ls {bucket}/{prefix}/data --recursive
```

This should return the following output:

```
2021-01-14 17:52:09    786285 demo-sagemaker-xgboost-adult-income-prediction/data/train.csv
2021-01-14 17:52:10    262122 demo-sagemaker-xgboost-adult-income-prediction/data/validation.csv
```

**Step 4: Train a Model**

The [Amazon SageMaker Python SDK](https://docs.aws.amazon.com/sagemaker/latest/dg/python-sdk-training.html) provides framework estimators and generic estimators to train your model while orchestrating the machine learning (ML) lifecycle accessing the SageMaker features for training and the AWS infrastructures, such as Amazon Elastic Container Registry (Amazon ECR), Amazon Elastic Compute Cloud (Amazon EC2), Amazon Simple Storage Service (Amazon S3). For more information about SageMaker built-in framework estimators, see [Frameworks](https://docs.aws.amazon.com/sagemaker/latest/dg/python-sdk-training.html) in the [Amazon SageMaker Python SDK](https://docs.aws.amazon.com/sagemaker/latest/dg/python-sdk-training.html) documentation. For more information about built-in algorithms, see [Use Amazon SageMaker Built-in Algorithms or Pre-trained Models](https://docs.aws.amazon.com/sagemaker/latest/dg/algorithms.html).

**Topics**

- [Choose the Training Algorithm](#)
- [Create and Run a Training Job](#)

**Choose the Training Algorithm**

To choose the right algorithm for your dataset, you typically need to evaluate different models to find the most suitable models to your data. For simplicity, the SageMaker [XGBoost Algorithm](https://docs.aws.amazon.com/sagemaker/latest/dg/xgboost.html) built-in algorithm is used throughout this tutorial without the pre-evaluation of models.

**Tip**

If you want SageMaker to find an appropriate model for your tabular dataset, use Amazon SageMaker Autopilot that automates a machine learning solution. For more information, see [SageMaker Autopilot](https://docs.aws.amazon.com/sagemaker/latest/dg/autopilot.html).
Create and Run a Training Job

After you figured out which model to use, start constructing a SageMaker estimator for training. This tutorial uses the XGBoost built-in algorithm for the SageMaker generic estimator.

To run a model training job

1. Import the Amazon SageMaker Python SDK and start by retrieving the basic information from your current SageMaker session.

   ```python
   import sagemaker
   region = sagemaker.Session().boto_region_name
   print("AWS Region: {}".format(region))
   role = sagemaker.get_execution_role()
   print("RoleArn: {}".format(role))
   ```

   This returns the following information:

   • `region` – The current AWS Region where the SageMaker notebook instance is running.
   • `role` – The IAM role used by the notebook instance.

   **Note**

   Check the SageMaker Python SDK version by running `sagemaker.__version__`. This tutorial is based on `sagemaker>=2.20`. If the SDK is outdated, install the latest version by running the following command:

   ```shell
   ! pip install -qU sagemaker
   ```

   If you run this installation in your exiting SageMaker Studio or notebook instances, you need to manually refresh the kernel to finish applying the version update.

2. Create an XGBoost estimator using the `sagemaker.estimator.Estimator` class. In the following example code, the XGBoost estimator is named `xgb_model`.

   ```python
   from sagemaker.debugger import Rule, ProfilerRule, rule_configs
   from sagemaker.session import TrainingInput
   ```
To construct the SageMaker estimator, specify the following parameters:

- **image_uri** – Specify the training container image URI. In this example, the SageMaker XGBoost training container URI is specified using `sagemaker.image_uris.retrieve`.

- **role** – The AWS Identity and Access Management (IAM) role that SageMaker uses to perform tasks on your behalf (for example, reading training results, call model artifacts from Amazon S3, and writing training results to Amazon S3).

- **instance_count** and **instance_type** – The type and number of Amazon EC2 ML compute instances to use for model training. For this training exercise, you use a single `ml.m4.xlarge` instance, which has 4 CPUs, 16 GB of memory, an Amazon Elastic Block Store (Amazon EBS) storage, and a high network performance. For more information about EC2 compute instance types, see Amazon EC2 Instance Types. For more information about billing, see Amazon SageMaker pricing.

- **volume_size** – The size, in GB, of the EBS storage volume to attach to the training instance. This must be large enough to store training data if you use File mode (File mode is on by default). If you don’t specify this parameter, its value defaults to 30.

- **output_path** – The path to the S3 bucket where SageMaker stores the model artifact and training results.
• **sagemaker_session** – The session object that manages interactions with SageMaker API operations and other AWS service that the training job uses.

• **rules** – Specify a list of SageMaker Debugger built-in rules. In this example, the `create_xgboost_report()` rule creates an XGBoost report that provides insights into the training progress and results, and the `ProfilerReport()` rule creates a report regarding the EC2 compute resource utilization. For more information, see [SageMaker Debugger XGBoost Training Report](#).

**Tip**

If you want to run distributed training of large sized deep learning models, such as convolutional neural networks (CNN) and natural language processing (NLP) models, use SageMaker Distributed for data parallelism or model parallelism. For more information, see [Distributed training in Amazon SageMaker](#).

3. Set the hyperparameters for the XGBoost algorithm by calling the `set_hyperparameters` method of the estimator. For a complete list of XGBoost hyperparameters, see [XGBoost Hyperparameters](#).

```python
xgb_model.set_hyperparameters(
    max_depth = 5,
    eta = 0.2,
    gamma = 4,
    min_child_weight = 6,
    subsample = 0.7,
    objective = "binary:logistic",
    num_round = 1000
)
```

**Tip**

You can also tune the hyperparameters using the SageMaker hyperparameter optimization feature. For more information, see [Perform Automatic Model Tuning with SageMaker](#).

4. Use the `TrainingInput` class to configure a data input flow for training. The following example code shows how to configure `TrainingInput` objects to use the training and
validation datasets you uploaded to Amazon S3 in the Split the Dataset into Train, Validation, and Test Datasets section.

```python
from sagemaker.session import TrainingInput

train_input = TrainingInput(
    "s3://{}/{}/{}".format(bucket, prefix, "data/train.csv"), content_type="csv"
)
validation_input = TrainingInput(
    "s3://{}/{}/{}".format(bucket, prefix, "data/validation.csv"),
    content_type="csv"
)
```

5. To start model training, call the estimator's fit method with the training and validation datasets. By setting wait=True, the fit method displays progress logs and waits until training is complete.

```python
xgb_model.fit({"train": train_input, "validation": validation_input}, wait=True)
```

For more information about model training, see Train a Model with Amazon SageMaker. This tutorial training job might take up to 10 minutes.

After the training job has done, you can download an XGBoost training report and a profiling report generated by SageMaker Debugger. The XGBoost training report offers you insights into the training progress and results, such as the loss function with respect to iteration, feature importance, confusion matrix, accuracy curves, and other statistical results of training. For example, you can find the following loss curve from the XGBoost training report which clearly indicates that there is an overfitting problem.
Run the following code to specify the S3 bucket URI where the Debugger training reports are generated and check if the reports exist.

```python
rule_output_path = xgb_model.output_path + "/" +
xgb_model.latest_training_job.job_name + "/rule-output"
! aws s3 ls {rule_output_path} --recursive
```
Download the Debugger XGBoost training and profiling reports to the current workspace:

```bash
! aws s3 cp {rule_output_path} ./ --recursive
```

Run the following IPython script to get the file link of the XGBoost training report:

```python
from IPython.display import FileLink, FileLinks
display("Click link below to view the XGBoost Training report",
    FileLink("CreateXgboostReport/xgboost_report.html"))
```

The following IPython script returns the file link of the Debugger profiling report that shows summaries and details of the EC2 instance resource utilization, system bottleneck detection results, and python operation profiling results:

```python
profiler_report_name = [rule["RuleConfigurationName"]
                       for rule in
                       xgb_model.latest_training_job.rule_job_summary()
                       if "Profiler" in rule["RuleConfigurationName"]][0]

display("Click link below to view the profiler report",
    FileLink(profiler_report_name+"/profiler-output/profiler-report.html"))
```

⚠️ **Tip**

If the HTML reports do not render plots in the JupyterLab view, you must choose **Trust HTML** at the top of the reports.

To identify training issues, such as overfitting, vanishing gradients, and other problems that prevents your model from converging, use SageMaker Debugger and take automated actions while prototyping and training your ML models. For more information, see [Use Amazon SageMaker Debugger to debug and improve model performance](#). To find a complete analysis of model parameters, see the [Explainability with Amazon SageMaker Debugger](#) example notebook.

You now have a trained XGBoost model. SageMaker stores the model artifact in your S3 bucket. To find the location of the model artifact, run the following code to print the model_data attribute of the `xgb_model` estimator:

---

*Use Notebook Instances to build models*
Tip

To measure biases that can occur during each stage of the ML lifecycle (data collection, model training and tuning, and monitoring of ML models deployed for prediction), use SageMaker Clarify. For more information, see Model Explainability. For an end-to-end example, see the Fairness and Explainability with SageMaker Clarify example notebook.

Step 5: Deploy the Model to Amazon EC2

To get predictions, deploy your model to Amazon EC2 using Amazon SageMaker.

Topics

- [Deploy the Model to SageMaker Hosting Services](#)
- (Optional) Use SageMaker Predictor to Reuse the Hosted Endpoint
- (Optional) Make Prediction with Batch Transform

Deploy the Model to SageMaker Hosting Services

To host a model through Amazon EC2 using Amazon SageMaker, deploy the model that you trained in [Create and Run a Training Job](#) by calling the deploy method of the xgb_model estimator. When you call the deploy method, you must specify the number and type of EC2 ML instances that you want to use for hosting an endpoint.

```python
import sagemaker
from sagemaker.serializers import CSVSerializer
xgb_predictor=xgb_model.deploy(
    initial_instance_count=1,
    instance_type='ml.t2.medium',
    serializer=CSVSerializer()
)
```

- initial_instance_count (int) – The number of instances to deploy the model.
- instance_type (str) – The type of instances that you want to operate your deployed model.
• **serializer (int)** – Serialize input data of various formats (a NumPy array, list, file, or buffer) to a CSV-formatted string. We use this because the XGBoost algorithm accepts input files in CSV format.

The deploy method creates a deployable model, configures the SageMaker hosting services endpoint, and launches the endpoint to host the model. For more information, see the [SageMaker generic Estimator's deploy class method](https://docs.aws.amazon.com/sagemaker/latest/dg/python-sdk-composing-models-deploy.html) in the Amazon SageMaker Python SDK. To retrieve the name of endpoint that's generated by the deploy method, run the following code:

```python
xgb_predictor.endpoint_name
```

This should return the endpoint name of the `xgb_predictor`. The format of the endpoint name is "sagemaker-xgboost-YYYY-MM-DD-HH-MM-SS-SSS". This endpoint stays active in the ML instance, and you can make instantaneous predictions at any time unless you shut it down later. Copy this endpoint name and save it to reuse and make real-time predictions elsewhere in SageMaker Studio or SageMaker notebook instances.

**Tip**

To learn more about compiling and optimizing your model for deployment to Amazon EC2 instances or edge devices, see [Compile and Deploy Models with Neo](https://docs.aws.amazon.com/sagemaker/latest/dg/compiling-models-neo.html).

**(Optional) Use SageMaker Predictor to Reuse the Hosted Endpoint**

After you deploy the model to an endpoint, you can set up a new SageMaker predictor by pairing the endpoint and continuously make real-time predictions in any other notebooks. The following example code shows how to use the SageMaker Predictor class to set up a new predictor object using the same endpoint. Re-use the endpoint name that you used for the `xgb_predictor`.

```python
import sagemaker
xgb_predictor_reuse=sagemaker.predictor.Predictor(
    endpoint_name="sagemaker-xgboost-YYYY-MM-DD-HH-MM-SS-SSS",
    sagemaker_session=sagemaker.Session(),
    serializer=sagemaker.serializers.CSVSerializer()
)
```
The `xgb_predictor_reuse` Predictor behaves exactly the same as the original `xgb_predictor`. For more information, see the SageMaker Predictor class in the Amazon SageMaker Python SDK.

(Optional) Make Prediction with Batch Transform

Instead of hosting an endpoint in production, you can run a one-time batch inference job to make predictions on a test dataset using the SageMaker batch transform. After your model training has completed, you can extend the estimator to a transformer object, which is based on the SageMaker Transformer class. The batch transformer reads in input data from a specified S3 bucket and makes predictions.

To run a batch transform job

1. Run the following code to convert the feature columns of the test dataset to a CSV file and uploads to the S3 bucket:

   ```
   X_test.to_csv('test.csv', index=False, header=False)
   boto3.Session().resource('s3').Bucket(bucket).Object(os.path.join(prefix, 'test/test.csv')).upload_file('test.csv')
   ```

2. Specify S3 bucket URIs of input and output for the batch transform job as shown following:

   ```
   # The location of the test dataset
   batch_input = 's3://{0}/{1}/test'.format(bucket, prefix)

   # The location to store the results of the batch transform job
   batch_output = 's3://{0}/{1}/batch-prediction'.format(bucket, prefix)
   ```

3. Create a transformer object specifying the minimal number of parameters: the `instance_count` and `instance_type` parameters to run the batch transform job, and the `output_path` to save prediction data as shown following:

   ```
   transformer = xgb_model.transformer(
       instance_count=1,
       instance_type='ml.m4.xlarge',
       output_path=batch_output
   )
   ```

4. Initiate the batch transform job by executing the `transform()` method of the `transformer` object as shown following:
5. When the batch transform job is complete, SageMaker creates the test.csv.out prediction data saved in the batch_output path, which should be in the following format: s3://sagemaker-<region>-111122223333/demo-sagemaker-xgboost-adult-income-prediction/batch-prediction. Run the following AWS CLI to download the output data of the batch transform job:

```
! aws s3 cp {batch_output} ./ --recursive
```

This should create the test.csv.out file under the current working directory. You'll be able to see the float values that are predicted based on the logistic regression of the XGBoost training job.

### Step 6: Evaluate the Model

Now that you have trained and deployed a model using Amazon SageMaker, evaluate the model to ensure that it generates accurate predictions on new data. For model evaluation, use the test dataset that you created in **Step 3: Download, Explore, and Transform a Dataset**.

**Evaluate the Model Deployed to SageMaker Hosting Services**

To evaluate the model and use it in production, invoke the endpoint with the test dataset and check whether the inferences you get returns a target accuracy you want to achieve.

**To evaluate the model**

1. Set up the following function to predict each line of the test set. In the following example code, the rows argument is to specify the number of lines to predict at a time. You can change the value of it to perform a batch inference that fully utilizes the instance's hardware resource.

```python
import numpy as np
def predict(data, rows=1000):
    transformer.transform(
        data=batch_input,
        data_type='S3Prefix',
        content_type='text/csv',
        split_type='Line'
    )
    transformer.wait()
```
2. Run the following code to make predictions of the test dataset and plot a histogram. You need to take only the feature columns of the test dataset, excluding the 0th column for the actual values.

```python
import matplotlib.pyplot as plt

predictions = predict(test.to_numpy()[:, 1:]
plt.hist(predictions)
plt.show()
```

3. The predicted values are float type. To determine True or False based on the float values, you need to set a cutoff value. As shown in the following example code, use the Scikit-learn library to return the output confusion metrics and classification report with a cutoff of 0.5.

```python
import sklearn

cutoff = 0.5
```
print(sklearn.metrics.confusion_matrix(test.iloc[:, 0], np.where(predictions > cutoff, 1, 0)))
print(sklearn.metrics.classification_report(test.iloc[:, 0], np.where(predictions > cutoff, 1, 0)))

This should return the following confusion matrix:

```
[[4670  356]
 [ 480 1007]]
```

<table>
<thead>
<tr>
<th></th>
<th>precision</th>
<th>recall</th>
<th>f1-score</th>
<th>support</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.91</td>
<td>0.93</td>
<td>0.92</td>
<td>5026</td>
</tr>
<tr>
<td>1</td>
<td>0.74</td>
<td>0.68</td>
<td>0.71</td>
<td>1487</td>
</tr>
</tbody>
</table>

4. To find the best cutoff with the given test set, compute the log loss function of the logistic regression. The log loss function is defined as the negative log-likelihood of a logistic model that returns prediction probabilities for its ground truth labels. The following example code numerically and iteratively calculates the log loss values \((-y \log(p) + (1-y)\log(1-p))\), where \(y\) is the true label and \(p\) is a probability estimate of the corresponding test sample. It returns a log loss versus cutoff graph.

```python
import matplotlib.pyplot as plt
cutoffs = np.arange(0.01, 1, 0.01)
log_loss = []
for c in cutoffs:
    log_loss.append(
        sklearn.metrics.log_loss(test.iloc[:, 0], np.where(predictions > c, 1, 0))
    )
plt.figure(figsize=(15,10))
plt.plot(cutoffs, log_loss)
plt.xlabel("Cutoff")
plt.ylabel("Log loss")
plt.show()
```
This should return the following log loss curve.

5. Find the minimum points of the error curve using the NumPy `argmin` and `min` functions:

    ```python
    print('Log loss is minimized at a cutoff of ', cutoffs[np.argmin(log_loss)],
          ', and the log loss value at the minimum is ', np.min(log_loss))
    ```

This should return:

    Log loss is minimized at a cutoff of 0.53, and the log loss value at the minimum is 4.348539186773897.

Instead of computing and minimizing the log loss function, you can estimate a cost function as an alternative. For example, if you want to train a model to perform a binary classification for a business problem such as a customer churn prediction problem, you can set weights to the elements of confusion matrix and calculate the cost function accordingly.

You have now trained, deployed, and evaluated your first model in SageMaker.
Tip
To monitor model quality, data quality, and bias drift, use Amazon SageMaker Model Monitor and SageMaker Clarify. To learn more, see Amazon SageMaker Model Monitor, Monitor Data Quality, Monitor Model Quality, Monitor Bias Drift, and Monitor Feature Attribution Drift.

Tip
To get human review of low confidence ML predictions or a random sample of predictions, use Amazon Augmented AI human review workflows. For more information, see Using Amazon Augmented AI for Human Review.

Step 7: Clean Up
To avoid incurring unnecessary charges, use the AWS Management Console to delete the endpoints and resources that you created while running the exercises.

Note
Training jobs and logs cannot be deleted and are retained indefinitely.

Note
If you plan to explore other exercises in this guide, you might want to keep some of these resources, such as your notebook instance, S3 bucket, and IAM role.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/ and delete the following resources:
   - The endpoint. Deleting the endpoint also deletes the ML compute instance or instances that support it.
     1. Under Inference, choose Endpoints.
2. Choose the endpoint that you created in the example, choose Actions, and then choose Delete.

- The endpoint configuration.
  1. Under Inference, choose Endpoint configurations.
  2. Choose the endpoint configuration that you created in the example, choose Actions, and then choose Delete.

- The model.
  1. Under Inference, choose Models.
  2. Choose the model that you created in the example, choose Actions, and then choose Delete.

- The notebook instance. Before deleting the notebook instance, stop it.
  1. Under Notebook, choose Notebook instances.
  2. Choose the notebook instance that you created in the example, choose Actions, and then choose Stop. The notebook instance takes several minutes to stop. When the Status changes to Stopped, move on to the next step.
  3. Choose Actions, and then choose Delete.

2. Open the Amazon S3 console at https://console.aws.amazon.com/s3/, and then delete the bucket that you created for storing model artifacts and the training dataset.

3. Open the Amazon CloudWatch console at https://console.aws.amazon.com/cloudwatch/, and then delete all of the log groups that have names starting with /aws/sagemaker/.

Amazon Linux 2 notebook instances

Amazon SageMaker notebook instances currently support Amazon Linux 2 (AL2) operating systems. You can select the operating system that your notebook instance is based on when you create the notebook instance.

SageMaker supports notebook instances based on the following Amazon Linux 2 operating systems.

- **notebook-al2-v1**: These notebook instances support JupyterLab version 1. For information about JupyterLab versions, see [JupyterLab versioning](#).
• **notebook-al2-v2**: These notebook instances support JupyterLab version 3. For information about JupyterLab versions, see [JupyterLab versioning](#).

Notebook instances created before 08/18/2021 automatically run on Amazon Linux (AL1). Notebook instances based on AL1 entered a maintenance phase as of 12/01/2022 and are no longer available for new notebook instance creation as of 02/01/2023. To replace AL1, you now have the option to create Amazon SageMaker notebook instances with AL2. For more information, see [AL1 Maintenance Phase Plan](#).

**Topics**

- [Supported instance types](#)
- [Available Kernels](#)
- [AL1 Maintenance Phase Plan](#)

**Supported instance types**

Amazon Linux 2 supports instance types listed under **Notebook Instances** in [Amazon SageMaker Pricing](#) with the exception that Amazon Linux 2 does not support ml.p2 instances.

**Available Kernels**

The following table gives information about the available kernels for SageMaker notebook instances. All of these images are supported on notebook instances based on both the notebook-al2-v1 and notebook-al2-v2 operating systems.

### SageMaker notebook instance kernels

<table>
<thead>
<tr>
<th>Kernel name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>R</td>
<td>A kernel used to perform data analysis and visualization using R code from a Jupyter notebook.</td>
</tr>
<tr>
<td>Sparkmagic (PySpark)</td>
<td>A kernel used to do data science with remote Spark clusters from Jupyter notebooks using the Python programming language. This kernel comes with Python 3.10.</td>
</tr>
<tr>
<td>Kernel name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Sparkmagic (Spark)</td>
<td>A kernel used to do data science with remote Spark clusters from Jupyter notebooks using the Scala programming language. This kernel comes with Python 3.10.</td>
</tr>
<tr>
<td>Sparkmagic (SparkR)</td>
<td>A kernel used to do data science with remote Spark clusters from Jupyter notebooks using the R programming language. This kernel comes with Python 3.10.</td>
</tr>
<tr>
<td>conda_python3</td>
<td>A conda environment that comes pre-installed with popular packages for data science and machine learning. This kernel comes with Python 3.10.</td>
</tr>
<tr>
<td>conda_pytorch_p310</td>
<td>A conda environment that comes pre-installed with PyTorch version 2.0.1, as well as popular data science and machine learning packages. This kernel comes with Python 3.10.</td>
</tr>
<tr>
<td>conda_tensorflow2_p310</td>
<td>A conda environment that comes pre-installed with TensorFlow version 2.13, as well as popular data science and machine learning packages. This kernel comes with Python 3.10.</td>
</tr>
</tbody>
</table>

**AL1 Maintenance Phase Plan**

The following table is a timeline for when AL1 entered its extended maintenance phase. The AL1 maintenance phase also coincides with the deprecation of Python 2 and Chainer. Notebooks based on AL2 do not have managed Python 2 and Chainer kernels.

<table>
<thead>
<tr>
<th>Date</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>08/18/2021</td>
<td>Notebook instances based on AL2 are launched. Newly launched notebook instances</td>
</tr>
<tr>
<td>Date</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>10/31/2022</td>
<td>The default platform identifier for SageMaker notebook instances changes from Amazon Linux (al1-v1) to Amazon Linux 2 (al2-v2). You can choose between the two operating systems when launching a new notebook instance.</td>
</tr>
<tr>
<td>12/01/2022</td>
<td>AL1 is no longer supported with non-critical security patches and updates. AL1 still receives fixes for critical security-related issues. You can still launch instances on AL1, but assume the risks associated with using an unsupported operating system.</td>
</tr>
<tr>
<td>02/01/2023</td>
<td>AL1 is no longer an available option for new notebook instance creation. After this date, customers can create notebook instances with the AL2 platform identifiers. Existing al1-v1 notebook instances are not affected.</td>
</tr>
</tbody>
</table>
AL1 reaches its end of life on notebook instances on March 31, 2024. After this date, AL1 will no longer receive any security updates, bug fixes, or be available for new notebook instance creation.

- Existing AL1 notebook instances with a STOPPED status cannot be restarted.
- AL1 notebook instances with the INSERVICE status are not affected until they are stopped.

**Migrating to Amazon Linux 2**

Your existing AL1 notebook instance is not automatically migrated to Amazon Linux 2. To upgrade your AL1 notebook instance to Amazon Linux 2, you must create a new notebook instance, replicate your code and environment, and delete your old notebook instance. For more information, see the [Amazon Linux 2 migration blog](https://aws.amazon.com/blogs/integer/al2-migration-guide/).

**JupyterLab versioning**

The Amazon SageMaker notebook instance interface is based on JupyterLab, which is a web-based interactive development environment for notebooks, code, and data. Notebooks now support using either JupyterLab 1 or JupyterLab 3. A single notebook instance can run a single instance of JupyterLab (at most). You can have multiple notebook instances with different JupyterLab versions.

You can configure your notebook to run your preferred JupyterLab version by selecting the appropriate platform identifier. Use either the AWS CLI or the SageMaker console when creating your notebook instance. For more information about platform identifiers, see [Amazon Linux 2 vs Amazon Linux notebook instances](https://docs.aws.amazon.com/sagemaker/latest/dg/jupyterlab-platform-identifiers.html). If you don’t explicitly configure a platform identifier, your notebook instance defaults to running JupyterLab 1.

**Topics**

- [JupyterLab 3](https://docs.aws.amazon.com/sagemaker/latest/dg/jupyterlab-platform-identifiers.html)
- [Creating a notebook with your JupyterLab version](https://docs.aws.amazon.com/sagemaker/latest/dg/jupyterlab-platform-identifiers.html)
• View the JupyterLab version of a notebook from the console

JupyterLab 3

JupyterLab 3 support is available only on the Amazon Linux 2 operating system platform. JupyterLab 3 includes the following features that are not available in JupyterLab 1. For more information about these features, see JupyterLab 3.0 is released!

• Visual debugger when using the following kernels:
  • conda_pytorch_p38
  • conda_tensorflow2_p38
  • conda_amazonei_pytorch_latest_p37
• File browser filter
• Table of Contents (TOC)
• Multi-language support
• Simple mode
• Single interface mode
• Live editing SVG files with updated rendering
• User interface for notebook cell tags

Important changes to JupyterLab 3

For information about important changes when using JupyterLab 3, see the following JupyterLab change logs:

• v2.0.0
• v3.0.0

Package version changes

JupyterLab 3 has the following package version changes from JupyterLab 1:

• JupyterLab has been upgraded from 1.x to 3.x.
• Jupyter notebook has been upgraded from 5.x to 6.x.
• jupyterlab-git has been updated to version 0.37.1.
• nbserverproxy 0.x (0.3.2) has been replaced with jupyter-server-proxy 3.x (3.2.1).

Creating a notebook with your JupyterLab version

You can select the JupyterLab version when creating your notebook instance from the console following the steps in Create a Notebook Instance.

You can also select the JupyterLab version by passing the platform-identifier parameter when creating your notebook instance using the AWS CLI as follows:

```
create-notebook-instance --notebook-instance-name <NEW_NOTEBOOK_NAME> \  
--instance-type <INSTANCE_TYPE> \  
--role-arn <YOUR_ROLE_ARN> \  
--platform-identifier <PLATFORM_TO_USE>
```

View the JupyterLab version of a notebook from the console

You can view the JupyterLab version of a notebook using the following procedure:

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation, select Notebook.
3. From the dropdown menu, select Notebook instances to navigate to the Notebook instances page.
4. From the list of notebook instances, select your notebook instance name.
5. On the Notebook instance settings page, view the Platform Identifier to see the JupyterLab version of the notebook.

Create a Notebook Instance

An Amazon SageMaker notebook instance is a ML compute instance running the Jupyter Notebook App. SageMaker manages creating the instance and related resources. Use Jupyter notebooks in your notebook instance to prepare and process data, write code to train models, deploy models to SageMaker hosting, and test or validate your models.

To create a notebook instance, use either the SageMaker console or the CreateNotebookInstance API.
The notebook instance type you choose depends on how you use your notebook instance. You want to ensure that your notebook instance is not bound by memory, CPU, or IO. If you plan to load a dataset into memory on the notebook instance for exploration or preprocessing, we recommend that you choose an instance type with enough RAM memory for your dataset. This would require an instance with at least 16 GB of memory (xlarge or larger). If you plan to use the notebook for compute intensive preprocessing, we recommend you choose a compute-optimized instance such as a c4 or c5.

A best practice when using a SageMaker notebook is to use the notebook instance to orchestrate other AWS services. For example, you can use the notebook instance to manage large dataset processing by making calls to AWS Glue for ETL (extract, transform, and load) services or Amazon EMR for mapping and data reduction using Hadoop. You can use AWS services as temporary forms of computation or storage for your data.

You can store and retrieve your training and test data using an Amazon S3 bucket. You can then use SageMaker to train and build your model, so the instance type of your notebook would have no bearing on the speed of your model training and testing.

After receiving the request, SageMaker does the following:

- **Creates a network interface**—If you choose the optional VPC configuration, SageMaker creates the network interface in your VPC. It uses the subnet ID that you provide in the request to determine which Availability Zone to create the subnet in. SageMaker associates the security group that you provide in the request with the subnet. For more information, see [Connect a Notebook Instance in a VPC to External Resources](#).

- **Launches an ML compute instance**—SageMaker launches an ML compute instance in a SageMaker VPC. SageMaker performs the configuration tasks that allow it to manage your notebook instance, and if you specified your VPC, it enables traffic between your VPC and the notebook instance.

- **Installs Anaconda packages and libraries for common deep learning platforms**—SageMaker installs all of the Anaconda packages that are included in the installer. For more information, see [Anaconda package list](#). In addition, SageMaker installs the TensorFlow and Apache MXNet deep learning libraries.

- **Attaches an ML storage volume**—SageMaker attaches an ML storage volume to the ML compute instance. You can use the volume as a working area to clean up the training dataset or to temporarily store validation, test, or other data. Choose any size between 5 GB and 16384 GB, in 1 GB increments, for the volume. The default is 5 GB. ML storage volumes are encrypted, so SageMaker can't determine the amount of available free space on the volume. Because of this,
you can increase the volume size when you update a notebook instance, but you can't decrease the volume size. If you want to decrease the size of the ML storage volume in use, create a new notebook instance with the desired size.

Only files and data saved within the `/home/ec2-user/SageMaker` folder persist between notebook instance sessions. Files and data that are saved outside this directory are overwritten when the notebook instance stops and restarts. Each notebook instance's `/tmp` directory provides a minimum of 10 GB of storage in an instance store. An instance store is temporary, block-level storage that isn't persistent. When the instance is stopped or restarted, SageMaker deletes the directory's contents. This temporary storage is part of the root volume of the notebook instance.

If the instance type used by the notebook instance has NVMe support, then customers can use the NVMe instance store volumes available for that instance type. For instances with NVMe store volumes, all instance store volumes are automatically attached to the instance at launch. For more information about instance types and their associated NVMe store volumes, see the Amazon Elastic Compute Cloud Instance Type Details.

To make the attached NVMe store volume available for your notebook instance, complete the steps in Make instance store volumes available on your instance with root access or using a lifecycle configuration script.

**Note**

NVMe instance store volumes are not persistent storage. This storage is short-lived with the instance and must be reconfigured every time an instance with this storage is launched.

- **Copies example Jupyter notebooks**— These Python code examples illustrate model training and hosting exercises using various algorithms and training datasets.

**To create a SageMaker notebook instance:**

2. Choose **Notebook instances**, then choose **Create notebook instance**.
3. On the **Create notebook instance** page, provide the following information:
   a. For **Notebook instance name**, type a name for your notebook instance.
b. For **Notebook instance type**, choose an instance size suitable for your use case. For a list of supported instance types and quotas, see [Amazon SageMaker Service Quotas](https://docs.aws.amazon.com/sagemaker/latest/dg/service-quotas.html).

c. For **Elastic Inference**, choose an inference accelerator type to associate with the notebook instance if you plan to conduct inferences from the notebook instance, or choose `none`. For information about elastic inference, see [Use Amazon SageMaker Elastic Inference (EI)](https://docs.aws.amazon.com/sagemaker/latest/dg/elastic-inference.html).

d. For **Platform Identifier**, choose a platform type to create the notebook instance on. This platform type dictates the Operating System and the JupyterLab version that your notebook instance is created with. For information about platform identifier type, see [Amazon Linux 2 notebook instances](https://docs.aws.amazon.com/sagemaker/latest/dg/what-is.html) and [JupyterLab versioning](https://docs.aws.amazon.com/sagemaker/latest/dg/jupyter-lab.html).

e. (Optional) **Additional configuration** lets advanced users create a shell script that can run when you create or start the instance. This script, called a lifecycle configuration script, can be used to set the environment for the notebook or to perform other functions. For information, see [Customize a Notebook Instance Using a Lifecycle Configuration Script](https://docs.aws.amazon.com/sagemaker/latest/dg/lifecycle-hooks.html).

f. (Optional) **Additional configuration** also lets you specify the size, in GB, of the ML storage volume that is attached to the notebook instance. You can choose a size between 5 GB and 16,384 GB, in 1 GB increments. You can use the volume to clean up the training dataset or to temporarily store validation or other data.

g. (Optional) For **Minimum IMDS Version**, select a version from the dropdown list. If this value is set to v1, both versions can be used with the notebook instance. If v2 is selected, then only IMDSv2 can be used with the notebook instance. For information about IMDSv2, see [Use IMDSv2](https://docs.aws.amazon.com/ NHS娃2/Windows/latest/HnsGuide/using-imds-windows.html).

h. For **IAM role**, choose either an existing IAM role in your account that has the necessary permissions to access SageMaker resources or choose Create a new role. If you choose Create a new role, SageMaker creates an IAM role named `AmazonSageMaker-ExecutionRole-YYYYMMDDTHmmSS`. The AWS managed policy

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**Note**

Starting October 31, 2022, the default minimum IMDS Version for SageMaker notebook instances changes from IMDSv1 to IMDSv2. Starting February 1, 2023, IMDSv1 is no longer be available for new notebook instance creation. After this date, you can create notebook instances with a minimum IMDS version of 2.
AmazonSageMakerFullAccess is attached to the role. The role provides permissions that allow the notebook instance to call SageMaker and Amazon S3.

i. For **Root access**, to enable root access for all notebook instance users, choose **Enable**. To disable root access for users, choose **Disable**. If you enable root access, all notebook instance users have administrator privileges and can access and edit all files on it.

j. (Optional) **Encryption key** lets you encrypt data on the ML storage volume attached to the notebook instance using an AWS Key Management Service (AWS KMS) key. If you plan to store sensitive information on the ML storage volume, consider encrypting the information.

k. (Optional) **Network** lets you put your notebook instance inside a Virtual Private Cloud (VPC). A VPC provides additional security and restricts access to resources in the VPC from sources outside the VPC. For more information on VPCs, see [Amazon VPC User Guide](https://docs.aws.amazon.com/vpc/latest/userguide/). To add your notebook instance to a VPC:

   i. Choose the **VPC** and a **SubnetId**.
   
   ii. For **Security Group**, choose your VPC's default security group.
   
   iii. If you need your notebook instance to have internet access, enable direct internet access. For **Direct internet access**, choose **Enable**. Internet access can make your notebook instance less secure. For more information, see [Connect a Notebook Instance in a VPC to External Resources](https://docs.aws.amazon.com/sagemaker/latest/dg/).  

l. (Optional) To associate Git repositories with the notebook instance, choose a default repository and up to three additional repositories. For more information, see [Associate Git Repositories with SageMaker Notebook Instances](https://docs.aws.amazon.com/sagemaker/latest/dg).  

m. Choose **Create notebook instance**.

   In a few minutes, Amazon SageMaker launches an ML compute instance—in this case, a notebook instance—and attaches an ML storage volume to it. The notebook instance has a preconfigured Jupyter notebook server and a set of Anaconda libraries. For more information, see the [CreateNotebookInstance API](https://docs.aws.amazon.com/sagemaker/latest/dg).

4. When the status of the notebook instance is **InService**, in the console, the notebook instance is ready to use. Choose **Open Jupyter** next to the notebook name to open the classic Jupyter dashboard.
Note

To augment the security of your Amazon SageMaker notebook instance, all regional `notebook.region.sagemaker.aws` domains are registered in the internet Public Suffix List (PSL). For further security, we recommend that you use cookies with a `__Host-` prefix if you ever need to set sensitive cookies for the domains of your SageMaker notebook instances. This helps to defend your domain against cross-site request forgery attempts (CSRF). For more information, see the `Set-Cookie` page in the [mozilla.org](http://mozilla.org) developer documentation website.

You can choose [Open JupyterLab](#) to open the JupyterLab dashboard. The dashboard provides access to your notebook instance and sample SageMaker notebooks that contain complete code walkthroughs. These walkthroughs show how to use SageMaker to perform common machine learning tasks. For more information, see [Example Notebooks](#). For more information, see [Control root access to a SageMaker notebook instance](#).

For more information about Jupyter notebooks, see [The Jupyter notebook](#).

## Access Notebook Instances

To access your Amazon SageMaker notebook instances, choose one of the following options:

- Use the console.

Choose **Notebook instances**. The console displays a list of notebook instances in your account. To open a notebook instance with a standard Jupyter interface, choose **Open Jupyter** for that instance. To open a notebook instance with a JupyterLab interface, choose **Open JupyterLab** for that instance.

The console uses your sign-in credentials to send a
CreatePresignedNotebookInstanceUrl API request to SageMaker. SageMaker returns the URL for your notebook instance, and the console opens the URL in another browser tab and displays the Jupyter notebook dashboard.

**Note**
The URL that you get from a call to CreatePresignedNotebookInstanceUrl is valid only for 5 minutes. If you try to use the URL after the 5-minute limit expires, you are directed to the AWS Management Console sign-in page.

- Use the API.

  To get the URL for the notebook instance, call the CreatePresignedNotebookInstanceUrl API and use the URL that the API returns to open the notebook instance.

Use the Jupyter notebook dashboard to create and manage notebooks and to write code. For more information about Jupyter notebooks, see http://jupyter.org/documentation.html.

### Update a Notebook Instance

After you create a notebook instance, you can update it using the SageMaker console and UpdateNotebookInstance API operation.

You can update the tags of a notebook instance that is InService. To update any other attribute of a notebook instance, its status must be Stopped.

**To update a notebook instance in the SageMaker console:**

2. Choose Notebook instances.
3. Choose the notebook instance that you want to update by selecting the notebook instance Name from the list.
4. If your notebook Status is not Stopped, select the Stop button to stop the notebook instance.

   When you do this, the notebook instance status changes to Stopping. Wait until the status changes to Stopped to complete the following steps.
5. Select the **Edit** button to open the **Edit notebook instance** page. For information about the notebook properties you can update, see [Create a Notebook Instance](#).

6. Update your notebook instance and select the **Update notebook instance** button at the bottom of the page when you are done to return to the notebook instances page. Your notebook instance status changes to **Updating**.

   When the notebook instance update is complete, the status changes to **Stopped**.

### Customize a Notebook Instance Using a Lifecycle Configuration Script

To install packages or sample notebooks on your notebook instance, configure networking and security for it, or otherwise use a shell script to customize it, use a lifecycle configuration. A *lifecycle configuration* provides shell scripts that run only when you create the notebook instance or whenever you start one. When you create a notebook instance, you can create a new lifecycle configuration and the scripts it uses or apply one that you already have.

You can also use a lifecycle configuration script to access AWS services from your notebook. For example, you can create a script that lets you use your notebook to control other AWS resources, such as an Amazon EMR instance.

We maintain a public repository of notebook lifecycle configuration scripts that address common use cases for customizing notebook instances at [https://github.com/aws-samples/amazon-sagemaker-notebook-instance-lifecycle-configuration-samples](https://github.com/aws-samples/amazon-sagemaker-notebook-instance-lifecycle-configuration-samples).

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**Note**

Each script has a limit of 16384 characters.

The value of the `PATH` environment variable that is available to both scripts is `/usr/local/sbin:/usr/local/bin:/usr/bin:/usr/sbin:/sbin:/bin`. The working directory, which is the value of the `PWD` environment variable, is `/`.


Scripts cannot run for longer than 5 minutes. If a script runs for longer than 5 minutes, it fails and the notebook instance is not created or started. To help decrease the run time of scripts, try the following:
You can see a list of notebook instance lifecycle configurations you previously created by choosing **Lifecyle configuration** in the SageMaker console. You can attach a notebook instance lifecycle configuration when you create a new notebook instance. For more information about creating a notebook instance, see [Create a Notebook Instance](#).

### To create a lifecycle configuration

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **Lifecycle configurations**.
4. From the **Lifecycle configurations** page, choose the **Notebook Instance** tab.
5. Choose **Create configuration**.
6. For **Name**, type a name using alphanumeric characters and "-", but no spaces. The name can have a maximum of 63 characters.
7. (Optional) To create a script that runs when you create the notebook and every time you start it, choose **Start notebook**.
8. In the **Start notebook** editor, type the script.
9. (Optional) To create a script that runs only once, when you create the notebook, choose **Create notebook**.
10. In the **Create notebook** editor, type the script configure networking.
11. Choose **Create configuration**.

### Lifecycle Configuration Best Practices

The following are best practices for using lifecycle configurations:

- Cut down on necessary steps. For example, limit which conda environments in which to install large packages.
- Run tasks in parallel processes.
- Use the nohup command in your script.
Important

We do not recommend storing sensitive information in your lifecycle configuration script.

- Lifecycle configurations run as the root user. If your script makes any changes within the /home/ec2-user/SageMaker directory, (for example, installing a package with pip), use the command `sudo -u ec2-user` to run as the ec2-user user. This is the same user that Amazon SageMaker runs as.

- SageMaker notebook instances use conda environments to implement different kernels for Jupyter notebooks. If you want to install packages that are available to one or more notebook kernels, enclose the commands to install the packages with conda environment commands that activate the conda environment that contains the kernel where you want to install the packages.

For example, if you want to install a package only for the python3 environment, use the following code:

```bash
#!/bin/bash
sudo -u ec2-user -i <<EOF
# This will affect only the Jupyter kernel called "conda_python3".
source activate python3

# Replace myPackage with the name of the package you want to install.
pip install myPackage
# You can also perform "conda install" here as well.

source deactivate
EOF
```

If you want to install a package in all conda environments in the notebook instance, use the following code:

```bash
#!/bin/bash
sudo -u ec2-user -i <<EOF
# Note that "base" is special environment name, include it there as well.
for env in base /home/ec2-user/anaconda3/envs/*; do
    source /home/ec2-user/anaconda3/bin/activate $(basename "$env")
EOF
```
# Installing packages in the Jupyter system environment can affect stability of your SageMaker Notebook Instance. You can remove this check if you'd like to install Jupyter extensions, etc.

```bash
if [ $env = 'JupyterSystemEnv' ]; then
  continue
fi

# Replace myPackage with the name of the package you want to install.
myPackage

# You can also perform "conda install" here as well.

source /home/ec2-user/anaconda3/bin/deactivate
done
EOF
```

- You must store all conda environments in the default environments folder (/home/user/anaconda3/envs).

⚠️ **Important**

When you create or change a script, we recommend that you use a text editor that provides Unix-style line breaks, such as the text editor available in the console when you create a notebook. Copying text from a non-Linux operating system might introduce incompatible line breaks and result in an unexpected error.

## Install External Libraries and Kernels in Notebook Instances

Amazon SageMaker notebook instances come with multiple environments already installed. These environments contain Jupyter kernels and Python packages including: scikit, Pandas, NumPy, TensorFlow, and MXNet. These environments, along with all files in the sample-notebooks folder, are refreshed when you stop and start a notebook instance. You can also install your own environments that contain your choice of packages and kernels.

The different Jupyter kernels in Amazon SageMaker notebook instances are separate conda environments. For information about conda environments, see [Managing environments](#) in the Conda documentation.
Install custom environments and kernels on the notebook instance's Amazon EBS volume. This ensures that they persist when you stop and restart the notebook instance, and that any external libraries you install are not updated by SageMaker. To do that, use a lifecycle configuration that includes both a script that runs when you create the notebook instance (on-create) and a script that runs each time you restart the notebook instance (on-start). For more information about using notebook instance lifecycle configurations, see Customize a Notebook Instance Using a Lifecycle Configuration Script. There is a GitHub repository that contains sample lifecycle configuration scripts at SageMaker Notebook Instance Lifecycle Config Samples.

The examples at https://github.com/aws-samples/amazon-sagemaker-notebook-instance-lifecycle-config-samples/blob/master/scripts/persistent-conda-ebs/on-create.sh and https://github.com/aws-samples/amazon-sagemaker-notebook-instance-lifecycle-config-samples/blob/master/scripts/persistent-conda-ebs/on-start.sh show the best practice for installing environments and kernels on a notebook instance. The on-create script installs the ipykernel library to create custom environments as Jupyter kernels, then uses pip install and conda install to install libraries. You can adapt the script to create custom environments and install libraries that you want. SageMaker does not update these libraries when you stop and restart the notebook instance, so you can ensure that your custom environment has specific versions of libraries that you want. The on-start script installs any custom environments that you create as Jupyter kernels, so that they appear in the dropdown list in the Jupyter New menu.

**Package installation tools**

SageMaker notebooks support the following package installation tools:

- conda install
- pip install

You can install packages using the following methods:

- Lifecycle configuration scripts.
  
  For example scripts, see SageMaker Notebook Instance Lifecycle Config Samples. For more information on lifecycle configuration, see Customize a Notebook Instance Using a Lifecycle Configuration Script.

- Notebooks – The following commands are supported.
  
  - %conda install
  - %pip install
• The Jupyter terminal – You can install packages using pip and conda directly.

From within a notebook you can use the system command syntax (lines starting with !) to install packages, for example, `!pip install` and `!conda install`. More recently, new commands have been added to IPython: `%pip` and `%conda`. These commands are the recommended way to install packages from a notebook as they correctly take into account the active environment or interpreter being used. For more information, see [Add `%pip` and `%conda magic functions](#).

**Conda**

Conda is an open source package management system and environment management system, which can install packages and their dependencies. SageMaker supports using Conda with either of the two main channels, the default channel, and the conda-forge channel. For more information, see [Conda channels](#). The conda-forge channel is a community channel where contributors can upload packages.

ℹ️ **Note**

Due to how Conda resolves the dependency graph, installing packages from conda-forge can take significantly longer (in the worst cases, upwards of 10 minutes).

The Deep Learning AMI comes with many conda environments and many packages preinstalled. Due to the number of packages preinstalled, finding a set of packages that are guaranteed to be compatible is difficult. You may see a warning "The environment is inconsistent, please check the package plan carefully". Despite this warning, SageMaker ensures that all the SageMaker provided environments are correct. SageMaker cannot guarantee that any user installed packages will function correctly.

ℹ️ **Note**

Users of SageMaker, AWS Deep Learning AMI and Amazon EMR can access the commercial Anaconda repository without taking a commercial license through February 1, 2024 when using Anaconda in those services. For any usage of the commercial Anaconda repository after February 1, 2024, customers are responsible for determining their own Anaconda license requirements.
Conda has two methods for activating environments: conda activate/deactivate, and source activate/deactivate. For more information, see Should I use 'conda activate' or 'source activate' in Linux.

SageMaker supports moving Conda environments onto the Amazon EBS volume, which is persisted when the instance is stopped. The environments aren't persisted when the environments are installed to the root volume, which is the default behavior. For an example lifecycle script, see persistent-conda-ebs.

**Supported conda operations (see note at the bottom of this topic)**

- conda install of a package in a single environment
- conda install of a package in all environments
- conda install of a R package in the R environment
- Installing a package from the main conda repository
- Installing a package from conda-forge
- Changing the Conda install location to use EBS
- Supporting both conda activate and source activate

**Pip**

Pip is the de facto tool for installing and managing Python packages. Pip searches for packages on the Python Package Index (PyPI) by default. Unlike Conda, pip doesn't have built in environment support, and is not as thorough as Conda when it comes to packages with native/system library dependencies. Pip can be used to install packages in Conda environments.

You can use alternative package repositories with pip instead of the PyPI. For an example lifecycle script, see on-start.sh.

**Supported pip operations (see note at the bottom of this topic)**

- Using pip to install a package without an active conda environment (install packages system wide)
- Using pip to install a package in a conda environment
- Using pip to install a package in all conda environments
- Changing the pip install location to use EBS
- Using an alternative repository to install packages with pip
Unsupported

SageMaker aims to support as many package installation operations as possible. However, if the packages were installed by SageMaker or DLAMI, and you use the following operations on these packages, it might make your notebook instance unstable:

- Uninstalling
- Downgrading
- Upgrading

We do not provide support for installing packages via yum install or installing R packages from CRAN.

Due to potential issues with network conditions or configurations, or the availability of Conda or PyPi, we cannot guarantee that packages will install in a fixed or deterministic amount of time.

Note

We cannot guarantee that a package installation will be successful. Attempting to install a package in an environment with incompatible dependencies can result in a failure. In such a case you should contact the library maintainer to see if it is possible to update the package dependencies. Alternatively you can attempt to modify the environment in such a way as to allow the installation. This modification however will likely mean removing or updating existing packages, which means we can no longer guarantee stability of this environment.

Notebook Instance Software Updates

Amazon SageMaker periodically tests and releases software that is installed on notebook instances. This includes:

- Kernel updates
- Security patches
- AWS SDK updates
- Amazon SageMaker Python SDK updates
- Open source software updates
To ensure that you have the most recent software updates, stop and restart your notebook instance, either in the SageMaker console or by calling

StopNotebookInstance.

You can also manually update software installed on your notebook instance while it is running by using update commands in a terminal or in a notebook.

Note

Updating kernels and some packages might depend on whether root access is enabled for the notebook instance. For more information, see Control root access to a SageMaker notebook instance.

You can check the Personal Health Dashboard or the security bulletin at Security Bulletins for updates.

Control an Amazon EMR Spark Instance Using a Notebook

You can use a notebook instance created with a custom lifecycle configuration script to access AWS services from your notebook. For example, you can create a script that lets you use your notebook with Sparkmagic to control other AWS resources, such as an Amazon EMR instance. You can then use the Amazon EMR instance to process your data instead of running the data analysis on your notebook. This allows you to create a smaller notebook instance because you won't use the instance to process data. This is helpful when you have large datasets that would require a large notebook instance to process the data.

The process requires three procedures using the Amazon SageMaker console:

- Create the Amazon EMR Spark instance
- Create the Jupyter Notebook
- Test the notebook-to-Amazon EMR connection

To create an Amazon EMR Spark instance that can be controlled from a notebook using Sparkmagic

1. Open the Amazon EMR console at https://console.aws.amazon.com/elasticmapreduce/.
2. In the navigation pane, choose Create cluster.
3. On the **Create Cluster - Quick Options** page, under **Software configuration**, choose **Spark: Spark 2.4.4 on Hadoop 2.8.5 YARN with Ganglia 3.7.2 and Zeppelin 0.8.2**.

4. Set additional parameters on the page and then choose **Create cluster**.

5. On the **Cluster** page, choose the cluster name that you created. Note the **Master Public DNS**, the **EMR master's security group**, and the VPC name and subnet ID where the EMR cluster was created. You will use these values when you create a notebook.

**To create a notebook that uses Sparkmagic to control an Amazon EMR Spark instance**


2. In the navigation pane, under **Notebook instances**, choose **Create notebook**.

3. Enter the notebook instance name and choose the instance type.

4. Choose **Additional configuration**, then, under **Lifecycle configuration**, choose **Create a new lifecycle configuration**.

5. Add the following code to the lifecycle configuration script:

```bash
# OVERVIEW
# This script connects an Amazon EMR cluster to an Amazon SageMaker notebook instance that uses Sparkmagic.
#
# Note that this script will fail if the Amazon EMR cluster's master node IP address is not reachable.
#   1. Ensure that the EMR master node IP is resolvable from the notebook instance. One way to accomplish this is to have the notebook instance and the Amazon EMR cluster in the same subnet.
#   2. Ensure the EMR master node security group provides inbound access from the notebook instance security group.
#      Type        - Protocol - Port - Source
#      Custom TCP  - TCP      - 8998 - $NOTEBOOK_SECURITY_GROUP
#   3. Ensure the notebook instance has internet connectivity to fetch the SparkMagic example config.
#
#
# PARAMETERS
EMR_MASTER_IP=your.emr.master.ip
```
cd /home/ec2-user/.sparkmagic

echo "Fetching Sparkmagic example config from GitHub..."
wget https://raw.githubusercontent.com/jupyter-incubator/sparkmagic/master/sparkmagic/example_config.json

echo "Replacing EMR master node IP in Sparkmagic config..."

sed -i -- "s/localhost/$EMR_MASTER_IP/g" example_config.json
mv example_config.json config.json

echo "Sending a sample request to Livy.."
curl "#$EMR_MASTER_IP:8998/sessions"

6. In the PARAMETERS section of the script, replace your.emr.master.ip with the Master Public DNS name for the Amazon EMR instance.

7. Choose Create configuration.

8. On the Create notebook page, choose Network - optional.

9. Choose the VPC and subnet where the Amazon EMR instance is located.

10. Choose the security group used by the Amazon EMR master node.

11. Choose Create notebook instance.

While the notebook instance is being created, the status is Pending. After the instance has been created and the lifecycle configuration script has successfully run, the status is InService.

Note
If the notebook instance can't connect to the Amazon EMR instance, SageMaker can't create the notebook instance. The connection can fail if the Amazon EMR instance and notebook are not in the same VPC and subnet, if the Amazon EMR master security group is not used by the notebook, or if the Master Public DNS name in the script is incorrect.

To test the connection between the Amazon EMR instance and the notebook

1. When the status of the notebook is InService, choose Open Jupyter to open the notebook.

2. Choose New, then choose Sparkmagic (PySpark).
3. In the code cell, enter `%%info` and then run the cell.

The output should be similar to the following

```
Current session configs: {'driverMemory': '1000M', 'executorCores': 2, 'kind': 'pyspark'}
No active sessions.
```

**Example Notebooks**

Your notebook instance contains example notebooks provided by Amazon SageMaker. The example notebooks contain code that shows how to apply machine learning solutions by using SageMaker. Notebook instances use the `nbexamples` Jupyter extension, which enables you to view a read-only version of an example notebook or create a copy of it that you can modify and run. For more information about the `nbexamples` extension, see [https://github.com/danielballan/nbexamples](https://github.com/danielballan/nbexamples).

For information about example notebooks for SageMaker Studio, see [Use Amazon SageMaker Studio Classic Notebooks](#).

**Note**

Example notebooks typically download datasets from the internet. If you disable SageMaker-provided internet access when you create your notebook instance, example notebooks might not work. For more information, see [Connect a Notebook Instance in a VPC to External Resources](#).

**Use or View Example Notebooks in Jupyter Classic**

To view or use the example notebooks in the classic Jupyter view, choose the **SageMaker Examples** tab.
To view a read-only version of an example notebook in the Jupyter classic view, on the **SageMaker Examples** tab, choose **Preview** for that notebook. To create a copy of an example notebook in the home directory of your notebook instance, choose **Use**. In the dialog box, you can change the notebook's name before saving it.

**Use or View Example Notebooks in Jupyterlab**

To view or use the example notebooks in the Jupyterlab view, choose the examples icon in the left navigation panel.
To view a read-only version of an example notebook, choose the name of the notebook. This opens the notebook as a tab in the main area. To create a copy of an example notebook in the home directory of your notebook instance, choose **Create a Copy** in the top banner. In the dialog box, type a name for the notebook and then choose **CREATE COPY**.

For more information about the example notebooks, see the [SageMaker examples GitHub repository](https://github.com/aws-sagemaker).
Set the Notebook Kernel

Amazon SageMaker provides several kernels for Jupyter that provide support for Python 2 and 3, Apache MXNet, TensorFlow, and PySpark. To set a kernel for a new notebook in the Jupyter notebook dashboard, choose New, and then choose the kernel from the list. For more information about the available kernels, see Available Kernels.

You can also create a custom kernel that you can use in your notebook instance. For information, see Install External Libraries and Kernels in Notebook Instances.

Associate Git Repositories with SageMaker Notebook Instances

Associate Git repositories with your notebook instance to save your notebooks in a source control environment that persists even if you stop or delete your notebook instance. You can associate one default repository and up to three additional repositories with a notebook instance. The repositories can be hosted in AWS CodeCommit, GitHub, or on any other Git server. Associating Git repositories with your notebook instance can be useful for:

- Persistence - Notebooks in a notebook instance are stored on durable Amazon EBS volumes, but they do not persist beyond the life of your notebook instance. Storing notebooks in a Git repository enables you to store and use notebooks even if you stop or delete your notebook instance.
- Collaboration - Peers on a team often work on machine learning projects together. Storing your notebooks in Git repositories allows peers working in different notebook instances to share notebooks and collaborate on them in a source-control environment.
- Learning - Many Jupyter notebooks that demonstrate machine learning techniques are available in publicly hosted Git repositories, such as on GitHub. You can associate your notebook instance with a repository to easily load Jupyter notebooks contained in that repository.

There are two ways to associate a Git repository with a notebook instance:
• Add a Git repository as a resource in your Amazon SageMaker account. Then, to access the repository, you can specify an AWS Secrets Manager secret that contains credentials. That way, you can access repositories that require authentication.

• Associate a public Git repository that is not a resource in your account. If you do this, you cannot specify credentials to access the repository.

Topics
• Add a Git Repository to Your Amazon SageMaker Account
• Create a Notebook Instance with an Associated Git Repository
• Associate a CodeCommit Repository in a Different AWS Account with a Notebook Instance
• Use Git Repositories in a Notebook Instance

Add a Git Repository to Your Amazon SageMaker Account

To manage your GitHub repositories, easily associate them with your notebook instances, and associate credentials for repositories that require authentication, add the repositories as resources in your Amazon SageMaker account. You can view a list of repositories that are stored in your account and details about each repository in the SageMaker console and by using the API.

You can add Git repositories to your SageMaker account in the SageMaker console or by using the AWS CLI.

⚠️ Note
You can use the SageMaker API CreateCodeRepository to add Git repositories to your SageMaker account, but step-by-step instructions are not provided here.

Add a Git Repository to Your SageMaker Account (Console)

To add a Git repository as a resource in your SageMaker account

2. Under Notebook, choose Git repositories, then choose Add repository.
3. To add an CodeCommit repository, choose **AWS CodeCommit**. To add a GitHub or other Git-based repository, choose **GitHub/Other Git-based repo**.

**To add an existing CodeCommit repository**

1. Choose **Use existing repository**.
2. For **Repository**, choose a repository from the list.
3. Enter a name to use for the repository in SageMaker. The name must be 1 to 63 characters. Valid characters are a-z, A-Z, 0-9, and - (hyphen).
4. Choose **Add repository**.

**To create a new CodeCommit repository**

1. Choose **Create new repository**.
2. Enter a name for the repository that you can use in both CodeCommit and SageMaker. The name must be 1 to 63 characters. Valid characters are a-z, A-Z, 0-9, and - (hyphen).
3. Choose **Create repository**.

**To add a Git repository hosted somewhere other than CodeCommit**

1. Choose **GitHub/Other Git-based repo**.
2. Enter a name of up to 63 characters. Valid characters include alpha-numeric characters, a hyphen (-), and 0-9.
3. Enter the URL for the repository. Do not provide a username in the URL. Add the sign-in credentials in AWS Secrets Manager as described in the next step.
4. For **Git credentials**, choose the credentials to use to authenticate to the repository. This is necessary only if the Git repository is private.

### Note

If you have two-factor authentication enabled for your Git repository, enter a personal access token generated by your Git service provider in the password field.
a. To use an existing AWS Secrets Manager secret, choose **Use existing secret**, and then choose a secret from the list. For information about creating and storing a secret, see [Creating a Basic Secret](https://docs.aws.amazon.com/secretsmanager/latest/userguide/create-a-basic-secret.html) in the *AWS Secrets Manager User Guide*. The name of the secret you use must contain the string `sagemaker`.

**Note**

The secret must have a staging label of `AWSCURRENT` and must be in the following format:

```
{"username": UserName, "password": Password}
```

For GitHub repositories, we recommend using a personal access token in the `password` field. For information, see [https://help.github.com/articles/creating-a-personal-access-token-for-the-command-line/](https://help.github.com/articles/creating-a-personal-access-token-for-the-command-line/).

b. To create a new AWS Secrets Manager secret, choose **Create secret**, enter a name for the secret, and then enter the sign-in credentials to use to authenticate to the repository. The name for the secret must contain the string `sagemaker`.

**Note**

The IAM role you use to create the secret must have the `secretsmanager:GetSecretValue` permission in its IAM policy. The secret must have a staging label of `AWSCURRENT` and must be in the following format:

```
{"username": UserName, "password": Password}
```

For GitHub repositories, we recommend using a personal access token.

c. To not use any credentials, choose **No secret**.

5. Choose **Create secret**.

**Add a Git Repository to Your Amazon SageMaker Account (CLI)**

Use the `create-code-repository` AWS CLI command. Specify a name for the repository as the value of the `code-repository-name` argument. The name must be 1 to 63 characters. Valid characters are a-z, A-Z, 0-9, and - (hyphen). Also specify the following:
• The default branch
• The URL of the Git repository

Note
Do not provide a username in the URL. Add the sign-in credentials in AWS Secrets Manager as described in the next step.

• The Amazon Resource Name (ARN) of an AWS Secrets Manager secret that contains the credentials to use to authenticate the repository as the value of the `git-config` argument

For information about creating and storing a secret, see [Creating a Basic Secret](#) in the *AWS Secrets Manager User Guide*. The following command creates a new repository named `MyRepository` in your Amazon SageMaker account that points to a Git repository hosted at `https://github.com/myprofile/my-repo`.

For Linux, OS X, or Unix:

```bash
aws sagemaker create-code-repository
   --code-repository-name "MyRepository"
```

For Windows:

```bash
aws sagemaker create-code-repository ^
   --code-repository-name "MyRepository" ^
```

Note
The secret must have a staging label of `AWSCURRENT` and must be in the following format:

```json
{"username": UserName, "password": Password}
```

For GitHub repositories, we recommend using a personal access token.
Create a Notebook Instance with an Associated Git Repository

You can associate Git repositories with a notebook instance when you create the notebook instance by using the AWS Management Console, or the AWS CLI. If you want to use a CodeCommit repository that is in a different AWS account than the notebook instance, set up cross-account access for the repository. For information, see Associate a CodeCommit Repository in a Different AWS Account with a Notebook Instance.

Topics

- Create a Notebook Instance with an Associated Git Repository (Console)
- Create a Notebook Instance with an Associated Git Repository (CLI)

Create a Notebook Instance with an Associated Git Repository (Console)

To create a notebook instance and associate Git repositories in the Amazon SageMaker console

1. Follow the instructions at Step 1: Create an Amazon SageMaker Notebook Instance.
2. For Git repositories, choose Git repositories to associate with the notebook instance.
   a. For Default repository, choose a repository that you want to use as your default repository. SageMaker clones this repository as a subdirectory in the Jupyter startup directory at /home/ec2-user/SageMaker. When you open your notebook instance, it opens in this repository. To choose a repository that is stored as a resource in your account, choose its name from the list. To add a new repository as a resource in your account, choose Add a repository to SageMaker (opens the Add repository flow in a new window) and then follow the instructions at Create a Notebook Instance with an Associated Git Repository (Console). To clone a public repository that is not stored in your account, choose Clone a public Git repository to this notebook instance only, and then specify the URL for that repository.
   b. For Additional repository 1, choose a repository that you want to add as an additional directory. SageMaker clones this repository as a subdirectory in the Jupyter startup directory at /home/ec2-user/SageMaker. To choose a repository that is stored as a resource in your account, choose its name from the list. To add a new repository as a resource in your account, choose Add a repository to SageMaker (opens the Add repository flow in a new window) and then follow the instructions at Create a Notebook Instance with an Associated Git Repository (Console). To clone a repository that is not
stored in your account, choose **Clone a public Git repository to this notebook instance only**, and then specify the URL for that repository.

Repeat this step up to three times to add up to three additional repositories to your notebook instance.

### Create a Notebook Instance with an Associated Git Repository (CLI)

To create a notebook instance and associate Git repositories by using the AWS CLI, use the `create-notebook-instance` command as follows:

- Specify the repository that you want to use as your default repository as the value of the `default-code-repository` argument. Amazon SageMaker clones this repository as a subdirectory in the Jupyter startup directory at `/home/ec2-user/SageMaker`. When you open your notebook instance, it opens in this repository. To use a repository that is stored as a resource in your SageMaker account, specify the name of the repository as the value of the `default-code-repository` argument. To use a repository that is not stored in your account, specify the URL of the repository as the value of the `default-code-repository` argument.

- Specify up to three additional repositories as the value of the `additional-code-repositories` argument. SageMaker clones this repository as a subdirectory in the Jupyter startup directory at `/home/ec2-user/SageMaker`, and the repository is excluded from the default repository by adding it to the `.git/info/exclude` directory of the default repository. To use repositories that are stored as resources in your SageMaker account, specify the names of the repositories as the value of the `additional-code-repositories` argument. To use repositories that are not stored in your account, specify the URLs of the repositories as the value of the `additional-code-repositories` argument.

For example, the following command creates a notebook instance that has a repository named `MyGitRepo`, that is stored as a resource in your SageMaker account, as a default repository, and an additional repository that is hosted on GitHub:

```bash
aws sagemaker create-notebook-instance \
  --notebook-instance-name "MyNotebookInstance" \
  --instance-type "ml.t2.medium" \
  --role-arn "arn:aws:iam::012345678901:role/service-role/AmazonSageMaker-ExecutionRole-20181129T121390" \
  --default-code-repository "MyGitRepo" \
```
Note
If you use an AWS CodeCommit repository that does not contain "SageMaker" in its name, add the codecommit:GitPull and codecommit:GitPush permissions to the role that you pass as the role-arn argument to the create-notebook-instance command. For information about how to add permissions to a role, see Adding and Removing IAM Policies in the AWS Identity and Access Management User Guide.

Associate a CodeCommit Repository in a Different AWS Account with a Notebook Instance

To associate a CodeCommit repository in a different AWS account with your notebook instance, set up cross-account access for the CodeCommit repository.

To set up cross-account access for a CodeCommit repository and associate it with a notebook instance:

1. In the AWS account that contains the CodeCommit repository, create an IAM policy that allows access to the repository from users in the account that contains your notebook instance. For information, see Step 1: Create a Policy for Repository Access in AccountA in the CodeCommit User Guide.

2. In the AWS account that contains the CodeCommit repository, create an IAM role, and attach the policy that you created in the previous step to that role. For information, see Step 2: Create a Role for Repository Access in AccountA in the CodeCommit User Guide.

3. Create a profile in the notebook instance that uses the role that you created in the previous step:
   a. Open the notebook instance.
   b. Open a terminal in the notebook instance.
   c. Edit a new profile by typing the following in the terminal:

   ```
   vi /home/ec2-user/.aws/config
   ```

   d. Edit the file with the following profile information:
[profile CrossAccountAccessProfile]
region = us-west-2
role_arn = arn:aws:iam::CodeCommitAccount:role/CrossAccountRepositoryContributorRole
credential_source=Ec2InstanceMetadata
output = json

Where CodeCommitAccount is the account that contains the CodeCommit repository, CrossAccountAccessProfile is the name of the new profile, and CrossAccountRepositoryContributorRole is the name of the role you created in the previous step.

4. On the notebook instance, configure git to use the profile you created in the previous step:
   a. Open the notebook instance.
   b. Open a terminal in the notebook instance.
   c. Edit the Git configuration file typing the following in the terminal:

   vi /home/ec2-user/.gitconfig

   d. Edit the file with the following profile information:

   [credential]
   helper = !aws codecommit credential-helper --
   profile CrossAccountAccessProfile $@
   UseHttpPath = true

   Where CrossAccountAccessProfile is the name of the profile that you created in the previous step.

Use Git Repositories in a Notebook Instance

When you open a notebook instance that has Git repositories associated with it, it opens in the default repository, which is installed in your notebook instance directly under /home/ec2-user/SageMaker. You can open and create notebooks, and you can manually run Git commands in a notebook cell. For example:

!git pull origin master
To open any of the additional repositories, navigate up one folder. The additional repositories are also installed as directories under `/home/ec2-user/SageMaker`.

If you open the notebook instance with a JupyterLab interface, the jupyter-git extension is installed and available to use. For information about the jupyter-git extension for JupyterLab, see https://github.com/jupyterlab/jupyterlab-git.

When you open a notebook instance in JupyterLab, you see the git repositories associated with it on the left menu:

You can use the jupyter-git extension to manage git visually, instead of using the command line:
Notebook Instance Metadata

When you create a notebook instance, Amazon SageMaker creates a JSON file on the instance at the location `/opt/ml/metadata/resource-metadata.json` that contains the ResourceName and ResourceArn of the notebook instance. You can access this metadata from anywhere within the notebook instance, including in lifecycle configurations. For information about notebook instance lifecycle configurations, see [Customize a Notebook Instance Using a Lifecycle Configuration Script](#).

### Note

The `resource-metadata.json` file can be modified with root access.

The `resource-metadata.json` file has the following structure:

```
{
    "ResourceArn": "NotebookInstanceArn",
    "ResourceName": "NotebookInstanceName"
}
```
You can use this metadata from within the notebook instance to get other information about the notebook instance. For example, the following commands get the tags associated with the notebook instance:

```
NOTEBOOK_ARN=$(jq '.ResourceArn' /opt/ml/metadata/resource-metadata.json --raw-output)
aws sagemaker list-tags --resource-arn $NOTEBOOK_ARN
```

The output looks like the following:

```
{
  "Tags": [
    {
      "Key": "test",
      "Value": "true"
    }
  ]
}
```

## Monitor Jupyter Logs in Amazon CloudWatch Logs

Jupyter logs include important information such as events, metrics, and health information that provide actionable insights when running Amazon SageMaker notebooks. By importing Jupyter logs into CloudWatch Logs, customers can use CloudWatch Logs to detect anomalous behaviors, set alarms, and discover insights to keep the SageMaker notebooks running more smoothly. You can access the logs even when the Amazon EC2 instance that hosts the notebook is unresponsive, and use the logs to troubleshoot the unresponsive notebook. Sensitive information such as AWS account IDs, secret keys, and authentication tokens in presigned URLs are removed so that customers can share logs without leaking private information.

### To view Jupyter logs for a notebook instance:

2. Choose **Notebook instances**.
3. In the list of notebook instances, choose the notebook instance for which you want to view Jupyter logs by selecting the Notebook instance **Name**.
This will bring you to the details page for that notebook instance.

4. Under **Monitor** on the notebook instance details page, choose **View logs**.

5. In the CloudWatch console, choose the log stream for your notebook instance. Its name is in the form `NotebookInstanceName/jupyter.log`.

For more information about monitoring CloudWatch logs for SageMaker, see [Log Amazon SageMaker Events with Amazon CloudWatch](#).

---

**Amazon SageMaker Studio Lab**

Amazon SageMaker Studio Lab is a free service that gives customers access to AWS compute resources, in an environment based on open-source JupyterLab. It is based on the same architecture and user interface as Amazon SageMaker Studio Classic, but with a subset of Studio Classic capabilities.

With Studio Lab, you can use AWS compute resources to create and run your Jupyter notebooks without signing up for an AWS account. Because Studio Lab is based on open-source JupyterLab, you can take advantage of open-source Jupyter extensions to run your Jupyter notebooks.

**Studio Lab compared to Amazon SageMaker Studio Classic**

While Studio Lab provides free access to AWS compute resources, Amazon SageMaker Studio Classic provides the following advanced machine learning capabilities that Studio Lab does not support.

- Continuous integration and continuous delivery (SageMaker Pipelines)
- Real-time predictions
- Large-scale distributed training
- Data preparation (Amazon SageMaker Data Wrangler)
- Data labeling (Amazon SageMaker Ground Truth)
- Feature Store
- Bias analysis (Clarify)
- Model deployment
- Model monitoring
Studio Classic also supports fine-grained access control and security by using AWS Identity and Access Management (IAM), Amazon Virtual Private Cloud (Amazon VPC), and AWS Key Management Service (AWS KMS). Studio Lab does not support these Studio Classic features, nor does it support the use of estimators and built-in SageMaker algorithms.

To export your Studio Lab projects for use with Studio Classic, see Export an Amazon SageMaker Studio Lab environment to Amazon SageMaker Studio Classic.

The following topics give information about Studio Lab and how to use it.

**Topics**

- Amazon SageMaker Studio Lab components overview
- Onboard to Amazon SageMaker Studio Lab
- Manage your account
- Launch your Amazon SageMaker Studio Lab project runtime
- Use Amazon SageMaker Studio Lab starter assets
- Studio Lab pre-installed environments
- Use the Amazon SageMaker Studio Lab project runtime
- Troubleshooting

**Amazon SageMaker Studio Lab components overview**

Amazon SageMaker Studio Lab consists of the following components. The following topics give more details about these components.

**Topics**

- Landing page
- Studio Lab account
- Project overview page
- Preview page
- Project
- Compute instance type
- Project runtime
Landing page

You can request an account and sign in to an existing account on your landing page. To navigate to the landing page, see the Amazon SageMaker Studio Lab website. For more information about creating a Studio Lab account, see Onboard to Amazon SageMaker Studio Lab.

The following screenshot shows the Studio Lab landing page interface for requesting a user account and signing in.

![Studio Lab landing page interface](image)

Learn and experiment with machine learning

Quickly create data analytics, scientific computing, and machine learning projects with notebooks in your browser.

Studio Lab account

Your Studio Lab account gives you access to Studio Lab. For more information about creating a user account, see Onboard to Amazon SageMaker Studio Lab.

Project overview page

You can launch a compute instance and view information about your project on this page. To navigate to this page, you must sign in from the Amazon SageMaker Studio Lab website. The URL takes the following format.

```
https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>
```
The following screenshot shows a project overview in the Studio Lab user interface.

![Studio Lab user interface screenshot](image)

**My Project**

<table>
<thead>
<tr>
<th>Status</th>
<th>Time remaining</th>
<th>Select compute type</th>
<th>Start runtime</th>
</tr>
</thead>
<tbody>
<tr>
<td>Idle</td>
<td></td>
<td>CPU</td>
<td>GPU</td>
</tr>
</tbody>
</table>

**Preview page**

On this page, you can access a read-only preview of a Jupyter notebook. You can not execute the notebook from preview, but you can copy that notebook into your project. For many customers, this may be the first Studio Lab page that customers see, as they may be opening a notebook from GitHub notebook. For more information on how to use GitHub resources, see [Use GitHub resources](#).

To copy the notebook preview to your Studio Lab project:

1. Sign in to your Studio Lab account. For more information about creating a Studio Lab account, see [Onboard to Amazon SageMaker Studio Lab](#).
2. Under **Notebook compute instance**, choose a compute instance type. For more information about compute instance types, see [Compute instance type](#).
3. Choose **Start runtime**. You might be asked to solve a CAPTCHA puzzle. For more information on CAPTCHA, see [What is a CAPTCHA puzzle?](#)
4. One time setup, for first time starting runtime using your Studio Lab account:
   a. Enter a mobile phone number to associate with your Amazon SageMaker Studio Lab account and choose **Continue**.
      
      For information on supported countries and regions, see [Supported countries and regions (SMS channel)](#).
   b. Enter the 6-digit code sent to the associated mobile phone number and choose **Verify**.
5. Choose **Copy to project**.
Project

Your project contains all of your files and folders, including your Jupyter notebooks. You have full control over the files in your project. Your project also includes the JupyterLab-based user interface. From this interface, you can interact with your Jupyter notebooks, edit your source code files, integrate with GitHub, and connect to Amazon S3. For more information, see Use the Amazon SageMaker Studio Lab project runtime.

The following screenshot shows a Studio Lab project with the file browser open and the Studio Lab Launcher displayed.

Compute instance type

Your Amazon SageMaker Studio Lab project runtime is based on an EC2 instance. You are allotted 15 GB of storage and 16 GB of RAM. Availability of compute instances is not guaranteed and is subject to demand. If you require additional storage or compute resources, consider switching to Studio.

Amazon SageMaker Studio Lab offers the choice of a CPU (Central Processing Unit) and a GPU (Graphical Processing Unit). The following sections give information about these two options, including selection guidance.

CPU
A central processing unit (CPU) is designed to handle a wide range of tasks efficiently, but is limited in how many tasks it can run concurrently. For machine learning, a CPU is recommended for compute intensive algorithms, such as time series, forecasting, and tabular data.

The CPU compute type has up to 4 hours at a time with a limit of 8 hours in a 24-hour period.

**GPU**

A graphics processing unit (GPU) is designed to render high-resolution images and video concurrently. A GPU is recommended for deep learning tasks, especially for transformers and computer vision.

The GPU compute type has up to 4 hours at a time with a limit of 4 hours in a 24-hour period.

**Compute time**

When compute time for Studio Lab reaches its time limit, the instance stops all running computations. Studio Lab does not support time limit increases.

Studio Lab automatically saves your environment when you update your environment and every time you create a new file. Custom-installed extensions and packages persist even after your runtime has ended.

File edits are periodically saved, but are not saved when your runtime ends. To ensure that you do not lose your progress, save your work manually. If you have content in your Studio Lab project that you don’t want to lose, we recommend that you back up your content elsewhere. For more information about exporting your environment and files, see Export an Amazon SageMaker Studio Lab environment to Amazon SageMaker Studio Classic.

During long computation, you do not need to keep your project open. For example, you can start training a model, then close your browser. The instance keeps running for up to the compute type limit in a 24-hour period. You can then sign in later to continue your work.

We recommend that you use checkpointing in your deep learning jobs. You can use saved checkpoints to restart a job from the previously saved checkpoint. For more information, see File I/O.

**Project runtime**

The project runtime is the period of time when your compute instance is running.
Session

A user session begins every time you launch your project.

Onboard to Amazon SageMaker Studio Lab

To onboard to Amazon SageMaker Studio Lab, follow the steps in this guide. In the following sections, you learn how to request a Studio Lab account, create your account, and sign in.

Topics

- Request a Studio Lab account
- Create a Studio Lab account
- Sign in to Studio Lab

Request a Studio Lab account

To use Studio Lab, you must first request approval to create a Studio Lab account. An AWS account cannot be used for onboarding to Studio Lab.

The following steps show how to request a Studio Lab account.

1. Navigate to the Studio Lab landing page.
2. Select Request account.
3. Enter the required information into the form.
4. Select Submit request.
5. If you receive an email to verify your email address, follow the instructions in the email to complete this step.

Your account request must be approved before you can register for a Studio Lab account. Your request will be reviewed within five business days. When your account request is approved, you receive an email with a link to the Studio Lab account registration page. This link expires seven days after your request is approved. If the link expires, you must submit a new account request.

Note: Your account request is denied if your email has been associated with activity that violates our Terms of Service or other agreements.
Referral codes

Studio Lab referral codes enable new account requests to be automatically approved to support machine learning events like workshops, hackathons, and classes. With a referral code, a trusted host can get their participants immediate access to Studio Lab. After an account has been created using a referral code, the account continues to exist after the expiration of the code.

To get a referral code, contact Sales Support. To use a referral code, enter the code as part of the account request form.

Create a Studio Lab account

After your request is approved, complete the following steps to create your Studio Lab account.

1. Select Create account in the account request approval email to open a new page.
2. From the new page, enter your Email, a Password, and a Username.
3. Select Create account.

You might be asked to solve a CAPTCHA puzzle. For more information on CAPTCHA, see What is a CAPTCHA puzzle?

Sign in to Studio Lab

After you register for your account, you can sign in to Studio Lab.

1. Navigate to the Studio Lab landing page.
2. Select Sign in to open a new page.
3. Enter your Email or Username and Password.
4. Select Sign in to open a new page to your project.

You might be asked to solve a CAPTCHA puzzle. For more information on CAPTCHA, see What is a CAPTCHA puzzle?

Manage your account

The following topic gives information about managing your account, including changing your password, deleting your account, and getting information that we have collected. These topics
require that you sign in to your Amazon SageMaker Studio Lab account. For more information, see Sign in to Studio Lab.

**Change your password**

Follow these steps to change your Amazon SageMaker Studio Lab password.

1. Navigate to the Studio Lab project overview page. The URL takes the following format.

   https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>

2. From the top-right corner, select your user name to open a dropdown menu.

3. From the dropdown menu, select **Change password** to open a new page.

4. Enter your current password into the **Enter your current password** field.

5. Enter your new password into the **Create a new password** and **Confirm your new password** fields.

6. Select **Submit**.

**Delete your account**

Follow these steps to delete your Studio Lab account.

1. Navigate to the Studio Lab project overview page. The URL takes the following format.

   https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>

2. From the top-right corner, select your user name to open a dropdown menu.

3. From the dropdown menu, select **Delete account** to open a new page.

4. Enter your password to confirm the deletion of your Studio Lab account.

5. Select **Delete**.

**Customer information**

Studio Lab collects your email address, user name, encrypted password, project files, and metadata. When requesting an account, you can optionally choose to provide your first and last name, country, organization name, occupation, and the reason for your interest in this product.
We protect all customer personal data with encryption. For more information about how your personal information is handled, see the Privacy Notice.

When you delete your account, all of your information is deleted immediately. If you have an inquiry about this, submit the Amazon SageMaker Studio Lab Form. For information and support related to AWS compliance, see Compliance support.

Launch your Amazon SageMaker Studio Lab project runtime

The Amazon SageMaker Studio Lab project runtime lets you write and run code directly from your browser. It is based on JupyterLab and has an integrated terminal and console. For more information about JupyterLab, see the JupyterLab Documentation.

The following topic gives information about how to manage your project runtime. These topics require that you sign in to your Amazon SageMaker Studio Lab account. For more information about signing in, see Sign in to Studio Lab. For more information about your project, see Amazon SageMaker Studio Lab components overview.

Topics

- Start your project runtime
- Stop your project runtime
- View remaining compute time
- Change your compute type

Start your project runtime

To use Studio Lab, you must start your project runtime. This runtime gives you access to the JupyterLab environment.

1. Navigate to the Studio Lab project overview page. The URL takes the following format.

   https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>

2. Under My Project, select a compute type. For more information about compute types, see Compute instance type.

3. Select Start runtime.
You might be asked to solve a CAPTCHA puzzle. For more information on CAPTCHA, see What is a CAPTCHA puzzle?

4. One time setup, for first time starting runtime using your Studio Lab account:
   a. Enter a mobile phone number to associate with your Amazon SageMaker Studio Lab account and choose Continue.
   
   For information on supported countries and regions, see Supported countries and regions (SMS channel).
   
   b. Enter the 6-digit code sent to the associated mobile phone number and choose Verify.

5. After the runtime is running, select Open project to open the project runtime environment in a new browser tab.

Stop your project runtime

When you stop your project runtime, your files are not automatically saved. To ensure that you don't lose your work, save all of your changes before stopping your project runtime.

- Under My Project, select Stop runtime.

View remaining compute time

Your project runtime has limited compute time based on the compute type that you select. For more information about compute time in Studio Lab, see Compute instance type.

- Under My Project, view Time remaining.

Change your compute type

You can switch your compute type based on your workflow. For more information about compute types, see Compute instance type.

1. Save any project files before changing the compute type.

2. Navigate to the Studio Lab project overview page. The URL takes the following format.

   https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>
3. Under **My Project**, select the desired compute type (CPU or GPU).

4. Confirm your choice by selecting **Restart** in the **Restart project runtime?** dialog box. Studio Lab stops your current project runtime, then starts a new project runtime with your updated compute type.

5. After your project runtime has started, select **Open project**. This opens your project runtime environment in a new browser tab. For information about using your project runtime environment, see **Use the Amazon SageMaker Studio Lab project runtime**.

---

## Use Amazon SageMaker Studio Lab starter assets

Amazon SageMaker Studio Lab supports the following assets to help machine learning (ML) practitioners get started. This guide shows you how to clone notebooks for your project.

### Getting started notebook

Studio Lab comes with a starter notebook that gives general information and guides you through key workflows. When you launch your project runtime for the first time, this notebook automatically opens.

### Dive into Deep Learning

Dive into Deep Learning (D2L) is an interactive, open-source book that teaches the ideas, mathematical theory, and code that power machine learning. With over 150 Jupyter notebooks, D2L provides a comprehensive overview of deep learning principles. For more information about D2L, see the [D2L website](#).

The following procedure shows how to clone the D2L Jupyter notebooks to your instance.

1. Start and open the Studio Lab project runtime environment by following **Start your project runtime**.

2. Once Studio Lab is open, choose the Git tab on the left sidebar.

3. Choose **Clone a Repository**. Under **Git repository URL (.git)** paste the MLU git repository D2L by following the steps below. If you do not see the **Clone a Repository** option because you are currently in a Git repository, return to the user directory to clone a new repository. You return to the user directory by choosing the Folder tab.
4. Navigate to the Studio Lab project overview page. The URL takes the following format.

https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>

5. Under **New to machine learning?**, choose **Dive into Deep Learning**.

6. From the new **Dive into Deep Learning** browser tab, choose **GitHub** to open a new page with the example notebooks.

7. Choose **Code** and copy the GitHub repository's URL in the **HTTPS** tab.

8. Return to the Studio Lab open project browser tab, paste the D2L repository URL, and clone the repository.

**AWS Machine Learning University**

The AWS Machine Learning University (MLU) provides access to the machine learning courses used to train Amazon's own developers. With AWS MLU, any developer can learn how to use machine learning with the learn-at-your-own-pace MLU Accelerator learning series. The MLU Accelerator series is designed to help developers begin their ML journey. It offers three-day foundational courses on these three subjects: Natural Language Processing, Tabular Data, and Computer Vision. For more information, see Machine Learning University.

The following procedure shows how to clone the AWS MLU Jupyter notebooks to your instance.

1. Start and open the Studio Lab project runtime environment by following **Start your project runtime**.

2. Once Studio Lab is open, choose the Git tab on the left sidebar.

3. Choose **Clone a Repository**. Under **Git repository URL (.git)** paste the MLU git repository URL by following the steps below. If you do not see the **Clone a Repository** option because you are currently in a Git repository, return to the user directory to clone a new repository. You return to the user directory by choosing the Folder tab on the left sidebar.
on the left sidebar. In the Folder tab beneath the file search bar choose the folder icon to the left of the currently open repository. Once you are in the user directory, choose the Git tab on the left sidebar and choose **Clone a Repository**.

4. Navigate to the Studio Lab project overview page. The URL takes the following format.

   https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>

5. Under **New to machine learning?**, choose **AWS Machine Learning University**.

6. From the new **AWS Machine Learning University** browser tab, find a course that interests you by reading the **Course Summary** for each course.

7. Choose the corresponding GitHub repository of interest under **Course Content**, to open a new page with the example notebooks.

8. Choose **Code** and copy the GitHub repository's URL in the **HTTPS** tab.

9. Return to the Studio Lab open project browser tab, paste the D2L repository URL, and choose **Clone** to clone the repository.

**Roboflow**

Roboflow gives you the tools to train, fine-tune, and label objects for computer vision applications. For more information, see [https://roboflow.com/](https://roboflow.com/).

The following procedure shows how to clone the Roboflow Jupyter notebooks to your instance.

1. Navigate to the Studio Lab project overview page. The URL takes the following format.

   https://studiolab.sagemaker.aws/users/<YOUR_USER_NAME>

2. Under **Resources and community**, find **Try Computer Vision**.

3. Under **Try Computer Vision** choose a Roboflow model. For more information, see [https://roboflow.com/](https://roboflow.com/).

4. Follow the tutorial under the Notebook preview.

**Studio Lab pre-installed environments**

Amazon SageMaker Studio Lab uses conda environments to contain your packages (or libraries). An environment is a folder that contains the packages you have installed. You can interact with an environment by using the terminal or your JupyterLab notebook. To use an environment and
the packages installed within, you must choose the corresponding kernel that contains the same name as the environment when opening your JupyterLab notebook. For a walkthrough on how to manage your environments, see Manage your environment. For more information on installing packages within your environment, see Customize your environment.

Studio Lab has various environments pre-installed for you. Any changes made to persistent memory environments will remain for your next session. Any changes to non-persistent memory environments will not remain for your next sessions, but the packages within will be updated and tested for compatibility by Amazon SageMaker. You will typically want to use the `sagemaker-distribution` non-persistent memory environment if you want to use a fully managed environment that already contains many popular packages used by machine learning (ML) engineers and data scientists. Otherwise you can use the default environment if you want to significantly customize your environment.

In the following we list the pre-installed environments and their use cases. To view the packages installed in an environment, see Customize your environment.

- **sagemaker-distribution**: Non-persistent memory environment that is regularly updated and tested for compatibility, fully managed by Amazon SageMaker. This environment contains popular packages used in ML, data science, and visualization. The `sagemaker-distribution` environment is closely related to the environment used in Amazon SageMaker Studio Classic, so after graduating from Studio Lab to Studio Classic the notebooks should run similarly. For information on exporting your environment from Studio Lab to Studio Classic, see Export an Amazon SageMaker Studio Lab environment to Amazon SageMaker Studio Classic.

- **default**: Persistent memory environment with very few packages pre-installed. Any installed packages or changes to this environment will continue on your next session.

- **studiolab**: Persistent memory environment where JupyterLab and other related packages are installed. This environment should only be used for JupyterLab and Jupyter server extensions, for configuring the JupyterLab user interface.

- **studiolab-safemode**: Non-persistent memory environment. This environment is automatically activated when there is an issue while starting your project runtime. Used for troubleshooting. For information on troubleshooting, see Troubleshooting.

- **base**: Non-persistent memory environment. This environment is only used for system tooling and should not be used by customers.

For information on SageMaker images and their versions, see Available Amazon SageMaker Images.
Use the Amazon SageMaker Studio Lab project runtime

The following topics give information about using the Amazon SageMaker Studio Lab project runtime. Before you can use the Studio Lab project runtime, you must onboard to Studio Lab by following the steps in Onboard to Amazon SageMaker Studio Lab.

Topics

- Amazon SageMaker Studio Lab UI overview
- Create or open an Amazon SageMaker Studio Lab notebook
- Use the Amazon SageMaker Studio Lab notebook toolbar
- Manage your environment
- Use external resources in Amazon SageMaker Studio Lab
- Get notebook differences
- Export an Amazon SageMaker Studio Lab environment to Amazon SageMaker Studio Classic
- Shut down resources

Amazon SageMaker Studio Lab UI overview

Amazon SageMaker Studio Lab extends the JupyterLab interface. Previous users of JupyterLab will notice similarities between the JupyterLab and Studio Lab UI, including the workspace. For an overview of the basic JupyterLab interface, see The JupyterLab Interface.

The following image shows Studio Lab with the file browser open and the Studio Lab Launcher displayed.
You will find the *menu bar* at the top of the screen. The *left sidebar* contains icons to open file browsers, resource browsers, and tools. The *status bar* is located at the bottom-left corner of Studio Lab.

The main work area is divided horizontally into two panes. The left pane is the *file and resource browser*. The right pane contains one or more tabs for resources, such as notebooks and terminals.

**Topics**

- *[Left sidebar]*
- *[File and resource browser]*
- *[Main work area]*

**Left sidebar**

The left sidebar includes the following icons. When you hover over an icon, a tooltip displays the icon name. When you choose an icon, the file and resource browser displays the described functionality. For hierarchical entries, a selectable breadcrumb at the top of the browser shows your location in the hierarchy.
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
</table>
| 📦 | **File Browser**  
Choose the **Upload Files** icon ( {}
  
) to add files to Studio Lab.  

Double-click a file to open the file in a new tab.  

To have adjacent files open, choose a tab that contains a notebook, Python, or text file, and then choose **New View for File**.  

Choose the plus (+) sign on the menu at the top of the file browser to open the Studio Lab Launcher. |
| 🟢 | **Running Terminals and Kernels**  
You can see a list of all of the running terminals and kernels in your project. For more information, see [Shut down resources](#). |
| 🗬️ | **Git**  
You can connect to a Git repository and then access a full range of Git tools and operations. For more information, see [Use external resources in Amazon SageMaker Studio Lab](#). |
| 📑 | **Table of Contents**  
You can access the Table of Contents for your current Jupyter notebook. |
| 🕵️‍♂️ | **Extension Manager**  
You can enable and manage third-party JupyterLab extensions. |
File and resource browser

The file and resource browser shows lists of your notebooks and files. On the menu at the top of the file browser, choose the plus (+) sign to open the Studio Lab Launcher. The Launcher allows you to create a notebook or open a terminal.

Main work area

The main work area has multiple tabs that contain your open notebooks and terminals.

Create or open an Amazon SageMaker Studio Lab notebook

When you create a notebook in Amazon SageMaker Studio Lab or open a notebook in Studio Lab, you must select a kernel for the notebook. The following topics describe how to create and open notebooks in Studio Lab.

For information about shutting down the notebook, see Shut down resources.

Topics

- Open a Studio Lab notebook
- Create a notebook from the file menu
- Create a notebook from the Launcher

Open a Studio Lab notebook

Studio Lab can only open notebooks listed in the Studio Lab file browser. To clone a notebook into your file browser from an external repository, see Use external resources in Amazon SageMaker Studio Lab.

To open a notebook

1. In the left sidebar, choose the File Browser icon ( ) to display the file browser.
2. Browse to a notebook file and double-click it to open the notebook in a new tab.
Create a notebook from the file menu

To create a notebook from the File menu

1. From the Studio Lab menu, choose File, choose New, and then choose Notebook.
2. To use the default kernel, in the Select Kernel dialog box, choose Select. Otherwise, to select a different kernel, use the dropdown menu.

Create a notebook from the Launcher

To create a notebook from the Launcher

1. Open the Launcher by using the keyboard shortcut Ctrl + Shift + L.

   Alternatively, you can open Launcher from the left sidebar: Choose the File Browser icon, and then choose the plus (+) icon.
2. To use the default kernel from the Launcher, under Notebook, choose default:Python. Otherwise, select a different kernel.

After you choose the kernel, your notebook launches and opens in a new Studio Lab tab.

To view the notebook's kernel session, in the left sidebar, choose the Running Terminals and Kernels icon ( ).

You can stop the notebook's kernel session from this view.

Use the Amazon SageMaker Studio Lab notebook toolbar

Amazon SageMaker Studio Lab notebooks extend the JupyterLab interface. For an overview of the basic JupyterLab interface, see The JupyterLab Interface.

The following image shows the toolbar and an empty cell from a Studio Lab notebook.

![Demo.ipynb](attachment:Demo.ipynb)

When you hover over a toolbar icon, a tooltip displays the icon function. You can find additional notebook commands in the Studio Lab main menu. The toolbar includes the following icons:
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="save" /></td>
<td><strong>Save and checkpoint</strong>&lt;br&gt;Saves the notebook and updates the checkpoint file.</td>
</tr>
<tr>
<td><img src="image" alt="add" /></td>
<td><strong>Insert cell</strong>&lt;br&gt;Inserts a code cell below the current cell. The current cell is noted by the blue vertical marker in the left margin.</td>
</tr>
<tr>
<td><img src="image" alt="cut-copy-paste" /></td>
<td><strong>Cut, copy, and paste cells</strong>&lt;br&gt;Cuts, copies, and pastes the selected cells.</td>
</tr>
<tr>
<td><img src="image" alt="run" /></td>
<td><strong>Run cells</strong>&lt;br&gt;Runs the selected cells. The cell that follows the last-selected cell becomes the new-selected cell.</td>
</tr>
<tr>
<td><img src="image" alt="interrupt" /></td>
<td><strong>Interrupt kernel</strong>&lt;br&gt;Interrupts the kernel, which cancels the currently-running operation. The kernel remains active.</td>
</tr>
<tr>
<td><img src="image" alt="re-run" /></td>
<td><strong>Restart kernel</strong>&lt;br&gt;Restarts the kernel. Variables are reset. Unsaved information is not affected.</td>
</tr>
<tr>
<td><img src="image" alt="re-run-all" /></td>
<td><strong>Restart kernel and re-run notebook</strong>&lt;br&gt;Restarts the kernel. Variables are reset. Unsaved information is not affected. Then re-runs the entire notebook.</td>
</tr>
<tr>
<td><img src="image" alt="code" /></td>
<td><strong>Cell type</strong>&lt;br&gt;Displays or changes the current cell type. The cell types are:&lt;br&gt;• Code – Code that the kernel runs.&lt;br&gt;• Markdown – Text rendered as markdown.</td>
</tr>
<tr>
<td><strong>Icon</strong></td>
<td><strong>Description</strong></td>
</tr>
<tr>
<td>----------</td>
<td>----------------</td>
</tr>
<tr>
<td></td>
<td>• Raw – Content, including Markdown markup, that's displayed as text.</td>
</tr>
</tbody>
</table>

### Checkpoint diff

Opens a new tab that displays the difference between the notebook and the checkpoint file. For more information, see [Get notebook differences](#).

### Git diff

Only enabled if the notebook is opened from a Git repository. Opens a new tab that displays the difference between the notebook and the last Git commit. For more information, see [Get notebook differences](#).

### default

**Kernel**

Displays or changes the kernel that processes the cells in the notebook.

No Kernel indicates that the notebook was opened without specifying a kernel. You can edit the notebook, but you can't run any cells.

### Kernel busy status

Displays a kernel's busy status by showing the circle's edge and its interior as the same color. The kernel is busy when it is starting and when it is processing cells. Additional kernel states are displayed in the status bar at the bottom-left corner of Studio Lab.

### Manage your environment

Amazon SageMaker Studio Lab provides pre-installed environments for your Studio Lab notebook instances. Environments allow you to start up a Studio Lab notebook instance with the packages you want to use. This is done by installing packages in the environment and then selecting the environment as a Kernel.

Studio Lab has various environments pre-installed for you. You will typically want to use the sagemaker-distribution environment if you want to use a fully managed environment that already contains many popular packages used for machine learning (ML) engineers and...
data scientists. Otherwise you can use the default environment if you want persistent customization for your environment. For more information on the available pre-installed Studio Lab environments, see Studio Lab pre-installed environments.

You can customize your environment by adding new packages (or libraries) to it. You can also create new environments from Studio Lab, import compatible environments, reset your environment to create space, and more.

The following commands are for running in a Studio Lab terminal. However, while installing packages it is highly recommended to install them within your Studio Lab Jupyter notebook to ensure that the packages are installed in the intended environment. To run the commands in a Jupyter notebook, prefix the command with a % before running the cell. For example, the code snippet `pip list` in a terminal is the same as `%pip list` in a Jupyter notebook.

The following sections give information about your default conda environment, how to customize it, and how to add and remove conda environments. For a list of sample environments that you can install into Studio Lab, see Creating Custom conda Environments. To use these sample environment YAML files with Studio Lab, see Step 4: Install your Studio Lab conda environments in Studio Classic.

Topics

- Your default environment
- View environments
- Create, activate, and use new conda environments
- Using sample Studio Lab environments
- Customize your environment
- Refresh Studio Lab

Your default environment

Studio Lab uses conda environments to encapsulate the software packages that are needed to run notebooks. Your project contains a default conda environment, named default, with the IPython kernel. This environment serves as the default kernel for your Jupyter notebooks.
View environments

To view the environments in Studio Lab you can use a terminal or Jupyter notebook. The following command will be for a Studio Lab terminal. If you wish to run the corresponding commands in a Jupyter notebook, see Manage your environment.

Open the Studio Lab terminal by opening the File Browser panel, choose the plus (+) sign on the menu at the top of the file browser to open the Launcher, then choose Terminal. From the Studio Lab terminal, list the conda environments by running the following.

```
conda env list
```

This command outputs a list of the conda environments and their locations in the file system. When you onboard to Studio Lab, you automatically activate the studiolab conda environment. The following is an example of listed environments after you onboard.

```
# conda environments:
#
default                  /home/studio-lab-user/.conda/envs/default
studiolab             *  /home/studio-lab-user/.conda/envs/studiolab
studiolab-safemode       /opt/amazon/sagemaker/safemode-home/.conda/envs/studiolab-safemode
base                     /opt/conda
sagemaker-distribution     /opt/conda/envs/sagemaker-distribution
```

The * marks the activated environment.

Create, activate, and use new conda environments

If you would like to maintain multiple environments for different use cases, you can create new conda environments in your project. The following sections show how to create and activate new conda environments. For a Jupyter notebook that shows how to create a custom environment, see Setting up a Custom Environment in SageMaker Studio Lab.

⚠️ Note

Maintaining multiple environments counts against your available Studio Lab memory.
Create conda environment

To create a conda environment, run the following conda command from your terminal. This example creates a new environment with Python 3.9.

```
conda create --name <ENVIRONMENT_NAME> python=3.9
```

Once the conda environment is created, you can view the environment in your environment list. For more information on how to view your environment list, see View environments.

Activate a conda environment

To activate any conda environment, run the following command in the terminal.

```
conda activate <ENVIRONMENT_NAME>
```

When you run this command, any packages installed using conda or pip are installed in the environment. For more information on installing packages, see Customize your environment.

Use a conda environment

To use your new conda environments with notebooks, make sure the ipykernel package is installed in the environment.

```
conda install ipykernel
```

Once the ipykernel package is installed in the environment, you can select the environment as the kernel for your notebook.

You may need to restart JupyterLab to see the environment available as a kernel. This can be done by choosing Amazon SageMaker Studio Lab in the top menu of Studio Lab and choosing Restart JupyterLab....

When you create a new notebook from the Studio Lab Launcher, you will have the option to choose the kernel under Notebook. For an overview of the Studio Lab UI, see Amazon SageMaker Studio Lab UI overview.

When a Jupyter notebook is open, you can choose the kernel by choosing Kernel from the top menu and choose Change Kernel....
Using sample Studio Lab environments

Studio Lab provides sample custom environments through the SageMaker Studio Lab Examples repository. The following shows how to clone and build these environments.

1. Clone the SageMaker Studio Lab Examples GitHub repository by following the instructions in Use GitHub resources.
2. In Studio Lab choose the File Browser icon on the left menu, so that the File Browser panel shows on the left.
3. Navigate to the studio-lab-examples/custom-environments directory in the File Browser.
4. Open the directory for the environment that you want to build.
5. Right click the .yml file in the folder, then select Build conda Environment.
6. You can now use the environment as a kernel after your conda environment has finished building. For instructions on how to use an existing environment as a kernel, see Create, activate, and use new conda environments

Customize your environment

You can customize your environment by installing and removing extensions and packages as needed. Studio Lab comes with environments with packages pre-installed and using an existing environment may save you time and memory, as pre-installed packages do not count against your available Studio Lab memory. For more information on the available pre-installed Studio Lab environments, see Studio Lab pre-installed environments.

Any installed extensions and packages installed on your default environment will persist in your project, so you do not need to install your packages for every project runtime session. However, extensions and packages installed on your sagemaker-distribution environment will not persist, so you will need to install new packages during your next session. Thus, it is highly recommended to install packages within your notebook to ensure that the packages are installed in the intended environment.

To view your environments, run the command conda env list.

To activate your environment, run the command conda activate <ENVIRONMENT_NAME>.

To view the packages in an environment, run the command conda list.
Install packages

It is highly recommended to install your packages within your Jupyter notebook to ensure that your packages are installed in the intended environment. To install additional packages to your environment from a Jupyter notebook, run one of the following commands in a cell within your Jupyter notebook. These commands install packages in the currently activated environment.

- `%conda install <PACKAGE>`
- `%pip install <PACKAGE>`

We don't recommend using the `!pip` or `!conda` commands because they can behave in unexpected ways when you have multiple environments.

After you install new packages to your environment, you may need to restart the kernel to ensure that the packages work in your notebook. This can be done by choosing Amazon SageMaker Studio Lab in the top menu of Studio Lab and choosing Restart JupyterLab.

Remove packages

To remove a package, run the command

```
%conda remove <PACKAGE_NAME>
```

This command will also remove any package that depends on `<PACKAGE_NAME>`, unless a replacement can be found without that dependency.

To remove all of the packages in an environment, run the command

```
conda deactivate
& & conda env remove --name <ENVIRONMENT_NAME>
```

Refresh Studio Lab

To refresh Studio Lab, remove all of your environments and files.

1. List all conda environments.

```
conda env list
```
2. Activate the base environment.

   conda activate base

3. Remove each environment in the list of conda environments, besides base.

   conda remove --name <ENVIRONMENT_NAME> --all

4. Delete all of the files on your Studio Lab.

   rm -rf *.*

---

**Use external resources in Amazon SageMaker Studio Lab**

With Amazon SageMaker Studio Lab, you can integrate external resources, such as Jupyter notebooks and data, from Git repositories and Amazon S3. You can also add an **Open in Studio Lab** button to your GitHub repo and notebooks. This button lets you clone your notebooks directly from Studio Lab.

The following topics show how to integrate external resources.

**Topics**

- Use GitHub resources
- Add an Open in Studio Lab button to your notebook
- Import files from your computer
- Connect to Amazon S3

**Use GitHub resources**

Studio Lab offers integration with GitHub. With this integration, you can clone notebooks and repositories directly to your Studio Lab project.

The following topics give information about how to use GitHub resources with Studio Lab.

**Studio Lab sample notebooks**

To get started with a repository of sample notebooks tailored for Studio Lab, see **Studio Lab Sample Notebooks**.
This repository provides notebooks for the following use cases and others.

- Computer vision
- Connecting to AWS
- Creating custom environments
- Geospatial data analysis
- Natural language processing
- Using R

**Clone a GitHub repo**

To clone a GitHub repo to your Studio Lab project, follow these steps.

1. Start your Studio Lab project runtime. For more information on launching Studio Lab project runtime, see [Start your project runtime](#).
2. In Studio Lab, choose the **File Browser** icon on the left menu, so that the **File Browser** panel shows on the left.
3. Navigate to your user directory by choosing the file icon beneath the file search bar.
4. Select the **Git** icon from the left menu to open a new dropdown menu.
5. Choose **Clone a Repository**.
6. Paste the repository's URL under **Git repository URL (.git)**.
7. Select **Clone**.

**Clone individual notebooks from GitHub**

To open a notebook in Studio Lab, you must have access to the repo that the notebook is in. The following examples describe Studio Lab permission-related behavior in various situations.

- If a repo is public, you can automatically clone the notebook into your project from the Studio Lab preview page.
- If a repo is private, you are prompted to sign in to GitHub from the Studio Lab preview page. If you have access to a private repo, you can clone the notebook into your project.
If you don't have access to a private repo, you cannot clone the notebook from the Studio Lab preview page.

The following sections show two options for you to copy a GitHub notebook in your Studio Lab project. These options depend on whether the notebook has an Open in Studio Lab button.

**Option 1: Copy notebook with an Open in Studio Lab button**

The following procedure shows how to copy a notebook that has an Open in Studio Lab button. If you want to add this button to your notebook, see Add an Open in Studio Lab button to your notebook.

1. Sign in to Studio Lab following the steps in Sign in to Studio Lab.
2. In a new browser tab, navigate to the GitHub notebook that you want to clone.
3. In the notebook, select the Open in Studio Lab button to open a new page in Studio Lab with a preview of the notebook.
4. If your project runtime is not already running, start it by choosing the Start runtime button at the top of the preview page. Wait for the runtime to start before proceeding to the next step.
5. After your project runtime has started, select Copy to project to open your project runtime in a new browser tab.
6. In the Copy from GitHub? dialog box, select Copy notebook only. This copies the notebook file to your project.

**Option 2: Clone any GitHub notebook**

The following procedure shows how to copy any notebook from GitHub.

1. Navigate to the notebook in GitHub.
2. In the browser's address bar, modify the notebook URL, as follows.

   # Original URL
   https://github.com/\(<PATH\_TO\_NOTEBOOK>\)

   # Modified URL
   https://studiolab.sagemaker.aws/import/github/\(<PATH\_TO\_NOTEBOOK>\)

3. Navigate to the modified URL. This opens a preview of the notebook in Studio Lab.
4. If your project runtime is not already running, start it by choosing the **Start runtime** button at the top of the preview page. Wait for the runtime to start before proceeding to the next step.

5. After your project runtime has started, select **Copy to project** to open your project runtime in a new browser tab.

6. In the **Copy from GitHub?** dialog box, select **Copy notebook only** to copy the notebook file to your project.

### Add an Open in Studio Lab button to your notebook

When you add the **Open in Studio Lab** button to your notebooks, others can clone your notebooks or repositories directly to their Studio Lab projects. If you are sharing your notebook within a public GitHub repository, your content will be publicly readable. Do not share private content, such as AWS access keys or AWS Identity and Access Management credentials, in your notebook.

To add the functional **Open in Studio Lab** button to your Jupyter notebook or repository, add the following markdown to the top of your notebook or repository.

```
[![Open In SageMaker Studio Lab](https://studiolab.sagemaker.aws/studiolab.svg)](https://studiolab.sagemaker.aws/import/github/<PATH_TO_YOUR_NOTEBOOK_ON_GITHUB>)
```

### Import files from your computer

The following steps show how to import files from your computer to your Studio Lab project.

1. Open the Studio Lab project runtime.
2. Open the **File Browser** panel.
3. In the actions bar of the **File Browser** panel, select the **Upload Files** button.
4. Select the files that you want to upload from your local machine.
5. Select **Open**.

Alternatively, you can drag and drop files from your computer into the **File Browser** panel.

### Connect to Amazon S3

The AWS CLI enables AWS integration in your Studio Lab project. With this integration, you can pull resources from Amazon S3 to use with your Jupyter notebooks.
To use AWS CLI with Studio Lab, complete the following steps. For a notebook that outlines this integration, see Using Studio Lab with AWS Resources.

1. Install the AWS CLI following the steps in Installing or updating the latest version of the AWS CLI.
2. Configure your AWS credentials by following the steps in Quick setup. The role for your AWS account must have permissions to access the Amazon S3 bucket that you are copying data from.
3. From your Jupyter notebook, clone resources from the Amazon S3 bucket, as needed. The following command shows how to clone all resources from an Amazon S3 path to your project. For more information, see the AWS CLI Command Reference.

```
aws s3 cp s3://<BUCKET_NAME>/<PATH_TO_RESOURCES>/ <PROJECT_DESTINATION_PATH>/ --recursive
```

Get notebook differences

You can display the difference between the current notebook and the last checkpoint, or the last Git commit, using the Amazon SageMaker Studio Lab project UI.

Topics

- Get the difference between the last checkpoint
- Get the difference between the last commit

Get the difference between the last checkpoint

When you create a notebook, a hidden checkpoint file that matches the notebook is created. You can view changes between the notebook and the checkpoint file, or revert the notebook to match the checkpoint file.

To save the Studio Lab notebook and update the checkpoint file to match: Choose the Save notebook and create checkpoint icon ( ).

This is located on the Studio Lab menu's left side. The keyboard shortcut for Save notebook and create checkpoint is Ctrl + s.
To view changes between the Studio Lab notebook and the checkpoint file: Choose the **Checkpoint diff** icon ( ), located in the center of the Studio Lab menu.

To revert the Studio Lab notebook to the checkpoint file: On the main Studio Lab menu, choose **File**, and then **Revert Notebook to Checkpoint**.

**Get the difference between the last commit**

If a notebook is opened from a Git repository, you can view the difference between the notebook and the last Git commit.

To view the changes in the notebook from the last Git commit: Choose the **Git diff** icon ( ) in the center of the notebook menu.

**Export an Amazon SageMaker Studio Lab environment to Amazon SageMaker Studio Classic**

Amazon SageMaker Studio Classic offers many features for machine learning and deep learning work flows that are unavailable in Amazon SageMaker Studio Lab. This page shows how to migrate a Studio Lab environment to Studio Classic to take advantage of more compute capacity, storage, and features. However, you may want to familiarize yourself with Studio Classic's prebuilt containers, which are optimized for the full MLOP pipeline. For more information, see [Amazon SageMaker Studio Lab](#).

To migrate your Studio Lab environment to Studio Classic, you must first onboard to Studio Classic following the steps in [Amazon SageMaker domain overview](#).

**Topics**

- **Step 1: Export your Studio Lab conda environment**
- **Step 2: Save your Studio Lab artifacts**
- **Step 3: Import your Studio Lab artifacts to Studio Classic**
- **Step 4: Install your Studio Lab conda environments in Studio Classic**
Step 1: Export your Studio Lab conda environment

You can export a conda environment and add libraries or packages to the environment by following the steps in Manage your environment. The following example demonstrates using the default environment to be exported to Studio Classic.

1. Open the Studio Lab terminal by opening the File Browser panel, choose the plus (+) sign on the menu at the top of the file browser to open the Launcher, then choose Terminal. From the Studio Lab terminal, list the conda environments by running the following.

   ```bash
   conda env list
   ```

   This command outputs a list of the conda environments and their locations in the file system. When you onboard to Studio Lab, you automatically activate the studiolab conda environment.

   ```
   # conda environments: #
   default                  /home/studio-lab-user/.conda/envs/default
   studiolab             *  /home/studio-lab-user/.conda/envs/studiolab
   studiolab-safemode       /opt/amazon/sagemaker/safemode-home/.conda/
   envs/studiolab-safemode
   base                     /opt/conda
   ```

   We recommend that you do not export the studiolab, studiolab-safemode, and base environments. These environments are not usable in Studio Classic for the following reasons:

   - **studiolab**: This sets up the JupyterLab environment for Studio Lab. Studio Lab runs a different major version of JupyterLab than Studio Classic, so it is not usable in Studio Classic.
   - **studiolab-safemode**: This also sets up the JupyterLab environment for Studio Lab. Studio Lab runs a different major version of JupyterLab than Studio Classic, so it is not usable in Studio Classic.
   - **base**: This environment comes with conda by default. The base environment in Studio Lab and the base environment in Studio Classic have incompatible versions of many packages.

2. For the conda environment that you want to migrate to Studio Classic, first activate the conda environment. The default environment is then changed when new libraries are installed or
removed from it. To get the exact state of the environment, export it into a YAML file using the command line. The following command lines export the default environment into a YAML file, creating a file called `myenv.yml`.

```bash
conda activate default
conda env export > ~/myenv.yml
```

**Step 2: Save your Studio Lab artifacts**

Now that you have saved your environment to a YAML file, you can move the environment file to any platform.

**Save to a local machine using Studio Lab GUI**

> Note

Downloading a directory from the Studio Lab GUI by right-clicking on the directory is currently unavailable. If you wish to export a directory, please follow the steps using the **Save to Git repository** tab.

One option is to save the environment onto your local machine. To do this, use the following procedure.

1. In Studio Lab, choose the **File Browser** icon on the left menu, so that the **File Browser** panel shows on the left.
2. Navigate to your user directory by choosing the file icon beneath the file search bar.
3. Choose (right-click) the `myenv.yml` file and then choose **Download**. You can repeat this process for other files you want to import to Studio Classic.

**Save to a Git repository**

Another option is to save your environment to a Git repository. This option uses GitHub as an example. These steps require a GitHub account and repository. For more information, visit [GitHub](https://github.com). The following procedure shows how to synchronize your content with GitHub using the Studio Lab terminal.
1. From the Studio Lab terminal, navigate to your user directory and make a new directory to contain the files you want to export.

   ```
cd ~
mkdir <NEW_DIRECTORY_NAME>
   ```

2. After you create a new directory, copy any file or directory you want to export to `<NEW_DIRECTORY_NAME>`.

   Copy a file using the following code format:

   ```
cp <FILE_NAME> <NEW_DIRECTORY_NAME>
   ```

   For example, replace `<FILE_NAME>` with `myenv.yml`.

   Copy any directory using the following code format:

   ```
cp -r <DIRECTORY_NAME> <NEW_DIRECTORY_NAME>
   ```

   For example, replace `<DIRECTORY_NAME>` with any directory name in your user directory.

3. Navigate to the new directory and initialize the directory as a Git repository using the following command. For more information, see the git-init documentation.

   ```
cd <NEW_DIRECTORY_NAME>
git init
   ```

4. Using Git, add all relevant files and then commit your changes.

   ```
git add .
git commit -m "<COMMIT_MESSAGE>"
   ```

   For example, replace `<COMMIT_MESSAGE>` with Add Amazon SageMaker Studio Lab artifacts to GitHub repository to migrate to Amazon SageMaker Studio Classic.

5. Push the commit to your remote repository. This repository has the format https://github.com/<GITHUB_USERNAME>/<REPOSITORY_NAME>.git where `<GITHUB_USERNAME>` is your GitHub user name and the `<REPOSITORY_NAME>` is your
remote repository name. Create a branch `<BRANCH_NAME>` to push the content to the GitHub repository.

```
git branch -M <BRANCH_NAME>
git remote add origin https://github.com/<GITHUB_USERNAME>/<REPOSITORY_NAME>.git
git push -u origin <BRANCH_NAME>
```

**Step 3: Import your Studio Lab artifacts to Studio Classic**

The following procedure shows how to import artifacts to Studio Classic. The instructions on using Feature Store through the console depends on if you have enabled Studio or Studio Classic as your default experience. For information on accessing Studio Classic through the console, see Launch Studio Classic if Studio is your default experience.

From Studio Classic, you can import files from your local machine or from a Git repository. You can do this using the Studio Classic GUI or terminal. The following procedure uses the examples from Step 2: Save your Studio Lab artifacts.

**Import using the Studio Classic GUI**

If you saved the files to your local machine, you can import the files to Studio Classic using the following steps.

1. Open the File Browser panel at the top left of Studio Classic.
2. Choose the Upload Files icon ( ) on the menu at the top of the File Browser panel.
3. Navigate to the file that you want to import, then choose Open.

**Note**

To import a directory into Studio Classic, first compress the directory on your local machine to a file. On a Mac, right-click the directory and choose Compress "<DIRECTORY_NAME>". In Windows, right-click the directory and choose Send to, and then choose Compressed (zipped) folder. After the directory is compressed, import the
imported file using the preceding steps. Unzip the compressed file by navigating to
the Studio Classic terminal and running the command `<DIRECTORY_NAME>.zip`

Import using a Git repository

This example provides two options for how to clone a GitHub repository
into Studio Classic. You can use the Studio Classic GUI by choosing the **Git**

) tab on the left side of Studio Classic. Choose **Clone a Repository**, then paste your GitHub
repository URL from **Step 2: Save your Studio Lab artifacts**. Another option is to use the Studio
Classic terminal by using the following procedure.

1. Open the Studio Classic **Launcher**. For more information on opening the **Launcher**, see
   Amazon SageMaker Studio Classic Launcher.
2. In the **Launcher**, in the **Notebooks and compute resources** section, choose **Change
   environment**.
3. In Studio Classic, open the **Launcher**. To open the **Launcher**, choose **Amazon SageMaker
   Studio Classic** at the top-left corner of Studio Classic.

   To learn about all the available ways to open the **Launcher**, see Use the Amazon SageMaker
   Studio Classic Launcher.
4. In the **Change environment** dialog, use the **Image** dropdown list to select the **Data Science**
   image and choose **Select**. This image comes with conda pre-installed.
5. In the Studio Classic **Launcher**, choose **Open image terminal**.
6. From the image terminal, run the following command to clone your repository. This
   command creates a directory named after `<REPOSITORY_NAME>` in your Studio Classic
   instance and clones your artifacts in that repository.

   
   ```
   git clone https://github.com/<GITHUB_USERNAME>/<REPOSITORY_NAME>.git
   ```

**Step 4: Install your Studio Lab conda environments in Studio Classic**

You can now recreate your conda environment by using your YAML file in your Studio Classic
instance. Open the Studio Classic **Launcher**. For more information on opening the **Launcher**, see
Amazon SageMaker Studio Classic Launcher. From the **Launcher**, choose **Open image terminal**.
In the terminal navigate to the directory that contains the YAML file, then run the following commands.

```bash
conda env create --file <ENVIRONMENT_NAME>.yml
conda activate <ENVIRONMENT_NAME>
```

After these commands are complete, you can select your environment as the kernel for your Studio Classic notebook instances. To view the available environment, run `conda env list`. To activate your environment, run `conda activate <ENVIRONMENT_NAME>`.

### Shut down resources

In this guide, you will learn how to shut down individual resources, including notebooks, terminals, and kernels. You can also shut down all resources in one of these categories at the same time.

**Topics**
- Shut down an open notebook
- Shut down resources

### Shut down an open notebook

You can shut down an open notebook from the Amazon SageMaker Studio Lab File menu or from the Running Terminals and Kernels pane.

**Note**

When you shut down a notebook, any unsaved information in the notebook is lost. The notebook is not deleted.

**To shut down an open notebook from the File menu**

1. Save the notebook contents by choosing the icon, located in the notebook menu.
2. Choose File then Close and Shutdown Notebook.
3. Choose OK.
Shut down resources

On the left sidebar of Studio Lab, you will find the Running Terminals and Kernels pane and icon. The Running Terminals and Kernels pane has three sections. Each section lists all of the resources of that type. You can shut down each resource individually, or shut down all resources in a section simultaneously.

When you shut down all resources in a section, the following occurs:

- **Kernels** – All kernels, notebooks, and consoles are shut down.
- **Terminals** – All terminals are shut down.

To shut down resources

1. In the left sidebar, choose the Running Terminals and Kernels icon ( ).

2. Do either of the following:
   - To shut down a specific resource: Choose the SHUT DOWN icon on the same row as the resource.
   - To shut down all resources in a section: Choose Shut Down All, which is located to the right of the section label. After a confirmation dialog box appears, choose Shut down all to proceed.

Troubleshooting

The guide shows common errors that might occur when using Amazon SageMaker Studio Lab. Each error contains a description, as well as a solution to the error.

**Note**

You cannot share your password with multiple users or use Studio Lab to mine cryptocurrency. We don’t recommend using Studio Lab for production tasks because of runtime limits.
Can’t access account

If you can’t access your account, verify that you are using the correct email and password. If you have forgotten your password, use the following steps to reset your password. If you still cannot access your account, you must request and register for a new account using the instructions in Onboard to Amazon SageMaker Studio Lab.

Forgot password

If you forget your password, you must reset it using the following steps.

1. Navigate to the Studio Lab landing page.
2. Select Sign in.
3. Select Forgot password? to open a new page.
4. Enter the email address that you used to sign up for an account.
5. Select Send reset link to send an email with a password reset link.
6. From the password reset email, select Reset your password.
7. Enter your new password.
8. Select Submit.

Can’t launch project runtime

If the Studio Lab project runtime does not launch, try launching it again. If this doesn't work, switch the instance type from CPU to GPU (or in reverse). For more information, see Change your compute type.

Runtime stopped running unexpectedly

If there is an issue with the environment used to run JupyterLab, then Studio Lab will automatically recreate the environment. Studio Lab does not support manual activation of this process.

Conflicting versions

Because you can add packages and modify your environment as needed, you may run into conflicts between packages in your environment. If there are conflicts between packages in your environment, you must remove the conflicting package.

Environment build fails
When you build an environment from a YAML file, a package-version conflict or file issue might cause a build to fail. To resolve this, remove the environment by running the following command. Do this before attempting to build it again.

```
conda remove --name <YOUR_ENVIRONMENT> --all
```

**Error message about allowing to download script from domain *.awswaf.com**

Studio Classic uses the web application firewall service AWS WAF to protect your resources, which uses JavaScript. If you are using a browser security plugin that prevents JavaScript from downloading, this error may pop up. To use Studio Classic, allow the JavaScript download from *.awswaf.com as a trusted domain. For more information on AWS WAF, see [AWS WAF](#) from the AWS WAF, AWS Firewall Manager, and AWS Shield Advanced. Developer Guide.

**Disk space is full**

If you run into a notification saying mentioning that your disk space is full or **File Load Error for <FILE_NAME>** while attempting to open a file, you can remove files, directories, libraries, or environments to increase space. For more information on managing your libraries and environments, see [Manage your environment](#).

**Project runtime is in safe mode notification**

If you run into a notification that **Project runtime is in safe mode**, you must free up some disk space to resume using the Studio Lab project runtime. Follow the instructions in the preceding troubleshoot item, **Disk space is full**. Once up to at least 500 MB of space has been cleared, you may restart the project runtime to use Studio Lab. This can be done by choosing [Amazon SageMaker Studio Lab](#) in the top menu of Studio Lab and choosing **Restart JupyterLab**.

**git Cannot import cv2**

If you run into an error when importing cv2 after installing opencv-python, you must uninstall opencv-python and install opencv-python-headless as follows.

```
%pip uninstall opencv-python --yes
%pip install opencv-python-headless
```

You can then import cv2 as expected.

**Studio Lab becomes unresponsive when opening large files**
The Studio Lab IDE may fail to render when large files are opened, resulting in blocked access to Studio Lab resources. To resolve this, reset the Studio Lab workspace using the following procedure.

1. After you open the IDE, copy the URL in your browser's address bar. This URL should be in the https://xxxxxx.studio.us-east-2.sagemaker.aws/studiolab/default/lab format. Close the tab.
2. In a new tab, paste the URL and remove anything after https://xxxxxx.studio.us-east-2.sagemaker.aws/studiolab/default/lab.
3. Add ?reset to the end of the URL, so it is in the https://xxxxxx.studio.us-east-2.sagemaker.aws/studiolab/default/lab?reset format.
4. Navigate to the updated URL. This resets the saved UI state and makes the Studio Lab IDE responsive.

Amazon SageMaker Canvas

Amazon SageMaker Canvas gives you the ability to use machine learning to generate predictions without needing to write any code. The following are some use cases where you can use SageMaker Canvas:

- Predict customer churn
- Plan inventory efficiently
- Optimize price and revenue
- Improve on-time deliveries
- Classify text or images based on custom categories
- Identify objects and text in images
- Extract information from documents

With Canvas, you can chat with popular large language models (LLMs), access Ready-to-use models, or build a custom model trained on your data.

Canvas chat is a functionality that leverages open-source and Amazon LLMs to help you boost your productivity. You can prompt the models to get assistance with tasks such as generating content, summarizing or categorizing documents, and answering questions. To learn more, see Use generative AI with foundation models.
The Ready-to-use models in Canvas can extract insights from your data for a variety of use cases. You don’t have to build a model to use Ready-to-use models because they are powered by Amazon AI services, including Amazon Rekognition, Amazon Textract, and Amazon Comprehend. You only have to import your data and start using a solution to generate predictions.

If you want a model that is customized to your use case and trained with your data, you can build a model. You can get predictions customized to your data by doing the following:

1. Import your data from one or more data sources.
2. Build a predictive model.
3. Evaluate the model's performance.
4. Generate predictions with the model.

Canvas supports the following types of custom models:

- Numeric prediction (also known as regression)
- Categorical prediction for 2 and 3+ categories (also known as binary and multi-class classification)
- Time series forecasting
- Single-label image prediction (also known as image classification)
- Multi-category text prediction (also known as multi-class text classification)

You can also bring your own models into Canvas from Amazon SageMaker Studio Classic.

To learn more about pricing, see the SageMaker Canvas pricing page. You can also see Manage billing and cost in SageMaker Canvas for more information.

SageMaker Canvas is currently available in the following Regions:

- US East (Ohio)
- US East (N. Virginia)
- US West (N. California)
- US West (Oregon)
- Asia Pacific (Mumbai)
- Asia Pacific (Seoul)
• Asia Pacific (Singapore)
• Asia Pacific (Sydney)
• Asia Pacific (Tokyo)
• Canada (Central)
• Europe (Frankfurt)
• Europe (Ireland)
• Europe (London)
• Europe (Paris)
• Europe (Stockholm)
• South America (São Paulo)

Topics

• Are you a first-time SageMaker Canvas user?
• Getting started with using Amazon SageMaker Canvas
• Setting Up and Managing Amazon SageMaker Canvas (for IT Administrators)
• Import data into Canvas
• Prepare data
• Use generative AI with foundation models
• Use Ready-to-use models
• Use custom models
• Logging out of Amazon SageMaker Canvas
• Limitations and troubleshooting
• Manage billing and cost in SageMaker Canvas

Are you a first-time SageMaker Canvas user?

If you are a first-time user of SageMaker Canvas, we recommend that you begin by reading the following sections:

• For IT administrators – Setting Up and Managing Amazon SageMaker Canvas (for IT Administrators)
• For analysts and individual users – Getting started with using Amazon SageMaker Canvas

Getting started with using Amazon SageMaker Canvas

This guide tells you how to get started with using SageMaker Canvas. If you’re an IT administrator and would like more in-depth details, see Setting Up and Managing Amazon SageMaker Canvas (for IT Administrators) to set up SageMaker Canvas for your users.

Topics

• Prerequisites for setting up Amazon SageMaker Canvas
• Step 1: Log in to SageMaker Canvas
• Step 2: Use SageMaker Canvas to get predictions

Prerequisites for setting up Amazon SageMaker Canvas

To set up a SageMaker Canvas application, you must first onboard to Amazon SageMaker domain, which supports the various machine learning (ML) environments such as Canvas and SageMaker Studio.

The following section describes how to set up an Amazon SageMaker domain and give yourself Canvas permissions.

⚠️ Important

For you to set up Amazon SageMaker Canvas, your version of Amazon SageMaker Studio must be 3.19.0 or later. For information about updating Amazon SageMaker Studio, see Shut down and Update SageMaker Studio Classic.

Onboard to domain

To set up your domain, first see Amazon SageMaker domain overview to learn more about domains.

Then, when you’re ready to set up your domain, choose one of the following setup methods:

1. (Quick) Quick onboard to Amazon SageMaker domain – Choose this option if you’d like to quickly set up your domain. This grants your user all of the default Canvas permissions and basic
functionality. Any additional features such as document querying can be enabled later by an admin. If you want to configure more granular permissions, we recommend that you choose option 2 or 3.

2. (Advanced) **Custom onboarding to Amazon SageMaker domain using IAM Identity Center** – Choose this option if you’d like to complete a more advanced setup of your domain. To use the IAM Identity Center method, you must be part of an organization in AWS Organizations.

3. (Advanced) **Custom onboarding to Amazon SageMaker domain using IAM** – Choose this option if you’d like to complete a more advanced setup of your domain and don’t require the organizational setup of the IAM Identity Center.

If you’re doing the Quick setup (option 1 in the preceding list), then you can skip the rest of this section and move on to **Step 1: Log in to SageMaker Canvas**.

If you’re doing the Advanced setup (options 2 or 3), then you can specify the Canvas features to which you’d like to grant your users access. Use the rest of this section as you complete the advanced domain setup to help you configure the permissions that are specific to Canvas.

In either the **Custom onboarding to Amazon SageMaker domain using IAM Identity Center** or the **Custom onboarding to Amazon SageMaker domain using IAM** setup instructions, for **Step 2: Users and ML Activities**, you must select the Canvas permissions that you want to grant. In the **ML activities** section, you can select the following permissions policies to grant access to Canvas features. You can only select up to 8 **ML activities** total when setting up your domain. The first two permissions in the following list are required to use Canvas, while the rest are for additional features.

- **Run Studio Applications** – These permissions are necessary to start up the Canvas application.
- **Canvas Core Access** – These permissions grant you access to the Canvas application and the basic functionality of Canvas, such as creating datasets, using basic data transforms, and building and analyzing models.
- (Optional) **Canvas Data Preparation (powered by Data Wrangler)** – These permissions grant you access to create data flows and use advanced transforms to prepare your data in Canvas. These permissions are also necessary for creating data processing jobs and data preparation job schedules.
- (Optional) **Canvas AI Services** – These permissions grant you access to the Ready-to-use models, foundation models, and Chat with Data features in Canvas.
• **Optional) Kendra access** – This permission grants you access to the document querying feature, where you can query documents stored in an Amazon Kendra index using foundation models in Canvas.

If you select this option, then in the **Canvas Kendra Access** section, enter the IDs for your Amazon Kendra indexes to which you want to grant access.

• **Optional) Canvas MLOps** – This permission grants you access to the model deployment feature in Canvas, where you can deploy models for use in production.

In the domain setup’s **Step 3: Applications** section, choose **Configure Canvas** and then do the following:

1. **For the Canvas storage configuration**, specify where you want Canvas to store the application data, such as model artifacts, batch predictions, datasets, and logs. SageMaker creates a Canvas/ folder inside this bucket to store the data. For more information, see **Configure your Amazon S3 storage**. For this section, do the following:
   a. Select **System managed** if you want to set the location to the default SageMaker-created bucket that follows the pattern `s3://sagemaker-{Region}-{your-account-id}`.
   b. Select **Custom S3** to specify your own Amazon S3 bucket as the storage location. Then, enter the Amazon S3 URI.
   c. **Optional) For Encryption key**, specify a KMS key for encrypting Canvas artifacts stored at the specified location.

2. **(Optional) For the Canvas Ready-to-use models configuration**, do the following:
   a. Leave the **Enable Canvas Ready-to-use models** option turned on to give your users permissions to generate predictions with Ready-to-use models in Canvas (it is turned on by default). This option also gives you permissions to chat with generative-AI powered models. For more information, see **Use generative AI with foundation models**.
   b. Leave the **Enable document query using Amazon Kendra** option turned on to give your users permissions to use foundation models for querying documents stored in an Amazon Kendra index. Then, from the dropdown menu, select the existing indexes to which you want to grant access. For more information, see **Use generative AI with foundation models**.

3. **(Optional) For the ML Ops permissions configuration** section, do the following:
a. Leave the **Enable direct deployment of Canvas models** option turned on to give your users permissions to deploy their models from Canvas to a SageMaker endpoint. For more information about model deployment in Canvas, see [Deploy your models to an endpoint](#).

b. Leave the **Enable Model Registry registration permissions for all users** option turned on to give your users permissions to register their model version to the SageMaker model registry (it is turned on by default). For more information, see [Register a model version in the SageMaker model registry](#).

c. If you left the **Enable Model Registry registration permissions for all users** option turned on, then select either **Register to Model Registry only** or **Register and approve model in Model Registry**.

4. (Optional) For the **Local file upload configuration** section, turn on the **Enable local file upload** option to give your users permissions to upload files to Canvas from their local machines. Turning this option on attaches a cross-origin resource sharing (CORS) policy to the Amazon S3 bucket specified in the **Canvas storage configuration** (and overrides any existing CORS policy). To learn more about local file upload permissions, see [Grant Your Users Permissions to Upload Local Files](#).

5. (Optional) For the **OAuth settings** section, do the following:

   a. Choose **Add OAuth configuration**.

   b. For **Data source**, select your data source.

   c. For **Secret setup**, select **Create a new secret** and enter the information you have from your identity provider. If you haven’t done the initial OAuth setup with your data source yet, see [Set up connections to data sources with OAuth](#).

6. (Optional) For the **Time series forecasting configuration**, leave the **Enable time series forecasting** option turned on to give your users permissions to do time series forecasting in SageMaker Canvas (it is turned on by default).

   - If you left **Enable time series forecasting** turned on, select **Create and use a new execution role**, or select **Use an existing execution role** if you already have an IAM role with the required Amazon Forecast permissions attached (for more information, see the [IAM role setup method](#)).

7. Finish configuring the rest of the domain settings using the [Custom onboarding to Amazon SageMaker domain using IAM Identity Center](#) or [Custom onboarding to Amazon SageMaker domain using IAM](#) procedures.
Note

If you encounter any issues with granting permissions through the console, such as permissions for Ready-to-use models, see the topic Troubleshooting issues with granting permissions through the SageMaker console.

You should now have a SageMaker domain set up and all of the Canvas permissions configured.

You can edit the Canvas permissions for a domain or a specific user after the initial domain setup. Individual user settings override the domain settings. To learn how to view or edit your Canvas permissions in the domain settings, see View and edit domains.

Give yourself permissions to use specific features in Canvas

The following information outlines the various permissions that you can grant to a Canvas user to allow the use of various features and functionalities within Canvas. Some of these permissions can be granted during the domain setup, but some require additional permissions or configuration. Refer to the specific permissions information for each feature that you want to enable:

- **Local file upload.** The permissions for local file upload are turned on by default in the Canvas base permissions when setting up your domain. If you can't upload local files from your machine to SageMaker Canvas, you can attach a CORS policy to the Amazon S3 bucket that you specified in the Canvas storage configuration. If you allowed SageMaker to use the default bucket, the bucket follows the naming pattern s3://sagemaker-{Region}-{your-account-id}. For more information, see Grant Your Users Permissions to Upload Local Files.

- **Custom image and text prediction models.** The permissions for building custom image and text prediction models are turned on by default in the Canvas base permissions when setting up your domain. However, if you have a custom IAM configuration and don't want to attach the AmazonSageMakerCanvasFullAccess policy to your user's IAM execution role, then you must explicitly grant your user the necessary permissions. For more information, see Grant Your Users Permissions to Build Custom Image and Text Prediction Models.

- **Ready-to-use models and foundation models.** You might want to use the Canvas Ready-to-use models to make predictions for your data. With the Ready-to-use models permissions, you can also chat with generative AI-powered models. The permissions are turned on by default when setting up your domain, or you can edit the permissions for a domain that you've already created. The Canvas Ready-to-use models permissions option adds the
Amazon SageMakerCanvasAIServicesAccess policy to your execution role. For more information, see the Get started section of the Ready-to-use models documentation.

For more information about getting started with generative AI foundation models, see Use generative AI with foundation models.

- **Fine-tune foundation models.** If you’d like to fine-tune foundation models in Canvas, you can either add the permissions when setting up your domain, or you can edit the permissions for the domain or user profile after creating your domain. You must add the AmazonSageMakerCanvasAIServicesAccess policy to the AWS IAM role you chose when setting up the user profile, and you must also add a trust relationship with Amazon Bedrock to the role. For instructions on how to add these permissions to your IAM role, see Grant Users Permissions to Fine-tune Foundation Models.

- **Time series forecasting.** If you’d like to perform forecasts on time series data, you can add time series forecasting permissions when setting up your domain, or you can edit the permissions for a domain or user profile after creating your domain. The required permissions are the AmazonSageMakerCanvasForecastAccess managed policy and a trust relationship with Amazon Forecast to the AWS IAM role you chose when setting up the user profile. For instructions on how to add these permissions to your IAM role, see Grant Your Users Permissions to Perform Time Series Forecasting.

- **Send batch predictions to Amazon QuickSight.** You might want to send batch predictions, or datasets of predictions you generate from a custom model, to Amazon QuickSight for analysis. In QuickSight, you can build and publish predictive dashboards with your prediction results. For instructions on how to add these permissions to your Canvas user’s IAM role, see Grant Your Users Permissions to Send Predictions to Amazon QuickSight.

- **Deploy Canvas models to a SageMaker endpoint.** SageMaker Hosting offers endpoints which you can use to deploy your model for use in production. You can deploy models built in Canvas to a SageMaker endpoint and then make predictions programmatically in a production environment. For more information, see Deploy your models to an endpoint.

- **Register model versions to the model registry.** You might want to register versions of your model to the SageMaker model registry, which is a repository for tracking the status of updated versions of your model. A data scientist or MLOps team working in the SageMaker model registry can view the versions of your model that you’ve built and approve or reject them. Then, they can deploy your model version to production or kick off an automated workflow. Model registration permissions are turned on by default for your domain. You can manage permissions at the user profile level and grant or remove permissions to specific users. For more information, see Register a model version in the SageMaker model registry.
• **Collaboration with data scientists.** If you want to collaborate with Studio Classic users and share models, you must add additional permissions to the AWS IAM role you chose when setting up the user profile. For instructions on how to add the policy to the role, see [Grant Users Permissions to Collaborate with Studio Classic](#).

• **Import data from Amazon Redshift.** If you want to import data from Amazon Redshift, you must give yourself additional permissions. You must add the `AmazonRedshiftFullAccess` managed policy to the AWS IAM role you chose when setting up the user profile. For instructions on how to add the policy to the role, see [Grant Users Permissions to Import Amazon Redshift Data](#).

**Note**

The necessary permissions to import through other data sources, such as Amazon Athena and SaaS platforms, are included in the `AmazonSageMakerFullAccess` and `AmazonSageMakerCanvasFullAccess` policies. If you followed the standard setup instructions, these policies should already be attached to your execution role. For more information about these data sources and their permissions, see [Connect to data sources](#).

**Step 1: Log in to SageMaker Canvas**

When the initial setup is complete, you can access SageMaker Canvas with any of the following methods, depending on your use case:

- In the [SageMaker console](https://console.aws.amazon.com), choose the **Canvas** in the left navigation pane. Then, on the **Canvas** page, select your user from the dropdown and launch the Canvas application.

- Open [SageMaker Studio](https://console.aws.amazon.com/studio/home), and in the Studio interface, go to the Canvas page and launch the Canvas application.

- Use your organization’s SAML 2.0-based SSO methods, such as Okta or the IAM Identity Center.

When you log into SageMaker Canvas for the first time, there is a welcome message with quick links to help you get started using the main features of Canvas.
Welcome to SageMaker Canvas
Build and use ML and generative AI models - no code required

If you close this message, you can revisit it at any time by choosing the Help button in the left navigation pane of the application.

Step 2: Use SageMaker Canvas to get predictions

After you’ve logged in to Canvas, you can start building models and generating predictions for your data.

You can either use Canvas Ready-to-use models to make predictions without building a model, or you can build a custom model for your specific business problem. Review the following information to decide whether Ready-to-use models or custom models are best for your use case.

- **Ready-to-use models.** With Ready-to-use models, you can use pre-built models to extract insights from your data. The Ready-to-use models cover a variety of use cases, such as language detection and document analysis. To get started making predictions with Ready-to-use models, see Use Ready-to-use models.

- **Custom models.** With custom models, you can build a variety of model types that are customized to make predictions for your data. Use custom models if you'd like to build a model that is trained on your business-specific data and if you'd like to use features such as collaborating with data scientists and evaluating your model's performance. To get started with building a custom model, see Use custom models.

You can also bring your own model (BYOM) from other features in SageMaker. An Amazon SageMaker Studio user can share their model with a Canvas user, and the Canvas user can generate predictions with the model. To learn more, see Bring your own model to SageMaker Canvas.

Setting Up and Managing Amazon SageMaker Canvas (for IT Administrators)

You can use the information in this section to help your users do the following:
• Optional: Grant your users permissions to upload their files locally.

• Set up Okta SSO for your users.

• Update SageMaker Canvas.

• Clean up or delete the installation of SageMaker Canvas.

• Optional: Set up Amazon Forecast so users can do time series forecasting.

• Optional: Set up an Amazon Virtual Private Cloud.

• Optional: Encrypt data using AWS Key Management Service.

• Optional: Grant your users permissions to import Amazon Redshift data.


**Topics**

• [Grant Your Users Permissions to Upload Local Files](#)

• [Set Up SageMaker Canvas for Your Users](#)

• [Configure your Amazon S3 storage](#)

• [Grant permissions for cross-account Amazon S3 storage](#)

• [Encrypt Your SageMaker Canvas Data with AWS KMS](#)

• [Grant Your Users Permissions to Build Custom Image and Text Prediction Models](#)

• [Grant Your Users Permissions to Perform Time Series Forecasting](#)

• [Grant Users Permissions to Fine-tune Foundation Models](#)

• [Update SageMaker Canvas for Your Users](#)

• [Request a Quota Increase](#)

• [Grant Users Permissions to Import Amazon Redshift Data](#)

• [Grant Users Permissions to Collaborate with Studio Classic](#)

• [Grant Your Users Permissions to Send Predictions to Amazon QuickSight](#)

• [Manage applications](#)

• [Configure Amazon SageMaker Canvas in a VPC without internet access](#)

• [Set up connections to data sources with OAuth](#)
Grant Your Users Permissions to Upload Local Files

If your users are uploading files from their local machines to SageMaker Canvas, you must attach a CORS (cross-origin resource sharing) configuration to the Amazon S3 bucket that they're using. When setting up the SageMaker domain or user profile, you can specify either a custom Amazon S3 location or the default location, which is a SageMaker created Amazon S3 bucket with a name that uses the following pattern: s3://sagemaker-{Region}-{your-account-id}. SageMaker Canvas adds your users' data to the bucket whenever they upload a file.

To grant users permissions to upload local files to the bucket, you can attach a CORS configuration to it using either of the following procedures. You can use the first method when setting up your domain or editing the existing domain settings, where you opt in to allow SageMaker to attach the CORS configuration to the bucket for you. The second method is the manual method, where you can attach the CORS configuration to the bucket yourself.

domain setup method

To grant your users permissions to upload local files, you can choose Enable Canvas permissions when setting up your domain. This attaches a Cross-Origin Resource Sharing (CORS) configuration to the Canvas storage configuration's Amazon S3 bucket and grants all users in the domain permission to upload local files into SageMaker Canvas. By default, the permissions option is turned on when you set up a domain, but you can turn off this option if you don't want to grant your users permission to upload files.

Nota

If you have an existing CORS configuration on the storage configuration Amazon S3 bucket, turning on Enable Canvas permissions overwrites the existing configuration with the new configuration.

The following procedure shows how you can turn on this option when doing a Quick setup for your domain in the console.

1. In the User profile section, enter a Name for the user.
2. Select an Execution role for the user.
3. Turn on Enable SageMaker Canvas permissions. (By default, this option is turned on.)
4. Finish setting up the domain.
If you are doing a **Standard setup** for your domain, then use the following procedure for the **Canvas settings** section to turn on local file upload.

1. For **Enable and configure Canvas permissions**, select **Local file upload**. (It's already checked by default.)
2. Choose **Next**.
3. Finish setting up the domain.

Your users can now upload local files into their SageMaker Canvas application.

You can also turn on or turn off local upload permissions for an existing domain by using the following procedure.

1. Go to the [Amazon SageMaker console](https://console.aws.amazon.com/sagemaker/).
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of **Domains**, choose your domain.
5. On the **Domain settings** page, choose the **Domain settings**, tab.
6. Choose **Edit**.
7. In the navigation pane, choose **Canvas settings**.
8. Select or deselect **Enable local file upload**.
9. Finish any other modifications you want to make to the domain, and then choose **Submit** to submit your changes.

**Amazon S3 bucket method**

If you want to manually attach the CORS configuration to the SageMaker Amazon S3 bucket, use the following procedure.

2. Choose your bucket. If your domain uses the default SageMaker created bucket, the bucket’s name uses the following pattern: s3://sagemaker-{Region}-{your-account-id}.
3. Choose **Permissions**.
4. Navigate to **Cross-origins resource sharing (CORS)**.
5. Choose **Edit**.

6. Add the following CORS policy:

```json
[
  {
    "AllowedHeaders": [
      "*"
    ],
    "AllowedMethods": [
      "POST"
    ],
    "AllowedOrigins": [
      "*"
    ],
    "ExposeHeaders": []
  }
]
```

7. Choose **Save changes**.

In the preceding procedure, the CORS policy must have "POST" listed under `AllowedMethods`.

After you've gone through the procedure, you should have:

- An IAM role assigned to each of your users.
- Amazon SageMaker Studio Classic runtime permissions for each of your users. SageMaker Canvas uses Studio Classic to run the commands from your users.
- If the users are uploading files from their local machines, a CORS policy attached to their Amazon S3 bucket.

If your users still can't upload the local files after you update the CORS policy, the browser might be caching the CORS settings from a previous upload attempt. If they're running into issues, instruct them to clear their browser cache and try again.

**Set Up SageMaker Canvas for Your Users**

To set up Amazon SageMaker Canvas, do the following:
• Create an Amazon SageMaker domain.
• Create user profiles for the domain
• Set up Okta Single Sign On (Okta SSO) for your users.
• Activate link sharing for models.

Use Okta Single-Sign On (Okta SSO) to grant your users access to Amazon SageMaker Canvas. SageMaker Canvas supports SAML 2.0 SSO methods. The following sections guide you through procedures to set up Okta SSO.

To set up a domain, see [Onboard to Amazon SageMaker Runtime Studio Using IAM](#). You can use the following information to help you complete the procedure in the section:

• You can ignore the step about creating projects.
• You don't need to provide access to additional Amazon S3 buckets. Your users can use the default bucket that we provide when we create a role.
• To grant your users access to share their notebooks with data scientists, turn on Notebook Sharing Configuration.
• Use Amazon SageMaker Studio Classic version 3.19.0 or later. For information about updating Amazon SageMaker Studio Classic, see [Shut down and Update SageMaker Studio Classic](#).

Use the following procedure to set up Okta. For all of the following procedures, you specify the same IAM role for `IAM-role`.

**Add the SageMaker Canvas application to Okta**

Set up the sign-on method for Okta.

1. Sign in to the Okta Admin dashboard.
2. Choose Add application. Search for AWS Account Federation.
3. Choose Add.
4. Optional: Change the name to Amazon SageMaker Canvas.
5. Choose Next.
6. Choose SAML 2.0 as the Sign-On method.
7. Choose Identity Provider Metadata to open the metadata XML file. Save the file locally.
8. Choose Done.

Set up ID federation in IAM

AWS Identity and Access Management (IAM) is the AWS service that you use to gain access to your AWS account. You gain access to AWS through an IAM account.

1. Sign in to the AWS console.
2. Choose AWS Identity and Access Management (IAM).
3. Choose Identity Providers.
4. Choose Create Provider.
5. For Configure Provider, specify the following:
   - Provider Type – From the dropdown list, choose SAML.
   - Provider Name – Specify Okta.
   - Metadata Document – Upload the XML document that you've saved locally from step 7 of Add the SageMaker Canvas application to Okta.
7. For Roles, choose the IAM role that you're using for Okta SSO access.
8. Under Trust Relationship for the IAM role, choose Edit Trust Relationship.
9. Modify the IAM trust relationship policy by specifying the Provider ARN value that you've copied and add the following policy:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Federated": "arn:aws:iam::123456789012:saml-provider/Okta"
            },
            "Action": [
                "sts:AssumeRoleWithSAML",
                "sts:SetSourceIdentity",
                "sts:TagSession"
            ]
        }]
}
```
10. For **Permissions**, add the following policy:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "AmazonSageMakerPresignedUrlPolicy",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreatePresignedDomainUrl",
                "sagemaker:CreatePresignedDomainUrlWithPrincipalTag"
            ],
            "Resource": "*"
        }
    ]
}
```

---

**Configure SageMaker Canvas in Okta**

Configure Amazon SageMaker Canvas in Okta using the following procedure.

To configure Amazon SageMaker Canvas to use Okta, follow the steps in this section. You must specify unique user names for each **SageMakerStudioProfileName** field. For example, you can use `user.login` as a value. If the username is different from the SageMaker Canvas profile name, choose a different uniquely identifying attribute. For example, you can use an employee's ID number for the profile name.

For an example of values that you can set for **Attributes**, see the code following the procedure.

1. **Under Directory**, choose **Groups**.
2. Add a group with the following pattern: sagemaker#canvas#IAM-role#AWS-account-id.

3. In Okta, open the AWS Account Federation application integration configuration.

4. Select Sign On for the AWS Account Federation application.

5. Choose Edit and specify the following:
   - SAML 2.0
   - Default Relay State – https://Region.conso...StudioId. You can find the Studio Classic ID in the console: https://console.aws.amazon.com/sagemaker/

6. Choose Attributes.

7. In the SageMakerStudioProfileName fields, specify unique values for each username. The usernames must match the usernames that you've created in the AWS console.

   Attribute 1:
   Name: https://aws.amazon.com/SAML/Attributes/
   PrincipalTag:SageMakerStudioUserProfileName
   Value: ${user.login}

   Attribute 2:
   Name: https://aws.amazon.com/SAML/Attributes/TransitiveTagKeys
   Value: {"SageMakerStudioUserProfileName"}

8. Select Environment Type. Choose Regular AWS.
   - If your environment type isn't listed, you can set your ACS URL in the ACS URL field. If your environment type is listed, you don’t need to enter your ACS URL

9. For Identity Provider ARN, specify the ARN you used in step 6 of the preceding procedure.

10. Specify a Session Duration.

11. Choose Join all roles.

12. Turn on Use Group Mapping by specifying the following fields:
   - App Filter – okta
   - Group Filter – ^aws\#S+\#(?IAM-role[\w\-]+)\#(?accountid\d+)$
13. Choose **Save/Next**.

14. Under **Assignments**, assign the application to the group that you've created.

### Add optional policies on access control in IAM

In IAM, you can apply the following policy to the administrator user who creates the user profiles.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "CreateSageMakerStudioUserProfilePolicy",
            "Effect": "Allow",
            "Action": "sagemaker:CreateUserProfile",
            "Resource": "/",
            "Condition": {
                "ForAnyValue:StringEquals": {
                    "aws:TagKeys": [
                        "studiouserid"
                    ]
                }
            }
        }
    ]
}
```

If you choose to add the preceding policy to the admin user, you must use the following permissions from **Set up ID federation in IAM**.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "AmazonSageMakerPresignedUrlPolicy",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreatePresignedDomainUrl",
                "sagemaker:CreatePresignedDomainUrlWithPrincipalTag"
            ],
        }
    ]
}
```
Configure your Amazon S3 storage

When you set up your SageMaker Canvas application, the default storage location for model artifacts, datasets, and other application data is an Amazon S3 bucket that Canvas creates. This default Amazon S3 bucket follows the naming pattern s3://sagemaker-{Region}-{your-account-id} and exists in the same Region as your Canvas application.

However, you can customize the storage location and specify your own Amazon S3 bucket for storing Canvas application data. You might want to use your own Amazon S3 bucket for storing application data for any of the following reasons:

- Your organization has internal naming conventions for Amazon S3 buckets.
- You want to enable cross-account access to model artifacts or other Canvas data.
- You want to be compliant with internal security guidelines, such as restricting users to specific Amazon S3 buckets or model artifacts.
- You want enhanced visibility and access to logs produced by Canvas, independent of the AWS console or SageMaker Studio Classic.

By specifying your own Amazon S3 bucket, you can have increased control over your own storage and be compliant with your organization.

To get started, you can either create a new SageMaker domain or user profile, or you can update an existing domain or user profile. Note that the user profile settings override the domain-level settings. For example, you can use the default bucket configuration at the domain level, but you can specify a custom Amazon S3 bucket for an individual user. After specifying your own Amazon S3 bucket for the domain or user profile, Canvas creates a subfolder called Canvas/
<UserProfileName> under the input Amazon S3 URI and saves all artifacts generated in the Canvas application under this subfolder.

⚠️ Important

If you update an existing domain or user profile, you no longer have access to your Canvas artifacts from the previous location. Your files are still in the old Amazon S3 location, but you can no longer view them from Canvas. The new configuration takes effect the next time you log into the application.

For more information about granting cross-account access to your Amazon S3 bucket, see Granting cross-account object permissions in the Amazon S3 User Guide.

The following sections describe how to specify a custom Amazon S3 bucket for your Canvas storage configuration. If you’re setting up a new SageMaker domain (or a new user in a domain), then use the New domain setup method or the New user profile setup method. If you have an existing Canvas user profile and would like to update the profile's storage configuration, use the Existing user method.

Before you begin

If you’re specifying an Amazon S3 URI from a different AWS account, or if you’re using a bucket that is encrypted with AWS KMS, then you must configure permissions before proceeding. You must grant AWS IAM permissions to ensure that Canvas can download and upload objects to and from your bucket. For detailed information on how to grant the required permissions, see Grant permissions for cross-account Amazon S3 storage.

Additionally, the final Amazon S3 URI for the training folder in your Canvas storage location must be 128 characters or less. The final Amazon S3 URI consists of your bucket path s3://<your-bucket-name>/<folder-name>/ plus the path that Canvas adds to your bucket: Canvas/<user-profile-name>/Training. For example, an acceptable path that is less than 128 characters is s3://<my-bucket>/<machine-learning>/Canvas/<user-1>/Training.

New domain setup method

If you’re setting up a new domain and Canvas application, use this section to configure the storage location at the domain level. This configuration applies to all new users you create in the domain, unless you specify a different storage location for individual user profiles.
When doing a **Standard setup** for your domain, use the following procedure for the **Canvas settings** section:

1. For the **Canvas storage configuration**, do the following:
   
   a. Select **System managed** if you want to set the location to the default SageMaker bucket that follows the pattern `s3://sagemaker-{Region}-{your-account-id}`.
   
   b. Select **Custom S3** to specify your own Amazon S3 bucket as the storage location. Then, enter the Amazon S3 URI.
   
   c. (Optional) For **Encryption key**, specify a KMS key for encrypting Canvas artifacts stored at the specified location.

2. Finish setting up the domain and choose **Submit**.

Your domain is now configured to use the Amazon S3 location you specified for SageMaker Canvas application storage.

**New user profile setup method**

If you're setting up a new user profile in your domain, use this section to configure the storage location for the user. This configuration overrides the domain-level configuration.

When adding a user profile to your domain, use the following procedure for the **Canvas settings** section:

1. For the **Canvas storage configuration**, do the following:
   
   a. Select **System managed** if you want to set the location to the default SageMaker created bucket that follows the pattern `s3://sagemaker-{Region}-{your-account-id}`.
   
   b. Select **Custom S3** to specify your own Amazon S3 bucket as the storage location. Then, enter the Amazon S3 URI.
   
   c. (Optional) For **Encryption key**, specify a KMS key for encrypting Canvas artifacts stored at the specified location.

2. Finish setting up the user profile and choose **Submit**.

Your user profile is now configured to use the Amazon S3 location you specified for SageMaker Canvas application storage.
Existing user method

If you have an existing Canvas user profile and would like to update the Amazon S3 storage location, you can edit the SageMaker domain or user profile settings. The change takes effect the next time you log into the Canvas application.

Note

When you change the storage location for an existing Canvas application, you lose access to your Canvas artifacts from the previous storage location. The artifacts are still stored in the old Amazon S3 location, but you can no longer view them from Canvas.

Remember that the user profile settings override the general domain settings, so you can update the Amazon S3 storage location for specific user profiles without changing it for all of the users. You can update the storage configuration for an existing domain or user by using the following procedures.

Update an existing domain

Use the following procedure to update the storage configuration for a domain.

2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, choose your domain.
5. On the domain settings page, choose the domain settings tab.
6. Choose Edit.
7. In the navigation pane, choose Canvas settings.
8. For the Canvas storage configuration, do the following:
   a. Select System managed if you want to set the location to the default SageMaker created bucket that follows the pattern s3://sagemaker-{Region}-{your-account-id}.
   b. Select Custom S3 to specify your own Amazon S3 bucket as the storage location. Then, enter the Amazon S3 URI.
Update an existing user profile

Use the following procedure to update the storage configuration for a user profile.

2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, choose your domain.
5. From the list of users in the domain, choose the user whose configuration you want to edit.
6. On the User Details page, choose Edit.
7. In the navigation pane, choose Canvas settings.
8. For the Canvas storage configuration, do the following:
   a. Select System managed if you want to set the location to the default SageMaker bucket that follows the pattern s3://sagemaker-{Region}-{your-account-id}.
   b. Select Custom S3 to specify your own Amazon S3 bucket as the storage location. Then, enter the Amazon S3 URI.
   c. (Optional) For Encryption key, specify a KMS key for encrypting Canvas artifacts stored at the specified location.
9. Finish any other modifications you want to make to the user profile, and then choose Submit to save your changes.

The storage location for your Canvas user profile should now be updated. The next time you log into the Canvas application, you receive a notification that the storage location has been updated. You lose access to any previous artifacts that you created in Canvas. You can still access the files in Amazon S3, but you can no longer view them in Canvas.
Grant permissions for cross-account Amazon S3 storage

When setting up your SageMaker domain or user profile for users to access SageMaker Canvas, you specify an Amazon S3 storage location for Canvas artifacts. These artifacts include saved copies of your input datasets, model artifacts, predictions, and other application data. You can either use the default SageMaker created Amazon S3 bucket, or you can customize the storage location and specify your own bucket for storing Canvas application data.

You can specify an Amazon S3 bucket in another AWS account for storing your Canvas data, but first you must grant cross-account permissions so that Canvas can access the bucket.

The following sections describe how to grant permissions to Canvas for uploading and downloading objects to and from an Amazon S3 bucket in another account. There are additional permissions for when your bucket is encrypted with AWS KMS.

Requirements

Before you begin, review the following requirements:

- Cross-account Amazon S3 buckets (and any associated AWS KMS keys) must be in the same AWS Region as the Canvas user domain or user profile.
- The final Amazon S3 URI for the training folder in your Canvas storage location must be 128 characters or less. The final S3 URI consists of your bucket path s3://<your-bucket-name>/<folder-name>/ plus the path that Canvas adds to your bucket: Canvas/<user-profile-name>/Training. For example, an acceptable path that is less than 128 characters is s3://<my-bucket>/<machine-learning>/Canvas/<user-1>/Training.

Permissions for cross-account Amazon S3 buckets

The following section outlines the basic steps for granting the necessary permissions so that Canvas can access your Amazon S3 bucket in another account. For more detailed instructions, see Example 2: Bucket owner granting cross-account bucket permissions in the Amazon S3 User Guide.

1. Create an Amazon S3 bucket, bucketA, in Account A.
2. The Canvas user exists in another account called Account B. In the following steps, we refer to the Canvas user's IAM role as roleB in Account B.

   Give the IAM role roleB in Account B permission to download (GetObject) and upload (PutObject) objects to and from bucketA in Account A by attaching an IAM policy.
To limit access to a specific bucket folder, define the folder name in the resource element, such as `arn:aws:s3::<bucketA>/FolderName/*`. For more information, see [How can I use IAM policies to grant user-specific access to specific folders?](#).

**Note**

Bucket-level actions, such as `GetBucketCors` and `GetBucketLocation`, should be added on bucket-level resources, not folders.

The following example IAM policy grants the required permissions for `roleB` to access objects in `bucketA`:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": ["s3:GetObject", "s3:PutObject", "s3:DeleteObject"],
      "Resource": ["arn:aws:s3::<bucketA>/FolderName/*", ]
    },
    {
      "Effect": "Allow",
      "Action": ["s3:ListBucket", "s3:GetBucketCors", "s3:GetBucketLocation"],
      "Resource": ["arn:aws:s3::<bucketA>", ]
    }
  ]
}
```
3. Configure the bucket policy for bucketA in Account A to grant permissions to the IAM role roleB in Account B.

⚠️ Note
Admins must also turn off **Block all public access** under the bucket **Permissions** section.

The following is an example bucket policy for bucketA to grant the necessary permissions to roleB:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::accountB:role/roleB"
      },
      "Action": ["s3:DeleteObject", "s3:GetObject", "s3:PutObject"],
      "Resource": "arn:aws:s3:::bucketA/FolderName/*"
    },
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::accountB:role/roleB"
      },
      "Action": ["s3:ListBucket", "s3:GetBucketCors", "s3:GetBucketLocation"],
      "Resource": "arn:aws:s3:::bucketA"
    }
  ]
}
```
After configuring the preceding permissions, your Canvas user profile in Account B can now use the Amazon S3 bucket in Account A as the storage location for Canvas artifacts.

Permissions for cross-account Amazon S3 buckets encrypted with AWS KMS

The following procedure shows you how to grant the necessary permissions so that Canvas can access your Amazon S3 bucket in another account that is encrypted with AWS KMS. The steps are similar to the procedure above, but with additional permissions. For more information about granting cross-account KMS key access, see Allowing users in other accounts to use a KMS key in the AWS KMS Developer Guide.

1. Create an Amazon S3 bucket, bucketA, and an Amazon S3 KMS key s3KmsInAccountA in Account A.

2. The Canvas user exists in another account called Account B. In the following steps, we refer to the Canvas user's IAM role as roleB in Account B.

Give the IAM role roleB in Account B permission to do the following:

- Download (GetObject) and upload (PutObject) objects to and from bucketA in Account A.
- Access the AWS KMS key s3KmsInAccountA in Account A.

The following example IAM policy grants the required permissions for roleB to access objects in bucketA and use the KMS key s3KmsInAccountA:

```json
{
    "Version": "2012-10-17",
    "Statement": [ 
        {
            "Effect": "Allow",
            "Action": [ 
                "s3:GetObject",
                "s3:PutObject",
                "s3:DeleteObject"
            ],
            "Resource": [ 
                "arn:aws:s3:::bucketA/FolderName/*"
            ]
        },
        {
            "Effect": "Allow",
            "Action": [ 
                "kms:Encrypt",
                "kms:Decrypt",
                "kms:ReEncrypt*,
                "kms:GenerateDataKey", 
                "kms:ListKeyVersions",
                "kms:DescribeKey",
                "kms:CreateGrant",
                "kms:TagResource",
                "kms:UntagResource"
            ],
            "Resource": [ 
                "arn:aws:kms:region:account-id:key/s3KmsInAccountA"
            ]
        }
    ]
}
```
3. Configure the bucket policy for bucketA and the key policy for s3KmsInAccountA in Account A to grant permissions to the IAM role roleB in Account B.

The following is an example bucket policy for bucketA to grant the necessary permissions to roleB:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::accountB:role/roleB"
            },
            "Action": [
                "s3:DeleteObject",
                "s3:GetObject",
                "s3:PutObject"
            ]
        }
    ]
}
```
The following example is a key policy that you attach to the KMS key s3KmsInAccountA in Account A to grant roleB access. For more information about how to create and attach a key policy statement, see Creating a key policy in the AWS KMS Developer Guide.

```json
{
    "Sid": "Allow use of the key",
    "Effect": "Allow",
    "Principal": {
        "AWS": [ "arn:aws:iam::accountB:role/roleB"
    ],
    "Action": [ "kms:DescribeKey", "kms:CreateGrant", "kms:RetireGrant", "kms:GenerateDataKey", "kms:GenerateDataKeyWithoutPlainText", "kms:Decrypt"
    ],
    "Resource": "*"
}
```

After configuring the preceding permissions, your Canvas user profile in Account B can now use the encrypted Amazon S3 bucket in Account A as the storage location for Canvas artifacts.
Encrypt Your SageMaker Canvas Data with AWS KMS

You might have data that you want to encrypt while using Amazon SageMaker Canvas, such as your private company information or customer data. SageMaker Canvas uses AWS Key Management Service to protect your data. AWS KMS is a service that you can use to create and manage cryptographic keys for encrypting your data. For more information about AWS KMS, see AWS Key Management Service in the AWS KMS Developer Guide.

Amazon SageMaker Canvas provides you with several options for encrypting your data. SageMaker Canvas provides default encryption within the application for tasks such as building your model and generating insights. You can also choose to encrypt data stored in Amazon S3 to protect your data at rest. SageMaker Canvas supports importing encrypted datasets, so you can establish an encrypted workflow. The following sections describe how you can use AWS KMS encryption to protect your data while building models with SageMaker Canvas.

Encrypt your data in SageMaker Canvas

With SageMaker Canvas, you can use two different AWS KMS encryption keys to encrypt your data in SageMaker Canvas, which you can specify when setting up your domain. These two keys can be the same or different. SageMaker Canvas uses one key for temporary application storage, visualizations, or compute purposes (such as building models). You can use either the default AWS managed key or specify your own. You can also specify an optional key that SageMaker Canvas uses for long-term storage of model objects and datasets, which are stored in the Region's default SageMaker S3 bucket for your account.

Prerequisites

To use your own KMS key for either of the previously described purposes, you must first grant your user's IAM role permission to use the key. Then, you can specify the KMS key when setting up your domain.

The simplest way to grant your role permission to use the key is to modify the key policy. Use the following procedure to grant your role the necessary permissions.

1. Open the AWS KMS console.
2. In the Key Policy section, choose Switch to policy view.
3. Modify the key's policy to grant permissions for the kms:GenerateDataKey and kms:Decrypt actions to the IAM role. You can add a statement that's similar to the following:
4. Choose **Save changes**.

The less preferred method is to modify the user’s IAM role to grant the user permissions to use or manage the KMS key. If you use this method, the KMS key policy must also allow access management through IAM. To learn how to grant permission to a KMS key through the user’s IAM role, see [Specifying KMS keys in IAM policy statements](https://docs.aws.amazon.com/kms/latest/developerguide/specifying-kms-keys-in-iam-policy-statements.html) in the *AWS KMS Developer Guide*.

**Prerequisites for time series forecasting**

To use your AWS KMS key to encrypt time series forecasting models in SageMaker Canvas, you must modify the key policy for the KMS key used to store objects to Amazon S3. Your key policy must grant permissions to the `AmazonSageMakerCanvasForecastRole`, which SageMaker creates when you [grant time series forecasting permissions for your users](https://docs.aws.amazon.com/forecast/latest/tutorials/iam-permissions.html). Amazon Forecast uses the `AmazonSageMakerCanvasForecastRole` to perform time series forecasting operations in SageMaker Canvas. Your KMS key must grant permissions to this role in order to ensure data is encrypted for time series forecasting.

To modify the permissions of your KMS key policy to allow encrypted time series forecasting, do the following.

1. Open the [AWS KMS console](https://console.aws.amazon.com/kms/home).
2. In the **Key Policy** section, choose **Switch to policy view**.
3. Modify the key's policy to have the permissions specified in the following example:

   ```json
   {
     "Sid": "Enable IAM Permissions for Amazon Forecast KMS access",
   }
   ```
4. Choose **Save changes**.

You can now use your KMS key to encrypt time series forecasting operations in SageMaker Canvas.

**Note**

The following permissions are only required if you are using the [IAM role setup method](#) to configure time series forecasting. Add the following permissions policy to your user's IAM role. You must also update the key policy with updated policies required for Amazon Forecast. For more information about the permissions required for time series forecasting, see [Grant Your Users Permissions to Perform Time Series Forecasting](#).

```json
{
   "Sid": "Enable IAM Permissions for Amazon Forecast KMS access",
   "Effect": "Allow",
   "Principal": {
      "AWS": "<arn:aws:iam::111122223333:role/AmazonSageMaker-111122223333>"
   },
   "Action": [ 
      "kms:Decrypt",
      "kms:DescribeKey",
      "kms:CreateGrant",
      "kms:RetireGrant",
      "kms:GenerateDataKey",
      "kms:GenerateDataKeyWithout Plaintext",
      "kms:Encrypt"
   ],
   "Resource": "*"
}
```
Encrypt your data in the SageMaker Canvas application

The first KMS key you can use in SageMaker Canvas is used for encrypting application data stored on Amazon Elastic Block Store (Amazon EBS) volumes and in the Amazon Elastic File System that SageMaker creates in your domain. SageMaker Canvas encrypts your data with this key in the underlying application and temporary storage systems created when using compute instances for building models and generating insights. SageMaker Canvas passes the key to other AWS services, such as Autopilot, whenever SageMaker Canvas initiates jobs with them to process your data.

You can specify this key by setting the KmsKeyId in the CreateDomain API call or while doing the Standard domain setup in the console. If you don't specify your own KMS key, SageMaker uses a default AWS managed KMS key to encrypt your data in the SageMaker Canvas application.

To specify your own KMS key for use in the SageMaker Canvas application through the console, first set up your Amazon SageMaker domain using the Standard setup. Use the following procedure to complete the Network and Storage Section for the domain.

1. Fill out your desired Amazon VPC settings.
2. For Encryption key, choose Enter a KMS key ARN.
3. For KMS ARN, enter the ARN for your KMS key, which should have a format similar to the following: arn:aws:kms:example-region-1:123456789098:key/111aa2bb-333c-4d44-5555-a111bb2c33dd

Encrypt your SageMaker Canvas data saved in Amazon S3

The second KMS key you can specify is used for data that SageMaker Canvas stores to Amazon S3. SageMaker Canvas saves duplicates of your input datasets, application and model data, and output data to the Region's default SageMaker S3 bucket for your account. The naming pattern for this bucket is s3://sagemaker-{Region}-{your-account-id}, and SageMaker Canvas stores data in the Canvas/ folder.

1. Turn on Enable notebook resource sharing.
2. For **S3 location for shareable notebook resources**, leave the default Amazon S3 path. Note that SageMaker Canvas does not use this Amazon S3 path; this Amazon S3 path is used for Studio Classic notebooks.

3. For **Encryption key**, choose **Enter a KMS key ARN**.

4. For **KMS ARN**, enter the ARN for your KMS key, which should have a format similar to the following: `arn:aws:kms:us-east-1:111122223333:key/111aa2bb-333c-4d44-5555-a111bb2c33dd`

**Import encrypted datasets from Amazon S3**

Your users might have datasets that have been encrypted with a KMS key. While the preceding section shows you how to encrypt data in SageMaker Canvas and data stored to Amazon S3, you must grant your user's IAM role additional permissions if you want to import data from Amazon S3 that is already encrypted with AWS KMS.

To grant your user permissions to import encrypted datasets from Amazon S3 into SageMaker Canvas, add the following permissions to the IAM execution role that you've used for the user profile:

```
"kms:Decrypt",
"kms:GenerateDataKey"
```

To learn how to edit the IAM permissions for a role, see [Adding and removing IAM identity permissions](#) in the *IAM User Guide*. For more information about KMS keys, see [Key policies in AWS Key Management Service](#) in the *AWS KMS Developer Guide*.

**FAQs**

Refer to the following FAQ items for answers to commonly asked questions about SageMaker Canvas AWS KMS support.

**Q: Does SageMaker Canvas retain my KMS key?**

A: No. SageMaker Canvas may temporarily cache your key or pass it on to other AWS services (such as Autopilot), but SageMaker Canvas does not retain your KMS key.
Q: I specified a KMS key when setting up my domain. Why did my dataset fail to import in SageMaker Canvas?

A: Your user's IAM role may not have permissions to use that KMS key. To grant your user permissions, see the Prerequisites. Another possible error is that you have a bucket policy on your Amazon S3 bucket that requires the use of a specific KMS key that doesn’t match the KMS key you specified in your domain. Make sure that you specify the same KMS key for your Amazon S3 bucket and your domain.

Q: How do I find the Region's default SageMaker Amazon S3 bucket for my account?

A: The default Amazon S3 bucket follows the naming pattern s3://sagemaker-{Region}-{your-account-id}. The Canvas/ folder in this bucket stores your SageMaker Canvas application data.

Q: Can I change the default SageMaker Amazon S3 bucket used to store SageMaker Canvas data?

A: No, SageMaker creates this bucket for you.

Q: What does SageMaker Canvas store in the default SageMaker Amazon S3 bucket?

A: SageMaker Canvas uses the default SageMaker Amazon S3 bucket to store duplicates of your input datasets, model artifacts, and model outputs.

Q: What use cases are supported for using KMS keys with SageMaker Canvas?

A: With SageMaker Canvas, you can use your own encryption keys with AWS KMS for building regression, binary and multi-class classification, and time series forecasting models, as well as for batch inference with your model.

Q: Can I encrypt time series forecasting models in SageMaker Canvas?

A: Yes. You must give your KMS key additional permissions in order to perform encrypted time series forecasting. For more information about how to modify your key's policy in order to grant time series forecasting permissions, see Prerequisites for time series forecasting.

Grant Your Users Permissions to Build Custom Image and Text Prediction Models

In Amazon SageMaker Canvas, you can build custom models to meet your specific business need. Two of these custom model types are single-label image prediction and multi-category text
prediction. The permissions to build these model types are included in the AWS Identity and Access Management (IAM) policy called AmazonSageMakerCanvasFullAccess, which SageMaker attaches by default to your user's IAM execution role if you leave the Canvas base permissions turned on.

However, if you are using a custom IAM configuration, then you must explicitly add permissions to your user's IAM execution role so that they can build custom image and text prediction model types. To grant the necessary permissions to build image and text prediction models, read the following section to learn how to attach a least-permissions policy to your role.

To add the permissions to the user's IAM role, do the following:

1. Go to the IAM console.
2. Choose Roles.
3. In the search box, search for the user's IAM role by name and select it.
4. On the page for the user's role, under Permissions, choose Add permissions.
5. Choose Create inline policy.
6. Select the JSON tab, and then paste the following least-permissions policy into the editor.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "sagemaker:CreateAutoMLJobV2",
            "sagemaker:DescribeAutoMLJobV2"
         ],
         "Resource": "*"
      }
   ]
}
```
7. Choose Review policy.
8. Enter a Name for the policy.
9. Choose Create policy.

For more information about AWS managed policies, see Managed policies and inline policies in the IAM User Guide.
Grant Your Users Permissions to Perform Time Series Forecasting

In order to perform time series forecasts in Amazon SageMaker Canvas, your users must have the necessary permissions. The preferred method to give your users these permissions is to turn on the time series forecasting option when setting up the Amazon SageMaker domain, or when editing the settings for a specific user profile. You can also use the manual method of attaching a policy and trust relationship for Amazon Forecast to the AWS Identity and Access Management (IAM) role.

If you want to encrypt your time series forecasts with your own key, you must use an AWS KMS key and modify your KMS key's policy to grant permissions to the role used by Amazon Forecast. For more information about setting up your KMS key and modifying the policy for time series forecasting, see Prerequisites for time series forecasting.

domain setup method

SageMaker provides you with the option to grant time series forecasting permissions to users through the domain settings. You can toggle the permissions for all of the users in your domain, and SageMaker manages attaching the required IAM policy and trust relationship for you.

If you are setting up your Amazon SageMaker domain for the first time and want to turn on time series forecasting permissions for all users in the domain, then use the following procedures.

Quick setup

Use the following procedure to turn on SageMaker Canvas time series forecasting permissions when doing a Quick setup for your domain.

1. In the Amazon SageMaker domain Quick setup, fill out the Name and Default execution role fields in the User profile section.
2. Leave the Enable SageMaker Canvas permissions option turned on. It is turned on by default.
3. Choose Submit to finish setting up your domain.

Standard setup

Use the following procedure to turn on SageMaker Canvas time series forecasting permissions when doing a Standard setup for your domain.

1. In the Amazon SageMaker domain Standard setup, fill out the General settings, Studio settings, and RStudio settings pages.
2. Choose the **Canvas settings** page.

3. For the **Canvas base permissions configuration**, leave the **Enable Canvas base permissions** option turned on. It is turned on by default. These permissions are required in order to turn on time series forecasting permissions.

4. For the **Time series forecasting configuration**, leave the **Enable time series forecasting** option turned on. It is turned on by default.

5. Select **Create and use a new execution role**, or select **Use an existing execution role** if you already have an IAM role with the required Amazon Forecast permissions attached. For more information, see the [IAM role setup method](#).

6. Finish making any other changes to your domain setup, and then choose **Submit**.

Your users should now have the necessary permissions to perform time series forecasting in SageMaker Canvas.

**User setup method**

You can configure time series forecasting permissions for individual users in an existing domain. The user profile settings override the general domain settings, so you can grant permissions to specific users without giving permissions to all of your users. To grant time series forecasting permissions to a specific user that doesn't already have permissions, use the following procedure.


2. On the left navigation pane, choose **Admin configurations**.

3. Under **Admin configurations**, choose **domains**.

4. On the **domains** page, choose your domain.

5. In the **User profiles** tab, select the name of the user whose permissions you want to edit.

6. On the **User Details** page, choose **Edit**.

7. Choose the **Canvas settings** page.

8. Turn on **Enable Canvas base permissions**. These permissions are required in order to turn on time series forecasting permissions.

9. Turn on the **Enable time series forecasting** option.

10. If you want to use a different execution role for the user than the role specified in the domain, select **Create and use a new execution role**, or **Use an existing execution role** if you already have an IAM role ready to use.
Note

If you want to use an existing IAM role, make sure that it has the IAM policy `AmazonSageMakerCanvasForecastAccess` attached and has a trust relationship that establishes Amazon Forecast as a service principal. For more information, see the section IAM role setup method.

11. The Canvas settings page should look like the following screenshot. Finish making any other changes to your user profile, and then choose Submit to save your changes.
Your user should now have permission to do time series forecasting in SageMaker Canvas.

You can also remove your user's permissions by using the preceding procedure and turning off the **Enable time series forecasting** option.

**IAM role setup method**

You can manually grant your users permissions to perform time series forecasting in Amazon SageMaker Canvas by adding additional permissions to the AWS Identity and Access Management (IAM) role specified for the user's profile. The IAM role must have a trust relationship with Amazon Forecast and an attached policy that gives permissions to Forecast.

The following section shows you how to create the trust relationship and attach the **AmazonSageMakerCanvasForecastAccess** managed policy to your IAM role, which grants the minimum permissions necessary for time series forecasting to work in SageMaker Canvas.

**Note**

The AmazonSageMakerCanvasForecastAccess policy grants permissions to access the SageMaker created Amazon S3 bucket, which is the default storage location for Canvas application data. If you've specified a custom Amazon S3 storage location for Canvas application data, you must update the permissions in the policy to your own Amazon S3 bucket. For more information about custom Amazon S3 storage locations for Canvas, see [Configure your Amazon S3 storage](#).

To configure an IAM role with the manual method, use the following procedure.

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. On the **Domains** page, choose your domain.
5. From the list of **User profiles**, select the profile of the user you to whom want to grant time series forecasting permissions.
6. Under **Details**, copy or make a note of the name of the user's **Execution role**. The name of the IAM role should be similar to the following: 111122223333.
7. Once you have the name of the user's IAM role, go to the IAM console.
8. Choose Roles.
9. Search for the user's IAM role by name from the list of roles and select it.
10. Under Permissions, choose Add permissions.
11. Choose Attach policies.
12. Search for the AmazonSageMakerCanvasForecastAccess managed policy and select it. Choose Attach policies to attach the policy to the role.

   After attaching the policy, the role's Permissions section should now include AmazonSageMakerCanvasForecastAccess.
13. Return to the IAM role's page, and under Trust relationships, choose Edit trust policy.
14. In the Edit trust policy editor, update the trust policy to add Forecast as a service principal. The policy should look like the following example.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
```
15. After editing the trust policy, choose **Update policy**.

You should now have an IAM role that has the policy `AmazonSageMakerCanvasForecastAccess` attached to it and a trust relationship established with Amazon Forecast, giving users permission to perform time series forecasting in SageMaker Canvas. For information about AWS managed policies, see [Managed policies and inline policies](#).

**Note**

If you use this method to set up time series forecasting and want to use AWS KMS encryption for your forecasts, then you must configure your KMS key’s policy to grant additional permissions. For more information, see [Prerequisites for time series forecasting](#).

---

**Grant Users Permissions to Fine-tune Foundation Models**

In order to grant permissions for fine-tuning foundation models in Amazon SageMaker Canvas, you must complete the permissions setup described on this page. You must grant the user permissions for **Ready-to-use models**, which attaches the `AmazonSageMakerCanvasAIServicesAccess` policy to your user’s AWS IAM execution role. You must also create a trust relationship between Amazon Bedrock and the IAM execution role.

**Grant Ready-to-use models permissions**

The permissions you need for fine-tuning foundation models are included in the Canvas Ready-to-use models permissions. You must turn on the **Canvas Ready-to-use models configuration** permissions when setting up your Amazon SageMaker domain. For more information, see [Prerequisites for setting up Amazon SageMaker Canvas](#).
You can edit your domain or user profile settings to turn on the **Canvas Ready-to-use models configuration** setting. For more information on editing your domain settings, see [View and Edit domains](#).

You can also manually attach the **AmazonSageMakerCanvasAlServicesAccess** policy to the user’s IAM execution role through the IAM console. For more information about attaching a policy to a role, see [Adding and removing IAM identity permissions](#) in the *AWS IAM User Guide*.

### Create a trust relationship with Amazon Bedrock

You must create a trust relationship with Amazon Bedrock so that Amazon Bedrock can assume your user’s IAM role while fine-tuning foundation models.

To add a trust relationship to your, take note of your domain or user profile’s IAM role. Then, do the following:

2. In the left navigation pane, choose **Roles**.
3. Search for the user’s IAM role by name from the list of roles and select it.
4. Choose the **Trust relationships** tab.
5. Choose **Edit trust policy**.
6. In the policy editor, find the **Add a principal option** in the right panel and choose **Add**.
7. In the dialog box, for **Principal type**, select **AWS services**.
8. For **ARN**, enter **bedrock.amazonaws.com**.
9. Choose **Add principal**.
10. Choose **Update policy**.

You should now have a trust relationship with Amazon Bedrock and the necessary permissions to fine-tune foundation models in SageMaker Canvas.

### Update SageMaker Canvas for Your Users

You can update to the latest version of Amazon SageMaker Canvas as either a user or an IT administrator. You can update Amazon SageMaker Canvas for a single user at a time.

To update the Amazon SageMaker Canvas application, you must delete the previous version.
Important
Deleting the previous version of Amazon SageMaker Canvas doesn’t delete the data or models that the users have created.

Use the following procedure to log in to AWS, open Amazon SageMaker domain, and update Amazon SageMaker Canvas. The users can start using the SageMaker Canvas application when they log back in.

1. Sign in to the Amazon SageMaker console at Amazon SageMaker Runtime.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. On the Domains page, choose your domain.
5. From the list of User profiles, choose a user profile.
6. For the list of Apps, find the Canvas application (the App type says Canvas) and choose Delete app.
7. Complete the dialog box and choose Confirm action.

The following image shows the user profile page and highlights the Delete app action from the preceding procedure.
Request a Quota Increase

Your users might use AWS resources in amounts that exceed those specified by their quotas. If your users are resource constrained and encounter errors in SageMaker Canvas, you can request a quota increase for them.

For more details about SageMaker quotas and how to request a quota increase, see [Quotas](#).

Amazon SageMaker Canvas uses the following services to process the requests of your users:

- Amazon SageMaker Autopilot
- Amazon SageMaker Studio Classic domain
- Amazon Forecast

For a list of the available quotas for SageMaker Canvas operations that aren't used to forecast time series data, see [Amazon SageMaker endpoints and quotas](#).

For a list of the available quotas for SageMaker Canvas operations that are used to forecast time series data, see [Amazon Forecast endpoints and quotas](#).
Request an increase for instances to build custom models

When building a custom model, if you encounter an error during post-building analysis that tells you to increase your quota for ml.m5.2xlarge instances, use the following information to resolve the issue.

You must increase the SageMaker Hosting endpoint quota for the ml.m5.2xlarge instance type to a non-zero value in your AWS account. After building a model, SageMaker Canvas hosts the model on a SageMaker Hosting endpoint and uses the endpoint to generate the post-building analysis. If you don't increase the default account quota of 0 for ml.m5.2xlarge instances, SageMaker Canvas cannot complete this step and generates an error during post-building analysis.

For the procedure to increase the quota, see Requesting a quota increase in the Service Quotas User Guide.

Grant Users Permissions to Import Amazon Redshift Data

Your users might have datasets stored in Amazon Redshift. Before users can import data from Amazon Redshift into SageMaker Canvas, you must add the AmazonRedshiftFullAccess managed policy to the IAM execution role that you've used for the user profile and add Amazon Redshift as a service principal to the role's trust policy. You must also associate the IAM execution role with your Amazon Redshift cluster. Complete the procedures in the following sections to give your users the required permissions to import Amazon Redshift data.

Add Amazon Redshift permissions to your IAM role

You must grant Amazon Redshift permissions to the IAM role specified in your user profile.

To add the AmazonRedshiftFullAccess policy to the user's IAM role, do the following.

2. Choose Roles.
3. In the search box, search for the user's IAM role by name and select it.
4. On the page for the user's role, under Permissions, choose Add permissions.
5. Choose Attach policies.
6. Search for the AmazonRedshiftFullAccess managed policy and select it.
7. Choose Attach policies to attach the policy to the role.
After attaching the policy, the role's Permissions section should now include AmazonRedshiftFullAccess.

To add Amazon Redshift as a service principal to the IAM role, do the following.

1. On the same page for the IAM role, under Trust relationships, choose Edit trust policy.
2. In the Edit trust policy editor, update the trust policy to add Amazon Redshift as a service principal. An IAM role that allows Amazon Redshift to access other AWS services on your behalf has a trust relationship as follows:

   ```json
   {
     "Version": "2012-10-17",
     "Statement": [
       {
         "Effect": "Allow",
         "Principal": {
           "Service": "redshift.amazonaws.com"
         },
         "Action": "sts:AssumeRole"
       }
     ]
   }
   ```

3. After editing the trust policy, choose Update policy.

You should now have an IAM role that has the policy AmazonRedshiftFullAccess attached to it and a trust relationship established with Amazon Redshift, giving users permission to import Amazon Redshift data into SageMaker Canvas. For more information about AWS managed policies, see Managed policies and inline policies in the IAM User Guide.

**Associate the IAM role with your Amazon Redshift cluster**

In the settings for your Amazon Redshift cluster, you must associate the IAM role that you granted permissions to in the preceding section.

To associate an IAM role with your cluster, do the following.

2. On the navigation menu, choose Clusters, and then choose the name of the cluster that you want to update.
3. In the **Actions** dropdown menu, choose **Manage IAM roles**. The **Cluster permissions** page appears.

4. For **Available IAM roles**, enter either the ARN or the name of the IAM role, or choose the IAM role from the list.

5. Choose **Associate IAM role** to add it to the list of **Associated IAM roles**.

6. Choose **Save changes** to associate the IAM role with the cluster.

Amazon Redshift modifies the cluster to complete the change, and the IAM role to which you previously granted Amazon Redshift permissions is now associated with your Amazon Redshift cluster. Your users now have the required permissions to import Amazon Redshift data into SageMaker Canvas.

### Grant Users Permissions to Collaborate with Studio Classic

**Note**

The functionality described on this page only applies to Amazon SageMaker Studio Classic. Currently, you can only share models to Canvas (or view shared Canvas models) in Studio Classic. If you’re currently using the latest version of Studio, you must run Studio Classic from within the latest version of Studio to share models to Canvas or view models shared from Canvas. For more information about accessing Studio Classic, see the [Studio Classic documentation](#).

Your Amazon SageMaker Canvas users might want to share their models with users in Amazon SageMaker Studio Classic to receive feedback and model updates, and Studio Classic users might want to share models with Canvas users so that they can generate predictions in Canvas. The following permissions grant Canvas users and Studio Classic users access to share models with each other.

For more information about how Canvas users can share models with Studio Classic users, see [Collaborate with data scientists](#). For more information about how Canvas users can bring a model shared from Studio Classic, see [Bring your own model to SageMaker Canvas](#).

Before Canvas and Studio Classic users can collaborate, the users must be in the same Amazon SageMaker domain. Add the following IAM permissions added to the same IAM execution role that you've used for their profiles.
To add the permissions to the users' IAM role, do the following:

1. Go to the IAM console.
2. Choose Roles.
3. In the search box, search for the user's IAM role by name and select it.
4. On the page for the user's role, under Permissions, choose Add permissions.
5. Choose Create inline policy.
6. In the Policy editor, choose JSON and enter the following IAM policy:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "sagemaker:CreateSharedModel",
            "sagemaker:DescribeSharedModel",
            "sagemaker:ListSharedModelEvents",
            "sagemaker:ListSharedModels",
            "sagemaker:ListSharedModelVersions",
            "sagemaker:SendSharedModelEvent",
            "sagemaker:UpdateSharedModel"
         ],
         "Resource": "*"
      }
   ]
}
```

7. Choose Next.
8. Enter a name for the policy in the Policy name field.
9. Choose Create policy to create the policy and attach it to the role.

For more information about AWS managed policies, see Managed policies and inline policies in the IAM User Guide.

**Grant Your Users Permissions to Send Predictions to Amazon QuickSight**

You must grant your SageMaker Canvas users permissions to send batch predictions to Amazon QuickSight. In Amazon QuickSight, users can create analyses and reports with a dataset and
prepare dashboards to share their results. For more information about sending prediction to QuickSight for analysis, see Send predictions to Amazon QuickSight.

To grant the necessary permissions to share batch predictions with users in QuickSight, you must add a permissions policy to the AWS Identity and Access Management (IAM) execution role that you’ve used for the user profile. The following section shows you how to attach a least-permissions policy to your role.

Add the permissions policy to your IAM role

To add the permissions policy, use the following procedure:

2. Choose Roles.
3. In the search box, search for the user's IAM role by name and select it.
4. On the page for the user's role, under Permissions, choose Add permissions.
5. Choose Create inline policy.
6. Select the JSON tab, and then paste the following least-permissions policy into the editor.
   Replace the placeholders <your-account-number> with your own AWS account number.

   ```json
   {
     "Version": "2012-10-17",
     "Statement": [
       {
         "Effect": "Allow",
         "Action": [
           "quicksight:CreateDataSet",
           "quicksight:ListUsers",
           "quicksight:ListNamespaces",
           "quicksight:CreateDataSource",
           "quicksight:PassDataSet",
           "quicksight:PassDataSource"
         ],
         "Resource": [
           "arn:aws:quicksight:*:<your-account-number>:datasource/*",
           "arn:aws:quicksight:*:<your-account-number>:user/*",
           "arn:aws:quicksight:*:<your-account-number>:namespace/*",
           "arn:aws:quicksight:*:<your-account-number>:dataset/*"
         ]
       }
     ]
   }
   ```
7. Choose **Review policy**.
8. Enter a **Name** for the policy.
9. Choose **Create policy**.

You should now have a customer-managed IAM policy attached to your execution role that grants your Canvas users the necessary permissions to send batch predictions to users in QuickSight.

**Manage applications**

The following sections describe how you can manage your SageMaker Canvas applications. You can view, delete, or relaunch your applications from the **Domains** section of the SageMaker console.

**Check for active applications**

To check if you have any actively running SageMaker Canvas applications, use the following procedure.

1. Open the [SageMaker console](#).
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. On the **Domains** page, choose your domain.
5. On the **Domain details** page, under **User profiles**, select the user profile name for the Canvas application that you want to view.
6. Under **Apps**, find the application that says **Canvas** in the **App type** column.

The **Status** column displays the status of the application, such as **Ready**, **Pending**, or **Deleted**. If the application is **Ready**, then your SageMaker Canvas workspace instance is active. You can delete the application from the console or log out from the SageMaker Canvas interface.

**Delete an application**

If you want to terminate your SageMaker Canvas workspace instance, you can either log out from the SageMaker Canvas application or delete your application from the SageMaker console. A **workspace instance** is dedicated for your use from when you start using SageMaker Canvas to the
point when you stop using it. Deleting the application only terminates the workspace instance and stops workspace instance charges. Models and datasets aren’t affected, but Quick build tasks automatically restart when you relaunch the application.

To delete your Canvas application through the AWS console, first close the browser tab in which your Canvas application was open. Then, use the following procedure to delete your SageMaker Canvas application.

1. Open the [SageMaker console](https://console.aws.amazon.com/sagemaker/).
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. On the **Domains** page, choose your domain.
5. On the **Domain details** page, under **User profiles**, select the user profile name for the Canvas application you want to view.
6. Under **Apps**, find the application that says **Canvas** in the **App type** column.
7. In the **Action** column, choose **Delete app**.
8. In the **Delete app** dialog box, select the **Yes, delete app** prompt, confirm the deletion by typing **delete** in the text field, and then choose **Delete**.

After you've successfully deleted the application, the **Status** column says **Deleted**. Otherwise, your application is still active.

You can also terminate the workspace instance by [logging out](https://console.aws.amazon.com/sagemaker/) from within the SageMaker Canvas application.

**Relaunch an application**

If you delete or log out of your SageMaker Canvas application and want to relaunch the application, use the following procedure.

1. Navigate to the [SageMaker console](https://console.aws.amazon.com/sagemaker/).
2. In the navigation pane, choose **Canvas**.
3. On the SageMaker Canvas landing page, in the **Get Started** box, select your user profile from the dropdown.
4. Choose **Open Canvas** to open the application.
SageMaker Canvas begins launching the application.

You can also use the following secondary procedure if you encounter any issues with the previous procedure.

1. Open the SageMaker console.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. On the Domains page, choose your domain.
5. On the Domain details page, under User profiles, select the user profile name for the SageMaker Canvas application you want to view.
6. Choose Launch and select Canvas from the dropdown list.

SageMaker Canvas begins launching the application.

**Configure Amazon SageMaker Canvas in a VPC without internet access**

The Amazon SageMaker Canvas application runs in a container in an AWS managed Amazon Virtual Private Cloud (VPC). If you want to further control access to your resources or run SageMaker Canvas without public internet access, you can configure your Amazon SageMaker domain and VPC settings. Within your own VPC, you can configure settings such as security groups (virtual firewalls that control inbound and outbound traffic from Amazon EC2 instances) and subnets (ranges of IP addresses in your VPC). To learn more about VPCs, see [How Amazon VPC works](#).

When the SageMaker Canvas application is running in the AWS managed VPC, it can interact with other AWS services using either an internet connection or through VPC endpoints created in a customer-managed VPC (without public internet access). SageMaker Canvas applications can access these VPC endpoints through a Studio Classic-created network interface that provides connectivity to the customer-managed VPC. The default behavior of the SageMaker Canvas application is to have internet access. When using an internet connection, the containers for the preceding jobs access AWS resources over the internet, such as the Amazon S3 buckets where you store training data and model artifacts.

However, if you have security requirements to control access to your data and job containers, we recommend that you configure SageMaker Canvas and your VPC so that your data and containers aren’t accessible over the internet. SageMaker uses the VPC configuration settings you specify when setting up your domain for SageMaker Canvas.
If you want to configure your SageMaker Canvas application without internet access, you must configure your VPC settings when you onboard to Amazon SageMaker domain, set up VPC endpoints, and grant the necessary AWS Identity and Access Management permissions. For information about configuring a VPC in Amazon SageMaker, see Choose an Amazon VPC. The following sections describe how to run SageMaker Canvas in a VPC without public internet access.

Configure Amazon SageMaker Canvas in a VPC without internet access

You can send traffic from SageMaker Canvas to other AWS services through your own VPC. If your own VPC doesn't have public internet access and you've set up your domain in VPC only mode, then SageMaker Canvas won't have public internet access as well. This includes all requests, such as accessing datasets in Amazon S3 or training jobs for standard builds, and the requests go through VPC endpoints in your VPC instead of the public internet. When you onboard to domain and Choose an Amazon VPC, you can specify your own VPC as the default VPC for the domain, along with your desired security group and subnet settings. Then, SageMaker creates a network interface in your VPC that SageMaker Canvas uses to access VPC endpoints in your VPC. Note that the security group and subnet settings are set after you finish onboarding to domain.

When onboarding to domain, if you choose Public internet only as the network access type, the VPC is SageMaker managed and allows internet access.

You can change this behavior by choosing VPC only so that SageMaker sends all traffic to a network interface that SageMaker creates in your specified VPC. When you choose this option, you must provide the subnets, security groups, and VPC endpoints that are necessary to communicate with the SageMaker API and SageMaker Runtime, and various AWS services, such as Amazon S3 and Amazon CloudWatch, that are used by SageMaker Canvas. Note that you can only import data from Amazon S3 buckets located in the same Region as your VPC.

The following procedures show how you can configure these settings to use SageMaker Canvas without the internet.

Step 1: Onboard to Amazon SageMaker domain

To send SageMaker Canvas traffic to a network interface in your own VPC instead of over the internet, specify the VPC you want to use when onboarding to Amazon SageMaker domain. You must also specify at least two subnets in your VPC that SageMaker can use. Choose Standard setup and do the following procedure when configuring the Network and Storage Section for the domain.

1. Select your desired VPC.
2. Choose two or more **Subnets**. If you don’t specify the subnets, SageMaker uses all of the subnets in the VPC.

3. Choose one or more **Security group(s)**.

4. Choose **VPC Only** to turn off direct internet access in the AWS managed VPC where SageMaker Canvas is hosted.

After disabling internet access, finish the onboarding process to set up your domain. For more information about the VPC settings for Amazon SageMaker domain, see [Choose an Amazon VPC](#).

---

**Step 2: Configure VPC endpoints and access**

---

**Note**

In order to configure Canvas in your own VPC, you must enable private DNS hostnames for your VPC endpoints. For more information, see [Connect to SageMaker Through a VPC Interface Endpoint](#).

---

SageMaker Canvas only accesses other AWS services to manage and store data for its functionality. For example, it connects to Amazon Redshift if your users access an Amazon Redshift database. It can connect to an AWS service such as Amazon Redshift using an internet connection or a VPC endpoint. Use VPC endpoints if you want to set up connections from your VPC to AWS services that don't use the public internet.

A VPC endpoint creates a private connection to an AWS service that uses a networking path that is isolated from the public internet. For example, if you set up access to Amazon S3 using a VPC endpoint from your own VPC, then the SageMaker Canvas application can access Amazon S3 by going through the network interface in your VPC and then through the VPC endpoint that connects to Amazon S3. The communication between SageMaker Canvas and Amazon S3 is private.

For more information about configuring VPC endpoints for your VPC, see [AWS PrivateLink](#). If you are using Amazon Bedrock models in Canvas with a VPC, for more information about controlling access to your data, see [Protect jobs using a VPC](#) in the Amazon Bedrock User Guide.

The following are the VPC endpoints for each service you can use with SageMaker Canvas:
<table>
<thead>
<tr>
<th>Service</th>
<th>Endpoint</th>
<th>Endpoint type</th>
</tr>
</thead>
<tbody>
<tr>
<td>AWS Application Auto Scaling</td>
<td>com.amazon.aws.Region.application-autoscaling</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Athena</td>
<td>com.amazon.aws.Region.athena</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon SageMaker</td>
<td>com.amazon.aws.Region.sagemaker.api</td>
<td>Interface</td>
</tr>
<tr>
<td></td>
<td>com.amazon.aws.Region.sagemaker.runtime</td>
<td></td>
</tr>
<tr>
<td></td>
<td>com.amazon.aws.Region.notebook</td>
<td></td>
</tr>
<tr>
<td>AWS Security Token Service</td>
<td>com.amazonaws.Region.sts</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Elastic Container Registry (Amazon ECR)</td>
<td>com.amazon.aws.Region.ecr.api</td>
<td>Interface</td>
</tr>
<tr>
<td></td>
<td>com.amazon.aws.Region.ecr.dkr</td>
<td></td>
</tr>
<tr>
<td>Amazon Elastic Compute Cloud (Amazon EC2)</td>
<td>com.amazonaws.Region.ec2</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Simple Storage Service (Amazon S3)</td>
<td>com.amazonaws.Region.s3</td>
<td>Gateway</td>
</tr>
<tr>
<td>Amazon Redshift</td>
<td>com.amazon.aws.Region.redshift-data</td>
<td>Interface</td>
</tr>
<tr>
<td>AWS Secrets Manager</td>
<td>com.amazon.aws.Region.secretsmanager</td>
<td>Interface</td>
</tr>
<tr>
<td>Service</td>
<td>Endpoint</td>
<td>Endpoint type</td>
</tr>
<tr>
<td>--------------------------------------------------</td>
<td>-----------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>AWS Systems Manager</td>
<td>com.amazonaws.Region.ssm</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon CloudWatch</td>
<td>com.amazonaws.Region.monitoring</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon CloudWatch Logs</td>
<td>com.amazonaws.Region.logs</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Forecast</td>
<td>com.amazonaws.Region.forecast</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Forecast</td>
<td>com.amazonaws.Region.forecastquery</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Textract</td>
<td>com.amazonaws.Region.textract</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Comprehend</td>
<td>com.amazonaws.Region.comprehend</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Rekognition</td>
<td>com.amazonaws.Region.rekognition</td>
<td>Interface</td>
</tr>
<tr>
<td>AWS Glue</td>
<td>com.amazonaws.Region.glue</td>
<td>Interface</td>
</tr>
<tr>
<td>AWS Application Auto Scaling</td>
<td>com.amazonaws.Region.application-autoscaling</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Relational Database Service (Amazon RDS)</td>
<td>com.amazonaws.Region.rds</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Bedrock</td>
<td>com.amazonaws.Region.bedrock-runtime</td>
<td>Interface</td>
</tr>
<tr>
<td>Amazon Kendra</td>
<td>com.amazonaws.Region.kendra</td>
<td>Interface</td>
</tr>
</tbody>
</table>
Note

For Amazon Bedrock, the interface endpoint service name `com.amazonaws.Region.bedrock` has been deprecated. Create a new VPC endpoint with the service name listed in the preceding table. Additionally, you can't fine-tune foundation models from Canvas VPCs with no internet access. This is because Amazon Bedrock doesn't support VPC endpoints for model customization APIs. To learn more about fine-tuning foundation models in Canvas, see Fine-tune foundation models.

You must also add an endpoint policy for Amazon S3 to control AWS principal access to your VPC endpoint. For information about how to update your VPC endpoint policy, see Control access to VPC endpoints using endpoint policies.

The following are two VPC endpoint policies that you can use. Use the first policy if you only want to grant access to the basic functionality of Canvas, such as importing data and creating models. Use the second policy if you want to grant access to the additional generative AI features in Canvas.

Basic VPC endpoint policy

The following policy grants the necessary access to your VPC endpoint for basic operations in Canvas.

```json
{
   "Effect": "Allow",
   "Action": [
      "s3:GetObject",
      "s3:GetObject",
      "s3:GetObject",
      "s3:GetObject",
      "s3:GetObject",
      "s3:GetObject",
      "s3:GetObject",
      "s3:GetObject"
   ],
   "Resource": [
      "arn:aws:s3:::*SageMaker*",
      "arn:aws:s3:::*Sagemaker*",
      "arn:aws:s3:::*sagemaker*"
   ]
}
```
Generative AI VPC endpoint policy

The following policy grants the necessary access to your VPC endpoint for basic operations in Canvas, as well as using generative AI foundation models.
Step 3: Grant IAM permissions

The SageMaker Canvas user must have the necessary AWS Identity and Access Management permissions to allow connection to the VPC endpoints. The IAM role to which you give permissions must be the same one you used when onboarding to Amazon SageMaker domain. You can attach the SageMaker managed AmazonSageMakerFullAccess policy to the IAM role for the user to give the user the required permissions. If you require more restrictive IAM permissions and use custom policies instead, then give the user's role the ec2:DescribeVpcEndpointServices permission. SageMaker Canvas requires these permissions to verify the existence of the required VPC endpoints for standard build jobs. If it detects these VPC endpoints, then standard build jobs run by default in your VPC. Otherwise, they will run in the default AWS managed VPC.

For instructions on how to attach the AmazonSageMakerFullAccess IAM policy to your user's IAM role, see Adding and removing IAM identity permissions.

To grant your user's IAM role the granular ec2:DescribeVpcEndpointServices permission, use the following procedure.

1. Sign in to the AWS Management Console and open the IAM console.
2. In the navigation pane, choose Roles.
3. In the list, choose the name of the role to which you want to grant permissions.
4. Choose the Permissions tab.
5. Choose Add permissions and then choose Create inline policy.
6. Choose the JSON tab and enter the following policy, which grants the ec2:DescribeVpcEndpointServices permission:

   ```json
   {
   "Version": "2012-10-17",
   "Statement": [
   {
   "Sid": "VisualEditor0",
   "Effect": "Allow",
   "Action": "ec2:DescribeVpcEndpointServices",
   "Resource": "*"
   }
   ]
   }
   ```
7. Choose **Review policy**, and then enter a **Name** for the policy (for example, VPCEndpointPermissions).

8. Choose **Create policy**.

The user’s IAM role should now have permissions to access the VPC endpoints configured in your VPC.

(Optional) Step 4: Override security group settings for specific users

If you are an administrator, you might want different users to have different VPC settings, or user-specific VPC settings. When you override the default VPC’s security group settings for a specific user, these settings are passed on to the SageMaker Canvas application for that user.

You can override the security groups that a specific user has access to in your VPC when you set up a new user profile in Studio Classic. You can use the **CreateUserProfile** SageMaker API call (or **create_user_profile** with the **AWS CLI**), and then in the **UserSettings**, you can specify the **SecurityGroups** for the user.

**Set up connections to data sources with OAuth**

The following section describes the steps you must take to set up OAuth connections to data sources from SageMaker Canvas. **OAuth** is a common authentication platform for granting access to resources without sharing passwords. With OAuth, you can quickly connect to your data from Canvas and import it for building models. Canvas currently supports OAuth for Snowflake and Salesforce Data Cloud.

Note

You can only establish one OAuth connection for each data source.

**Set up OAuth for Salesforce Data Cloud**

To set up OAuth for Salesforce Data Cloud, follow these general steps:

1. Sign in to Salesforce Data Cloud.
2. In Salesforce Data Cloud, create a new app connection and do the following:
   a. Enable OAuth settings.
b. When prompted for a callback URL (or the URL of the resource accessing your data), specify the URL for your Canvas application. The Canvas application URL follows this format: https://<domain-id>.studio.<region>.sagemaker.aws/canvas/default

c. Copy the consumer key and secret.
d. Copy your authorization URL and token URL.

For more detailed instructions about performing the preceding tasks in Salesforce Data Cloud, see Import data from Salesforce Data Cloud in the Data Wrangler documentation for importing data from Salesforce Data Cloud.

After enabling access from Salesforce Data Cloud and getting your connection information, you must create an AWS Secrets Manager secret to store the information and add it to your Amazon SageMaker domain or user profile. Note that you can add a secret to both a domain and user profile, but Canvas looks for secrets in the user profile first.

To add a secret to your domain or user profile, do the following:

1. Go to the Amazon SageMaker console.
2. Choose domains in the navigation pane.
3. From the list of domains, choose your domain.
   a. If adding your secret to your domain, do the following:
      i. Choose the domain.
      ii. On the domain settings page, choose the domain settings tab.
      iii. Choose Edit.
   b. If adding the secret to your user profile, do the following:
      i. Choose the user's domain.
      ii. On the domain settings page, choose the user profile.
      iii. On the User Details page, choose Edit.
4. In the navigation pane, choose Canvas settings.
5. For OAuth settings, choose Add OAuth configuration.
6. For Data source, select Salesforce Data Cloud.
7. For **Secret Setup**, select **Create a new secret**. Alternatively, if you already created an AWS Secrets Manager secret with your credentials, enter the ARN for the secret. If creating a new secret, do the following:

   a. For **Identity Provider**, select **SALESFORCE**.
   b. For **Client ID**, **Client Secret**, **Authorization URL**, and **Token URL**, enter all of the information you gathered from Salesforce Data Cloud in the previous procedure.

8. Save your domain or user profile settings.

You should now be able to create a connection to your data in Salesforce Data Cloud from Canvas.

**Set up OAuth for Snowflake**

To set up authentication for Snowflake, Canvas supports identity providers that you can use instead of having users directly enter their credentials into Canvas.

The following are links to the Snowflake documentation for the identity providers that Canvas supports:

- [Azure AD](#)
- [Okta](#)
- [Ping Federate](#)

The following process describes the general steps you must take. For more detailed instructions about performing these steps, you can refer to the Setting up Snowflake OAuth Access section in the Data Wrangler documentation for importing data from Snowflake.

To set up OAuth for Snowflake, do the following:

1. Register Canvas as an application with the identity provider. This requires specifying a redirect URL to Canvas, which should follow this format: https://<domain-id>.studio.<region>.sagemaker.aws/canvas/default

2. Within the identity provider, create a server or API that sends OAuth tokens to Canvas so that Canvas can access Snowflake. When setting up the server, use the authorization code and refresh token grant types, specify the access token lifetime, and set a refresh token policy. Additionally, within the External OAuth Security Integration for Snowflake, enable `external_oauth_any_role_mode`.


3. Get the following information from the identity provider: token URL, authorization URL, client ID, client secret. For Azure AD, also retrieve the OAuth scope credentials.

4. Store the information retrieved in the previous step in an AWS Secrets Manager secret.

   a. For Okta and Ping Federate, the secret should look like the following format:

   ```json
   {"token_url":"https://identityprovider.com/oauth2/example-portion-of-URL-path/v2/token",
   "client_id":"example-client-id", "client_secret":"example-client-secret",
   "identity_provider":"OKTA"|"PING_FEDERATE",
   "authorization_url":"https://identityprovider.com/oauth2/example-portion-of-URL-path/v2/authorize"}
   
   b. For Azure AD, the secret should also include the OAuth scope credentials as the datasource_oauth_scope field.

After configuring the identity provider and the secret, you must create an AWS Secrets Manager secret to store the information and add it to your Amazon SageMaker domain or user profile. Note that you can add a secret to both a domain and user profile, but Canvas looks for secrets in the user profile first.

To add a secret to your domain or user profile, do the following:

1. Go to the Amazon SageMaker console.
2. Choose domains in the navigation pane.
3. From the list of domains, choose your domain.

   a. If adding your secret to your domain, do the following:

      i. Choose the domain.
      ii. On the domain settings page, choose the domain settings tab.
      iii. Choose Edit.

   b. If adding the secret to your user profile, do the following:

      i. Choose the user's domain.
      ii. On the domain settings page, choose the user profile.
      iii. On the User Details page, choose Edit.

4. In the navigation pane, choose Canvas settings.
5. For **OAuth settings**, choose **Add OAuth configuration**.

6. For **Data source**, select **Snowflake**.

7. For **Secret Setup**, select **Create a new secret**. Alternatively, if you already created an AWS Secrets Manager secret with your credentials, enter the ARN for the secret. If creating a new secret, do the following:
   a. For **Identity Provider**, select **SNOWFLAKE**.
   b. For **Client ID**, **Client Secret**, **Authorization URL**, and **Token URL**, enter all of the information you gathered from the identity provider in the previous procedure.

8. Save your domain or user profile settings.

You should now be able to create a connection to your data in Snowflake from Canvas.

**Import data into Canvas**

Amazon SageMaker Canvas supports importing tabular, image, and document data. You can import data from both local and external data sources into Canvas. Use the datasets that you import to build models and make predictions for other datasets.

Each use case for which you can build a custom model accepts different types of input. For example, if you want to build a single-label image classification model, then you should import image data. For more information about the different model types and the data they accept, see [Build a custom model](#). You can import data and build custom models in SageMaker Canvas for the following data types:

- **Tabular** (CSV, Parquet, or tables)
  - Categorical – Use categorical data to build custom categorical prediction models for 2 and 3+ category prediction.
  - Numeric – Use numeric data to build custom numeric prediction models.
  - Text – Use text data to build custom multi-category text prediction models.
  - Timeseries – Use timeseries data to build custom time series forecasting models.
- **Image** (JPG or PNG) – Use image data to build custom single-label image prediction models.
- **Document** (PDF, JPG, PNG, TIFF) – Document data is only supported for SageMaker Canvas Ready-to-use models. To learn more about Ready-to-use models that can make predictions for document data, see [Use Ready-to-use models](#).
You can import data into Canvas from the following data sources:

- Local files on your computer
- Amazon S3 buckets
- Amazon Redshift provisioned clusters (not Amazon Redshift Serverless)
- AWS Glue Data Catalog through Amazon Athena
- Amazon Aurora
- Amazon Relational Database Service (Amazon RDS)
- Salesforce Data Cloud
- Snowflake
- Databricks, SQLServer, MariaDB, and other popular databases through JDBC connectors
- Over 40 external SaaS platforms, such as SAP OData

For a full list of data sources from which you can import, see the following table:

<table>
<thead>
<tr>
<th>Source</th>
<th>Type</th>
<th>Supported data types</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local file upload</td>
<td>Local</td>
<td>Tabular, Image, Document</td>
</tr>
<tr>
<td>Amazon Aurora</td>
<td>Amazon internal</td>
<td>Tabular</td>
</tr>
<tr>
<td>Amazon S3 bucket</td>
<td>Amazon internal</td>
<td>Tabular, Image, Document</td>
</tr>
<tr>
<td>Amazon RDS</td>
<td>Amazon internal</td>
<td>Tabular</td>
</tr>
<tr>
<td>Amazon Redshift provisioned clusters (not Redshift Serverless)</td>
<td>Amazon internal</td>
<td>Tabular</td>
</tr>
<tr>
<td>AWS Glue Data Catalog (through Amazon Athena)</td>
<td>Amazon internal</td>
<td>Tabular</td>
</tr>
<tr>
<td>Databricks</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td>Snowflake</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td>Salesforce Data Cloud</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td>Source</td>
<td>Type</td>
<td>Supported data types</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>SQLServer</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td>MySQL</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td>PostgreSQL</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td>MariaDB</td>
<td>External</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Amplitude</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>CircleCI</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>DocuSign Monitor</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Domo</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Datadog</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Dynatrace</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Facebook Ads</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Facebook Page Insights</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Google Ads</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Google Analytics 4</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Google Search Console</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>GitHub</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>GitLab</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Infor Nexus</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Instagram Ads</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td><strong>Jira Cloud</strong></td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Source</td>
<td>Type</td>
<td>Supported data types</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>LinkedIn Ads</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>LinkedIn Ads</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Mailchimp</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Marketo</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Microsoft Teams</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Mixpanel</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Okta</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Salesforce</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Salesforce Marketing Cloud</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Salesforce Pardot</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>SAP OData</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>SendGrid</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>ServiceNow</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Singular</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Slack</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Stripe</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Trend Micro</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Typeform</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Veeva</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Zendesk</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Source</td>
<td>Type</td>
<td>Supported data types</td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>Zendesk Chat</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Zendesk Sell</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Zendesk Sunshine</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
<tr>
<td>Zoom Meetings</td>
<td>External SaaS platform</td>
<td>Tabular</td>
</tr>
</tbody>
</table>

For instructions on how to import data and information regarding input data requirements, such as the maximum file size for images, see [Create a dataset](#).

Canvas also provides several sample datasets in your application to help you get started. To learn more about the SageMaker-provided sample datasets you can experiment with, see [Use sample datasets](#).

After you import a dataset into Canvas, you can update the dataset at any time. You can do a manual update or you can set up a schedule for automatic dataset updates. For more information, see [Update a dataset](#).

For more information specific to each dataset type, see the following sections:

**Tabular**

To import data from an external data source (such as a Snowflake database or a SaaS platform), you must authenticate and connect to the data source in the Canvas application. For more information, see [Connect to data sources](#).

After creating datasets in Canvas, you can join multiple datasets into a single dataset. Joining datasets is only supported for tabular datasets. As long as your data is arranged into tables, you can join datasets from various sources, such as Amazon Redshift, Amazon Athena, or Snowflake. For information about joining datasets, see [Join data that you’ve imported into SageMaker Canvas](#).

**Image**

For information about how to edit an image dataset and perform tasks such as assigning or reassigning labels, adding images, or deleting images, see [Edit an image dataset](#).
Create a dataset

The following sections describe how to create a dataset in Amazon SageMaker Canvas. For custom models, you can create datasets for tabular and image data. For Ready-to-use models, you can use tabular and image datasets as well as document datasets. Choose your workflow based on the following information:

- For categorical, numeric, text, and timeseries data, see [Import tabular data](#).
- For image data, see [Import image data](#).
- For document data, see [Import document data](#).

**Note**

For information about how to import a document dataset for Ready-to-use models that accept document data, see the [Import document data](#) workflow in the Ready-to-use models documentation.

A dataset can consist of multiple files. For example, you might have multiple files of inventory data in CSV format. You can upload these files together as a dataset as long as the schema (or column names and data types) of the files match.

Canvas also supports managing multiple versions of your dataset. When you create a dataset, the first version is labeled as V1. You can create a new version of your dataset by updating your dataset. You can do a manual update, or you can set up an automated schedule for updating your dataset with new data. For more information, see [Update a dataset](#).

When you import your data into Canvas, make sure that it meets the requirements in the following table. The limitations are specific to the type of model you’re building.

<table>
<thead>
<tr>
<th>Limit</th>
<th>2 category, 3+ category, numeric, and time series models</th>
<th>Text prediction models</th>
<th>Image prediction models</th>
<th>*Document data for Ready-to-use models</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supported file types</td>
<td>CSV and Parquet (local upload,)</td>
<td>CSV and Parquet (local upload,)</td>
<td>JPG, PNG</td>
<td>PDF, JPG, PNG, TIFF</td>
</tr>
<tr>
<td>Limit</td>
<td>2 category, 3+ category, numeric, and time series models</td>
<td>Text prediction models</td>
<td>Image prediction models</td>
<td>*Document data for Ready-to-use models</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>---------------------------------------------------------</td>
<td>------------------------</td>
<td>-------------------------</td>
<td>----------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Amazon S3, or databases</td>
<td>Amazon S3, or databases</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>JSON (databases)</td>
<td>JSON (databases)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum file size</td>
<td>5 GB (for all files in the dataset)</td>
<td>5 MB (for all files in the dataset)</td>
<td>30 MB per image</td>
<td>5 MB per document</td>
</tr>
<tr>
<td>Maximum number of files in tabular datasets</td>
<td>50</td>
<td>50</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Maximum number of files in tabular datasets for a single manual upload</td>
<td>20</td>
<td>20</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Maximum number of columns</td>
<td>1000</td>
<td>1000</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Maximum number of entries (rows, images, or documents) for Quick builds</td>
<td>50,000 rows</td>
<td>7500 rows</td>
<td>5000 images</td>
<td>N/A</td>
</tr>
<tr>
<td>Limit</td>
<td>2 category, 3+ category, numeric, and time series models</td>
<td>Text prediction models</td>
<td>Image prediction models</td>
<td>*Document data for Ready-to-use models</td>
</tr>
<tr>
<td>-------</td>
<td>------------------------------------------------------</td>
<td>------------------------</td>
<td>------------------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td>Maximum number of entries (rows, images, or documents) for <strong>Standard builds</strong></td>
<td>N/A</td>
<td>150,000 rows</td>
<td>180,000 images</td>
<td>N/A</td>
</tr>
<tr>
<td>Minimum number of entries (rows) for <strong>Quick builds</strong></td>
<td>2 category: 500 rows 3+ category, numeric, time series: N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Minimum number of entries (rows, images, or documents) for <strong>Standard builds</strong></td>
<td>250 rows</td>
<td>50 rows</td>
<td>50 images</td>
<td>N/A</td>
</tr>
<tr>
<td>Minimum number of entries (rows or images) per label</td>
<td>N/A</td>
<td>25 rows</td>
<td>25 rows</td>
<td>N/A</td>
</tr>
<tr>
<td>Limit</td>
<td>2 category, 3+ category, numeric, and time series models</td>
<td>Text prediction models</td>
<td>Image prediction models</td>
<td>*Document data for Ready-to-use models</td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------------------------------------------------------</td>
<td>------------------------</td>
<td>------------------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td>Minimum number of labels</td>
<td>2 category: 2</td>
<td>2</td>
<td>2</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>3+ category: 3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Numeric, time series: N/A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minimum sample size for random sampling</td>
<td>500</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Maximum sample size for random sampling</td>
<td>40,000</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Maximum number of labels</td>
<td>2 category: 2</td>
<td>1000</td>
<td>1000</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>3+ category, numeric, time series: N/A</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Document data is currently only supported for Ready-to-use models that accept document data. You can't build a custom model with document data.

Also note the following restrictions:

- For tabular data, Canvas disallows selecting any file with extensions other than .csv, .parquet, .parq, and .pqt for both local upload and Amazon S3 import. CSV files must be comma delimited and not have newline characters except when denoting a new row.
- For tabular data using Parquet files, note the following:
• Parquet files can’t include complex types like maps and lists.
• The column names of Parquet files can’t contain spaces.
• If using compression, Parquet files must use either gzip or snappy compression types. For more information about the preceding compression types, see the [gzip documentation](#) and the [snappy documentation](#).

For image data, if you have any unlabeled images, you must label them before building your model. For information about how to assign labels to images within the Canvas application, see [Edit an image dataset](#).

If you set up automatic dataset updates or automatic batch prediction configurations, you can only create a total of 20 configurations in your Canvas application. For more information, see [Manage automations](#).

After you import a dataset, you can view your datasets on the **Datasets** page at any time.

**Import tabular data**

With tabular datasets, you can build categorical, numeric, time series forecasting, and text prediction models. Review the limitations table in the preceding **Import a dataset** section to ensure that your data meets the requirements for tabular data (note that the sample size limits only apply when previewing your data before building your model).

Use the following procedure to import a tabular dataset into Canvas:

1. Open your SageMaker Canvas application.
2. In the left navigation pane, choose **Datasets**.
3. Choose **Import**.
4. In the popup dialog box, in the **Dataset name** field, enter a name for the dataset and choose **Create**.
5. On the **Import** page, open the **Data Source** dropdown menu.
6. Choose your data source:

   - To upload files from your computer, choose **Local upload**.
   - To import data from another source, such as an Amazon S3 bucket or a Snowflake database, search for your data source in the **Search data source bar**. Then, choose the tile for your desired data source.
Note

You can only import data from the tiles that have an active connection. If you want to connect to a data source that is unavailable to you, contact your administrator. If you’re an administrator, see Connect to data sources.

The following screenshot shows the Data Source dropdown menu.

7. (Optional) If you’re connecting to an Amazon Redshift or Snowflake database for the first time, a dialog box appears to create a connection. Fill out the dialog box with your credentials and choose Create connection. If you already have a connection, choose your connection.

8. From your data source, select your files to import. For local upload and importing from Amazon S3, you can select files. For Amazon S3 only, you also have the option to directly enter the S3 URI or ARN of your bucket in the Input S3 endpoint field and then choose files to import. For database sources, you can drag-and-drop data tables from the left navigation pane.

9. (Optional) For tabular data sources that support SQL querying (such as Amazon Redshift, Amazon Athena, or Snowflake), you can choose Edit in SQL to make SQL queries and join tables before importing them. For more information, see Join data that you’ve imported into SageMaker Canvas.
The following screenshot shows the **Edit SQL** view for an Amazon Athena data source.

10. (Optional) You can choose **Preview** to preview your dataset before importing. For tabular datasets, this shows you up to the first 100 rows of your dataset. The following screenshot shows you the **Import preview** screen.

11. When you’re ready to import your data, choose **Import data**.

While your dataset is importing into Canvas, you can see your datasets listed on the **Datasets** page. From this page, you can **View your dataset details**.

When the **Status** of your dataset shows as Ready, Canvas successfully imported your data and you can proceed with **building a model**.

If you have a connection to a data source, such as an Amazon Redshift database or a SaaS connector, you can return to that connection. For Amazon Redshift and Snowflake, you can add another connection by creating another dataset, returning to the **Import data** page, and choosing the **Data Source** tile for that connection. From the dropdown menu, you can open the previous connection or choose **Add connection**.

⚠️ **Note**

For SaaS platforms, you can only have one connection per data source.
Import image data

With image datasets, you can build single-label image prediction custom models, which predict a label for an image. Review the limitations in the preceding Import a dataset section to ensure that your image dataset meets the requirements for image data.

ℹ️ Note
You can only import image datasets from local file upload or an Amazon S3 bucket. Also, for image datasets, you must have at least 25 images per label.

Use the following procedure to import an image dataset into Canvas:

1. Open your SageMaker Canvas application.
2. In the left navigation pane, choose Datasets.
3. Choose Create.
4. From the dropdown menu, choose Image.
5. In the popup dialog box, in the Dataset name field, enter a name for the dataset and choose Create.
6. On the Import page, open the Data Source dropdown menu.
7. Choose your data source. To upload files from your computer, choose Local upload. To import files from Amazon S3, choose Amazon S3.
8. From your computer or Amazon S3 bucket, select the images or folders of images that you want to upload.
9. When you’re ready to import your data, choose Import data.

While your dataset is importing into Canvas, you can see your datasets listed on the Datasets page. From this page, you can View your dataset details.

When the Status of your dataset shows as Ready, Canvas successfully imported your data and you can proceed with building a model.

When you are building your model, you can edit your image dataset, and you can assign or re-assign labels, add images, or delete images from your dataset. For more information about how to edit your image dataset, see Edit an image dataset.
Import document data

The Ready-to-use models for expense analysis, identity document analysis, document analysis, and document queries support document data. You can’t build a custom model with document data.

With document datasets, you can generate predictions for expense analysis, identity document analysis, document analysis, and document queries Ready-to-use models. Review the limitations table in the Create a dataset section to ensure that your document dataset meets the requirements for document data.

Note
You can only import document datasets from local file upload or an Amazon S3 bucket.

Use the following procedure to import a document dataset into Canvas:

1. Open your SageMaker Canvas application.
2. In the left navigation pane, choose Datasets.
3. Choose Create.
4. From the dropdown menu, choose Document.
5. In the popup dialog box, in the Dataset name field, enter a name for the dataset and choose Create.
6. On the Import page, open the Data Source dropdown menu.
7. Choose your data source. To upload files from your computer, choose Local upload. To import files from Amazon S3, choose Amazon S3.
8. From your computer or Amazon S3 bucket, select the document files that you want to upload.
9. When you’re ready to import your data, choose Import data.

While your dataset is importing into Canvas, you can see your datasets listed on the Datasets page. From this page, you can View your dataset details.

When the Status of your dataset shows as Ready, Canvas has successfully imported your data.

On the Datasets page, you can choose your dataset to preview it, which shows you up to the first 100 documents of your dataset.
View your dataset details

For each of your datasets, you can view all of the files in a dataset, the dataset's version history, and any auto update configurations for the dataset. From the Datasets page, you can also initiate actions such as Update a dataset or Build a custom model.

To view the details for a dataset, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose Datasets.
3. From the list of datasets, choose your dataset.

On the Data tab, you can see a preview of your data. If you choose Dataset details, you can see all of the files that are part of your dataset. Choose a file to see only the data from that file in the preview. For image datasets, the preview only shows you the first 100 images of your dataset.

On the Version history tab, you can see a list of all of the versions of your dataset. A new version is made whenever you update a dataset. To learn more about updating a dataset, see Update a dataset. The following screenshot shows the Version history tab in the Canvas application.
On the **Auto updates** tab, you can enable auto updates for the dataset and set up a configuration to update your dataset on a regular schedule. To learn more about setting up auto updates for a dataset, see [Configure automatic updates for a dataset](#). The following screenshot shows the **Auto updates** tab with auto updates turned on and a list of auto update jobs that have been performed on the dataset.
Update a dataset

After importing your initial dataset into Amazon SageMaker Canvas, you might have additional data that you want to add to your dataset. For example, you might get inventory data at the end of every week that you want to add to your dataset. Instead of importing your data multiple times, you can update your existing dataset and add or remove files from it.

Note

You can only update datasets that you have imported through local upload or Amazon S3.
You can update your dataset either manually or automatically. With automatic updates, you specify a location where Canvas checks for files at a frequency you specify. If you import new files during the update, the schema of the files must match the existing dataset exactly.

Every time you update your dataset, Canvas creates a new version of your dataset. You can only use the latest version of your dataset to build a model or generate predictions. For more information about viewing the version history of your dataset, see View your dataset details.

You can also use dataset updates with automated batch predictions, which starts a batch prediction job whenever you update your dataset. For more information, see Make batch predictions.

The following sections describe how to do manual and automatic updates to your dataset.

**Manually update a dataset**

To do a manual update, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose Datasets.
3. From the list of datasets, choose the dataset you want to update.
4. Choose the Update dataset dropdown menu and choose Manual update. You are taken to the import data workflow.
5. From the Data source dropdown menu, choose either Local upload or Amazon S3.
6. The page shows you a preview of your data. From here, you can add or remove files from the dataset. If you’re importing tabular data, the schema of the new files (column names and data types) must match the schema of the existing files. Additionally, your new files must not exceed the maximum dataset size or file size. For more information about these limitations, see Import a dataset.

    **Note**
    
    If you add a file with the same name as an existing file in your dataset, the new file overwrites the old version of the file.

7. When you’re ready to save your changes, choose Update dataset.

You should now have a new version of your dataset.
On the **Datasets** page, you can choose the **Version history** tab to see all of the versions of your dataset and the history of both manual and automatic updates you’ve made.

**Configure automatic updates for a dataset**

An automatic update is when you set up a configuration for Canvas to update your dataset at a given frequency. We recommend that you use this option if you regularly receive new files of data that you want to add to your dataset.

When you set up the auto update configuration, you specify an Amazon S3 location where you upload your files and a frequency at which Canvas checks the location and imports files. Each instance of Canvas updating your dataset is referred to as a *job*. For each job, Canvas imports all of the files in the Amazon S3 location. If you have new files with the same names as existing files in your dataset, Canvas overwrites the old files with the new files.

For automatic dataset updates, Canvas doesn’t perform schema validation. If the schema of files imported during an automatic update don’t match the schema of the existing files or exceed the size limitations (see [Import a dataset](#) for a table of file size limitations), then you get errors when your jobs run.

### Note

You can only set up a maximum of 20 automatic configurations in your Canvas application. Additionally, Canvas only does automatic updates while you’re logged in to your Canvas application. If you log out of your Canvas application, automatic updates pause until you log back in.

To configure automatic updates for your dataset, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **Datasets**.
3. From the list of datasets, choose the dataset you want to update.
4. Choose the **Update dataset** dropdown menu and choose **Automatic update**. You are taken to the **Auto updates** tab for the dataset.
5. Turn on the **Auto update enabled** toggle.
6. For **Specify a data source**, enter the Amazon S3 path to a folder where you plan to regularly upload files.
7. For **Choose a frequency**, select **Hourly**, **Weekly**, or **Daily**.

8. For **Specify a starting time**, use the calendar and time picker to select when you want the first auto update job to start.

9. When you’re ready to create the auto update configuration, choose **Save**.

Canvas begins the first job of your auto update cadence at the specified starting time.

For more information about viewing your auto update job history or making changes to your auto update configuration through the **Automations** page in the Canvas application, see [Manage automations](#).

The following sections describe how to view, update, and delete your automatic update configuration through the **Datasets** page in the Canvas application.

**View your automatic dataset update jobs**

To view the job history for your automatic dataset updates, on your dataset details page, choose the **Auto updates** tab.

Each automatic update to a dataset shows as a job in the **Auto updates** tab under the **Job history** section. For each job, you can see the following:

- **Job created** – The timestamp for when Canvas started updating the dataset.
- **Files** – The number of files in the dataset.
- **Cells (Columns x Rows)** – The number of columns and rows in the dataset.
- **Status** – The status of the dataset after the update. If the job was successful, the status is **Ready**. If the job failed for any reason, the status is **Failed**, and you can hover over the status for more details.

**Edit your automatic dataset update configuration**

You might want to make changes to your auto update configuration for a dataset, such as changing the frequency of the updates. You might also want to turn off your automatic update configuration to pause the updates to your dataset.

To make changes to your auto update configuration for a dataset, go to the **Auto updates** tab of your dataset and choose **Edit** to make changes to the configuration.
To pause your dataset updates, turn off your automatic configuration. You can turn off auto updates by going to the **Auto updates** tab of your dataset and turning the **Enable auto updates** toggle off. You can turn this toggle back on at any time to resume the update schedule.

**Delete your automatic dataset update configuration**

To learn how to delete your configuration, see [Delete an automatic configuration](#).

**Connect to data sources**

In Amazon SageMaker Canvas, you can import data from a location outside of your local file system through an AWS service, a SaaS platform, or other databases using JDBC connectors. For example, you might want to import tables from a data warehouse in Amazon Redshift, or you might want to import Google Analytics data.

When you go through the **Import** workflow to import data in the Canvas application, you can choose your data source and then select the data that you want to import. For certain data sources, like Snowflake and Amazon Redshift, you must specify your credentials and add a connection to the data source.

The following screenshot shows the data sources toolbar in the **Import** workflow, with all of the available data sources highlighted. You can only import data from the data sources that are available to you. Contact your administrator if your desired data source isn't available.

![Data Sources Toolbar](image)

*How to connect to data sources*
The following sections provide information about establishing connections to external data sources and importing data from them. Review the following section first to determine what permissions you need to import data from your data source.

Permissions

Review the following information to ensure that you have the necessary permissions to import data from your data source:

- **Amazon S3**: You can import data from any Amazon S3 bucket as long as your user has permissions to access the bucket. For more information about using AWS IAM to control access to Amazon S3 buckets, see Identity and access management in Amazon S3 in the Amazon S3 User Guide.

- **Amazon Athena**: If you have the AmazonSageMakerFullAccess policy and the AmazonSageMakerCanvasFullAccess policy attached to your user's execution role, then you can query your AWS Glue Data Catalog with Amazon Athena. If you're part of an Athena workgroup, make sure that the Canvas user has permissions to run Athena queries on the data. For more information, see Using workgroups for running queries in the Amazon Athena User Guide.

- **Amazon DocumentDB**: You can import data from any Amazon DocumentDB database as long as you have the credentials (username and password) to connect to the database and have the minimum base Canvas permissions attached to your user's execution role. For more information about Canvas permissions, see the Prerequisites for setting up Amazon SageMaker Canvas.

- **Amazon Redshift**: To give yourself the necessary permissions to import data from Amazon Redshift, see Grant Users Permissions to Import Amazon Redshift Data.

- **Amazon RDS**: If you have the AmazonSageMakerCanvasFullAccess policy attached to your user's execution role, then you'll be able to access your Amazon RDS databases from Canvas.

- **SaaS platforms**: If you have the AmazonSageMakerFullAccess policy and the AmazonSageMakerCanvasFullAccess policy attached to your user's execution role, then you have the necessary permissions to import data from SaaS platforms. See Use SaaS connectors with Canvas for more information about connecting to a specific SaaS connector.

- **JDBC connectors**: For database sources such as Databricks, MySQL or MariaDB, you must enable username and password authentication on the source database before attempting to connect from Canvas. If you're connecting to a Databricks database, you must have the JDBC URL that contains the necessary credentials.
Connect to a database stored in AWS

You might want to import data that you’ve stored in AWS. You can import data from Amazon S3, use Amazon Athena to query a database in the AWS Glue Data Catalog, import data from Amazon RDS, or make a connection to a provisioned Amazon Redshift database (not Redshift Serverless).

You can create multiple connections to Amazon Redshift. For Amazon Athena, you can access any databases that you have in your AWS Glue Data Catalog. For Amazon S3, you can import data from a bucket as long as you have the necessary permissions.

Review the following sections for more detailed information.

Connect to data in Amazon S3, Amazon Athena, or Amazon RDS

For Amazon S3, you can import data from an Amazon S3 bucket as long as you have permissions to access the bucket.

For Amazon Athena, you can access databases in your AWS Glue Data Catalog as long as you have permissions through your Amazon Athena workgroup.

For Amazon RDS, if you have the AmazonSageMakerCanvasFullAccess policy attached to your user's role, then you'll be able to import data from your Amazon RDS databases into Canvas.

To import data from an Amazon S3 bucket, or to run queries and import data tables with Amazon Athena, see Create a dataset. You can only import tabular data from Amazon Athena, and you can import tabular and image data from Amazon S3.

Connect to an Amazon DocumentDB database

Amazon DocumentDB is a fully managed, serverless, document database service. You can import unstructured document data stored in an Amazon DocumentDB database into SageMaker Canvas as a tabular dataset, and then you can build machine learning models with the data.

⚠️ Important

Your SageMaker domain must be configured in VPC only mode to add connections to Amazon DocumentDB. You can only access Amazon DocumentDB clusters in the same Amazon VPC as your Canvas application. Additionally, Canvas can only connect to TLS-enabled Amazon DocumentDB clusters. For more information about how to set up Canvas in VPC only mode, see Configure Amazon SageMaker Canvas in a VPC without internet access.
To import data from Amazon DocumentDB databases, you must have credentials to access the Amazon DocumentDB database and specify the username and password when creating a database connection. You can configure more granular permissions and restrict access by modifying the Amazon DocumentDB user permissions. To learn more about access control in Amazon DocumentDB, see Database Access Using Role-Based Access Control in the Amazon DocumentDB Developer Guide.

When you import from Amazon DocumentDB, Canvas converts your unstructured data into a tabular dataset by mapping the fields to columns in a table. Additional tables are created for each complex field (or nested structure) in the data, where the columns correspond to the sub-fields of the complex field. For more detailed information about this process and examples of schema conversion, see the Amazon DocumentDB JDBC Driver Schema Discovery GitHub page.

Canvas can only make a connection to a single database in Amazon DocumentDB. To import data from a different database, you must create a new connection.

You can import data from Amazon DocumentDB into Canvas by using the following methods:

- **Create a dataset.** You can import your Amazon DocumentDB data and create a tabular dataset in Canvas. If you choose this method, make sure that you follow the Import tabular data procedure.
- **Create a Data Flow.** You can create a data preparation pipeline in Canvas and add your Amazon DocumentDB database as a data source.

To proceed with importing your data, follow the procedure for one of the methods linked in the preceding list.

When you reach the step in either workflow to choose a data source (Step 5 for creating a dataset, or Step 6 for creating a data flow), do the following:

1. For Data Source, open the dropdown menu and choose DocumentDB.
2. Choose Add connection.
3. In the dialog box, specify your Amazon DocumentDB credentials:
   a. Enter a Connection name. This is a name used by Canvas to identify this connection.
   b. For Cluster, select the cluster in Amazon DocumentDB that stores your data. Canvas automatically populates the dropdown menu with Amazon DocumentDB clusters in the same VPC as your Canvas application.
c. Enter the **Username** for your Amazon DocumentDB cluster.

d. Enter the **Password** for your Amazon DocumentDB cluster.

e. Enter the name of the **Database** to which you want to connect.

f. The **Read preference** option determines which types of instances on your cluster Canvas reads the data from. Select one of the following:

   - **Secondary preferred** – Canvas defaults to reading from the cluster's secondary instances, but if a secondary instance isn't available, then Canvas reads from a primary instance.
   - **Secondary** – Canvas only reads from the cluster's secondary instances, which prevents the read operations from interfering with the cluster's regular read and write operations.

g. Choose **Add connection**. The following image shows the dialog box with the preceding fields for an Amazon DocumentDB connection.
You should now have an Amazon DocumentDB connection, and you can use your Amazon DocumentDB data in Canvas to create either a dataset or a data flow.

**Connect to an Amazon Redshift database**

You can import data from Amazon Redshift, a data warehouse where your organization keeps its data. Before you can import data from Amazon Redshift, the AWS IAM role you use must have the AmazonRedshiftFullAccess managed policy attached. For instructions on how to attach this policy, see [Grant Users Permissions to Import Amazon Redshift Data](#).

To import data from Amazon Redshift, you do the following:

1. Create a connection to an Amazon Redshift database.
2. Choose the data that you're importing.
3. Import the data.

You can use the Amazon Redshift editor to drag datasets onto the import pane and import them into SageMaker Canvas. For more control over the values returned in the dataset, you can use the following:

- SQL queries
- Joins

With SQL queries, you can customize how you import the values in the dataset. For example, you can specify the columns returned in the dataset or the range of values for a column.

You can use joins to combine multiple datasets from Amazon Redshift into a single dataset. You can drag your datasets from Amazon Redshift into the panel that gives you the ability to join the datasets.

You can use the SQL editor to edit the dataset that you've joined and convert the joined dataset into a single node. You can join another dataset to the node. You can import the data that you've selected into SageMaker Canvas.

Use the following procedure to import data from Amazon Redshift.

1. In the SageMaker Canvas application, go to the **Datasets** page.
2. Choose **Create**, and from the dropdown menu, choose **Tabular**.
3. Enter a name for the dataset and choose **Create**.

4. For **Data Source**, open the dropdown menu and choose **Redshift**.

5. Choose **Add connection**.

6. In the dialog box, specify your Amazon Redshift credentials:
   
a. For **Authentication method**, choose **IAM**.
   
b. Enter the **Cluster identifier** to specify to which cluster you want to connect. Enter only the cluster identifier and not the full endpoint of the Amazon Redshift cluster.
   
c. Enter the **Database name** of the database to which you want to connect.
   
d. Enter a **Database user** to identify the user you want to use to connect to the database.
   
e. For **ARN**, enter the IAM role ARN of the role that the Amazon Redshift cluster should assume to move and write data to Amazon S3. For more information about this role, see [Authorizing Amazon Redshift to access other AWS services on your behalf](#) in the *Amazon Redshift Management Guide*.
   
f. Enter a **Connection name**. This is a name used by Canvas to identify this connection.

7. From the tab that has the name of your connection, drag the .csv file that you're importing to the **Drag and drop table to import** pane.

8. Optional: Drag additional tables to the import pane. You can use the GUI to join the tables. For more specificity in your joins, choose **Edit in SQL**.

9. Optional: If you're using SQL to query the data, you can choose **Context** to add context to the connection by specifying values for the following:
   
   - **Warehouse**
   - **Database**
   - **Schema**

10. Choose **Import data**.

The following image shows an example of fields specified for an Amazon Redshift connection.
The following image shows the page used to join datasets in Amazon Redshift.

The following image shows an SQL query being used to edit a join in Amazon Redshift.
Connect to your data with JDBC connectors

With JDBC, you can connect to your databases from sources such as Databricks, SQLServer, MySQL, PostgreSQL, MariaDB, Amazon RDS, and Amazon Aurora.

You must make sure that you have the necessary credentials and permissions to create the connection from Canvas.

- For Databricks, you must provide a JDBC URL. The URL formatting can vary between Databricks instances. For information about finding the URL and the specifying the parameters within it, see JDBC configuration and connection parameters in the Databricks documentation. The following is an example of how a URL can be formatted:
  jdbc:spark://aws-sagemaker-datawrangler.cloud.databricks.com:443/default;transportMode=http;ssl=1;httpPath=sql/protocolv1:o/3122619508517275/0909-200301-cut318;AuthMech=3;UID=token;PWD=personal-access-token

- For other database sources, you must set up username and password authentication, and then specify those credentials when connecting to the database from Canvas.

Additionally, your data source must either be accessible through the public internet, or if your Canvas application is running in VPC only mode, then the data source must run in the same VPC.
For more information about configuring an Amazon RDS database in a VPC, see Amazon VPC VPCs and Amazon RDS in the Amazon RDS User Guide.

After you’ve configured your data source credentials, you can sign in to the Canvas application and create a connection to the data source. Specify your credentials (or, for Databricks, the URL) when creating the connection.

**Connect to data sources with OAuth**

Canvas supports using OAuth as an authentication method for connecting to your data in Snowflake and Salesforce Data Cloud. OAuth is a common authentication platform for granting access to resources without sharing passwords.

⚠️ **Note**

You can only establish one OAuth connection for each data source.

To authorize the connection, you must following the initial setup described in Set up connections to data sources with OAuth.

After setting up the OAuth credentials, you can do the following to add a Snowflake or Salesforce Data Cloud connection with OAuth:

1. Sign in to the Canvas application.
2. Create a tabular dataset. When prompted to upload data, choose Snowflake or Salesforce Data Cloud as your data source.
3. Create a new connection to your Snowflake or Salesforce Data Cloud data source. Specify OAuth as the authentication method and enter your connection details.

You should now be able to import data from your databases in Snowflake or Salesforce Data Cloud.

**Connect to a SaaS platform**

You can import data from Snowflake and over 40 other external SaaS platforms. For a full list of the connectors, see the table on Import data into Canvas.
Use Snowflake with Canvas

Snowflake is a data storage and analytics service, and you can import your data from Snowflake into SageMaker Canvas. For more information about Snowflake, see the Snowflake documentation.

You can import data from your Snowflake account by doing the following:

1. Create a connection to the Snowflake database.
2. Choose the data that you're importing by dragging and dropping the table from the left navigation menu into the editor.
3. Import the data.

You can use the Snowflake editor to drag datasets onto the import pane and import them into SageMaker Canvas. For more control over the values returned in the dataset, you can use the following:

- SQL queries
- Joins

With SQL queries, you can customize how you import the values in the dataset. For example, you can specify the columns returned in the dataset or the range of values for a column.

You can join multiple Snowflake datasets into a single dataset before you import into Canvas using SQL or the Canvas interface. You can drag your datasets from Snowflake into the panel that gives you the ability to join the datasets, or you can edit the joins in SQL and convert the SQL into a single node. You can then combine the datasets that you've joined into a single node and join the nodes to a different Snowflake dataset. Finally, you can import the data that you've selected into Canvas.

Use the following procedure to import data from Snowflake to Amazon SageMaker Canvas.

1. In the SageMaker Canvas application, go to the Datasets page.
2. Choose **Import**.
3. For **Data Source**, open the dropdown menu and choose **Snowflake**.
4. Choose **Add connection**.
5. In the **Add a new Snowflake connection** dialog box, specify your Snowflake credentials. For the **Authentication method**, you can choose **Basic - username password**, **ARN** or **OAuth**. OAuth lets you authenticate without providing a password but requires additional setup. For more information about setting up OAuth credentials for Snowflake, see [Set up connections to data sources with OAuth](#).
6. Choose **Add connection**.
7. From the tab that has the name of your connection, drag the `.csv` file that you're importing to the **Drag and drop table to import** pane.
8. Optional: Drag additional tables to the import pane. You can use the user interface to join the tables. For more specificity in your joins, choose **Edit in SQL**.
9. Optional: If you're using SQL to query the data, you can choose **Context** to add context to the connection by specifying values for the following:
   - **Warehouse**
   - **Database**
   - **Schema**

   Adding context to a connection makes it easier to specify future queries.
10. Choose **Import data**.

   The following image shows an example of fields specified for a Snowflake connection.
The following image shows the page used to add context to a connection.
The following image shows the page used to join datasets in Snowflake.
The following image shows a SQL query being used to edit a join in Snowflake.
Use SaaS connectors with Canvas

Note
For SaaS platforms besides Snowflake, you can only have one connection per data source.

Before you can import data from a SaaS platform, your administrator must authenticate and create a connection to the data source. For more information about how administrators can create a connection with a SaaS platform, see Managing Amazon AppFlow connections in the Amazon AppFlow User Guide.

If you’re an administrator getting started with Amazon AppFlow for the first time, see Getting started in the Amazon AppFlow User Guide.
To import data from a SaaS platform, you can follow the standard Import tabular data procedure, which shows you how to import tabular datasets into Canvas.

**Join data that you've imported into SageMaker Canvas**

*Note*

You can only make joins for tabular datasets in SageMaker Canvas.

You can use Amazon SageMaker Canvas to join multiple datasets into a single dataset. A join combines the two datasets. By default, SageMaker Canvas automatically joins the datasets on their matching column names. The option to combine multiple datasets might give you the ability to get more insight from the models that you build.

You can make the following joins for your datasets:

- **Inner** – Returns a dataset with matching values in both datasets.
- **Left** – Returns a dataset that has:
  - All the rows from the dataset to the left of the join.
  - All the rows from the dataset to the right of the join that have matching values with the columns to the left of the join.
- **Right** – Returns a dataset that has:
  - All the rows from the dataset to the right of the join.
  - All the rows from the dataset to the left of the join that have matching values with the columns to the right of the join.
- **Outer** – Returns all the rows when there is a match in either the left or the right dataset. The dataset from an outer join might have null values that SageMaker Canvas might impute when you build a model.

Use the following procedure to join your datasets.

To join datasets, do the following.

1. Navigate to the **Datasets** page.
2. Choose **Join data**.
3. Drag and drop the datasets that you're joining into the **Drag and drop datasets to join** box.

4. Configure the join. Amazon SageMaker Canvas shows you a preview of the joined data after you configure it.

5. Choose **Save joined data** to save the output of the join.

The following images show the workflow of the preceding procedure.
Join Datasets

Datasets

- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv

Join Preview

<table>
<thead>
<tr>
<th>(selecteddataset1).csv</th>
<th>(selecteddataset2).csv</th>
<th>(selecteddataset3).csv</th>
<th>(selecteddataset4).csv</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sold</td>
<td>ABC</td>
<td>ABC</td>
<td>ABC</td>
</tr>
<tr>
<td>2 categories</td>
<td>2</td>
<td>200.99</td>
<td>ABC</td>
</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Southwest</td>
<td>23</td>
</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Southwest</td>
<td>23</td>
</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Southwest</td>
<td>23</td>
</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Southwest</td>
<td>23</td>
</tr>
</tbody>
</table>

Previewsing first 100 rows

Save joined data

Close
Join Datasets

**Datasets**
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv
- sales-data-May2020.csv

**Join Preview**

<table>
<thead>
<tr>
<th>(selecteddataset1).csv</th>
<th>(selecteddataset2).csv</th>
<th>(selecteddataset2).csv</th>
<th>(selecteddataset1).csv</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sold</strong></td>
<td><strong>Price</strong></td>
<td><strong>Age</strong></td>
<td><strong>Age</strong></td>
</tr>
<tr>
<td>2 categories</td>
<td>2</td>
<td>12 categories</td>
<td>4 categories</td>
</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>Yes</td>
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</tr>
<tr>
<td>Yes</td>
<td>29.99</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

**Join Options**
- Inner
- Left
- Right
- Full

**Key**
- (Left Data Set)  
- (Right Data Set)  

Show dropped columns

Close  Save joined data
Use sample datasets

SageMaker Canvas provides sample datasets addressing unique use cases so you can start building, training, and validating models quickly without writing any code. The use cases associated with these datasets highlight the capabilities of SageMaker Canvas, and you can leverage these datasets to get started with building models. You can find the sample datasets in the **Datasets** page of your SageMaker Canvas application.

Sample datasets

The following datasets are the samples that SageMaker Canvas provides by default. These datasets cover use cases such as predicting house prices, loan defaults, and readmission for diabetic patients; forecasting sales; predicting machine failures to streamline predictive maintenance in manufacturing units; and generating supply chain predictions for transportation and logistics.
The datasets are stored in the `sample_dataset` folder in the default Amazon S3 bucket that SageMaker creates for your account in a Region.

- **canvas-sample-diabetic-readmission.csv**: This dataset contains historical data including over fifteen features with patient and hospital outcomes. You can use this dataset to predict whether high-risk diabetic patients are likely to get readmitted to the hospital within 30 days of discharge, after 30 days, or not at all. Use the `redadmitted` column as the target column, and use the 3+ category prediction model type with this dataset. To learn more about how to build a model with this dataset, see the [SageMaker Canvas workshop page](https://sagemaker.aws). This dataset was obtained from the [UCI Machine Learning Repository](https://archive.ics.uci.edu/ml).  

- **canvas-sample-housing.csv**: This dataset contains data on the characteristics tied to a given housing price. You can use this dataset to predict housing prices. Use the `median_house_value` column as the target column, and use the numeric prediction model type with this dataset. To learn more about building a model with this dataset, see the [SageMaker Canvas workshop page](https://sagemaker.aws). This is the California housing dataset obtained from the [StatLib repository](http://lib.stat.cmu.edu).  

- **canvas-sample-loans.csv**: This dataset contains complete loan data for all loans issued from 2007–2011, including the current loan status and latest payment information. You can use this dataset to predict whether a customer will repay a loan. Use the `loan_status` column as the target column, and use the 3+ category prediction model type with this dataset. To learn more about how to build a model with this dataset, see the [SageMaker Canvas workshop page](https://sagemaker.aws). This data uses the LendingClub data obtained from [Kaggle](https://www.kaggle.com).  

- **canvas-sample-maintenance.csv**: This dataset contains data on the characteristics tied to a given maintenance failure type. You can use this dataset to predict which failure will occur in the future. Use the `Failure Type` column as the target column, and use the 3+ category prediction model type with this dataset. To learn more about how to build a model with this dataset, see the [SageMaker Canvas workshop page](https://sagemaker.aws). This dataset was obtained from the [UCI Machine Learning Repository](https://archive.ics.uci.edu/ml).  

- **canvas-sample-shipping-logs.csv**: This dataset contains complete shipping data for all products delivered, including estimated time shipping priority, carrier, and origin. You can use this dataset to predict the estimated time of arrival of the shipment in number of days. Use the `ActualShippingDays` column as the target column, and use the numeric prediction model type with this dataset. To learn more about how to build a model with this data, see the [SageMaker Canvas workshop page](https://sagemaker.aws). This is a synthetic dataset created by Amazon.  

- **canvas-sample-sales-forecasting.csv**: This dataset contains historical time series sales data for retail stores. You can use this dataset to forecast sales for a particular retail store. Use the `sales` column as the target column, and use the time series forecasting model type with this
dataset. To learn more about how to build a model with this dataset, see the SageMaker Canvas workshop page. This is a synthetic dataset created by Amazon.

Re-import a deleted sample dataset

If you no longer wish to use the sample datasets, you can delete them from the Datasets page of your SageMaker Canvas application. However, these datasets are still stored in the Amazon S3 bucket that you specified as the Canvas storage location, so you can always access them later.

If you used the default Amazon S3 bucket, the bucket name follows the pattern sagemaker-{region}-{account ID}. You can find the sample datasets in the directory path Canvas/sample_dataset.

If you delete a sample dataset from your SageMaker Canvas application and want to access the sample dataset again, use the following procedure.

1. Navigate to the Datasets page in your SageMaker Canvas application.
2. Choose Import data.
3. From the list of Amazon S3 buckets, select the bucket that is your Canvas storage location. If using the default SageMaker-created Amazon S3 bucket, it follows the naming pattern sagemaker-{region}-{account ID}.
4. Select the Canvas folder.
5. Select the sample_dataset folder, which contains all of the sample datasets for SageMaker Canvas.
6. Select the dataset you want to import, and then choose Import data.

Prepare data

Note

Previously, Amazon SageMaker Data Wrangler was part of the SageMaker Studio Classic experience. Now, if you update to using the new Studio experience, you must use SageMaker Canvas to access Data Wrangler and receive the latest feature updates. If you have been using Data Wrangler in Studio Classic until now and want to migrate to Data Wrangler in Canvas, you might have to grant additional permissions so that you can create
and use a Canvas application. For more information, see Migrate from Data Wrangler in Studio Classic to SageMaker Canvas.

Use Amazon SageMaker Data Wrangler in Amazon SageMaker Canvas to prepare, featurize and analyze your data. You can integrate a Data Wrangler data preparation flow into your machine learning (ML) workflows to simplify and streamline data pre-processing and feature engineering using little to no coding. You can also add your own Python scripts and transformations to customize workflows.

- **Data Flow** – Create a data flow to define a series of ML data prep steps. You can use a flow to combine datasets from different data sources, identify the number and types of transformations you want to apply to datasets, and define a data prep workflow that can be integrated into an ML pipeline.

- **Transform** – Clean and transform your dataset using standard transforms like string, vector, and numeric data formatting tools. Featurize your data using transforms like text and date/time embedding and categorical encoding.

- **Generate Data Insights** – Automatically verify data quality and detect abnormalities in your data with Data Wrangler Data Quality and Insights Report.

- **Analyze** – Analyze features in your dataset at any point in your flow. Data Wrangler includes built-in data visualization tools like scatter plots and histograms, as well as data analysis tools like target leakage analysis and quick modeling to understand feature correlation.

- **Export** – Export your data preparation workflow to a different location. The following are example locations:
  - Amazon Simple Storage Service (Amazon S3) bucket
  - Amazon SageMaker Feature Store – Store the features and their data in a centralized store.

- **Automate data preparation** – Create machine learning workflows from your data flow.
  - Amazon SageMaker Model Building Pipelines – Build workflows that manage your SageMaker data preparation, model training, and model deployment jobs.
  - Serial inference pipeline – Create a serial inference pipeline from your data flow. Use it to make predictions on new data.
  - Python script – Store the data and their transformations in a Python script for your custom workflows.
Create a Data Flow

Use a Data Wrangler flow in SageMaker Canvas, or data flow, to create and modify a data preparation pipeline. The datasets, transformations, and analyses that you use in the data flow are represented as steps.

Import data into a data flow

To get started with using a data flow, import your data into it. To use datasets larger than 5 GB, you must import your data directly from the data source instead of using a SageMaker Canvas dataset.

Use the following procedure to import your data into a data flow.

To import your data into a data flow

1. Open SageMaker Canvas.
2. On the left-hand navigation, choose .
3. Choose Data flows.
4. Choose Create.
5. (Optional) For Data flow name, specify a name for the data flow.
6. • To use a SageMaker Canvas dataset that you've already imported into SageMaker Canvas, choose Select existing dataset.
   a. Select the dataset type.
   b. Select the SageMaker Canvas dataset.
   • To import your data directly from a data source, choose Import data.
      a. For Data Source, choose a data source.
      b. Connect to a data source to browse through data and import a dataset. For information about connecting to a data source or importing data, see the following pages:
         • Import data into Canvas
         • Connect to data sources
      c. Choose Import data
d. (Optional) If the first row of your dataset is the header, choose **Use first row as header**.

e. Choose **Import data**.

**The Data Flow UI**

When you import a dataset, the original dataset appears on the data flow and is named **Source**. SageMaker Canvas automatically infers the types of each column in your dataset and creates a new dataframe named **Data types**. You can select this frame to update the inferred data types.

Each time you add a transform step, you create a new dataframe. When multiple transform steps (other than **Join** or **Concatenate**) are added to the same dataset, they are stacked.

**Join** and **Concatenate** create standalone steps that contain the new joined or concatenated dataset.

**Add a Step to Your Data Flow**

Select + next to any dataset or previously added step and then select one of the following options:

- **Edit data types** (For a **Data types** step only): If you have not added any transforms to a **Data types** step, you can select **Edit data types** to update the data types Data Wrangler inferred when importing your dataset.

- **Add transform**: Adds a new transform step. See **Transform data** to learn more about the data transformations you can add.

- **Add analysis**: Adds an analysis. You can use this option to analyze your data at any point in the data flow. See **Perform exploratory data analysis (EDA)** to learn more about the analyses you can add.

- **Join**: Joins two datasets and adds the resulting dataset to the data flow. To learn more, see **Join Datasets**.

- **Concatenate**: Concatenates two datasets and adds the resulting dataset to the data flow. To learn more, see **Concatenate Datasets**.

**Delete a Step from Your Data Flow**

To delete a step, select the + next to the step and select **Delete**. If the node is a node that has a single input, you delete only the step that you select. Deleting a step that has a single input doesn't
delete the steps that follow it. If you’re deleting a step for a source, join, or concatenate node, all the steps that follow it are also deleted.

To delete a step from a stack of steps, select the stack and then select the step you want to delete.

You can use one of the following procedures to delete a step without deleting the downstream steps.

Delete a step in the Data Wrangler flow

You can delete an individual step for nodes in your data flow that have a single input. You can’t delete individual steps for source, join, and concatenate nodes.

Use the following procedure to delete a step in the Data Wrangler flow.

1. Choose the group of steps that has the step that you're deleting.
2. Choose the icon next to the step.
3. Choose Delete step.

Delete a step in the table view

You can delete an individual step for nodes in your data flow that have a single input. You can’t delete individual steps for source, join, and concatenate nodes.

Use the following procedure to delete a step in the table view.

1. Choose the step and open the table view for the step.
2. Move your cursor over the step so the ellipsis icon appears.
3. Choose the icon next to the step.
4. Choose Delete.

Perform exploratory data analysis (EDA)

Data Wrangler includes built-in analyses that help you generate visualizations and data analyses in a few clicks. You can also create custom analyses using your own code.

You add an analysis to a dataframe by selecting a step in your data flow, and then choosing Add analysis. To access an analysis you’ve created, select the step that contains the analysis, and select the analysis.
All analyses are generated using 20,000 rows of your dataset.

You can add the following analysis to a dataframe:

- Data visualizations, including histograms and scatter plots.
- A quick summary of your dataset, including number of entries, minimum and maximum values (for numeric data), and most and least frequent categories (for categorical data).
- A quick model of the dataset, which can be used to generate an importance score for each feature.
- A target leakage report, which you can use to determine if one or more features are strongly correlated with your target feature.
- A custom visualization using your own code.

Use the following sections to learn more about these options.

**Get insights on data and data quality**

Use the [Data Quality and Insights Report](#) to perform an analysis of the data that you've imported into Data Wrangler. We recommend that you create the report after you import your dataset. You can use the report to help you clean and process your data. It gives you information such as the number of missing values and the number of outliers. If you have issues with your data, such as target leakage or imbalance, the insights report can bring those issues to your attention.

Use the following procedure to create a Data Quality and Insights report. It assumes that you've already imported a dataset into your Data Wrangler flow.

**To create a Data Quality and Insights report**

1. Choose a + next to a node in your Data Wrangler flow.
2. Select [Get data insights](#).
3. For Analysis name, specify a name for the insights report.
4. (Optional) For Target column, specify the target column.
5. For Problem type, specify [Regression](#) or [Classification](#).
6. For Data size, specify one of the following:
   - 20 K – Uses the first 20000 rows of the dataset that you've imported to create the report.
• **Entire dataset** – Uses the entire dataset that you've imported to create the report.

**Note**

Creating a Data Quality and Insights report on the entire dataset uses an Amazon SageMaker processing job. A SageMaker processing job provisions the additional compute resources required to get insights for all of your data. For more information about SageMaker processing jobs, see [Process data](#).

7. Choose **Create**.

The following topics show the sections of the report:

**Topics**

- Summary
- Target column
- Quick model
- Feature summary
- Samples
- Definitions

You can either download the report or view it online. To download the report, choose the download button at the top right corner of the screen.

**Summary**

The insights report has a brief summary of the data that includes general information such as missing values, invalid values, feature types, outlier counts, and more. It can also include high severity warnings that point to probable issues with the data. We recommend that you investigate the warnings.

**Target column**

When you create the Data Quality and Insights Report, Data Wrangler gives you the option to select a target column. A target column is a column that you're trying to predict. When you choose a target column, Data Wrangler automatically creates a target column analysis. It also ranks the
features in the order of their predictive power. When you select a target column, you must specify whether you're trying to solve a regression or a classification problem.

For classification, Data Wrangler shows a table and a histogram of the most common classes. A class is a category. It also presents observations, or rows, with a missing or invalid target value.

For regression, Data Wrangler shows a histogram of all the values in the target column. It also presents observations, or rows, with a missing, invalid, or outlier target value.

**Quick model**

The **Quick model** provides an estimate of the expected predicted quality of a model that you train on your data.

Data Wrangler splits your data into training and validation folds. It uses 80% of the samples for training and 20% of the values for validation. For classification, the sample is stratified split. For a stratified split, each data partition has the same ratio of labels. For classification problems, it's important to have the same ratio of labels between the training and classification folds. Data Wrangler trains the XGBoost model with the default hyperparameters. It applies early stopping on the validation data and performs minimal feature preprocessing.

For classification models, Data Wrangler returns both a model summary and a confusion matrix.

To learn more about the information that the classification model summary returns, see **Definitions**.

A confusion matrix gives you the following information:

- The number of times the predicted label matches the true label.
- The number of times the predicted label doesn't match the true label.

The true label represents an actual observation in your data. For example, if you're using a model to detect fraudulent transactions, the true label represents a transaction that is actually fraudulent or non-fraudulent. The predicted label represents the label that your model assigns to the data.

You can use the confusion matrix to see how well the model predicts the presence or the absence of a condition. If you're predicting fraudulent transactions, you can use the confusion matrix to get a sense of both the sensitivity and the specificity of the model. The sensitivity refers to the model's ability to detect fraudulent transactions. The specificity refers to the model's ability to avoid detecting non-fraudulent transactions as fraudulent.
**Feature summary**

When you specify a target column, Data Wrangler orders the features by their prediction power. Prediction power is measured on the data after it is split into 80% training and 20% validation folds. Data Wrangler fits a model for each feature separately on the training fold. It applies minimal feature preprocessing and measures prediction performance on the validation data.

It normalizes the scores to the range \([0, 1]\). Higher prediction scores indicate columns that are more useful for predicting the target on their own. Lower scores point to columns that aren’t predictive of the target column.

It’s uncommon for a column that isn’t predictive on its own to be predictive when it’s used in tandem with other columns. You can confidently use the prediction scores to determine whether a feature in your dataset is predictive.

A low score usually indicates the feature is redundant. A score of 1 implies perfect predictive abilities, which often indicates target leakage. Target leakage usually happens when the dataset contains a column that isn’t available at the prediction time. For example, it could be a duplicate of the target column.

**Samples**

Data Wrangler provides information about whether your samples are anomalous or if there are duplicates in your dataset.

Data Wrangler detects anomalous samples using the isolation forest algorithm. The isolation forest associates an anomaly score with each sample (row) of the dataset. Low anomaly scores indicate anomalous samples. High scores are associated with non-anomalous samples. Samples with a negative anomaly score are usually considered anomalous and samples with positive anomaly score are considered non-anomalous.

When you look at a sample that might be anomalous, we recommend that you pay attention to unusual values. For example, you might have anomalous values that result from errors in gathering and processing the data. The following is an example of the most anomalous samples according to the Data Wrangler’s implementation of the isolation forest algorithm. We recommend using domain knowledge and business logic when you examine the anomalous samples.

Data Wrangler detects duplicate rows and calculates the ratio of duplicate rows in your data. Some data sources could include valid duplicates. Other data sources could have duplicates that point
to problems in data collection. Duplicate samples that result from faulty data collection could interfere with machine learning processes that rely on splitting the data into independent training and validation folds.

The following are elements of the insights report that can be impacted by duplicated samples:

- Quick model
- Prediction power estimation
- Automatic hyperparameter tuning

You can remove duplicate samples from the dataset using the **Drop duplicates** transform under **Manage rows**. Data Wrangler shows you the most frequently duplicated rows.

**Definitions**

The following are definitions for the technical terms that are used in the data insights report.

**Feature types**

The following are the definitions for each of the feature types:

- **Numeric** – Numeric values can be either floats or integers, such as age or income. The machine learning models assume that numeric values are ordered and a distance is defined over them. For example, 3 is closer to 4 than to 10 and 3 < 4 < 10.

- **Categorical** – The column entries belong to a set of unique values, which is usually much smaller than the number of entries in the column. For example, a column of length 100 could contain the unique values Dog, Cat, and Mouse. The values could be numeric, text, or a combination of both. Horse, House, 8, Love, and 3.1 would all be valid values and could be found in the same categorical column. The machine learning model does not assume order or distance on the values of categorical features, as opposed to numeric features, even when all the values are numbers.

- **Binary** – Binary features are a special categorical feature type in which the cardinality of the set of unique values is 2.

- **Text** – A text column contains many non-numeric unique values. In extreme cases, all the elements of the column are unique. In an extreme case, no two entries are the same.

- **Datetime** – A datetime column contains information about the date or time. It can have information about both the date and time.
Feature statistics

The following are definitions for each of the feature statistics:

- **Prediction power** – Prediction power measures how useful the column is in predicting the target.
- **Outliers** (in numeric columns) – Data Wrangler detects outliers using two statistics that are robust to outliers: median and robust standard deviation (RSTD). RSTD is derived by clipping the feature values to the range [5 percentile, 95 percentile] and calculating the standard deviation of the clipped vector. All values larger than median + 5 * RSTD or smaller than median - 5 * RSTD are considered to be outliers.
- **Skew** (in numeric columns) – Skew measures the symmetry of the distribution and is defined as the third moment of the distribution divided by the third power of the standard deviation. The skewness of the normal distribution or any other symmetric distribution is zero. Positive values imply that the right tail of the distribution is longer than the left tail. Negative values imply that the left tail of the distribution is longer than the right tail. As a rule of thumb, a distribution is considered skewed when the absolute value of the skew is larger than 3.
- **Kurtosis** (in numeric columns) – Pearson's kurtosis measures the heaviness of the tail of the distribution. It's defined as the fourth moment of the distribution divided by the square of the second moment. The kurtosis of the normal distribution is 3. Kurtosis values lower than 3 imply that the distribution is concentrated around the mean and the tails are lighter than the tails of the normal distribution. Kurtosis values higher than 3 imply heavier tails or outliers.
- **Missing values** – Null-like objects, empty strings and strings composed of only white spaces are considered missing.
- **Valid values for numeric features or regression target** – All values that you can cast to finite floats are valid. Missing values are not valid.
- **Valid values for categorical, binary, or text features, or for classification target** – All values that are not missing are valid.
- **Datetime features** – All values that you can cast to a datetime object are valid. Missing values are not valid.
- **Invalid values** – Values that are either missing or you can't properly cast. For example, in a numeric column, you can't cast the string "six" or a null value.

Quick model metrics for regression

The following are the definitions for the quick model metrics:
• **R2 or coefficient of determination** – R2 is the proportion of the variation in the target that is predicted by the model. R2 is in the range of \([-\infty, 1]\). 1 is the score of the model that predicts the target perfectly and 0 is the score of the trivial model that always predicts the target mean.

• **MSE or mean squared error** – MSE is in the range \([0, \infty]\). 0 is the score of the model that predicts the target perfectly.

• **MAE or mean absolute error** – MAE is in the range \([0, \infty]\) where 0 is the score of the model that predicts the target perfectly.

• **RMSE or root mean square error** – RMSE is in the range \([0, \infty]\) where 0 is the score of the model that predicts the target perfectly.

• **Max error** – The maximum absolute value of the error over the dataset. Max error is in the range \([0, \infty]\). 0 is the score of the model that predicts the target perfectly.

• **Median absolute error** – Median absolute error is in the range \([0, \infty]\). 0 is the score of the model that predicts the target perfectly.

### Quick model metrics for classification

The following are the definitions for the quick model metrics:

• **Accuracy** – Accuracy is the ratio of samples that are predicted accurately. Accuracy is in the range \([0, 1]\). 0 is the score of the model that predicts all samples incorrectly and 1 is the score of the perfect model.

• **Balanced accuracy** – Balanced accuracy is the ratio of samples that are predicted accurately when the class weights are adjusted to balance the data. All classes are given the same importance, regardless of their frequency. Balanced accuracy is in the range \([0, 1]\). 0 is the score of the model that predicts all samples wrong. 1 is the score of the perfect model.

• **AUC (binary classification)** – This is the area under the receiver operating characteristic curve. AUC is in the range \([0, 1]\) where a random model returns a score of 0.5 and the perfect model returns a score of 1.

• **AUC (OVR)** – For multiclass classification, this is the area under the receiver operating characteristic curve calculated separately for each label using one versus rest. Data Wrangler reports the average of the areas. AUC is in the range \([0, 1]\) where a random model returns a score of 0.5 and the perfect model returns a score of 1.

• **Precision** – Precision is defined for a specific class. Precision is the fraction of true positives out of all the instances that the model classified as that class. Precision is in the range \([0, 1]\).
1 is the score of the model that has no false positives for the class. For binary classification, Data Wrangler reports the precision of the positive class.

- **Recall** – Recall is defined for a specific class. Recall is the fraction of the relevant class instances that are successfully retrieved. Recall is in the range \([0, 1]\). 1 is the score of the model that classifies all the instances of the class correctly. For binary classification, Data Wrangler reports the recall of the positive class.

- **F1** – F1 is defined for a specific class. It's the harmonic mean of the precision and recall. F1 is in the range \([0, 1]\). 1 is the score of the perfect model. For binary classification, Data Wrangler reports the F1 for classes with positive values.

**Textual patterns**

**Patterns** describe the textual format of a string using an easy to read format. The following are examples of textual patterns:

- "\{digits:4-7\}\" describes a sequence of digits that have a length between 4 and 7.
- "\{alnum:5\}\" describes an alpha-numeric string with a length of exactly 5.

Data Wrangler infers the patterns by looking at samples of non-empty strings from your data. It can describe many of the commonly used patterns. The **confidence** expressed as a percentage indicates how much of the data is estimated to match the pattern. Using the textual pattern, you can see which rows in your data you need to correct or drop.

The following describes the patterns that Data Wrangler can recognize:

<table>
<thead>
<tr>
<th>Pattern</th>
<th>Textual Format</th>
</tr>
</thead>
<tbody>
<tr>
<td>{alnum}</td>
<td>Alphanumeric strings</td>
</tr>
<tr>
<td>{any}</td>
<td>Any string of word characters</td>
</tr>
<tr>
<td>{digits}</td>
<td>A sequence of digits</td>
</tr>
<tr>
<td>{lower}</td>
<td>A lowercase word</td>
</tr>
<tr>
<td>{mixed}</td>
<td>A mixed-case word</td>
</tr>
<tr>
<td>{name}</td>
<td>A word beginning with a capital letter</td>
</tr>
<tr>
<td>Pattern</td>
<td>Textual Format</td>
</tr>
<tr>
<td>-------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>{upper}</td>
<td>An uppercase word</td>
</tr>
<tr>
<td>{whitespace}</td>
<td>Whitespace characters</td>
</tr>
</tbody>
</table>

A word character is either an underscore or a character that might appear in a word in any language. For example, the strings 'Hello_word' and 'écoute' both consist of word characters. 'H' and 'é' are both examples of word characters.

**Histogram**

Use histograms to see the counts of feature values for a specific feature. You can inspect the relationships between features using the **Color by** option.

You can use the **Facet by** feature to create histograms of one column, for each value in another column.

**Scatter plot**

Use the **Scatter Plot** feature to inspect the relationship between features. To create a scatter plot, select a feature to plot on the **X axis** and the **Y axis**. Both of these columns must be numeric typed columns.

You can color scatter plots by an additional column.

Additionally, you can facet scatter plots by features.

**Table summary**

Use the **Table Summary** analysis to quickly summarize your data.

For columns with numerical data, including log and float data, a table summary reports the number of entries (count), minimum (min), maximum (max), mean, and standard deviation (stddev) for each column.

For columns with non-numerical data, including columns with string, Boolean, or date/time data, a table summary reports the number of entries (count), least frequent value (min), and most frequent value (max).
Quick model

Use the **Quick Model** visualization to quickly evaluate your data and produce importance scores for each feature. A **feature importance score** score indicates how useful a feature is at predicting a target label. The feature importance score is between [0, 1] and a higher number indicates that the feature is more important to the whole dataset. On the top of the quick model chart, there is a model score. A classification problem shows an F1 score. A regression problem has a mean squared error (MSE) score.

When you create a quick model chart, you select a dataset you want evaluated, and a target label against which you want feature importance to be compared. Data Wrangler does the following:

- Infers the data types for the target label and each feature in the dataset selected.
- Determines the problem type. Based on the number of distinct values in the label column, Data Wrangler determines if this is a regression or classification problem type. Data Wrangler sets a categorical threshold to 100. If there are more than 100 distinct values in the label column, Data Wrangler classifies it as a regression problem; otherwise, it is classified as a classification problem.
- Pre-processes features and label data for training. The algorithm used requires encoding features to vector type and encoding labels to double type.
- Trains a random forest algorithm with 70% of data. Spark’s **RandomForestRegressor** is used to train a model for regression problems. The **RandomForestClassifier** is used to train a model for classification problems.
- Evaluates a random forest model with the remaining 30% of data. Data Wrangler evaluates classification models using an F1 score and evaluates regression models using an MSE score.
- Calculates feature importance for each feature using the Gini importance method.

Target leakage

Target leakage occurs when there is data in a machine learning training dataset that is strongly correlated with the target label, but is not available in real-world data. For example, you may have a column in your dataset that serves as a proxy for the column you want to predict with your model.

When you use the **Target Leakage** analysis, you specify the following:

- **Target**: This is the feature about which you want your ML model to be able to make predictions.
• **Problem type**: This is the ML problem type on which you are working. Problem type can either be **classification** or **regression**.

• *(Optional)* **Max features**: This is the maximum number of features to present in the visualization, which shows features ranked by their risk of being target leakage.

For classification, the target leakage analysis uses the area under the receiver operating characteristic, or AUC - ROC curve for each column, up to **Max features**. For regression, it uses a coefficient of determination, or R2 metric.

The AUC - ROC curve provides a predictive metric, computed individually for each column using cross-validation, on a sample of up to around 1000 rows. A score of 1 indicates perfect predictive abilities, which often indicates target leakage. A score of 0.5 or lower indicates that the information on the column could not provide, on its own, any useful information towards predicting the target. Although it can happen that a column is uninformative on its own but is useful in predicting the target when used in tandem with other features, a low score could indicate the feature is redundant.

**Multicollinearity**

Multicollinearity is a circumstance where two or more predictor variables are related to each other. The predictor variables are the features in your dataset that you're using to predict a target variable. When you have multicollinearity, the predictor variables are not only predictive of the target variable, but also predictive of each other.

You can use the **Variance Inflation Factor (VIF)**, **Principal Component Analysis (PCA)**, or **Lasso feature selection** as measures for the multicollinearity in your data. For more information, see the following.

**Variance Inflation Factor (VIF)**

The Variance Inflation Factor (VIF) is a measure of collinearity among variable pairs. Data Wrangler returns a VIF score as a measure of how closely the variables are related to each other. A VIF score is a positive number that is greater than or equal to 1.

A score of 1 means that the variable is uncorrelated with the other variables. Scores greater than 1 indicate higher correlation.

Theoretically, you can have a VIF score with a value of infinity. Data Wrangler clips high scores to 50. If you have a VIF score greater than 50, Data Wrangler sets the score to 50.
You can use the following guidelines to interpret your VIF scores:

- A VIF score less than or equal to 5 indicates that the variables are moderately correlated with the other variables.
- A VIF score greater than or equal to 5 indicates that the variables are highly correlated with the other variables.

Principle Component Analysis (PCA)

Principal Component Analysis (PCA) measures the variance of the data along different directions in the feature space. The feature space consists of all the predictor variables that you use to predict the target variable in your dataset.

For example, if you're trying to predict who survived on the *RMS Titanic* after it hit an iceberg, your feature space can include the passengers' age, gender, and the fare that they paid.

From the feature space, PCA generates an ordered list of variances. These variances are also known as singular values. The values in the list of variances are greater than or equal to 0. We can use them to determine how much multicollinearity there is in our data.

When the numbers are roughly uniform, the data has very few instances of multicollinearity. When there is a lot of variability among the values, we have many instances of multicollinearity. Before it performs PCA, Data Wrangler normalizes each feature to have a mean of 0 and a standard deviation of 1.

*Note*

PCA in this circumstance can also be referred to as Singular Value Decomposition (SVD).

Lasso feature selection

Lasso feature selection uses the L1 regularization technique to only include the most predictive features in your dataset.

For both classification and regression, the regularization technique generates a coefficient for each feature. The absolute value of the coefficient provides an importance score for the feature. A higher importance score indicates that it is more predictive of the target variable. A common feature selection method is to use all the features that have a non-zero lasso coefficient.
Detect anomalies in time series data

You can use the anomaly detection visualization to see outliers in your time series data. To understand what determines an anomaly, you need to understand that we decompose the time series into a predicted term and an error term. We treat the seasonality and trend of the time series as the predicted term. We treat the residuals as the error term.

For the error term, you specify a threshold as the number of standard of deviations the residual can be away from the mean for it to be considered an anomaly. For example, you can specify a threshold as being 3 standard deviations. Any residual greater than 3 standard deviations away from the mean is an anomaly.

You can use the following procedure to perform an Anomaly detection analysis.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add analysis.
3. For Analysis type, choose Time Series.
4. For Visualization, choose Anomaly detection.
5. For Anomaly threshold, choose the threshold that a value is considered an anomaly.
6. Choose Preview to generate a preview of the analysis.
7. Choose Add to add the transform to the Data Wrangler data flow.

Seasonal trend decomposition in time series data

You can determine whether there's seasonality in your time series data by using the Seasonal Trend Decomposition visualization. We use the STL (Seasonal Trend decomposition using LOESS) method to perform the decomposition. We decompose the time series into its seasonal, trend, and residual components. The trend reflects the long term progression of the series. The seasonal component is a signal that recurs in a time period. After removing the trend and the seasonal components from the time series, you have the residual.

You can use the following procedure to perform a Seasonal-Trend decomposition analysis.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add analysis.
3. For Analysis type, choose Time Series.
4. For Visualization, choose Seasonal-Trend decomposition.
5. For **Anomaly threshold**, choose the threshold that a value is considered an anomaly.

6. Choose **Preview** to generate a preview of the analysis.

7. Choose **Add** to add the transform to the Data Wrangler data flow.

### Create custom visualizations

You can add an analysis to your Data Wrangler flow to create a custom visualization. Your dataset, with all the transformations you've applied, is available as a **Pandas DataFrame**. Data Wrangler uses the `df` variable to store the dataframe. You access the dataframe by calling the variable.

You must provide the output variable, `chart`, to store an **Altair** output chart. For example, you can use the following code block to create a custom histogram using the Titanic dataset.

```python
import altair as alt
df = df.iloc[:30]
df = df.rename(columns={'Age': 'value'})
df = df.assign(count=df.groupby('value').value.transform('count'))
df = df[['value', 'count']]  
base = alt.Chart(df)  
bar = base.mark_bar().encode(x=alt.X('value', bin=True, axis=None), y=alt.Y('count'))
rule = base.mark_rule(color='red').encode(
    x='mean(value):Q',
    size=alt.value(5))
chart = bar + rule
```

To create a custom visualization:

1. Next to the node containing the transformation that you'd like to visualize, choose the +.

2. Choose **Add analysis**.

3. For **Analysis type**, choose **Custom Visualization**.

4. For **Analysis name**, specify a name.

5. Enter your code in the code box.

6. Choose **Preview** to preview your visualization.

7. Choose **Save** to add your visualization.

If you don’t know how to use the Altair visualization package in Python, you can use custom code snippets to help you get started.
Data Wrangler has a searchable collection of visualization snippets. To use a visualization snippet, choose **Search example snippets** and specify a query in the search bar.

The following example uses the **Binned scatterplot** code snippet. It plots a histogram for 2 dimensions.

The snippets have comments to help you understand the changes that you need to make to the code. You usually need to specify the column names of your dataset in the code.

```python
import altair as alt

# Specify the number of top rows for plotting
rows_number = 1000
df = df.head(rows_number)
# You can also choose bottom rows or randomly sampled rows
# df = df.tail(rows_number)
# df = df.sample(rows_number)

chart = (alt.Chart(df)
 .mark_circle()
 .encode(
     x=alt.X("col1:Q", bin=alt.Bin(maxbins=20)),
     y=alt.Y("col2:Q", bin=alt.Bin(maxbins=20)),
     size="count()",
 )

# :Q specifies that label column has quantitative type.
# For more details on Altair typing refer to
# https://altair-viz.github.io/user_guide/encoding.html#encoding-data-types
```

**Transform data**

Amazon SageMaker Data Wrangler provides numerous ML data transforms to streamline cleaning, transforming, and featurizing your data. When you add a transform, it adds a step to the data flow. Each transform you add modifies your dataset and produces a new dataframe. All subsequent transforms apply to the resulting dataframe.
Data Wrangler includes built-in transforms, which you can use to transform columns without any code. You can also add custom transformations using PySpark, Python (User-Defined Function), pandas, and PySpark SQL. Some transforms operate in place, while others create a new output column in your dataset.

You can apply transforms to multiple columns at once. For example, you can delete multiple columns in a single step.

You can apply the **Process numeric** and **Handle missing** transforms only to a single column.

Use this page to learn more about these built-in and custom transforms.

**Transform UI**

Most of the built-in transforms are located in the **Prepare** tab of the Data Wrangler UI. You can access the join and concatenate transforms through the data flow view. Use the following table to preview these two views.

**Transform**

You can add a transform to any step in your data flow. Use the following procedure to add a transform to your data flow.

To add a step to your data flow, do the following.

1. Choose the + next to the step in the data flow.
2. Choose **Add transform**.
3. Choose **Add step**.
4. Choose a transform.
5. (Optional) You can search for the transform that you want to use. Data Wrangler highlights the query in the results.

**Join View**

To join two datasets, select the first dataset in your data flow and choose **Join**. When you choose **Join**, your left and right datasets are displayed in the left panel. The main panel displays your data flow, with the newly joined dataset added.

When you choose **Configure** to configure your join, you see results similar to those shown in the following image. Your join configuration is displayed in the left panel. You can use this panel to
choose the joined dataset name, join type, and columns to join. The main panel displays three tables. The top two tables display the left and right datasets on the left and right respectively. Under this table, you can preview the joined dataset.

See Join Datasets to learn more.

Concatenate View

To concatenate two datasets, you select the first dataset in your data flow and choose Concatenate. Your left and right datasets are displayed in the left panel. The main panel displays your data flow, with the newly concatenated dataset added.

When you choose Configure to configure your concatenation, you see results similar to those shown in the following image. Your concatenate configuration displays in the left panel. You can use this panel to choose the concatenated dataset's name, and choose to remove duplicates after concatenation and add columns to indicate the source dataframe. The main panel displays three tables. The top two tables display the left and right datasets on the left and right respectively. Under this table, you can preview the concatenated dataset.

See Concatenate Datasets to learn more.

Join Datasets

You join dataframes directly in your data flow. When you join two datasets, the resulting joined dataset appears in your flow. The following join types are supported by Data Wrangler.

- **Left Outer** – Include all rows from the left table. If the value for the column joined on a left table row does not match any right table row values, that row contains null values for all right table columns in the joined table.
- **Left Anti** – Include rows from the left table that do not contain values in the right table for the joined column.
- **Left semi** – Include a single row from the left table for all identical rows that satisfy the criteria in the join statement. This excludes duplicate rows from the left table that match the criteria of the join.
- **Right Outer** – Include all rows from the right table. If the value for the joined column in a right table row does not match any left table row values, that row contains null values for all left table columns in the joined table.
- **Inner** – Include rows from left and right tables that contain matching values in the joined column.
• **Full Outer** – Include all rows from the left and right tables. If the row value for the joined column in either table does not match, separate rows are created in the joined table. If a row doesn’t contain a value for a column in the joined table, null is inserted for that column.

• **Cartesian Cross** – Include rows which combine each row from the first table with each row from the second table. This is a *Cartesian product* of rows from tables in the join. The result of this product is the size of the left table times the size of the right table. Therefore, we recommend caution in using this join between very large datasets.

Use the following procedure to join two dataframes.

1. Select + next to the left dataframe that you want to join. The first dataframe you select is always the left table in your join.
2. Choose **Join**.
3. Select the right dataframe. The second dataframe you select is always the right table in your join.
4. Choose **Configure** to configure your join.
5. Give your joined dataset a name using the **Name** field.
6. Select a **Join type**.
7. Select a column from the left and right tables to join.
8. Choose **Apply** to preview the joined dataset on the right.
9. To add the joined table to your data flow, choose **Add**.

**Concatenate Datasets**

**Concatenate two datasets:**

1. Choose + next to the left dataframe that you want to concatenate. The first dataframe you select is always the left table in your concatenate.
2. Choose **Concatenate**.
3. Select the right dataframe. The second dataframe you select is always the right table in your concatenate.
4. Choose **Configure** to configure your concatenate.
5. Give your concatenated dataset a name using the **Name** field.
6. (Optional) Select the checkbox next to **Remove duplicates after concatenation** to remove duplicate columns.

7. (Optional) Select the checkbox next to **Add column to indicate source dataframe** if, for each column in the new dataset, you want to add an indicator of the column's source.

8. Choose **Apply** to preview the new dataset.

9. Choose **Add** to add the new dataset to your data flow.

**Balance Data**

You can balance the data for datasets with an underrepresented category. Balancing a dataset can help you create better models for binary classification.

**Note**

You can't balance datasets containing column vectors.

You can use the **Balance data** operation to balance your data using one of the following operators:

- **Random oversampling** – Randomly duplicates samples in the minority category. For example, if you're trying to detect fraud, you might only have cases of fraud in 10% of your data. For an equal proportion of fraudulent and non-fraudulent cases, this operator randomly duplicates fraud cases within the dataset 8 times.

- **Random undersampling** – Roughly equivalent to random oversampling. Randomly removes samples from the overrepresented category to get the proportion of samples that you desire.

- **Synthetic Minority Oversampling Technique (SMOTE)** – Uses samples from the underrepresented category to interpolate new synthetic minority samples. For more information about SMOTE, see the following description.

You can use all transforms for datasets containing both numeric and non-numeric features. SMOTE interpolates values by using neighboring samples. Data Wrangler uses the R-squared distance to determine the neighborhood to interpolate the additional samples. Data Wrangler only uses numeric features to calculate the distances between samples in the underrepresented group.

For two real samples in the underrepresented group, Data Wrangler interpolates the numeric features by using a weighted average. It randomly assigns weights to those samples in the range of
[0, 1]. For numeric features, Data Wrangler interpolates samples using a weighted average of the samples. For samples A and B, Data Wrangler could randomly assign a weight of 0.7 to A and 0.3 to B. The interpolated sample has a value of 0.7A + 0.3B.

Data Wrangler interpolates non-numeric features by copying from either of the interpolated real samples. It copies the samples with a probability that it randomly assigns to each sample. For samples A and B, it can assign probabilities 0.8 to A and 0.2 to B. For the probabilities it assigned, it copies A 80% of the time.

Custom Transforms

The **Custom Transforms** group allows you to use Python (User-Defined Function), PySpark, pandas, or PySpark (SQL) to define custom transformations. For all three options, you use the variable `df` to access the dataframe to which you want to apply the transform. To apply your custom code to your dataframe, assign the dataframe with the transformations that you've made to the `df` variable. If you're not using Python (User-Defined Function), you don't need to include a return statement. Choose **Preview** to preview the result of the custom transform. Choose **Add** to add the custom transform to your list of **Previous steps**.

You can import the popular libraries with an `import` statement in the custom transform code block, such as the following:

- NumPy version 1.19.0
- scikit-learn version 0.23.2
- SciPy version 1.5.4
- pandas version 1.0.3
- PySpark version 3.0.0

⚠️ **Important**

**Custom transform** doesn't support columns with spaces or special characters in the name. We recommend that you specify column names that only have alphanumeric characters and underscores. You can use the **Rename column** transform in the **Manage columns** transform group to remove spaces from a column's name. You can also add a **Python (Pandas) Custom transform** similar to the following to remove spaces from multiple columns in a single step. This example changes columns named `A column` and `B column` to `A_column` and `B_column` respectively.
If you include print statements in the code block, the result appears when you select **Preview**. You can resize the custom code transformer panel. Resizing the panel provides more space to write code.

The following sections provide additional context and examples for writing custom transform code.

**Python (User-Defined Function)**

The Python function gives you the ability to write custom transformations without needing to know Apache Spark or pandas. Data Wrangler is optimized to run your custom code quickly. You get similar performance using custom Python code and an Apache Spark plugin.

To use the Python (User-Defined Function) code block, you specify the following:

- **Input column** – The input column where you're applying the transform.
- **Mode** – The scripting mode, either pandas or Python.
- **Return type** – The data type of the value that you're returning.

Using the pandas mode gives better performance. The Python mode makes it easier for you to write transformations by using pure Python functions.

**PySpark**

The following example extracts date and time from a timestamp.

```python
from pyspark.sql.functions import from_unixtime, to_date, date_format
df = df.withColumn('DATE_TIME', from_unixtime('TIMESTAMP'))
df = df.withColumn('EVENT_DATE', to_date('DATE_TIME')).withColumn('EVENT_TIME', date_format('DATE_TIME', 'HH:mm:ss'))
```

**pandas**

The following example provides an overview of the dataframe to which you are adding transforms.

```python
df.info()
```
**PySpark (SQL)**

The following example creates a new dataframe with four columns: *name, fare, pclass, survived*.

```sql
SELECT name, fare, pclass, survived FROM df
```

If you don't know how to use PySpark, you can use custom code snippets to help you get started.

Data Wrangler has a searchable collection of code snippets. You can use code snippets to perform tasks such as dropping columns, grouping by columns, or modelling.

To use a code snippet, choose **Search example snippets** and specify a query in the search bar. The text you specify in the query doesn’t have to match the name of the code snippet exactly.

The following example shows a **Drop duplicate rows** code snippet that can delete rows with similar data in your dataset. You can find the code snippet by searching for one of the following:

- Duplicates
- Identical
- Remove

The following snippet has comments to help you understand the changes that you need to make. For most snippets, you must specify the column names of your dataset in the code.

```python
# Specify the subset of columns
# all rows having identical values in these columns will be dropped

subset = ["col1", "col2", "col3"]
df = df.dropDuplicates(subset)

# to drop the full-duplicate rows run
# df = df.dropDuplicates()
```

To use a snippet, copy and paste its content into the **Custom transform** field. You can copy and paste multiple code snippets into the custom transform field.
Amazon SageMaker

Developer Guide

Custom Formula
Use Custom formula to deﬁne a new column using a Spark SQL expression to query data in the
current dataframe. The query must use the conventions of Spark SQL expressions.

Important
Custom formula doesn't support columns with spaces or special characters in the name.
We recommend that you specify column names that only have alphanumeric characters
and underscores. You can use the Rename column transform in the Manage columns
transform group to remove spaces from a column's name. You can also add a Python
(Pandas) Custom transform similar to the following to remove spaces from multiple
columns in a single step. This example changes columns named A column and B column
to A_column and B_column respectively.
df.rename(columns={"A column": "A_column", "B column": "B_column"})

You can use this transform to perform operations on columns, referencing the columns by name.
For example, assuming the current dataframe contains columns named col_a and col_b, you
can use the following operation to produce an Output column that is the product of these two
columns with the following code:
col_a * col_b

Other common operations include the following, assuming a dataframe contains col_a and col_b
columns:
• Concatenate two columns: concat(col_a, col_b)
• Add two columns: col_a + col_b
• Subtract two columns: col_a - col_b
• Divide two columns: col_a / col_b
• Take the absolute value of a column: abs(col_a)
For more information, see the Spark documentation on selecting data.
Prepare data

869


Reduce Dimensionality within a Dataset

Reduce the dimensionality in your data by using Principal Component Analysis (PCA). The dimensionality of your dataset corresponds to the number of features. When you use dimensionality reduction in Data Wrangler, you get a new set of features called components. Each component accounts for some variability in the data.

The first component accounts for the largest amount of variation in the data. The second component accounts for the second largest amount of variation in the data, and so on.

You can use dimensionality reduction to reduce the size of the data sets that you use to train models. Instead of using the features in your dataset, you can use the principal components instead.

To perform PCA, Data Wrangler creates axes for your data. An axis is an affine combination of columns in your dataset. The first principal component is the value on the axis that has the largest amount of variance. The second principal component is the value on the axis that has the second largest amount of variance. The nth principal component is the value on the axis that has the nth largest amount of variance.

You can configure the number of principal components that Data Wrangler returns. You can either specify the number of principal components directly or you can specify the variance threshold percentage. Each principal component explains an amount of variance in the data. For example, you might have a principal component with a value of 0.5. The component would explain 50% of the variation in the data. When you specify a variance threshold percentage, Data Wrangler returns the smallest number of components that meet the percentage that you specify.

The following are example principal components with the amount of variance that they explain in the data.

- Component 1 – 0.5
- Component 2 – 0.45
- Component 3 – 0.05

If you specify a variance threshold percentage of 94 or 95, Data Wrangler returns Component 1 and Component 2. If you specify a variance threshold percentage of 96, Data Wrangler returns all three principal components.

You can use the following procedure to run PCA on your dataset.
To run PCA on your dataset, do the following.

1. Open your Data Wrangler data flow.
2. Choose the +, and select **Add transform**.
3. Choose **Add step**.
4. Choose **Dimensionality Reduction**.
5. For **Input Columns**, choose the features that you're reducing into the principal components.
6. (Optional) For **Number of principal components**, choose the number of principal components that Data Wrangler returns in your dataset. If specify a value for the field, you can't specify a value for **Variance threshold percentage**.
7. (Optional) For **Variance threshold percentage**, specify the percentage of variation in the data that you want explained by the principal components. Data Wrangler uses the default value of 95 if you don't specify a value for the variance threshold. You can't specify a variance threshold percentage if you've specified a value for **Number of principal components**.
8. (Optional) Deselect **Center** to not use the mean of the columns as the center of the data. By default, Data Wrangler centers the data with the mean before scaling.
9. (Optional) Deselect **Scale** to not scale the data with the unit standard deviation.
10. (Optional) Choose **Columns** to output the components to separate columns. Choose **Vector** to output the components as a single vector.
11. (Optional) For **Output column**, specify a name for an output column. If you're outputting the components to separate columns, the name that you specify is a prefix. If you're outputting the components to a vector, the name that you specify is the name of the vector column.
12. (Optional) Select **Keep input columns**. We don't recommend selecting this option if you plan on only using the principal components to train your model.
13. Choose **Preview**.
14. Choose **Add**.

**Encode Categorical**

Categorical data is usually composed of a finite number of categories, where each category is represented with a string. For example, if you have a table of customer data, a column that indicates the country a person lives in is categorical. The categories would be Afghanistan, Albania, Algeria, and so on. Categorical data can be **nominal or ordinal**. Ordinal categories have an inherent
order, and nominal categories do not. The highest degree obtained (High school, Bachelors, Masters, and so on) is an example of ordinal categories.

Encoding categorical data is the process of creating a numerical representation for categories. For example, if your categories are Dog and Cat, you may encode this information into two vectors, [1, 0] to represent Dog, and [0, 1] to represent Cat.

When you encode ordinal categories, you may need to translate the natural order of categories into your encoding. For example, you can represent the highest degree obtained with the following map: {"High school": 1, "Bachelors": 2, "Masters": 3}.

Use categorical encoding to encode categorical data that is in string format into arrays of integers.

The Data Wrangler categorical encoders create encodings for all categories that exist in a column at the time the step is defined. If new categories have been added to a column when you start a Data Wrangler job to process your dataset at time $t$, and this column was the input for a Data Wrangler categorical encoding transform at time $t-1$, these new categories are considered missing in the Data Wrangler job. The option you select for Invalid handling strategy is applied to these missing values. Examples of when this can occur are:

- When you use a .flow file to create a Data Wrangler job to process a dataset that was updated after the creation of the data flow. For example, you may use a data flow to regularly process sales data each month. If that sales data is updated weekly, new categories may be introduced into columns for which an encode categorical step is defined.
- When you select Sampling when you import your dataset, some categories may be left out of the sample.

In these situations, these new categories are considered missing values in the Data Wrangler job.

You can choose from and configure an ordinal and a one-hot encode. Use the following sections to learn more about these options.

Both transforms create a new column named Output column name. You specify the output format of this column with Output style:

- Select Vector to produce a single column with a sparse vector.
- Select Columns to create a column for every category with an indicator variable for whether the text in the original column contains a value that is equal to that category.
Ordinal Encode

Select **Ordinal encode** to encode categories into an integer between 0 and the total number of categories in the **Input column** you select.

**Invalid handing strategy:** Select a method to handle invalid or missing values.

- **Choose Skip** if you want to omit the rows with missing values.
- **Choose Keep** to retain missing values as the last category.
- **Choose Error** if you want Data Wrangler to throw an error if missing values are encountered in the **Input column**.
- **Choose Replace with NaN** to replace missing with NaN. This option is recommended if your ML algorithm can handle missing values. Otherwise, the first three options in this list may produce better results.

One-Hot Encode

Select **One-hot encode** for **Transform** to use one-hot encoding. Configure this transform using the following:

- **Drop last category:** If **True**, the last category does not have a corresponding index in the one-hot encoding. When missing values are possible, a missing category is always the last one and setting this to **True** means that a missing value results in an all zero vector.
- **Invalid handing strategy:** Select a method to handle invalid or missing values.
  - **Choose Skip** if you want to omit the rows with missing values.
  - **Choose Keep** to retain missing values as the last category.
  - **Choose Error** if you want Data Wrangler to throw an error if missing values are encountered in the **Input column**.
- **Is input ordinal encoded:** Select this option if the input vector contains ordinal encoded data. This option requires that input data contain non-negative integers. If **True**, input $i$ is encoded as a vector with a non-zero in the $i$th location.

Similarity encode

Use similarity encoding when you have the following:

- A large number of categorical variables
Noisy data

The similarity encoder creates embeddings for columns with categorical data. An embedding is a mapping of discrete objects, such as words, to vectors of real numbers. It encodes similar strings to vectors containing similar values. For example, it creates very similar encodings for "California" and "Calfornia".

Data Wrangler converts each category in your dataset into a set of tokens using a 3-gram tokenizer. It converts the tokens into an embedding using min-hash encoding.

The similarity encodings that Data Wrangler creates:

- Have low dimensionality
- Are scalable to a large number of categories
- Are robust and resistant to noise

For the preceding reasons, similarity encoding is more versatile than one-hot encoding.

To add the similarity encoding transform to your dataset, use the following procedure.

To use similarity encoding, do the following.

1. Sign in to the Amazon SageMaker Console.
2. Choose Open Studio Classic.
3. Choose Launch app.
4. Choose Studio.
5. Specify your data flow.
6. Choose a step with a transformation.
7. Choose Add step.
8. Choose Encode categorical.
9. Specify the following:
   - Transform – Similarity encode
   - Input column – The column containing the categorical data that you're encoding.
• **Target dimension** – (Optional) The dimension of the categorical embedding vector. The default value is 30. We recommend using a larger target dimension if you have a large dataset with many categories.

• **Output style** – Choose **Vector** for a single vector with all of the encoded values. Choose **Column** to have the encoded values in separate columns.

• **Output column** – (Optional) The name of the output column for a vector encoded output. For a column-encoded output, this is the prefix of the column names followed by listed number.

**Featurize Text**

Use the **Featurize Text** transform group to inspect string-typed columns and use text embedding to featurize these columns.

This feature group contains two features, **Character statistics** and **Vectorize**. Use the following sections to learn more about these transforms. For both options, the **Input column** must contain text data (string type).

**Character Statistics**

Use **Character statistics** to generate statistics for each row in a column containing text data.

This transform computes the following ratios and counts for each row, and creates a new column to report the result. The new column is named using the input column name as a prefix and a suffix that is specific to the ratio or count.

• **Number of words**: The total number of words in that row. The suffix for this output column is -stats_word_count.

• **Number of characters**: The total number of characters in that row. The suffix for this output column is -stats_char_count.

• **Ratio of upper**: The number of uppercase characters, from A to Z, divided by all characters in the column. The suffix for this output column is -stats_capital_ratio.

• **Ratio of lower**: The number of lowercase characters, from a to z, divided by all characters in the column. The suffix for this output column is -stats_lower_ratio.

• **Ratio of digits**: The ratio of digits in a single row over the sum of digits in the input column. The suffix for this output column is -stats_digit_ratio.
• **Special characters ratio**: The ratio of non-alphanumeric (characters like #$&%:@) characters to over the sum of all characters in the input column. The suffix for this output column is `-stats_special_ratio`.

**Vectorize**

Text embedding involves mapping words or phrases from a vocabulary to vectors of real numbers. Use the Data Wrangler text embedding transform to tokenize and vectorize text data into term frequency–inverse document frequency (TF-IDF) vectors.

When TF-IDF is calculated for a column of text data, each word in each sentence is converted to a real number that represents its semantic importance. Higher numbers are associated with less frequent words, which tend to be more meaningful.

When you define a **Vectorize** transform step, Data Wrangler uses the data in your dataset to define the count vectorizer and TF-IDF methods. Running a Data Wrangler job uses these same methods.

You configure this transform using the following:

• **Output column name**: This transform creates a new column with the text embedding. Use this field to specify a name for this output column.

• **Tokenizer**: A tokenizer converts the sentence into a list of words, or *tokens*.

  Choose **Standard** to use a tokenizer that splits by white space and converts each word to lowercase. For example, "Good dog" is tokenized to ["good", "dog"].

  Choose **Custom** to use a customized tokenizer. If you choose **Custom**, you can use the following fields to configure the tokenizer:

  • **Minimum token length**: The minimum length, in characters, for a token to be valid. Defaults to 1. For example, if you specify 3 for minimum token length, words like a, at, in are dropped from the tokenized sentence.

  • **Should regex split on gaps**: If selected, regex splits on gaps. Otherwise, it matches tokens. Defaults to True.

  • **Regex pattern**: Regex pattern that defines the tokenization process. Defaults to \s+.

  • **To lowercase**: If chosen, Data Wrangler converts all characters to lowercase before tokenization. Defaults to True.

  To learn more, see the Spark documentation on [Tokenizer](#).
• **Vectorizer**: The vectorizer converts the list of tokens into a sparse numeric vector. Each token corresponds to an index in the vector and a non-zero indicates the existence of the token in the input sentence. You can choose from two vectorizer options, *Count* and *Hashing*.

  • **Count vectorize** allows customizations that filter infrequent or too common tokens. **Count vectorize parameters** include the following:

    • **Minimum term frequency**: In each row, terms (tokens) with smaller frequency are filtered. If you specify an integer, this is an absolute threshold (inclusive). If you specify a fraction between 0 (inclusive) and 1, the threshold is relative to the total term count. Defaults to 1.

    • **Minimum document frequency**: Minimum number of rows in which a term (token) must appear to be included. If you specify an integer, this is an absolute threshold (inclusive). If you specify a fraction between 0 (inclusive) and 1, the threshold is relative to the total term count. Defaults to 1.

    • **Maximum document frequency**: Maximum number of documents (rows) in which a term (token) can appear to be included. If you specify an integer, this is an absolute threshold (inclusive). If you specify a fraction between 0 (inclusive) and 1, the threshold is relative to the total term count. Defaults to 0.999.

    • **Maximum vocabulary size**: Maximum size of the vocabulary. The vocabulary is made up of all terms (tokens) in all rows of the column. Defaults to 262144.

    • **Binary outputs**: If selected, the vector outputs do not include the number of appearances of a term in a document, but rather are a binary indicator of its appearance. Defaults to False.

To learn more about this option, see the Spark documentation on [CountVectorizer](https://spark.apache.org/docs/latest/ml-features.html#countvectorizer).

• **Hashing** is computationally faster. **Hash vectorize parameters** includes the following:

  • **Number of features during hashing**: A hash vectorizer maps tokens to a vector index according to their hash value. This feature determines the number of possible hash values. Large values result in fewer collisions between hash values but a higher dimension output vector.

To learn more about this option, see the Spark documentation on [FeatureHasher](https://spark.apache.org/docs/latest/ml-features.html#feature hasher).

• **Apply IDF** applies an IDF transformation, which multiplies the term frequency with the standard inverse document frequency used for TF-IDF embedding. **IDF parameters** include the following:

  • **Minimum document frequency**: Minimum number of documents (rows) in which a term (token) must appear to be included. If count_vectorize is the chosen vectorizer, we recommend that you keep the default value and only modify the min_doc_freq field in Count vectorize parameters. Defaults to 5.
• **Output format**: The output format of each row.
  - Select **Vector** to produce a single column with a sparse vector.
  - Select **Flattened** to create a column for every category with an indicator variable for whether the text in the original column contains a value that is equal to that category. You can only choose flattened when **Vectorizer** is set as **Count vectorizer**.

**Transform Time Series**

In Data Wrangler, you can transform time series data. The values in a time series dataset are indexed to specific time. For example, a dataset that shows the number of customers in a store for each hour in a day is a time series dataset. The following table shows an example of a time series dataset.

**Hourly number of customers in a store**

<table>
<thead>
<tr>
<th>Number of customers</th>
<th>Time (hour)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>09:00</td>
</tr>
<tr>
<td>10</td>
<td>10:00</td>
</tr>
<tr>
<td>14</td>
<td>11:00</td>
</tr>
<tr>
<td>25</td>
<td>12:00</td>
</tr>
<tr>
<td>20</td>
<td>13:00</td>
</tr>
<tr>
<td>18</td>
<td>14:00</td>
</tr>
</tbody>
</table>

For the preceding table, the **Number of Customers** column contains the time series data. The time series data is indexed on the hourly data in the **Time (hour)** column.

You might need to perform a series of transformations on your data to get it in a format that you can use for your analysis. Use the **Time series** transform group to transform your time series data. For more information about the transformations that you can perform, see the following sections.

**Topics**

- [Group by a Time Series](#)
- [Resample Time Series Data](#)
Group by a Time Series

You can use the group by operation to group time series data for specific values in a column.

For example, you have the following table that tracks the average daily electricity usage in a household.

**Average daily household electricity usage**

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Daily timestamp</th>
<th>Electricity usage (kWh)</th>
<th>Number of household occupants</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>1/1/2020</td>
<td>30</td>
<td>2</td>
</tr>
<tr>
<td>household_0</td>
<td>1/2/2020</td>
<td>40</td>
<td>2</td>
</tr>
<tr>
<td>household_0</td>
<td>1/4/2020</td>
<td>35</td>
<td>3</td>
</tr>
<tr>
<td>household_1</td>
<td>1/2/2020</td>
<td>45</td>
<td>3</td>
</tr>
<tr>
<td>household_1</td>
<td>1/3/2020</td>
<td>55</td>
<td>4</td>
</tr>
</tbody>
</table>

If you choose to group by ID, you get the following table.

**Electricity usage grouped by household ID**

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
<th>Number of household occupants series</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35]</td>
<td>[2, 2, 3]</td>
</tr>
</tbody>
</table>
Each entry in the time series sequence is ordered by the corresponding timestamp. The first element of the sequence corresponds to the first timestamp of the series. For household_0, 30 is the first value of the Electricity Usage Series. The value of 30 corresponds to the first timestamp of 1/1/2020.

You can include the starting timestamp and ending timestamp. The following table shows how that information appears.

### Electricity usage grouped by household ID

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
<th>Number of household occupants series</th>
<th>Start_time</th>
<th>End_time</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35]</td>
<td>[2, 2, 3]</td>
<td>1/1/2020</td>
<td>1/4/2020</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, 55]</td>
<td>[3, 4]</td>
<td>1/2/2020</td>
<td>1/3/2020</td>
</tr>
</tbody>
</table>

You can use the following procedure to group by a time series column.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add transform.
3. Choose Add step.
5. Under Transform, choose Group by.
6. Specify a column in Group by this column.
7. For Apply to columns, specify a value.
8. Choose Preview to generate a preview of the transform.
9. Choose Add to add the transform to the Data Wrangler data flow.
Resample Time Series Data

Time series data usually has observations that aren't taken at regular intervals. For example, a dataset could have some observations that are recorded hourly and other observations that are recorded every two hours.

Many analyses, such as forecasting algorithms, require the observations to be taken at regular intervals. Resampling gives you the ability to establish regular intervals for the observations in your dataset.

You can either upsample or downsample a time series. Downsampling increases the interval between observations in the dataset. For example, if you downsample observations that are taken either every hour or every two hours, each observation in your dataset is taken every two hours. The hourly observations are aggregated into a single value using an aggregation method such as the mean or median.

Upsampling reduces the interval between observations in the dataset. For example, if you upsample observations that are taken every two hours into hourly observations, you can use an interpolation method to infer hourly observations from the ones that have been taken every two hours. For information on interpolation methods, see pandas.DataFrame.interpolate.

You can resample both numeric and non-numeric data.

Use the Resample operation to resample your time series data. If you have multiple time series in your dataset, Data Wrangler standardizes the time interval for each time series.

The following table shows an example of downsampling time series data by using the mean as the aggregation method. The data is downsamples from every two hours to every hour.

**Hourly temperature readings over a day before downsampling**

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Temperature (Celsius)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:00</td>
<td>30</td>
</tr>
<tr>
<td>1:00</td>
<td>32</td>
</tr>
<tr>
<td>2:00</td>
<td>35</td>
</tr>
<tr>
<td>3:00</td>
<td>32</td>
</tr>
<tr>
<td>4:00</td>
<td>30</td>
</tr>
</tbody>
</table>
Temperature readings downsampled to every two hours

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Temperature (Celsius)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:00</td>
<td>30</td>
</tr>
<tr>
<td>2:00</td>
<td>33.5</td>
</tr>
<tr>
<td>4:00</td>
<td>35</td>
</tr>
</tbody>
</table>

You can use the following procedure to resample time series data.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add transform.
3. Choose Add step.
5. For Timestamp, choose the timestamp column.
6. For Frequency unit, specify the frequency that you're resampling.
7. (Optional) Specify a value for Frequency quantity.
8. Configure the transform by specifying the remaining fields.
9. Choose Preview to generate a preview of the transform.
10. Choose Add to add the transform to the Data Wrangler data flow.

Handle Missing Time Series Data

If you have missing values in your dataset, you can do one of the following:

- For datasets that have multiple time series, drop the time series that have missing values that are greater than a threshold that you specify.
- Impute the missing values in a time series by using other values in the time series.

Imputing a missing value involves replacing the data by either specifying a value or by using an inferential method. The following are the methods that you can use for imputation:

- Constant value – Replace all the missing data in your dataset with a value that you specify.
• Most common value – Replace all the missing data with the value that has the highest frequency in the dataset.

• Forward fill – Use a forward fill to replace the missing values with the non-missing value that precedes the missing values. For the sequence: [2, 4, 7, NaN, NaN, NaN, 8], all of the missing values are replaced with 7. The sequence that results from using a forward fill is [2, 4, 7, 7, 7, 7, 8].

• Backward fill – Use a backward fill to replace the missing values with the non-missing value that follows the missing values. For the sequence: [2, 4, 7, NaN, NaN, NaN, 8], all of the missing values are replaced with 8. The sequence that results from using a backward fill is [2, 4, 7, 8, 8, 8, 8].

• Interpolate – Uses an interpolation function to impute the missing values. For more information on the functions that you can use for interpolation, see `pandas.DataFrame.interpolate`.

Some of the imputation methods might not be able to impute of all the missing value in your dataset. For example, a Forward fill can't impute a missing value that appears at the beginning of the time series. You can impute the values by using either a forward fill or a backward fill.

You can either impute missing values within a cell or within a column.

The following example shows how values are imputed within a cell.

### Electricity usage with missing values

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35, NaN, NaN]</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, NaN, 55]</td>
</tr>
</tbody>
</table>

### Electricity usage with values imputed using a forward fill

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35, 35, 35]</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, 45, 55]</td>
</tr>
</tbody>
</table>

The following example shows how values are imputed within a column.
Average daily household electricity usage with missing values

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>30</td>
</tr>
<tr>
<td>household_0</td>
<td>40</td>
</tr>
<tr>
<td>household_0</td>
<td>NaN</td>
</tr>
<tr>
<td>household_1</td>
<td>NaN</td>
</tr>
<tr>
<td>household_1</td>
<td>NaN</td>
</tr>
</tbody>
</table>

Average daily household electricity usage with values imputed using a forward fill

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>30</td>
</tr>
<tr>
<td>household_0</td>
<td>40</td>
</tr>
<tr>
<td>household_0</td>
<td>40</td>
</tr>
<tr>
<td>household_1</td>
<td>40</td>
</tr>
<tr>
<td>household_1</td>
<td>40</td>
</tr>
</tbody>
</table>

You can use the following procedure to handle missing values.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add transform.
3. Choose Add step.
4. Choose Handle missing.
5. For Time series input type, choose whether you want to handle missing values inside of a cell or along a column.
6. For Impute missing values for this column, specify the column that has the missing values.
7. For Method for imputing values, select a method.
8. Configure the transform by specifying the remaining fields.

9. Choose Preview to generate a preview of the transform.

10. If you have missing values, you can specify a method for imputing them under Method for imputing values.

11. Choose Add to add the transform to the Data Wrangler data flow.

Validate the Timestamp of Your Time Series Data

You might have time stamp data that is invalid. You can use the Validate time stamp function to determine whether the timestamps in your dataset are valid. Your timestamp can be invalid for one or more of the following reasons:

- Your timestamp column has missing values.
- The values in your timestamp column are not formatted correctly.

If you have invalid timestamps in your dataset, you can't perform your analysis successfully. You can use Data Wrangler to identify invalid timestamps and understand where you need to clean your data.

The time series validation works in one of the two ways:

You can configure Data Wrangler to do one of the following if it encounters missing values in your dataset:

- Drop the rows that have the missing or invalid values.
- Identify the rows that have the missing or invalid values.
- Throw an error if it finds any missing or invalid values in your dataset.

You can validate the timestamps on columns that either have the timestamp type or the string type. If the column has the string type, Data Wrangler converts the type of the column to timestamp and performs the validation.

You can use the following procedure to validate the timestamps in your dataset.

1. Open your Data Wrangler data flow.

2. In your data flow, under Data types, choose the +, and select Add transform.
3. Choose **Add step**.

4. Choose **Validate timestamps**.

5. For **Timestamp Column**, choose the timestamp column.

6. For **Policy**, choose whether you want to handle missing timestamps.

7. (Optional) For **Output column**, specify a name for the output column.

8. If the date time column is formatted for the string type, choose **Cast to datetime**.

9. Choose **Preview** to generate a preview of the transform.

10. Choose **Add** to add the transform to the Data Wrangler data flow.

**Standardizing the Length of the Time Series**

If you have time series data stored as arrays, you can standardize each time series to the same length. Standardizing the length of the time series array might make it easier for you to perform your analysis on the data.

You can standardize your time series for data transformations that require the length of your data to be fixed.

Many ML algorithms require you to flatten your time series data before you use them. Flattening time series data is separating each value of the time series into its own column in a dataset. The number of columns in a dataset can't change, so the lengths of the time series need to be standardized between you flatten each array into a set of features.

Each time series is set to the length that you specify as a quantile or percentile of the time series set. For example, you can have three sequences that have the following lengths:

- 3
- 4
- 5

You can set the length of all of the sequences as the length of the sequence that has the 50th percentile length.

Time series arrays that are shorter than the length you've specified have missing values added. The following is an example format of standardizing the time series to a longer length: [2, 4, 5, NaN, NaN, NaN].
You can use different approaches to handle the missing values. For information on those approaches, see Handle Missing Time Series Data.

The time series arrays that are longer than the length that you specify are truncated.

You can use the following procedure to standardize the length of the time series.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add transform.
3. Choose Add step.
4. Choose Standardize length.
5. For Standardize the time series length for the column, choose a column.
6. (Optional) For Output column, specify a name for the output column. If you don’t specify a name, the transform is done in place.
7. If the datetime column is formatted for the string type, choose Cast to datetime.
8. Choose Cutoff quantile and specify a quantile to set the length of the sequence.
9. Choose Flatten the output to output the values of the time series into separate columns.
10. Choose Preview to generate a preview of the transform.
11. Choose Add to add the transform to the Data Wrangler data flow.

Extract Features from Your Time Series Data

If you’re running a classification or a regression algorithm on your time series data, we recommend extracting features from the time series before running the algorithm. Extracting features might improve the performance of your algorithm.

Use the following options to choose how you want to extract features from your data:

- Use Minimal subset to specify extracting 8 features that you know are useful in downstream analyses. You can use a minimal subset when you need to perform computations quickly. You can also use it when your ML algorithm has a high risk of overfitting and you want to provide it with fewer features.
- Use Efficient subset to specify extracting the most features possible without extracting features that are computationally intensive in your analyses.
- Use All features to specify extracting all features from the tune series.
• Use **Manual subset** to choose a list of features that you think explain the variation in your data well.

Use the following the procedure to extract features from your time series data.

1. Open your Data Wrangler data flow.
2. In your data flow, under **Data types**, choose the +, and select **Add transform**.
3. Choose **Add step**.
4. Choose **Extract features**.
5. For **Extract features for this column**, choose a column.
6. (Optional) Select **Flatten** to output the features into separate columns.
7. For **Strategy**, choose a strategy to extract the features.
8. Choose **Preview** to generate a preview of the transform.
9. Choose **Add** to add the transform to the Data Wrangler data flow.

**Use Lagged Features from Your Time Series Data**

For many use cases, the best way to predict the future behavior of your time series is to use its most recent behavior.

The most common uses of lagged features are the following:

• Collecting a handful of past values. For example, for time, t + 1, you collect t, t - 1, t - 2, and t - 3.

• Collecting values that correspond to seasonal behavior in the data. For example, to predict the occupancy in a restaurant at 1:00 PM, you might want to use the features from 1:00 PM on the previous day. Using the features from 12:00 PM or 11:00 AM on the same day might not be as predictive as using the features from previous days.

1. Open your Data Wrangler data flow.
2. In your data flow, under **Data types**, choose the +, and select **Add transform**.
3. Choose **Add step**.
4. Choose **Lag features**.
5. For **Generate lag features for this column**, choose a column.
6. For **Timestamp Column**, choose the column containing the timestamps.
7. For **Lag**, specify the duration of the lag.
8. (Optional) Configure the output using one of the following options:
   - **Include the entire lag window**
   - **Flatten the output**
   - **Drop rows without history**
9. Choose **Preview** to generate a preview of the transform.
10. Choose **Add** to add the transform to the Data Wrangler data flow.

**Create a Datetime Range In Your Time Series**

You might have time series data that don’t have timestamps. If you know that the observations were taken at regular intervals, you can generate timestamps for the time series in a separate column. To generate timestamps, you specify the value for the start timestamp and the frequency of the timestamps.

For example, you might have the following time series data for the number of customers at a restaurant.

**Time series data on the number of customers at a restaurant**

<table>
<thead>
<tr>
<th>Number of customers</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
</tr>
<tr>
<td>14</td>
</tr>
<tr>
<td>24</td>
</tr>
<tr>
<td>40</td>
</tr>
<tr>
<td>30</td>
</tr>
<tr>
<td>20</td>
</tr>
</tbody>
</table>

If you know that the restaurant opened at 5:00 PM and that the observations are taken hourly, you can add a timestamp column that corresponds to the time series data. You can see the timestamp column in the following table.
Time series data on the number of customers at a restaurant

<table>
<thead>
<tr>
<th>Number of customers</th>
<th>Timestamp</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>1:00 PM</td>
</tr>
<tr>
<td>14</td>
<td>2:00 PM</td>
</tr>
<tr>
<td>24</td>
<td>3:00 PM</td>
</tr>
<tr>
<td>40</td>
<td>4:00 PM</td>
</tr>
<tr>
<td>30</td>
<td>5:00 PM</td>
</tr>
<tr>
<td>20</td>
<td>6:00 PM</td>
</tr>
</tbody>
</table>

Use the following procedure to add a datetime range to your data.

1. Open your Data Wrangler data flow.
2. In your data flow, under **Data types**, choose the +, and select **Add transform**.
3. Choose **Add step**.
4. Choose **Datetime range**.
5. For **Frequency type**, choose the unit used to measure the frequency of the timestamps.
6. For **Starting timestamp**, specify the start timestamp.
7. For **Output column**, specify a name for the output column.
8. (Optional) Configure the output using the remaining fields.
9. Choose **Preview** to generate a preview of the transform.
10. Choose **Add** to add the transform to the Data Wrangler data flow.

**Use a Rolling Window In Your Time Series**

You can extract features over a time period. For example, for time, $t$, and a time window length of 3, and for the row that indicates the $t$th timestamp, we append the features that are extracted from the time series at times $t - 3$, $t - 2$, and $t - 1$. For information on extracting features, see [Extract Features from Your Time Series Data](#).

You can use the following procedure to extract features over a time period.
1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add transform.
3. Choose Add step.
4. Choose Rolling window features.
5. For Generate rolling window features for this column, choose a column.
6. For Timestamp Column, choose the column containing the timestamps.
7. (Optional) For Output Column, specify the name of the output column.
8. For Window size, specify the window size.
9. For Strategy, choose the extraction strategy.
10. Choose Preview to generate a preview of the transform.
11. Choose Add to add the transform to the Data Wrangler data flow.

**Featurize Datetime**

Use Featurize date/time to create a vector embedding representing a datetime field. To use this transform, your datetime data must be in one of the following formats:

- Strings describing datetime: For example, "January 1st, 2020, 12:44pm".
- A Unix timestamp: A Unix timestamp describes the number of seconds, milliseconds, microseconds, or nanoseconds from 1/1/1970.

You can choose to Infer datetime format and provide a Datetime format. If you provide a datetime format, you must use the codes described in the Python documentation. The options you select for these two configurations have implications for the speed of the operation and the final results.

- The most manual and computationally fastest option is to specify a Datetime format and select No for Infer datetime format.
- To reduce manual labor, you can choose Infer datetime format and not specify a datetime format. It is also a computationally fast operation; however, the first datetime format encountered in the input column is assumed to be the format for the entire column. If there are other formats in the column, these values are NaN in the final output. Inferring the datetime format can give you unparsed strings.
• If you don't specify a format and select No for Infer datetime format, you get the most robust results. All the valid datetime strings are parsed. However, this operation can be an order of magnitude slower than the first two options in this list.

When you use this transform, you specify an Input column which contains datetime data in one of the formats listed above. The transform creates an output column named Output column name. The format of the output column depends on your configuration using the following:

• **Vector**: Outputs a single column as a vector.
• **Columns**: Creates a new column for every feature. For example, if the output contains a year, month, and day, three separate columns are created for year, month, and day.

Additionally, you must choose an **Embedding mode**. For linear models and deep networks, we recommend choosing cyclic. For tree-based algorithms, we recommend choosing ordinal.

**Format String**

The **Format string** transforms contain standard string formatting operations. For example, you can use these operations to remove special characters, normalize string lengths, and update string casing.

This feature group contains the following transforms. All transforms return copies of the strings in the Input column and add the result to a new, output column.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left pad</td>
<td>Left-pad the string with a given Fill character to the given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Right pad</td>
<td>Right-pad the string with a given Fill character to the given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Center (pad on either side)</td>
<td>Center-pad the string (add padding on both sides of the string) with a given Fill character</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Prepend zeros</td>
<td>Left-fill a numeric string with zeros, up to a given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Strip left and right</td>
<td>Returns a copy of the string with the leading and trailing characters removed.</td>
</tr>
<tr>
<td>Strip characters from left</td>
<td>Returns a copy of the string with leading characters removed.</td>
</tr>
<tr>
<td>Strip characters from right</td>
<td>Returns a copy of the string with trailing characters removed.</td>
</tr>
<tr>
<td>Lower case</td>
<td>Convert all letters in text to lowercase.</td>
</tr>
<tr>
<td>Upper case</td>
<td>Convert all letters in text to uppercase.</td>
</tr>
<tr>
<td>Capitalize</td>
<td>Capitalize the first letter in each sentence.</td>
</tr>
<tr>
<td>Swap case</td>
<td>Converts all uppercase characters to lowercase and all lowercase characters to uppercase characters of the given string, and returns it.</td>
</tr>
<tr>
<td>Add prefix or suffix</td>
<td>Adds a prefix and a suffix the string column. You must specify at least one of Prefix and Suffix.</td>
</tr>
<tr>
<td>Remove symbols</td>
<td>Removes given symbols from a string. All listed characters are removed. Defaults to white space.</td>
</tr>
</tbody>
</table>
Handle Outliers

Machine learning models are sensitive to the distribution and range of your feature values. Outliers, or rare values, can negatively impact model accuracy and lead to longer training times. Use this feature group to detect and update outliers in your dataset.

When you define a Handle outliers transform step, the statistics used to detect outliers are generated on the data available in Data Wrangler when defining this step. These same statistics are used when running a Data Wrangler job.

Use the following sections to learn more about the transforms this group contains. You specify an Output name and each of these transforms produces an output column with the resulting data.

Robust standard deviation numeric outliers

This transform detects and fixes outliers in numeric features using statistics that are robust to outliers.

You must define an Upper quantile and a Lower quantile for the statistics used to calculate outliers. You must also specify the number of Standard deviations from which a value must vary from the mean to be considered an outlier. For example, if you specify 3 for Standard deviations, a value must fall more than 3 standard deviations from the mean to be considered an outlier.

The Fix method is the method used to handle outliers when they are detected. You can choose from the following:

- **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.
- **Remove**: Use this option to remove rows with outliers from the dataframe.
- **Invalidate**: Use this option to replace outliers with invalid values.

Standard Deviation Numeric Outliers

This transform detects and fixes outliers in numeric features using the mean and standard deviation.

You specify the number of Standard deviations a value must vary from the mean to be considered an outlier. For example, if you specify 3 for Standard deviations, a value must fall more than 3 standard deviations from the mean to be considered an outlier.

The Fix method is the method used to handle outliers when they are detected. You can choose from the following:
• **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.

• **Remove**: Use this option to remove rows with outliers from the dataframe.

• **Invalidate**: Use this option to replace outliers with invalid values.

**Quantile Numeric Outliers**

Use this transform to detect and fix outliers in numeric features using quantiles. You can define an **Upper quantile** and a **Lower quantile**. All values that fall above the upper quantile or below the lower quantile are considered outliers.

The **Fix method** is the method used to handle outliers when they are detected. You can choose from the following:

• **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.

• **Remove**: Use this option to remove rows with outliers from the dataframe.

• **Invalidate**: Use this option to replace outliers with invalid values.

**Min-Max Numeric Outliers**

This transform detects and fixes outliers in numeric features using upper and lower thresholds. Use this method if you know threshold values that demark outliers.

You specify a **Upper threshold** and a **Lower threshold**, and if values fall above or below those thresholds respectively, they are considered outliers.

The **Fix method** is the method used to handle outliers when they are detected. You can choose from the following:

• **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.

• **Remove**: Use this option to remove rows with outliers from the dataframe.

• **Invalidate**: Use this option to replace outliers with invalid values.

**Replace Rare**

When you use the **Replace rare** transform, you specify a threshold and Data Wrangler finds all values that meet that threshold and replaces them with a string that you specify. For example, you may want to use this transform to categorize all outliers in a column into an "Others" category.
• **Replacement string**: The string with which to replace outliers.

• **Absolute threshold**: A category is rare if the number of instances is less than or equal to this absolute threshold.

• **Fraction threshold**: A category is rare if the number of instances is less than or equal to this fraction threshold multiplied by the number of rows.

• **Max common categories**: Maximum not-rare categories that remain after the operation. If the threshold does not filter enough categories, those with the top number of appearances are classified as not rare. If set to 0 (default), there is no hard limit to the number of categories.

**Handle Missing Values**

Missing values are a common occurrence in machine learning datasets. In some situations, it is appropriate to impute missing data with a calculated value, such as an average or categorically common value. You can process missing values using the **Handle missing values** transform group. This group contains the following transforms.

**Fill Missing**

Use the **Fill missing** transform to replace missing values with a **Fill value** you define.

**Impute Missing**

Use the **Impute missing** transform to create a new column that contains imputed values where missing values were found in input categorical and numerical data. The configuration depends on your data type.

For numeric data, choose an imputing strategy, the strategy used to determine the new value to impute. You can choose to impute the mean or the median over the values that are present in your dataset. Data Wrangler uses the value that it computes to impute the missing values.

For categorical data, Data Wrangler imputes missing values using the most frequent value in the column. To impute a custom string, use the **Fill missing** transform instead.

**Add Indicator for Missing**

Use the **Add indicator for missing** transform to create a new indicator column, which contains a Boolean "false" if a row contains a value, and "true" if a row contains a missing value.

**Drop Missing**

Use the **Drop missing** option to drop rows that contain missing values from the **Input column**.
Manage Columns

You can use the following transforms to quickly update and manage columns in your dataset:

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drop Column</td>
<td>Delete a column.</td>
</tr>
<tr>
<td>Duplicate Column</td>
<td>Duplicate a column.</td>
</tr>
<tr>
<td>Rename Column</td>
<td>Rename a column.</td>
</tr>
<tr>
<td>Move Column</td>
<td>Move a column's location in the dataset.</td>
</tr>
<tr>
<td></td>
<td>Choose to move your column to the start or</td>
</tr>
<tr>
<td></td>
<td>end of the dataset, before or after a reference</td>
</tr>
<tr>
<td></td>
<td>column, or to a specific index.</td>
</tr>
</tbody>
</table>

Manage Rows

Use this transform group to quickly perform sort and shuffle operations on rows. This group contains the following:

- **Sort**: Sort the entire dataframe by a given column. Select the check box next to *Ascending order* for this option; otherwise, deselect the check box and descending order is used for the sort.
- **Shuffle**: Randomly shuffle all rows in the dataset.

Manage Vectors

Use this transform group to combine or flatten vector columns. This group contains the following transforms.

- **Assemble**: Use this transform to combine Spark vectors and numeric data into a single column. For example, you can combine three columns: two containing numeric data and one containing vectors. Add all the columns you want to combine in *Input columns* and specify a *Output column name* for the combined data.
- **Flatten**: Use this transform to flatten a single column containing vector data. The input column must contain PySpark vectors or array-like objects. You can control the number of columns created by specifying a *Method to detect number of outputs*. For example, if you select *Length*
of first vector, the number of elements in the first valid vector or array found in the column determines the number of output columns that are created. All other input vectors with too many items are truncated. Inputs with too few items are filled with NaNs.

You also specify an Output prefix, which is used as the prefix for each output column.

Process Numeric

Use the Process Numeric feature group to process numeric data. Each scalar in this group is defined using the Spark library. The following scalars are supported:

- **Standard Scaler**: Standardize the input column by subtracting the mean from each value and scaling to unit variance. To learn more, see the Spark documentation for `StandardScaler`.
- **Robust Scaler**: Scale the input column using statistics that are robust to outliers. To learn more, see the Spark documentation for `RobustScaler`.
- **Min Max Scaler**: Transform the input column by scaling each feature to a given range. To learn more, see the Spark documentation for `MinMaxScaler`.
- **Max Absolute Scaler**: Scale the input column by dividing each value by the maximum absolute value. To learn more, see the Spark documentation for `MaxAbsScaler`.

Sampling

After you've imported your data, you can use the Sampling transformer to take one or more samples of it. When you use the sampling transformer, Data Wrangler samples your original dataset.

You can choose one of the following sample methods:

- **Limit**: Samples the dataset starting from the first row up to the limit that you specify.
- **Randomized**: Takes a random sample of a size that you specify.
- **Stratified**: Takes a stratified random sample.

You can stratify a randomized sample to make sure that it represents the original distribution of the dataset.

You might be performing data preparation for multiple use cases. For each use case, you can take a different sample and apply a different set of transformations.
The following procedure describes the process of creating a random sample.

To take a random sample from your data.

1. Choose the + to the right of the dataset that you've imported. The name of your dataset is located below the +.
2. Choose Add transform.
3. Choose Sampling.
4. For Sampling method, choose the sampling method.
5. For Approximate sample size, choose the approximate number of observations that you want in your sample.
6. (Optional) Specify an integer for Random seed to create a reproducible sample.

The following procedure describes the process of creating a stratified sample.

To take a stratified sample from your data.

1. Choose the + to the right of the dataset that you've imported. The name of your dataset is located below the +.
2. Choose Add transform.
3. Choose Sampling.
4. For Sampling method, choose the sampling method.
5. For Approximate sample size, choose the approximate number of observations that you want in your sample.
6. For Stratify column, specify the name of the column that you want to stratify on.
7. (Optional) Specify an integer for Random seed to create a reproducible sample.

**Search and Edit**

Use this section to search for and edit specific patterns within strings. For example, you can find and update strings within sentences or documents, split strings by delimiters, and find occurrences of specific strings.

The following transforms are supported under Search and edit. All transforms return copies of the strings in the Input column and add the result to a new output column.
<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Find substring</td>
<td>Returns the index of the first occurrence of the Substring for which you searched. You can start and end the search at Start and End respectively.</td>
</tr>
<tr>
<td>Find substring (from right)</td>
<td>Returns the index of the last occurrence of the Substring for which you searched. You can start and end the search at Start and End respectively.</td>
</tr>
<tr>
<td>Matches prefix</td>
<td>Returns a Boolean value if the string contains a given Pattern. A pattern can be a character sequence or regular expression. Optionally, you can make the pattern case sensitive.</td>
</tr>
<tr>
<td>Find all occurrences</td>
<td>Returns an array with all occurrences of a given pattern. A pattern can be a character sequence or regular expression.</td>
</tr>
<tr>
<td>Extract using regex</td>
<td>Returns a string that matches a given Regex pattern.</td>
</tr>
<tr>
<td>Extract between delimiters</td>
<td>Returns a string with all characters found between Left delimiter and Right delimiter.</td>
</tr>
<tr>
<td>Extract from position</td>
<td>Returns a string, starting from Start position in the input string, that contains all characters up to the start position plus Length.</td>
</tr>
<tr>
<td>Find and replace substring</td>
<td>Returns a string with all matches of a given Pattern (regular expression) replaced by Replacement string.</td>
</tr>
<tr>
<td>Replace between delimiters</td>
<td>Returns a string with the substring found between the first appearance of a Left delimiter and the last appearance of a Right delimiter.</td>
</tr>
</tbody>
</table>
### Name | Function
--- | ---
**delimiter** replaced by **Replacement string**. If no match is found, nothing is replaced.

**Replace from position**

Returns a string with the substring between **Start position** and **Start position** plus **Length** replaced by **Replacement string**. If **Start position** plus **Length** is greater than the length of the replacement string, the output contains ....

**Convert regex to missing**

Converts a string to None if invalid and returns the result. Validity is defined with a regular expression in **Pattern**.

**Split string by delimiter**

Returns an array of strings from the input string, split by **Delimiter**, with up to **Max number of splits** (optional). The delimiter defaults to white space.

---

### Split data

Use the **Split data** transform to split your dataset into two or three datasets. For example, you can split your dataset into a dataset used to train your model and a dataset used to test it. You can determine the proportion of the dataset that goes into each split. For example, if you’re splitting one dataset into two datasets, the training dataset can have 80% of the data while the testing dataset has 20%.

Splitting your data into three datasets gives you the ability to create training, validation, and test datasets. You can see how well the model performs on the test dataset by dropping the target column.

Your use case determines how much of the original dataset each of your datasets get and the method you use to split the data. For example, you might want to use a stratified split to make sure that the distribution of the observations in the target column are the same across datasets. You can use the following split transforms:
• Randomized split — Each split is a random, non-overlapping sample of the original dataset. For larger datasets, using a randomized split might be computationally expensive and take longer than an ordered split.

• Ordered split – Splits the dataset based on the sequential order of the observations. For example, for an 80/20 train-test split, the first observations that make up 80% of the dataset go to the training dataset. The last 20% of the observations go to the testing dataset. Ordered splits are effective in keeping the existing order of the data between splits.

• Stratified split – Splits the dataset to make sure that the number of observations in the input column have proportional representation. For an input column that has the observations 1, 1, 1, 1, 2, 2, 2, 2, 2, 3, 3, 3, 3, 3, 3, 3, 3, an 80/20 split on the column would mean that approximately 80% of the 1s, 80% of the 2s, and 80% of the 3s go to the training set. About 20% of each type of observation go to the testing set.

• Split by key – Avoids data with the same key occurring in more than one split. For example, if you have a dataset with the column 'customer_id' and you’re using it as a key, no customer id is in more than one split.

After you split the data, you can apply additional transformations to each dataset. For most use cases, they aren't necessary.

Data Wrangler calculates the proportions of the splits for performance. You can choose an error threshold to set the accuracy of the splits. Lower error thresholds more accurately reflect the proportions that you specify for the splits. If you set a higher error threshold, you get better performance, but lower accuracy.

For perfectly split data, set the error threshold to 0. You can specify a threshold between 0 and 1 for better performance. If you specify a value greater than 1, Data Wrangler interprets that value as 1.

If you have 10000 rows in your dataset and you specify an 80/20 split with an error of 0.001, you would get observations approximating one of the following results:

• 8010 observations in the training set and 1990 in the testing set
• 7990 observations in the training set and 2010 in the testing set

The number of observations for the testing set in the preceding example is in the interval between 8010 and 7990.
By default, Data Wrangler uses a random seed to make the splits reproducible. You can specify a different value for the seed to create a different reproducible split.

Randomized split

Use the following procedure to perform a randomized split on your dataset.

To split your dataset randomly, do the following

1. Choose the + next to the node containing the dataset that you're splitting.
2. Choose **Add transform**.
3. Choose **Split data**.
4. (Optional) For Splits, specify the names and proportions of each split. The proportions must sum to 1.
5. (Optional) Choose the + to create an additional split.
   - Specify the names and proportions of all the splits. The proportions must sum to 1.
6. (Optional) Specify a value for **Error threshold** other than the default value.
7. (Optional) Specify a value for **Random seed**.
8. Choose **Preview**.
9. Choose **Add**.

Ordered split

Use the following procedure to perform an ordered split on your dataset.

To make an ordered split in your dataset, do the following.

1. Choose the + next to the node containing the dataset that you're splitting.
2. Choose **Add transform**.
3. For **Transform**, choose **Ordered split**.
4. Choose **Split data**.
5. (Optional) For Splits, specify the names and proportions of each split. The proportions must sum to 1.
6. (Optional) Choose the + to create an additional split.
• Specify the names and proportions of all the splits. The proportions must sum to 1.

7. (Optional) Specify a value for **Error threshold** other than the default value.

8. (Optional) For **Input column**, specify a column with numeric values. Uses the values of the columns to infer which records are in each split. The smaller values are in one split with the larger values in the other splits.

9. (Optional) Select **Handle duplicates** to add noise to duplicate values and create a dataset of entirely unique values.

10. (Optional) Specify a value for **Random seed**.

11. Choose **Preview**.

12. Choose **Add**.

**Stratified split**

Use the following procedure to perform a stratified split on your dataset.

To make a stratified split in your dataset, do the following.

1. Choose the + next to the node containing the dataset that you're splitting.

2. Choose **Add transform**.

3. Choose **Split data**.

4. For **Transform**, choose **Stratified split**.

5. (Optional) For **Splits**, specify the names and proportions of each split. The proportions must sum to 1.

6. (Optional) Choose the + to create an additional split.

   • Specify the names and proportions of all the splits. The proportions must sum to 1.

7. For **Input column**, specify a column with up to 100 unique values. Data Wrangler can't stratify a column with more than 100 unique values.

8. (Optional) Specify a value for **Error threshold** other than the default value.

9. (Optional) Specify a value for **Random seed** to specify a different seed.

10. Choose **Preview**.

11. Choose **Add**.
Split by column keys

Use the following procedure to split by the column keys in your dataset.

To split by the column keys in your dataset, do the following.

1. Choose the + next to the node containing the dataset that you're splitting.
2. Choose **Add transform**.
3. Choose **Split data**.
4. For **Transform**, choose **Split by key**.
5. (Optional) For **Splits**, specify the names and proportions of each split. The proportions must sum to 1.
6. (Optional) Choose the + to create an additional split.
   - Specify the names and proportions of all the splits. The proportions must sum to 1.
7. For **Key columns**, specify the columns with values that you don't want to appear in both datasets.
8. (Optional) Specify a value for **Error threshold** other than the default value.
9. Choose **Preview**.
10. Choose **Add**.

**Parse Value as Type**

Use this transform to cast a column to a new type. The supported Data Wrangler data types are:

- Long
- Float
- Boolean
- Date, in the format dd-MM-yyyy, representing day, month, and year respectively.
- String

**Validate String**

Use the **Validate string** transforms to create a new column that indicates that a row of text data meets a specified condition. For example, you can use a **Validate string** transform to verify that a
string only contains lowercase characters. The following transforms are supported under **Validate string**.

The following transforms are included in this transform group. If a transform outputs a Boolean value, **True** is represented with a 1 and **False** is represented with a 0.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>String length</td>
<td>Returns <strong>True</strong> if a string length equals specified length. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Starts with</td>
<td>Returns <strong>True</strong> if a string starts will a specified prefix. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Ends with</td>
<td>Returns <strong>True</strong> if a string length equals specified length. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is alphanumeric</td>
<td>Returns <strong>True</strong> if a string only contains numbers and letters. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is alpha (letters)</td>
<td>Returns <strong>True</strong> if a string only contains letters. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is digit</td>
<td>Returns <strong>True</strong> if a string only contains digits. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is space</td>
<td>Returns <strong>True</strong> if a string only contains numbers and letters. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is title</td>
<td>Returns <strong>True</strong> if a string contains any white spaces. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is lowercase</td>
<td>Returns <strong>True</strong> if a string only contains lower case letters. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Is uppercase</td>
<td>Returns <strong>True</strong> if a string only contains upper case letters. Otherwise, returns <strong>False</strong>.</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>--------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Is numeric</td>
<td>Returns True if a string only contains numbers. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is decimal</td>
<td>Returns True if a string only contains decimal numbers. Otherwise, returns False.</td>
</tr>
</tbody>
</table>

**Unnest JSON Data**

If you have a .csv file, you might have values in your dataset that are JSON strings. Similarly, you might have nested data in columns of either a Parquet file or a JSON document.

Use the **Flatten structured** operator to separate the first level keys into separate columns. A first level key is a key that isn't nested within a value.

For example, you might have a dataset that has a `person` column with demographic information on each person stored as JSON strings. A JSON string might look like the following.

```
{"seq": 1,"name": {"first": "Nathaniel","last": "Ferguson"},"age": 59,"city": "Posbotno","state": "WV"}
```

The **Flatten structured** operator converts the following first level keys into additional columns in your dataset:

- seq
- name
- age
- city
- state

Data Wrangler puts the values of the keys as values under the columns. The following shows the column names and values of the JSON.
For each value in your dataset containing JSON, the **Flatten structured** operator creates columns for the first-level keys. To create columns for nested keys, call the operator again. For the preceding example, calling the operator creates the columns:

- name_first
- name_last

The following example shows the dataset that results from calling the operation again.

```
seq, name, age, city, state, name_first, name_last
1, {"first": "Nathaniel","last": "Ferguson"}, 59, Posbotno, WV, Nathaniel, Ferguson
```

Choose **Keys to flatten on** to specify the first-level keys that want to extract as separate columns. If you don't specify any keys, Data Wrangler extracts all the keys by default.

**Explode Array**

Use **Explode array** to expand the values of the array into separate output rows. For example, the operation can take each value in the array, `[[1, 2, 3], [4, 5, 6], [7, 8, 9]]` and create a new column with the following rows:

```
[1, 2, 3]
[4, 5, 6]
[7, 8, 9]
```

Data Wrangler names the new column, `input_column_name_flatten`.

You can call the **Explode array** operation multiple times to get the nested values of the array into separate output columns. The following example shows the result of calling the operation multiple times on a dataset with a nested array.
Putting the values of a nested array into separate columns

<table>
<thead>
<tr>
<th>id</th>
<th>array</th>
<th>id</th>
<th>array_items</th>
<th>id</th>
<th>array_items</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>[ [cat, dog], [bat, frog] ]</td>
<td>1</td>
<td>[cat, dog]</td>
<td>1</td>
<td>cat</td>
</tr>
<tr>
<td>2</td>
<td>[[rose, petunia], [lily, daisy]]</td>
<td>1</td>
<td>[bat, frog]</td>
<td>1</td>
<td>dog</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>[rose, petunia]</td>
<td>1</td>
<td>bat</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>[lily, daisy]</td>
<td>1</td>
<td>frog</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>rose</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>petunia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>lily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>daisy</td>
</tr>
</tbody>
</table>

Transform Image Data

Use Data Wrangler to import and transform the images that you're using for your machine learning (ML) pipelines. After you've prepared your image data, you can export it from your Data Wrangler flow to your ML pipeline.

You can use the information provided here to familiarize yourself with importing and transforming image data in Data Wrangler. Data Wrangler uses OpenCV to import images. For more information about supported image formats, see Image file reading and writing.

After you've familiarized yourself with the concepts of transforming your image data, go through the following tutorial, Prepare image data with Amazon SageMaker Data Wrangler.

The following industries and use cases are examples where applying machine learning to transformed image data can be useful:
• Manufacturing – Identifying defects in items from the assembly line
• Food – Identifying spoiled or rotten food
• Medicine – Identifying lesions in tissues

When you work with image data in Data Wrangler, you go through the following process:

1. Import – Select the images by choosing the directory containing them in your Amazon S3 bucket.
2. Transform – Use the built-in transformations to prepare the images for your machine learning pipeline.
3. Export – Export the images that you’ve transformed to a location that can be accessed from the pipeline.

Use the following procedure to import your image data.

**To import your image data**

1. Navigate to the Create connection page.
2. Choose Amazon S3.
3. Specify the Amazon S3 file path that contains the image data.
4. For File type, choose Image.
5. (Optional) Choose Import nested directories to import images from multiple Amazon S3 paths.
6. Choose Import.

Data Wrangler uses the open-source imgaug library for its built-in image transformations. You can use the following built-in transformations:

• ResizImage
• EnhancelImage
• CorruptImage
• SplitImage
• DropCorruptedImages
• DropImageDuplicates
• Brightness
• ColorChannels
• Grayscale
• Rotate

Use the following procedure to transform your images without writing code.

To transform the image data without writing code

1. From your Data Wrangler flow, choose the + next to the node representing the images that you've imported.
2. Choose Add transform.
3. Choose Add step.
4. Choose the transform and configure it.
5. Choose Preview.
6. Choose Add.

In addition to using the transformations that Data Wrangler provides, you can also use your own custom code snippets. For more information about using custom code snippets, see Custom Transforms. You can import the OpenCV and imgaug libraries within your code snippets and use the transforms associated with them. The following is an example of a code snippet that detects edges within the images.

```python
# A table with your image data is stored in the `df` variable
import cv2
import numpy as np
from pyspark.sql.functions import column
from sagemaker_dataprep.compute.operators.transforms.image.constants import DEFAULT_IMAGE_COLUMN, IMAGE_COLUMN_TYPE
from sagemaker_dataprep.compute.operators.transforms.image.decorators import BasicImageOperationDecorator, PandasUDFOperationDecorator

@BasicImageOperationDecorator
def my_transform(image: np.ndarray) -> np.ndarray:
    Prepare data
```
# To use the code snippet on your image data, modify the following lines within the function

```python
HYST_THRLD_1, HYST_THRLD_2 = 100, 200
edges = cv2.Canny(image, HYST_THRLD_1, HYST_THRLD_2)
return edges
```

```python
@PandasUDFOperationDecorator(IMAGE_COLUMN_TYPE)
def custom_image_udf(image_row):
    return my_transform(image_row)
```

```python
df = df.withColumn(DEFAULT_IMAGE_COLUMN,
custom_image_udf(column(DEFAULT_IMAGE_COLUMN)))
```

When apply transformations in your Data Wrangler flow, Data Wrangler only applies them to a sample of the images in your dataset. To optimize your experience with the application, Data Wrangler doesn't apply the transforms to all of your images.

**Filter data**

Use Data Wrangler to filter the data in your columns. When you filter the data in a column, you specify the following fields:

- **Column name** – The name of the column that you're using to filter the data.
- **Condition** – The type of filter that you're applying to values in the column.
- **Value** – The value or category in the column to which you're applying the filter.

You can filter on the following conditions:

- `=` – Returns values that match the value or category that you specify.
- `!=` – Returns values that don't match the value or category that you specify.
- `>=` – For **Long** or **Float** data, filters for values that are greater than or equal to the value that you specify.
- `<=` – For **Long** or **Float** data, filters for values that are less than or equal to the value that you specify.
- `>` – For **Long** or **Float** data, filters for values that are greater than the value that you specify.
• `<` – For **Long** or **Float** data, filters for values that are less than the value that you specify.

For a column that has the categories, male and female, you can filter out all the male values. You could also filter for all the female values. Because there are only male and female values in the column, the filter returns a column that only has female values.

You can also add multiple filters. The filters can be applied across multiple columns or the same column. For example, if you're creating a column that only has values within a certain range, you add two different filters. One filter specifies that the column must have values greater than the value that you provide. The other filter specifies that the column must have values less than the value that you provide.

Use the following procedure to add the filter transform to your data.

**To filter your data**

1. From your Data Wrangler flow, choose the + next to the node with the data that you're filtering.
2. Choose **Add transform**.
3. Choose **Add step**.
4. Choose **Filter data**.
5. Specify the following fields:
   - **Column name** – The column that you're filtering.
   - **Condition** – The condition of the filter.
   - **Value** – The value or category in the column to which you're applying the filter.
6. (Optional) Choose + following the filter that you've created.
7. Configure the filter.
8. Choose **Preview**.
9. Choose **Add**.

**Chat for data prep**

⚠️ **Important**

For administrators:
Chat for data prep requires the AmazonSageMakerCanvasAIServicesAccess policy. For more information, see AWS managed policy: AmazonSageMakerCanvasAIServicesAccess.

Chat for data prep requires access to Amazon Bedrock and the Anthropic Claude model within it. For more information, see Add model access.

You must run SageMaker Canvas data prep in the same AWS Region as the Region where you're running your model. Chat for data prep is available in the US East (N. Virginia), US West (Oregon), and Europe (Frankfurt) AWS Regions.

In addition to using the built-in transforms and analyses, you can use natural language to explore, visualize, and transform your data in a conversational interface. Within the conversational interface, you can use natural language queries to understand and prepare your data to build ML models.

The following are examples of some prompts that you can use:

- Summarize my data
- Drop column example-column-name
- Replace missing values with median
- Plot histogram of prices
- What is the most expensive item sold?
- How many distinct items were sold?
- Sort data by region

When you’re transforming your data using your prompts, you can view a preview that shows how data is being transformed. You can choose to add it as step in your Data Wrangler flow based on what you see in the preview.

The responses to your prompts generate code for your transformations and analyses. You can modify the code to update the output from the prompt. For example, you can modify the code for an analysis to change the values of the axes of a graph.

Use the following procedure to start chatting with your data:
To chat with your data

1. Open the SageMaker Canvas data flow.
2. Choose the speech bubble.
3. Specify a prompt.
4. (Optional) If an analysis has been generated by your query, choose Add to analyses to reference it for later.
5. (Optional) If you've transformed your data using a prompt, do the following.

   a. Choose **Preview** to view the results.
   
   b. (Optional) Modify the code in the transform and choose **Update**.
   
   c. (Optional) If you're happy with the results of the transform, choose **Add to steps** to add it to the steps panel on the right-hand navigation.

After you've prepared your data using natural language, you can create a model using your transformed data. For more information about creating a model, see [Build a custom model](#).

**Process data**

You can process or export your data to a location that is suitable for your machine learning workflows. For example, you can export the transformed data as SageMaker Canvas dataset and create a machine learning model from it.

After you've exported your data you can choose **Create model** to create a machine learning model from your data. For more information about creating a model, see [Build a custom model](#).
Important

Note that SageMaker Canvas datasets have a 5 GB limit. If you're processing more than 5 GB of data, you can use a sampling transform to reduce the size of the dataset before you export it. Alternatively, you can export data to Amazon S3. For more information about importing datasets, see Create a dataset.

Within the Data Wrangler flow (data flow), in SageMaker Canvas only applies the transformations to the SageMaker Canvas dataset. To process all of the data that you've imported, see Export data using a processing job. You can use any of the following methods to process your data:

- Exporting data – Exports the dataset with the transformations that you've made.
- Exporting your data flow – Exports your data flow using a Jupyter notebook. You can modify the code and use it in your machine learning workflows.
- Running a processing job from your data flow – Exports the dataset with the transformations that you've made in a highly performant manner. We recommend running a processing job on datasets larger than 5 GB.

Export data

Export your data to create a dataset with transformations from your flow applied to it. You can export any node in your data flow to the following locations:

- SageMaker Canvas dataset
- Amazon S3

For Amazon S3, you can export your data as one of the following file types:

- CSV
- Parquet

You can export your dataset as a SageMaker Canvas dataset to create a model using your transformed data. Use the following procedure to export a SageMaker Canvas dataset from a node in your data flow.
To export a node in your flow as a SageMaker Canvas dataset

1. Navigate to your data flow.
2. Choose the + next to the node that you're exporting.
3. Select Export data.
4. Select Canvas dataset.
5. Choose Export.

Export your dataset to Amazon S3 to use your transformed data in machine learning workflows external to SageMaker Canvas.

To export a SageMaker Canvas dataset to Amazon S3

1. Navigate to your data flow.
2. Choose the + next to the node that you’re exporting.
3. Select Export data.
4. Select Amazon S3.
5. Specify values for the following fields:
   - **Amazon S3 location** – the S3 location where you're exporting the file.
   - **File type** – The format of the file that you're exporting.
   - **Delimiter** – the value used to separate values in the file.
   - **Compression** (Optional) – The compression method used to reduce the file size. You can use the following compression methods:
     - None
     - bzip2
     - deflate
     - gzip
   - **KMS Key ID or ARN** (Optional) – An ARN or ID of an AWS KMS key. A KMS key is a cryptographic key. You can use the key to encrypt the output data from the job. For more information about KMS keys, see [AWS Key Management Service](#).
6. Choose Export.
Export data using a processing job

A SageMaker Canvas dataset is a sample of your data. Use an Amazon SageMaker processing job to apply transformations in your data flow to all of your data.

To create a SageMaker processing job, do the following:

1. Create a destination node
2. Create a job

A destination node tells SageMaker Canvas where to store the data that it processed. You create a processing job to output the transformed data to the location specified by the destination node. Creating a processing job runs the computational resources needed to output the transformed data to Amazon S3.

You can use a destination node to export some of the transformations or all of the transformations that you've made in your Data Wrangler flow.

You can use multiple destination nodes to export different transformations or sets of transformations. The following example shows two destination nodes in a single Data Wrangler flow.

Use the following procedure to create a destination node.

To create destination nodes

1. Choose the + next to the nodes that represent the transformations that you want to export.
2. Choose Add destination.
3. Choose Amazon S3.
4. Specify the following fields.
   - **Dataset name** – The name that you specify for the dataset that you're exporting.
   - **File type** – The format of the file that you're exporting.
   - **Delimiter** – The value used to separate other values.
   - **Compression** (CSV and Parquet files only) – The compression method used to reduce the file size. You can use the following compression methods:
     - bzip2
• deflate
• gzip
• Amazon S3 location – The S3 location that you're using to output the files.
• (Optional) Number of partitions – The number of datasets that you're writing as the output of the processing job.
• (Optional) Partition by column – Writes all data with the same unique value from the column.

5. Choose Add destination.

You can create multiple destination nodes within the same data flow. When you create a processing job, it can simultaneously perform different sets of transformations on your data and save them to different Amazon S3 locations.

**Export data flow**

Exporting your data flow translates the operations that you've made in Data Wrangler and exports it into a Jupyter notebook that you can modify and run.

A data flow is the series of data preparation steps that you've performed on your data. In your data preparation, you perform one or more transformations to your data. Each transformation is done using a transform step. The flow has a series of nodes that represent the import of your data and the transformations that you've performed. For an example of nodes, see the following image.

As an alternative to using a destination node, you can use the Export data flow option to export your Data Wrangler flow to Amazon S3 using a Jupyter notebook. You can integrate the output code into your machine learning pipelines. You can choose any data node in your data flow and export it. Exporting the data node exports the transformation that the node represents and the transformations that precede it.

Use the following procedure to generate a Jupyter notebook and run it to export your data flow to Amazon S3.

1. Choose the + next to the node that you want to export.

2. Choose Export data flow.

3. Choose one of the following:

   • Save to Amazon S3 (via Jupyter notebook).
• Amazon Personalize.

4. Select one of the following:
   • Download a local copy
   • Export to S3 location

5. If you're exporting to Amazon S3, specify the S3 location where you're export the notebook.
6. Choose Export.

Create a schedule to automatically process new data

If you're processing data periodically, you can create a schedule to run the processing job automatically. For example, you can create a schedule that runs a processing job automatically when you get new data. For more information about processing jobs, see Export data using a processing job.

When you create a job, you must specify an IAM role that has permissions to create the job. You can use the AmazonSageMakerCanvasDataPrepFullAccess policy to add permissions.

Add the following trust policy to the role to allow EventBridge to assume it.

```
{
   "Effect": "Allow",
   "Principal": {
      "Service": "events.amazonaws.com"
   },
   "Action": "sts:AssumeRole"
}
```

⚠️ Important

When you create a schedule, Data Wrangler creates an eventRule in EventBridge. You incur charges for both the event rules that you create and the instances used to run the processing job.

For information about EventBridge pricing, see Amazon EventBridge pricing. For information about processing job pricing, see Amazon SageMaker Pricing.
You can set a schedule using one of the following methods:

- **CRON expressions**

  ![Note](image)
  
  Data Wrangler doesn't support the following expressions:
  
  - LW#
  
  - Abbreviations for days
  
  - Abbreviations for months

- **RATE expressions**

  - Recurring – Set an hourly or daily interval to run the job.
  
  - Specific time – Set specific days and times to run the job.

The following sections provide procedures on creating jobs.

**CRON**

Use the following procedure to create a schedule with a CRON expression.

To specify a schedule with a CRON expression, do the following.

1. Open your Data Wrangler flow.
2. Choose Create job.
3. (Optional) For Output KMS key, specify an AWS KMS key to configure the output of the job.
4. Choose Next.
5. Select Associate Schedules.
6. Choose Create a new schedule.
7. For Schedule Name, specify the name of the schedule.
8. For Run Frequency, choose CRON.
9. Specify a valid CRON expression.
10. Choose Create.
11. (Optional) Choose Add another schedule to run the job on an additional schedule.
Note
You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

12. Choose one of the following:

- **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.
- **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

13. Choose **Run**

RATE

Use the following procedure to create a schedule with a RATE expression.

To specify a schedule with a RATE expression, do the following.

1. Open your Data Wrangler flow.
2. Choose **Create job**.
3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.
4. Choose **Next, 2. Configure job**.
5. Select **Associate Schedules**.
6. Choose **Create a new schedule**.
7. For **Schedule Name**, specify the name of the schedule.
8. For **Run Frequency**, choose **Rate**.
9. For **Value**, specify an integer.
10. For **Unit**, select one of the following:
    - **Minutes**
    - **Hours**
    - **Days**
11. Choose **Create**.
12. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

**Note**
You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

13. Choose one of the following:

- **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.
- **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

14. Choose **Run**

**Recurring**

Use the following procedure to create a schedule that runs a job on a recurring basis.

To specify a schedule with a CRON expression, do the following.

1. Open your Data Wrangler flow.
2. Choose **Create job**.
3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.
4. Choose **Next, 2. Configure job**.
5. Select **Associate Schedules**.
6. Choose **Create a new schedule**.
7. For **Schedule Name**, specify the name of the schedule.
8. For **Run Frequency**, make sure **Recurring** is selected by default.
9. For **Every x hours**, specify the hourly frequency that the job runs during the day. Valid values are integers in the inclusive range of 1 and 23.
10. For **On days**, select one of the following options:

- **Every Day**
- **Weekends**
• **Weekdays**

• **Select Days**

• (Optional) If you've selected **Select Days**, choose the days of the week to run the job.

**Note**

The schedule resets every day. If you schedule a job to run every five hours, it runs at the following times during the day:

• 00:00
• 05:00
• 10:00
• 15:00
• 20:00

11. Choose **Create**.

12. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

**Note**

You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

13. Choose one of the following:

• **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.

• **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

14. Choose **Run**

**Specific time**

Use the following procedure to create a schedule that runs a job at specific times.

To specify a schedule with a CRON expression, do the following.
1. Open your Data Wrangler flow.

2. Choose **Create job**.

3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.

4. Choose **Next, 2. Configure job**.

5. Select **Associate Schedules**.

6. Choose **Create a new schedule**.

7. For **Schedule Name**, specify the name of the schedule.

8. Choose **Create**.

9. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

   **Note**
   
   You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

10. Choose one of the following:

    - **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.
    - **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

11. Choose **Run**

You can use the SageMaker AWS Management Console to view the jobs that are scheduled to run. Your processing jobs run within SageMaker Pipelines. Each processing job has its own pipeline. It runs as a processing step within the pipeline. You can view the schedules that you've created within a pipeline. For information about viewing a pipeline, see **View a Pipeline**.

Use the following procedure to view the jobs that you've scheduled.

To view the jobs you've scheduled, do the following.

1. Open Amazon SageMaker Studio Classic.

2. Open SageMaker Pipelines

3. View the pipelines for the jobs that you've created.
The pipeline running the job uses the job name as a prefix. For example, if you've created a job named `housing-data-feature-engineering`, the name of the pipeline is `canvas-data-prep-housing-data-feature-engineering`.

4. Choose the pipeline containing your job.

5. View the status of the pipelines. Pipelines with a **Status** of **Succeeded** have run the processing job successfully.

To stop the processing job from running, do the following:

To stop a processing job from running, delete the event rule that specifies the schedule. Deleting an event rule stops all the jobs associated with the schedule from running. For information about deleting a rule, see [Disabling or deleting an Amazon EventBridge rule](#).

You can stop and delete the pipelines associated with the schedules as well. For information about stopping a pipeline, see [StopPipelineExecution](#). For information about deleting a pipeline, see [DeletePipeline](#).

**Refit transforms to the entire dataset and export them**

When you import data, Data Wrangler uses a sample of the data to apply the encodings. Data Wrangler uses the first 20,000 rows as a sample.

The following transformations can use your data to create a column in the dataset:

- **Encode Categorical**
- **Featurize Text**
- **Handle Outliers**
- **Handle Missing Values**

If you used sampling to import your data, the preceding transforms only use the data from the sample to create the column. The transform might not have used all of the relevant data. For example, if you use the **Encode Categorical** transform, there might have been a category in the entire dataset that wasn't present in the sample.

You can either use a destination node or a Jupyter notebook to refit the transformations to the entire dataset. When Data Wrangler exports the transformations in the flow, it creates a SageMaker...
processing job. When the processing job finishes, Data Wrangler saves the following files in either the default Amazon S3 location or an S3 location that you specify:

- The Data Wrangler flow file that specifies the transformations that are refit to the dataset
- The dataset with the refit transformations applied to it

You can open a Data Wrangler flow file within SageMaker Canvas and apply the transformations to a different dataset. For example, if you've applied the transformations to a training dataset, you can open and use the Data Wrangler flow file to apply the transformations to a dataset used for inference.

Use the following procedure to run a Jupyter notebook to refit the transformations and export the data.

To run a Jupyter notebook and to refit the transformations and export your Data Wrangler flow, do the following.

1. Choose the + next to the node that you want to export.
2. Choose Export to.
3. Choose the location to which you're exporting the data.
4. For the refit_trained_params object, set refit to True.
5. For the output_flow field, specify the name of the output flow file with the refit transformations.
6. Run the Jupyter notebook.

Automate data preparation in SageMaker Canvas

After you transform your data in data flow, you can export the transforms to your machine learning workflows. When you export your transforms, SageMaker Canvas creates a Jupyter notebook. You must run the notebook within Amazon SageMaker Studio Classic. For information about getting started with Studio Classic, contact your administrator.

Automate data preparation using SageMaker Pipelines

When you want to build and deploy large-scale machine learning (ML) workflows, you can use SageMaker Pipelines to create workflows that manage and deploy SageMaker jobs. With SageMaker Pipelines, you can build workflows that manage your SageMaker data preparation,
model training, and model deployment jobs. You can use the first-party algorithms that SageMaker offers by using SageMaker Pipelines. For more information on SageMaker Pipelines, see SageMaker Pipelines.

When you export one or more steps from your data flow to SageMaker Pipelines, Data Wrangler creates a Jupyter notebook that you can use to define, instantiate, run, and manage a pipeline.

Use a Jupyter Notebook to Create a Pipeline

Use the following procedure to create a Jupyter notebook to export your Data Wrangler flow to SageMaker Pipelines.

1. Choose the + next to the node that you want to export.
2. Choose Export data flow.
4. Download the Jupyter notebook or copy it to an Amazon S3 location. We recommend copying it to an Amazon S3 location that you can access within Studio Classic. Contact your administrator if you need guidance on a suitable location.
5. Run the Jupyter notebook.

You can use the Jupyter notebook that Data Wrangler produces to define a pipeline. The pipeline includes the data processing steps that are defined by your Data Wrangler flow.

You can add additional steps to your pipeline by adding steps to the steps list in the following code in the notebook:

```python
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[instance_type, instance_count],
    steps=[step_process],  #Add more steps to this list to run in your Pipeline
)
```

For more information on defining pipelines, see Define SageMaker Pipeline.
Automate data preparation using an inference endpoint

Use your Data Wrangler flow to process data at the time of inference by creating a SageMaker serial inference pipeline from your Data Wrangler flow. An inference pipeline is a series of steps that results in a trained model making predictions on new data. A serial inference pipeline within Data Wrangler transforms the raw data and provides it to the machine learning model for a prediction. You create, run, and manage the inference pipeline from a Jupyter notebook within Studio Classic. For more information about accessing the notebook, see Use a Jupyter notebook to create an inference endpoint.

Within the notebook, you can either train a machine learning model or specify one that you've already trained. You can either use Amazon SageMaker Autopilot or XGBoost to train the model using the data that you've transformed in your Data Wrangler flow.

The pipeline provides the ability to perform either batch or real-time inference. You can also add the Data Wrangler flow to SageMaker Model Registry. For more information about hosting models, see Host multiple models in one container behind one endpoint.

⚠️ Important

You can't export your Data Wrangler flow to an inference endpoint if it has the following transformations:

- Join
- Concatenate
- Group by

If you must use the preceding transforms to prepare your data, use the following procedure.

To prepare your data for inference with unsupported transforms

1. Create a Data Wrangler flow.
2. Apply the preceding transforms that aren't supported.
3. Export the data to an Amazon S3 bucket.
4. Create a separate Data Wrangler flow.
5. Import the data that you've exported from the preceding flow.
6. Apply the remaining transforms.
7. Create a serial inference pipeline using the Jupyter notebook that we provide.

For information about exporting your data to an Amazon S3 bucket see Export data. For information about opening the Jupyter notebook used to create the serial inference pipeline, see Use a Jupyter notebook to create an inference endpoint.

Data Wrangler ignores transforms that remove data at the time of inference. For example, Data Wrangler ignores the Handle Missing Values transform if you use the Drop missing configuration.

If you've refit transforms to your entire dataset, the transforms carry over to your inference pipeline. For example, if you used the median value to impute missing values, the median value from refitting the transform is applied to your inference requests. You can either refit the transforms from your Data Wrangler flow when you're using the Jupyter notebook or when you're exporting your data to an inference pipeline. For information about refitting transforms, see Refit transforms to the entire dataset and export them.

The serial inference pipeline supports the following data types for the input and output strings. Each data type has a set of requirements.

**Supported datatypes**

- **text/csv** – the datatype for CSV strings
  - The string can't have a header.
  - Features used for the inference pipeline must be in the same order as features in the training dataset.
  - There must be a comma delimiter between features.
  - Records must be delimited by a newline character.

  The following is an example of a validly formatted CSV string that you can provide in an inference request.

  `abc,0.0,"Doe, John",12345\ndef,1.1,"Doe, Jane",67890`

- **application/json** – the datatype for JSON strings
The features used in the dataset for the inference pipeline must be in the same order as the features in the training dataset.

The data must have a specific schema. You define schema as a single `instances` object that has a set of `features`. Each `features` object represents an observation.

The following is an example of a validly formatted JSON string that you can provide in an inference request.

```json
{
  "instances": [
    {
      "features": ["abc", 0.0, "Doe, John", 12345]
    },
    {
      "features": ["def", 1.1, "Doe, Jane", 67890]
    }
  ]
}
```

Use a Jupyter notebook to create an inference endpoint

Use the following procedure to export your Data Wrangler flow to create an inference pipeline.

To create an inference pipeline using a Jupyter notebook, do the following.

1. Choose the + next to the node that you want to export.
2. Choose Export data flow.
4. Download the Jupyter notebook or copy it to an Amazon S3 location. We recommend copying it to an Amazon S3 location that you can access within Studio Classic. Contact your administrator if you need guidance on a suitable location.
5. Run the Jupyter notebook.

When you run the Jupyter notebook, it creates an inference flow artifact. An inference flow artifact is a Data Wrangler flow file with additional metadata used to create the serial inference pipeline. The node that you're exporting encompasses all of the transforms from the preceding nodes.
Important
Data Wrangler needs the inference flow artifact to run the inference pipeline. You can't use your own flow file as the artifact. You must create it by using the preceding procedure.

Automate data preparation using Python Code

To export all steps in your data flow to a Python file that you can manually integrate into any data processing workflow, use the following procedure.

Use the following procedure to generate a Jupyter notebook and run it to export your Data Wrangler flow to Python code.

1. Choose the + next to the node that you want to export.
2. Choose Export data flow.
4. Download the Jupyter notebook or copy it to an Amazon S3 location. We recommend copying it to an Amazon S3 location that you can access within Studio Classic. Contact your administrator if you need guidance on a suitable location.
5. Run the Jupyter notebook.

You might need to configure the Python script to make it run in your pipeline. For example, if you're running a Spark environment, make sure that you are running the script from an environment that has permission to access AWS resources.

Use generative AI with foundation models

Amazon SageMaker Canvas provides generative AI foundation models that you can use to start conversational chats. These content generation models are trained on large amounts of text data to learn the statistical patterns and relationships between words, and they can produce coherent text that is statistically similar to the text on which they were trained. You can use this capability to increase your productivity by doing the following:

- Generate content, such as document outlines, reports, and blogs
- Summarize text from large corpuses of text, such as earnings call transcripts, annual reports, or chapters of user manuals
• Extract insights and key takeaways from large passages of text, such as meeting notes or narratives

• Improve text and catch grammatical errors or typos

The foundation models are a combination of Amazon SageMaker JumpStart and Amazon Bedrock large language models (LLMs). Canvas offers the following models:

<table>
<thead>
<tr>
<th>Model</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amazon Titan</td>
<td>Amazon Bedrock model</td>
<td>Amazon Titan is a powerful, general-purpose language model that you can use for tasks such as summarization, text generation (such as creating a blog post), classification, open-ended Q&amp;A, and information extraction. It is pretrained on large datasets, making it suitable for complex tasks and reasoning. To continue supporting best practices in the responsible use of AI, Amazon Titan foundation models are built to detect and remove harmful content in the data, reject inappropriate content in the user input, and filter model outputs that contain inappropriate content (such as hate speech, profanity, and violence).</td>
</tr>
<tr>
<td>Anthropic Claude Instant</td>
<td>Amazon Bedrock model</td>
<td>Anthropic's Claude Instant is a faster and more cost-effective yet still very capable</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>--------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Anthropic Claude-2</td>
<td>Amazon Bedrock model</td>
<td>Claude-2 is Anthropic's most powerful model, which excels at a wide range of tasks from sophisticated dialogue and creative content generation to detailed instruction following. Claude-2 can take up to 100,000 tokens in each prompt, equivalent to about 200 pages of information. It can generate longer responses compared to its prior version. It supports use cases such as question answering, information extraction, removing PII, content generation, multiple-choice classification, roleplay, comparing text, summarization, and document Q&amp;A with citation.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Falcon-7B-Instruct</td>
<td>SageMaker JumpStart model</td>
<td>Falcon-7B-Instruct has 7 billion parameters and was fine-tuned on a mixture of chat and instruct datasets. It is suitable as a virtual assistant and performs best when following instructions or engaging in conversation. Since the model was trained on large amounts of English-language web data, it carries the stereotypes and biases commonly found online and is not suitable for languages other than English. Compared to Falcon-40B-Instruct, Falcon-7B-Instruct is a slightly smaller and more compact model.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>--------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Falcon-40B-Instruct</td>
<td>SageMaker JumpStart model</td>
<td>Falcon-40B-Instruct has 40 billion parameters and was fine-tuned on a mixture of chat and instruct datasets. It is suitable as a virtual assistant and performs best when following instructions or engaging in conversation. Since the model was trained on large amounts of English-language web data, it carries the stereotypes and biases commonly found online and is not suitable for languages other than English. Compared to Falcon-7B-Instruct, Falcon-40B-Instruct is a slightly larger and more powerful model.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Jurassic-2 Mid</td>
<td>Amazon Bedrock model</td>
<td>Jurassic-2 Mid is a high-performance text generation model trained on a massive corpus of text (current up to mid 2022). It is highly versatile, general-purpose, and capable of composing human-like text and solving complex tasks such as question answering, text classification, and many others. This model offers zero-shot instruction capabilities, allowing it to be directed with only natural language and without the use of examples. It performs up to 30% faster than its predecessor, the Jurassic-1 model. Jurassic-2 Mid is AI21’s mid-sized model, carefully designed to strike the right balance between exceptional quality and affordability.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
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<tr>
<td>---------------</td>
<td>---------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Jurassic-2 Ultra</td>
<td>Amazon Bedrock model</td>
<td>Jurassic-2 Ultra is a high-performance text generation model trained on a massive corpus of text (current up to mid 2022). It is highly versatile, general-purpose, and capable of composing human-like text and solving complex tasks such as question answering, text classification, and many others. This model offers zero-shot instruction capabilities, allowing it to be directed with only natural language and without the use of examples. It performs up to 30% faster than its predecessor, the Jurassic-1 model. Compared to Jurassic-2 Mid, Jurassic-2 Ultra is a slightly larger and more powerful model.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
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<td>---------------</td>
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<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Llama-2-7b-Chat</td>
<td>SageMaker JumpStart model</td>
<td>Llama-2-7b-Chat is a foundation model by Meta that is suitable for engaging in meaningful and coherent conversations, generating new content, and extracting answers from existing notes. Since the model was trained on large amounts of English-language internet data, it carries the biases and limitations commonly found online and is best-suited for tasks in English.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>---------------</td>
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<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Llama-2-13B-Chat</td>
<td>Amazon Bedrock model</td>
<td>Llama-2-13B-Chat by Meta was fine-tuned on conversational data after initial training on internet data. It is optimized for natural dialog and engaging chat abilities, making it well-suited as a conversational agent. Compared to the smaller Llama-2-7b-Chat, Llama-2-13B-Chat has nearly twice as many parameters, allowing it to remember more context and produce more nuanced conversational responses. Like Llama-2-7b-Chat, Llama-2-13B-Chat was trained on English-language data and is best-suited for tasks in English.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------</td>
<td>---------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Llama-2-70B-Chat</td>
<td>Amazon Bedrock model</td>
<td>Like Llama-2-7b-Chat and Llama-2-13B-Chat, the Llama-2-70B-Chat model by Meta is optimized for engaging in natural and meaningful dialog. With 70 billion parameters, this large conversational model can remember more extensive context and produce highly coherent responses when compared to the more compact model versions. However, this comes at the cost of slower responses and higher resource requirements. Llama-2-70B-Chat was trained on large amounts of English-language internet data and is best-suited for tasks in English.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>-----------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Mistral-7B</td>
<td>SageMaker JumpStart model</td>
<td>Mistral-7B by Mistral.AI is an excellent general purpose language model suitable for a wide range of natural language (NLP) tasks like text generation, summarization, and question answering. It utilizes grouped-query attention (GQA) which allows for faster inference speeds, making it perform comparably to models with twice or three times as many parameters. It was trained on a mixture of text data including books, websites, and scientific papers in the English language, so it is best-suited for tasks in English.</td>
</tr>
<tr>
<td>Model</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Mistral-7B-Chat</td>
<td>SageMaker JumpStart model</td>
<td>Mistral-7B-Chat is a conversational model by Mistral.AI based on Mistral-7B. While Mistral-7B is best for general NLP tasks, Mistral-7B-Chat has been further fine-tuned on conversational data to optimize its abilities for natural, engaging chat. As a result, Mistral-7B-Chat generates more human-like responses and remembers the context of previous responses. Like Mistral-7B, this model is best-suited for English language tasks.</td>
</tr>
<tr>
<td>MPT-7B-Instruct</td>
<td>SageMaker JumpStart model</td>
<td>MPT-7B-Instruct is a model for long-form instruction following tasks and can assist you with writing tasks including text summarization and question-answering to save you time and effort. This model was trained on large amounts of fine-tuned data and can handle larger inputs, such as complex documents. Use this model when you want to process large bodies of text or want the model to generate long responses.</td>
</tr>
</tbody>
</table>
The foundation models from Amazon Bedrock are currently only available in the US East (N. Virginia) and US West (Oregon) Regions. Additionally, when using foundation models from Amazon Bedrock, you are charged based on the volume of input tokens and output tokens, as specified by each model provider. For more information, see the Amazon Bedrock pricing page. The SageMaker JumpStart foundation models are deployed on SageMaker Hosting instances, and you are charged for the duration of usage based on the instance type used. For more information about the cost of different instance types, see the Amazon SageMaker Hosting: Real-Time Inference section on the SageMaker pricing page.

Document querying is an additional feature that you can use to query and get insights from documents stored in indexes using Amazon Kendra. With this functionality, you can generate content from the context of those documents and receive responses that are specific to your business use case, as opposed to responses that are generic to the large amounts of data on which the foundation models were trained. For more information about indexes in Amazon Kendra, see the Amazon Kendra Developer Guide.

If you would like to get responses from any of the foundation models that are customized to your data and use case, you can fine-tune foundation models. To learn more, see Fine-tune foundation models.

To get started, see the following sections.

Prerequisites

The following sections outline the prerequisites for interacting with foundation models and using the document query feature in Canvas. The rest of the content on this page assumes that you’ve met the prerequisites for foundation models. The document query feature requires additional permissions.

Prerequisites for foundation models

The permissions you need for interacting with models are included in the Canvas Ready-to-use models permissions. To use the generative AI-powered models in Canvas, you must turn on the Canvas Ready-to-use models configuration permissions when setting up your Amazon SageMaker domain. For more information, see Prerequisites for setting up Amazon SageMaker Canvas. The Canvas Ready-to-use models configuration attaches the AmazonSageMakerCanvasAIServicesAccess policy to your Canvas user’s AWS Identity and Access Management (IAM) execution role. If you encounter any issues with granting permissions, see the topic Troubleshooting issues with granting permissions through the SageMaker console.
If you've already set up your domain, you can edit your domain settings and turn on the permissions. For instructions on how to edit your domain settings, see View and edit domains. When editing the settings for your domain, go to the Canvas settings and turn on the Enable Canvas Ready-to-use models option.

Certain SageMaker JumpStart foundation models also require that you request a SageMaker instance quota increase. Canvas hosts the models that you're currently interacting with on these instances, but the default quota for your account may be insufficient. If you run into an error while running any of the following models, request a quota increase for the associated instance types:

- Falcon-40B – ml.g5.12xlarge, ml.g5.24xlarge
- Falcon-13B – ml.g5.2xlarge, ml.g5.4xlarge, ml.g5.8xlarge
- MPT-7B-Instruct – ml.g5.2xlarge, ml.g5.4xlarge, ml.g5.8xlarge

For the preceding instances types, request an increase from 0 to 1 for the endpoint usage quota. For more information about how to increase an instance quota for your account, see Requesting a quota increase in the Service Quotas User Guide.

Prerequisites for document querying

Note

Document querying is supported in the following AWS Regions: US East (N. Virginia), US East (Ohio), US West (Oregon), Europe (Ireland), Asia Pacific (Singapore), Asia Pacific (Sydney), Asia Pacific (Tokyo), and Asia Pacific (Mumbai).

The document querying feature requires that you already have an Amazon Kendra index that stores your documents and document metadata. For more information about Amazon Kendra, see the Amazon Kendra Developer Guide. To learn more about the quotas for querying indexes, see Quotas in the Amazon Kendra Developer Guide.

You must also make sure that your Canvas user profile has the necessary permissions for document querying. The AmazonSageMakerCanvasFullAccess policy must be attached to the AWS IAM execution role for the SageMaker domain that hosts your Canvas application (this policy is attached by default to all new and existing Canvas user profiles). You must also specifically grant document querying permissions and specify access to one or more Amazon Kendra indexes.
If your Canvas administrator is setting up a new domain or user profile, have them set up the domain by following the instructions in Prerequisites for setting up Amazon SageMaker Canvas. While setting up the domain, they can turn on the document querying permissions through the Canvas Ready-to-use models configuration.

The Canvas administrator can manage document querying permissions at the user profile level as well. For example, if the administrator wants to grant document querying permissions to some user profiles but remove permissions for others, they can edit the permissions for a specific user.

The following procedure shows how to turn on document querying permissions for a specific user profile:

2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the user profile’s domain.
5. On the domain details page, choose the User profile whose permissions you want to edit.
6. On the User Details page, choose Edit.
7. In the left navigation pane, choose Canvas settings.
8. In the Canvas Ready-to-use models configuration section, turn on the Enable document query using Amazon Kendra toggle.
9. In the dropdown, select one or more Amazon Kendra indexes to which you want to grant access.
10. Choose Submit to save the changes to your domain settings.

You should now be able to use Canvas foundation models to query documents in the specified Amazon Kendra indexes.

Start a new conversation to generate, extract, or summarize content

To get started with generative AI foundation models in Canvas, you can initiate a new chat session with one of the models. For SageMaker JumpStart models, you are charged while the model is active, so you must start up models when you want to use them and shut them down when you are done interacting. If you do not shut down a SageMaker JumpStart model, Canvas shuts it down after 2 hours of inactivity. For Amazon Bedrock models (such as Amazon Titan), you are charged
by prompt; the models are already active and don’t need to be started up or shut down. You are charged directly for use of these models by Amazon Bedrock.

To open a chat with a model, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **Ready-to-use models**.
3. Choose **Generate, extract and summarize content**.
4. On the welcome page, you’ll receive a recommendation to start up the default model. You can start the recommended model, or you can choose **Select another model** from the dropdown to choose a different one.
5. If you selected a SageMaker JumpStart foundation model, you have to start it up before it is available for use. Choose **Start up the model**, and then the model is deployed to a SageMaker instance. It might take several minutes for this to complete. When the model is ready, you can enter prompts and ask the model questions.

   If you selected a foundation model from Amazon Bedrock, you can start using it instantly by entering a prompt and asking questions.

Depending on the model, you can perform various tasks. For example, you can enter a passage of text and ask the model to summarize it. Or, you can ask the model to come up with a short summary of the market trends in your domain.

The model's responses in a chat are based on the context of your previous prompts. If you want to ask a new question in the chat that is unrelated to the previous conversation topic, we recommend that you start a new chat with the model.

**Extract information from documents with document querying**

⚠️ **Note**

This section assumes that you’ve completed the section above [Prerequisites for document querying](#).

Document querying is a feature that you can use while interacting with foundation models in Canvas. With document querying, you can access a corpus of documents stored in an Amazon Kendra *index*, which holds the contents of your documents and is structured in a way to make
documents searchable. You can ask specific questions that are targeted to the data in your Amazon Kendra index, and the foundation model returns answers to your questions. For example, you can query an internal knowledge base of IT information and ask questions such as “How do I connect to my company’s network?” For more information about setting up an index, see the Amazon Kendra Developer Guide.

When using the document query feature, the foundation models restrict their responses to the content of the documents in your index with a technique called Retrieval Augmented Generation (RAG). This technique bundles the most relevant information from the index along with the user's prompt and sends it to the foundation model to get a response. Responses are limited to what can be found in your index, preventing the model from giving you incorrect responses based on external data. For more information about this process, see the blog post Quickly build high-accuracy Generative AI applications on enterprise data.

To get started, in a chat with a foundation model in Canvas, turn on the Document query toggle at the top of the page. From the dropdown, select the Amazon Kendra index that you want to query. Then, you can begin asking questions related to the documents in your index.

⚠️ Important
Document querying supports the Compare model outputs feature. Any existing chat history is overwritten when you start a new chat to compare model outputs.

Model management

ℹ️ Note
The following section describes starting up and shutting down models, which only applies to the SageMaker JumpStart foundation models, such as Falcon-40B-Instruct. You can access Amazon Bedrock models, such as Amazon Titan, instantly at any time.

You can start up as many SageMaker JumpStart models as you like. Each active SageMaker JumpStart model incurs charges on your account, so we recommend that you don’t start up more models than you are currently using.

To start up another model, you can do the following:
1. On the **Generate, extract and summarize content** page, choose **New chat**.
2. Choose the model from the dropdown menu. If you want to choose a model not displayed in the dropdown, choose **Start up another model**, and then select the model that you want to start up.
3. Choose **Start up model**.

The model should begin starting up, and within a few minutes you can chat with the model.

We highly recommend that you shut down models that you aren’t using. The models automatically shut down after 2 hours of inactivity. However, to manually shut down a model, you can do the following:

1. On the **Generate, extract and summarize content** page, open the chat for the model that you want to shut down.
2. On the chat page, choose the **More options** icon.
3. Choose **Shut down model**.
4. In the **Shut down model** confirmation box, choose **Shut down**.

The model begins shutting down. If your chat compares two or more models, you can shut down an individual model from the chat page by choosing the model's **More options** icon and then choosing **Shut down model**.

**Compare model outputs**

You might want to compare the output of different models side by side to see which model output you prefer. This can help you decide which model is best suited to your use case. You can compare up to three models in chats.

**Note**

Each individual model incurs charges on your account.

You must start a new chat to add models for comparison. To compare the output of models side by side in a chat, do the following:
1. In a chat, choose **New chat**.

2. Choose **Compare**, and use the dropdown menu to select the model that you want to add. To add a third model, choose **Compare** again to add another model.

### Note

If you want to use a SageMaker JumpStart model that isn’t currently active, you are prompted to start up the model.

When the models are active, you see the two models side by side in the chat. You can submit your prompt, and each model responds in the same chat, as shown in the following screenshot.

When you’re done interacting, make sure to shut down any SageMaker JumpStart models individually to avoid incurring further charges.
Fine-tune foundation models

The foundation models that you can access through Amazon SageMaker Canvas can help you with a range of general purpose tasks. However, if you have a specific use case and would like to customized responses based on your own data, you can fine-tune a foundation model.

To fine-tune a foundation model, you provide a dataset that consists of sample prompts and model responses. Then, you train the foundation model on the data. Finally, the fine-tuned foundation model is able to provide you with more specific responses.

This topic describes how to fine-tune foundation models in Canvas.

Before you begin

Before fine-tuning a foundation model, make sure that you have the permissions for Ready-to-use models in Canvas and a trust relationship with Amazon Bedrock. For more information, see the Grant Users Permissions to Fine-tune Foundation Models.

You must also have a dataset that is formatted for fine-tuning large language models (LLMs). The following is a list of requirements for your dataset:

- The dataset must be tabular and contain at least two columns of text data—one input column (which contains example prompts to the model) and one output column (which contains example responses from the model).

An example is the following:

<table>
<thead>
<tr>
<th>Input</th>
<th>Output</th>
</tr>
</thead>
<tbody>
<tr>
<td>What are your shipping terms?</td>
<td>We offer free shipping on all orders over $50. Orders under $50 have a shipping fee of $5.99.</td>
</tr>
<tr>
<td>How can I return an item?</td>
<td>To return an item, please visit our returns center and follow the instructions. You must provide your order number and the reason for the return.</td>
</tr>
</tbody>
</table>
I'm having trouble with my product. What can I do?

Please contact our customer support team and we will be happy to help you troubleshoot the issue.

- We recommend that the dataset has at least 100 text pairs (rows of corresponding input and output items). This ensures that the foundation model has enough data for fine-tuning and increases the accuracy of its responses.

- Each input and output item should contain a maximum of 512 characters. Anything longer is reduced to 512 characters when fine-tuning the foundation model.

When fine-tuning an Amazon Bedrock model, you must adhere to the Amazon Bedrock quotas. For more information, see Model customization quotas in the Amazon Bedrock User Guide.

For more information about general dataset requirements and limitations in Canvas, see Create a dataset.

Fine-tune a foundation model

You can fine-tune a foundation model by using any of the following methods in the Canvas application:

- While in a Generate, extract and summarize content chat with a foundation model, choose the Fine-tune model icon.

- While in a chat with a foundation model, if you’ve re-generated the response two or more times, then Canvas offers you the option to Fine-tune model. The following screenshot shows you what this looks like.

Not happy with the model's response? You can fine-tune it to get the responses you want.

Learn more about fine-tuning a model.

- On the My models page, you can create a new model by choosing New model, and then select Fine-tune foundation model.

- On the Ready-to-use models home page, you can choose Create your own model, and then in the Create new model dialog box, choose Fine-tune foundation model.
• While browsing your datasets in the Data Wrangler tab, you can select a dataset and choose Create a model. Then, choose Fine-tune foundation model.

After you’ve begun to fine-tune a model, do the following:

Select a dataset

On the Select tab of fine-tuning a model, you choose the data on which you’d like to train the foundation model.

Either select an existing dataset or create a new dataset that meets the requirements listed in the Before you begin section. For more information about how to create a dataset, see Create a dataset.

When you’ve selected or created a dataset and you’re ready to move on, choose Select dataset.

Fine-tune the model

After selecting your data, you’re now ready to begin training and fine-tune the model.

On the Fine-tune tab, do the following:

1. For Select up to 3 base models, open the dropdown menu and check up to 3 foundation models (up to 2 SageMaker JumpStart models and 1 Amazon Bedrock model) that you’d like to fine-tune during the training job. By fine-tuning multiple foundation models, you can compare their performance and ultimately choose the one best suited to your use case as the default model. For more information about default models, see View model candidates in the model leaderboard.

2. For Select Input column, select the column of text data in your dataset that contains the example model prompts.

3. For Select Output column, select the column of text data in your dataset that contains the example model responses.

4. (Optional) To configure advanced settings for the training job, choose Configure model. For more information about the advanced model building settings, see Advanced model building configurations.

    In the Configure model pop-up window, do the following:

    a. For Hyperparameters, you can adjust the Epoch count, Batch size, Learning rate, and Learning rate warmup steps for each model you selected. For more information
about these parameters, see the [Hyperparameters section in the SageMaker JumpStart documentation](https://docs.aws.amazon.com/sagemaker/latest/dg/basics-hyperparameters.html).

b. For **Data split**, you can specify percentages for how to divide your data between the **Training set** and **Validation set**.

c. For **Max job runtime**, you can set the maximum amount of time that Canvas runs the build job. This feature is only available for SageMaker JumpStart foundation models.

d. After configuring the settings, choose **Save**.

5. Choose **Fine-tune** to begin training the foundation models you selected.

After the fine-tuning job begins, you can leave the page. When the model shows as **Ready** on the **My models** page, it’s ready for use, and you can now analyze the performance of your fine-tuned model.

### Analyze the fine-tuned model

On the **Analyze** tab of your fine-tuned model, you can see the model's performance.

The **Overview** tab on this page shows you the perplexity and loss scores, along with analyses that visualize the model's improvement over time during training. The following screenshot shows the **Overview** tab.

On this page, you can see the following visualizations:
• **The Perplexity Curve** measures how well the model predicts the next word in a sequence, or how grammatical the model's output is. Ideally, as the model improves during training, the score decreases and results in a curve that lowers and flattens over time.

• **The Loss Curve** quantifies the difference between the correct output and the model's predicted output. A loss curve that decreases and flattens over time indicates that the model is improving its ability to make accurate predictions.

The **Advanced metrics** tab shows you the hyperparameters and additional metrics for your model.

It looks like the following screenshot:

The **Advanced metrics** tab contains the following information:

• The **Explainability** section contains the **Hyperparameters**, which are the values set before the job to guide the model's fine-tuning. If you didn't specify custom hyperparameters in the model's advanced settings in the **Fine-tune the model** section, then Canvas selects default hyperparameters for you.

For SageMaker JumpStart models, you can also see the advanced metric **ROUGE (Recall-Oriented Understudy for Gisting Evaluation)**, which evaluates the quality of summaries generated by the model. It measures how well the model can summarize the main points of a passage.
• The **Artifacts** section provides you with links to artifacts generated during the fine-tuning job. You can access the training and validation data saved in Amazon S3, as well as the link to the model evaluation report (to learn more, see the following paragraph).

To get more model evaluation insights, you can download a report that is generated using SageMaker Clarify, which is a feature that can help you detect bias in your model and data. First, generate the report by choosing **Generate evaluation report** at the bottom of the page. After the report has generated, you can download the full report by choosing **Download report** or by returning to the **Artifacts** section.

You can also access a Jupyter notebook that shows you how to replicate your fine-tuning job in Python code. You can use this to replicate or make programmatic changes to your fine-tuning job or get a deeper understanding of how Canvas fine-tunes your model. To learn more about model notebooks and how to access them, see [Download a model notebook](#).

For more information about how to interpret the information in the **Analyze** tab of your fine-tuned model, see the topic [Evaluate Your Model's Performance in Amazon SageMaker Canvas](#).

After analyzing the **Overview** and **Advanced metrics** tabs, you can also choose to open the **Model leaderboard**, which shows you the list of the base models trained during the build. The model with the lowest loss score is considered the best performing model and is selected as the **Default model**, which is the model whose analysis you see in the **Analyze** tab. You can only test and deploy the default model. For more information about the model leaderboard and how to change the default model, see [View model candidates in the model leaderboard](#).

**Test a fine-tuned model in a chat**

After analyzing the performance of a fine-tuned model, you might want to test it out or compare its responses with the base model. You can test a fine-tuned model in a chat in the **Generate, extract and summarize content** feature.

Start a chat with a fine-tuned model by choosing one of the following methods:

• On the fine-tuned model's **Analyze** tab, choose **Test in Ready-to-use foundation models**.

• On the Canvas **Ready-to-use models** page, choose **Generate, extract and summarize content**. Then, choose **New chat** and select the version of your fine-tuned model that you want to test.
The model starts up in a chat, and you can interact with it like any other foundation model. You can add more models to the chat and compare their outputs. For more information about the functionality of chats, see Use generative AI with foundation models.

Operationalize fine-tuned models

After fine-tuning your model in Canvas, you can register the model to the SageMaker model registry for integration into your organizations MLOps processes. For more information, see Register a model version in the SageMaker model registry.

Important

You can only register SageMaker JumpStart based fine-tuned models, not Amazon Bedrock based models.

Use Ready-to-use models

With Amazon SageMaker Canvas Ready-to-use models, you can make predictions on your data without writing a single line of code or having to build a model—all you have to bring is your data. The Ready-to-use models use pre-built models to generate predictions without requiring you to spend the time, expertise, or cost required to build a model, and you can choose from a variety of use cases ranging from language detection to expense analysis.

Canvas integrates with existing AWS services, such as Amazon Textract, Amazon Rekognition, and Amazon Comprehend, to analyze your data and make predictions or extract insights. You can use the predictive power of these services from within the Canvas application to get high quality predictions for your data.

Canvas supports the following Ready-to-use models types:

<table>
<thead>
<tr>
<th>Ready-to-use model</th>
<th>Description</th>
<th>Supported data type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sentiment analysis</td>
<td>Detect sentiment in lines of text, which can be positive, negative, neutral, or mixed. Currently, you can only do sentiment analysis for English language text.</td>
<td>Plain text or tabular (CSV, Parquet)</td>
</tr>
<tr>
<td>Ready-to-use model</td>
<td>Description</td>
<td>Supported data type</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------------------------------------</td>
</tr>
<tr>
<td>Entities extraction</td>
<td>Extract entities, which are real-world objects such as people, places, and</td>
<td>Plain text or tabular (CSV, Parquet)</td>
</tr>
<tr>
<td></td>
<td>commercial items, or units such as dates and quantities, from text.</td>
<td></td>
</tr>
<tr>
<td>Language detection</td>
<td>Determine the dominant language in text such as English, French, or German.</td>
<td>Plain text or tabular (CSV, Parquet)</td>
</tr>
<tr>
<td>Personal information detection</td>
<td>Detect personal information that could be used to identify an individual,</td>
<td>Plain text or tabular (CSV, Parquet)</td>
</tr>
<tr>
<td></td>
<td>such as addresses, bank account numbers, and phone numbers, from text.</td>
<td></td>
</tr>
<tr>
<td>Object detection in images</td>
<td>Detect objects, concepts, scenes, and actions in your images.</td>
<td>Image (JPG, PNG)</td>
</tr>
<tr>
<td>Text detection in images</td>
<td>Detect text in your images.</td>
<td>Image (JPG, PNG)</td>
</tr>
<tr>
<td>Expense analysis</td>
<td>Extract information from invoices and receipts, such as date, number, item</td>
<td>Document (PDF, JPG, PNG, TIFF)</td>
</tr>
<tr>
<td></td>
<td>prices, total amount, and payment terms.</td>
<td></td>
</tr>
<tr>
<td>Identity document analysis</td>
<td>Extract information from passports, driver licenses, and other identity</td>
<td>Document (PDF, JPG, PNG, TIFF)</td>
</tr>
<tr>
<td></td>
<td>documentation issued by the US Government.</td>
<td></td>
</tr>
<tr>
<td>Ready-to-use model</td>
<td>Description</td>
<td>Supported data type</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>---------------------------</td>
</tr>
<tr>
<td>Document analysis</td>
<td>Analyze documents and forms for relationships among detected text.</td>
<td>Document (PDF, JPG, PNG, TIFF)</td>
</tr>
<tr>
<td>Document queries</td>
<td>Extract information from structured documents such as paystubs, bank statements, W-2s, and mortgage application forms by asking questions using natural language.</td>
<td>Document (PDF)</td>
</tr>
</tbody>
</table>

**Get started**

To get started with Ready-to-use models, review the following information.

**Prerequisites**

To use Ready-to-use models in Canvas, you must turn on the **Canvas Ready-to-use models configuration** permissions when setting up your Amazon SageMaker domain. The **Canvas Ready-to-use models configuration** attaches the **AmazonSageMakerCanvasAIServicesAccess** policy to your Canvas user's AWS Identity and Access Management (IAM) execution role. If you encounter any issues with granting permissions, see the topic **Troubleshooting issues with granting permissions through the SageMaker console**.

If you’ve already set up your domain, you can edit your domain settings and turn on the permissions. For instructions on how to edit your domain settings, see **View and Edit domains**. When editing the settings for your domain, go to the **Canvas settings** and turn on the **Enable Canvas Ready-to-use models** option.

**(Optional) Opt out of AI services data storage**

Certain AWS AI services store and use your data to make improvements to the service. You can opt out of having your data stored or used for service improvements. To learn more about how to opt out, see **AI services opt-out policies** in the **AWS Organizations User Guide**.

**How to use Ready-to-use models**

To get started with Ready-to-use models, do the following:
1. **(Optional) Import your data.** You can import a tabular, image, or document dataset to generate batch predictions, or a dataset of predictions, with Ready-to-use models. To get started with importing a dataset, see [Import data into a data flow](#).

2. **Generate predictions.** You can generate single or batch predictions with your chosen Ready-to-use model. To get started with making predictions, see [Make predictions with Ready-to-use models](#).

---

**Make predictions with Ready-to-use models**

Ready-to-use models are available for text, image, and document data. Each data type has Ready-to-use models that are designed to work best for each use case. Use the following guide to determine which Ready-to-use models you can use with your input data:

- **Text data:** Sentiment analysis, entities extraction, language detection, personal information detection
- **Image data:** Object detection in images, text detection in images
- **Document data:** Expense analysis, identity document analysis, document analysis, document queries

The following screenshot shows you the landing page for Ready-to-use models, which showcases all of the different solutions.
Each Ready-to-use model supports both **Single predictions** and **Batch predictions** for your dataset. A **Single prediction** is when you only need to make one prediction. For example, you have one image from which you want to extract text, or one paragraph of text for which you want to detect the dominant language. A **Batch prediction** is when you'd like to make predictions for an entire dataset. For example, you might have a CSV file of customer reviews for which you'd like to analyze the customer sentiment, or you might have image files in which you'd like to detect objects.

When you have your data and have identified your use case, choose one of the following workflows to make predictions for your data.

**Make predictions for text data**

The following procedures describe how to make both single and batch predictions for text datasets. You can use the procedures for the following Ready-to-use model types: sentiment analysis, entities extraction, language detection, and personal information detection.

**Note**

For sentiment analysis, you can only use English language text.

**Single predictions**

To make a single prediction for Ready-to-use models that accept text data, do the following:

1. In the left navigation pane of the Canvas application, choose **Ready-to-use models**.
2. On the **Ready-to-use models** page, choose the Ready-to-use model for your use case. For text data, it should be one of the following: **Sentiment analysis**, **Entities extraction**, **Language detection**, or **Personal information detection**.
3. On the **Run predictions** page for your chosen Ready-to-use model, choose **Single prediction**.
4. For **Text field**, enter the text for which you’d like to get a prediction.
5. Choose **Generate prediction results** to get your prediction.

In the right pane **Prediction results**, you receive an analysis of your text in addition to a **Confidence** score for each result or label. For example, if you chose language detection and entered a passage of text in French, you might get French with a 95% confidence score and traces of other languages, like English, with a 5% confidence score.
The following screenshot shows the results for a single prediction using language detection where the model is 100% confident that the passage is English.

Batch predictions

To make batch predictions for Ready-to-use models that accept text data, do the following:

1. In the left navigation pane of the Canvas application, choose Ready-to-use models.
2. On the Ready-to-use models page, choose the Ready-to-use model for your use case. For text data, it should be one of the following: Sentiment analysis, Entities extraction, Language detection, or Personal information detection.
3. On the Run predictions page for your chosen Ready-to-use model, choose Batch prediction.
4. Choose Select dataset if you've already imported your dataset. If not, choose Import new dataset, and then you are directed through the import data workflow.
5. From the list of available datasets, select your dataset and choose Generate predictions to get your predictions.

After the prediction job finishes running, on the Run predictions page, you see an output dataset listed under Predictions. This dataset contains your results, and if you select the More options icon, you can Preview the output data. Then, you can choose Download to download the results.
Make predictions for image data

The following procedures describe how to make both single and batch predictions for image datasets. You can use the procedures for the following Ready-to-use model types: object detection images and text detection in images.

Single predictions

To make a single prediction for Ready-to-use models that accept image data, do the following:

1. In the left navigation pane of the Canvas application, choose **Ready-to-use models**.
2. On the **Ready-to-use models** page, choose the Ready-to-use model for your use case. For image data, it should be one of the following: **Object detection images** or **Text detection in images**.
3. On the **Run predictions** page for your chosen Ready-to-use model, choose **Single prediction**.
4. Choose **Upload image**.
5. You are prompted to select an image to upload from your local computer. Select the image from your local files, and then the prediction results generate.

In the right pane **Prediction results**, you receive an analysis of your image in addition to a **Confidence** score for each object or text detected. For example, if you chose object detection in images, you receive a list of objects in the image along with a confidence score of how certain the model is that each object was accurately detected, such as 93%.

The following screenshot shows the results for a single prediction using the object detection in images solution, where the model predicts objects such as a clock tower and bus with 100% confidence.
Batch predictions

To make batch predictions for Ready-to-use models that accept image data, do the following:

1. In the left navigation pane of the Canvas application, choose **Ready-to-use models**.

2. On the **Ready-to-use models** page, choose the Ready-to-use model for your use case. For image data, it should be one of the following: **Object detection images** or **Text detection in images**.

3. On the **Run predictions** page for your chosen Ready-to-use model, choose **Batch prediction**.

4. Choose **Select dataset** if you’ve already imported your dataset. If not, choose **Import new dataset**, and then you are directed through the import data workflow.

5. From the list of available datasets, select your dataset and choose **Generate predictions** to get your predictions.

After the prediction job finishes running, on the **Run predictions** page, you see an output dataset listed under **Predictions**. This dataset contains your results, and if you select the **More options** icon ( ), you can choose **View prediction results** to preview the output data. Then, you can choose **Download prediction** and download the results as a CSV or a ZIP file.
Make predictions for document data

The following procedures describe how to make both single and batch predictions for document datasets. You can use the procedures for the following Ready-to-use model types: expense analysis, identity document analysis, and document analysis.

**Note**
For document queries, only single predictions are currently supported.

**Single predictions**

To make a single prediction for Ready-to-use models that accept document data, do the following:

1. In the left navigation pane of the Canvas application, choose **Ready-to-use models**.
2. On the **Ready-to-use models** page, choose the Ready-to-use model for your use case. For document data, it should be one of the following: Expense analysis, Identity document analysis, or Document analysis.
3. On the **Run predictions** page for your chosen Ready-to-use model, choose **Single prediction**.
4. If your Ready-to-use model is identity document analysis or document analysis, complete the following actions. If you’re doing expense analysis or document queries, skip this step and go to Step 5 or Step 6, respectively.
   a. Choose **Upload document**.
   b. You are prompted to upload a PDF, JPG, or PNG file from your local computer. Select the document from your local files, and then the prediction results will generate.
5. If your Ready-to-use model is expense analysis, do the following:
   a. Choose **Upload invoice or receipt**.
   b. You are prompted to upload a PDF, JPG, PNG, or TIFF file from your local computer. Select the document from your local files, and then the prediction results will generate.
6. If your Ready-to-use model is document queries, do the following:
   a. Choose **Upload document**.
   b. You are prompted to upload a PDF file from your local computer. Select the document from your local files. Your PDF must be 1–100 pages long.
c. In the right side pane, enter queries to search for information in the document. The number of characters you can have in a single query is from 1–200. You can add up to 15 queries at a time.

d. Choose Submit queries, and then the results generate with answers to your queries. You are billed once for each submissions of queries you make.

In the right pane Prediction results, you'll receive an analysis of your document.

The following information describes the results for each type of solution:

• For expense analysis, the results are categorized into Summary fields, which include fields such as the total on a receipt, and Line item fields, which include fields such as individual items on a receipt. The identified fields are highlighted on the document image in the output.

• For identity document analysis, the output shows you the fields that the Ready-to-use model identified, such as first and last name, address, or date of birth. The identified fields are highlighted on the document image in the output.

• For document analysis, the results are categorized into Raw text, Forms, Tables, and Signatures. Raw text includes all of the extracted text, while Forms, Tables, and Signatures only include information on the form that falls into those categories. For example, Tables only includes information extracted from tables in the document. The identified fields are highlighted on the document image in the output.

• For document queries, Canvas returns answers to each of your queries. You can open the collapsible query dropdown to view a result, along with a confidence score for the prediction. If Canvas finds multiple answers in the document, then you might have more than one result for each query.

The following screenshot shows the results for a single prediction using the document analysis solution.
Batch predictions

To make batch predictions for Ready-to-use models that accept document data, do the following:

1. In the left navigation pane of the Canvas application, choose **Ready-to-use models**.
2. On the **Ready-to-use models** page, choose the Ready-to-use model for your use case. For image data, it should be one of the following: **Expense analysis**, **Identity document analysis**, or **Document analysis**.
3. On the **Run predictions** page for your chosen Ready-to-use model, choose **Batch prediction**.
4. Choose **Select dataset** if you’ve already imported your dataset. If not, choose **Import new dataset**, and then you are directed through the import data workflow.
5. From the list of available datasets, select your dataset and choose **Generate predictions**. If your use case is document analysis, continue to Step 6.
6. (Optional) If your use case is Document analysis, another dialog box called **Select features to include in batch prediction** appears. You can select **Forms**, **Tables**, and **Signatures** to group the results by those features. Then, choose **Generate predictions**.

After the prediction job finishes running, on the **Run predictions** page, you see an output dataset listed under **Predictions**. This dataset contains your results, and if you select the **More options** icon...
you can choose **View prediction results** to preview the analysis of your document data.

The following information describes the results for each type of solution:

- For expense analysis, the results are categorized into **Summary fields**, which include fields such as the total on a receipt, and **Line item fields**, which include fields such as individual items on a receipt. The identified fields are highlighted on the document image in the output.

- For identity document analysis, the output shows you the fields that the Ready-to-use model identified, such as first and last name, address, or date of birth. The identified fields are highlighted on the document image in the output.

- For document analysis, the results are categorized into **Raw text**, **Forms**, **Tables**, and **Signatures**. **Raw text** includes all of the extracted text, while **Forms**, **Tables**, and **Signatures** only include information on the form that falls into those categories. For example, **Tables** only includes information extracted from tables in the document. The identified fields are highlighted on the document image in the output.

After previewing your results, you can choose **Download prediction** and download the results as a ZIP file.

### Use custom models

With Amazon SageMaker Canvas, you can build a custom model that is trained with your data. By training a custom model on your data, you are able to capture characteristics and trends that are specific and most representative of your data. For example, you might want to create a custom time series forecasting model that you train on inventory data from your warehouse to manage your logistics operations.

You can train a Canvas custom model on the following types of datasets:

- Tabular (including numeric, categorical, timeseries, and text data)
- Image

The following table shows the types of custom models that you can build in Canvas, along with their supported data types and data sources.
<table>
<thead>
<tr>
<th>Model type</th>
<th>Example use case</th>
<th>Supported data types</th>
<th>Supported data sources</th>
</tr>
</thead>
<tbody>
<tr>
<td>Numeric prediction</td>
<td>Predicting house prices based on features like square footage</td>
<td>Numeric</td>
<td>Local upload, Amazon S3, SaaS connectors</td>
</tr>
<tr>
<td>2 category prediction</td>
<td>Predicting whether or not a customer is likely to churn</td>
<td>Binary or categorical</td>
<td>Local upload, Amazon S3, SaaS connectors</td>
</tr>
<tr>
<td>3+ category prediction</td>
<td>Predicting patient outcomes after being discharged from the hospital</td>
<td>Categorical</td>
<td>Local upload, Amazon S3, SaaS connectors</td>
</tr>
<tr>
<td>Time series forecasting</td>
<td>Predicting your inventory for the next quarter</td>
<td>Timeseries</td>
<td>Local upload, Amazon S3, SaaS connectors</td>
</tr>
<tr>
<td>Single-label image prediction</td>
<td>Predicting types of manufacturing defects in images</td>
<td>Image (JPG, PNG)</td>
<td>Local upload, Amazon S3</td>
</tr>
<tr>
<td>Multi-category text prediction</td>
<td>Predicting categories of products, such as clothing, electronics, or household goods, based on product descriptions</td>
<td>Source column: text</td>
<td>Local upload, Amazon S3</td>
</tr>
</tbody>
</table>

**Get started**

To get started with building and generating predictions from a custom model, do the following:
• Determine your use case and type of model that you want to build. For more information about the custom model types, see Build a custom model. For more information about the data types and sources supported for custom models, see Import data into Canvas.

• Import your data into Canvas. You can build a custom model with any tabular or image dataset that meets the input requirements. For more information about the input requirements, see Create a dataset.

To learn more about sample datasets provided by SageMaker with which you can experiment, see Use sample datasets.

• Build your custom model. You can do a Quick build to get your model and start making predictions more quickly, or you can do a Standard build for greater accuracy.

For numeric, categorical, and time series forecasting model types, you can clean and prepare your data with features such as advanced transforms and joins. For image prediction models, you can Edit an image dataset to update your labels or add and delete images. Note that you can't use these features for multi-category text prediction models.

• Evaluate your model's performance and determine how well it might perform on real-world data.

• (Optional) For certain model types, you can collaborate with data scientists in Amazon SageMaker Studio Classic who can help review and improve your model.

• Make single or batch predictions with your model.

Note
If you already have a trained model in Amazon SageMaker Studio Classic that you'd like to share with Canvas, you can bring your own model to SageMaker Canvas. Review the BYOM prerequisites to determine whether your model is eligible for sharing.

Build a custom model

Use Amazon SageMaker Canvas to build a custom model on the dataset that you've imported. Use the model that you've built to make predictions on new data. SageMaker Canvas uses the information in the dataset to build up to 250 models and choose the one that performs the best.

When you begin building a model, Canvas automatically recommends one or more model types. Model types fall into one of the following categories:
• **Numeric prediction** – This is known as *regression* in machine learning. Use the numeric prediction model type when you want to make predictions for numeric data. For example, you might want to predict the price of houses based on features such as the house's square footage.

• **Categorical prediction** – This is known as *classification* in machine learning. When you want to categorize data into groups, use the categorical prediction model types:
  
  • **2 category prediction** – Use the 2 category prediction model type (also known as *binary classification* in machine learning) when you have two categories that you want to predict for your data. For example, you might want to determine whether a customer is likely to churn.

  • **3+ category prediction** – Use the 3+ category prediction model type (also known as *multi-class classification* in machine learning) when you have three or more categories that you want to predict for your data. For example, you might want to predict a customer's loan status based on features such as previous payments.

• **Time series forecasting** – Use time series forecasts when you want to make predictions over a period of time. For example, you might want to predict the number of items you'll sell in the next quarter. For information about time series forecasts, see [Time Series Forecasts in Amazon SageMaker Canvas](#).

• **Image prediction** – Use the single-label image prediction model type (also known as *single-label image classification* in machine learning) when you want to assign labels to images. For example, you might want to classify different types of manufacturing defects in images of your product.

• **Text prediction** – Use the multi-category text prediction model type (also known as *multi-class text classification* in machine learning) when you want to assign labels to passages of text. For example, you might have a dataset of customer reviews for a product, and you want to determine whether customers liked or disliked the product. You might have your model predict whether a given passage of text is Positive, Negative, or Neutral.

For a table of the supported input data types for each model type, see [Use custom models](#).

For each tabular data model that you build (which includes numeric, categorical, time series forecasting, and text prediction models), you choose the **Target column**. The **Target column** is the column that contains the information that you want to predict. For example, if you're building a model to predict whether people have cancelled their subscriptions, the **Target column** contains data points that are either a yes or a no about someone's cancellation status.

For image prediction models, you build the model with a dataset of images that have been assigned labels. For the unlabeled images that you provide, the model predicts a label. For example, if you're building a model to predict whether an image is a cat or a dog, you provide
images labeled as cats or dogs when building the model. Then, the model can accept unlabeled images and predict them as either cats or dogs.

**What happens when you build a model**

To build your model, you can choose either a **Quick build** or a **Standard build**. The **Quick build** has a shorter build time, but the **Standard build** generally has a higher accuracy. The following table outlines the average build times for each model and build type, along with the minimum and maximum number of data points you should have for each build type.

<table>
<thead>
<tr>
<th>Limit</th>
<th>Numeric and categorical prediction</th>
<th>Time series forecasting</th>
<th>Image prediction</th>
<th>Text prediction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quick build time</td>
<td>2-20 minutes</td>
<td>2-20 minutes</td>
<td>15-30 minutes</td>
<td>15-30 minutes</td>
</tr>
<tr>
<td>Standard build time</td>
<td>2-4 hours</td>
<td>2-4 hours</td>
<td>2-5 hours</td>
<td>2-5 hours</td>
</tr>
<tr>
<td>Maximum number of entries (rows or images) for Quick builds</td>
<td>50,000</td>
<td>50,000</td>
<td>5000</td>
<td>7500</td>
</tr>
</tbody>
</table>

If you log out while running a **Quick build**, your build might be interrupted until you log in again. When you log in again, Canvas resumes the **Quick build**.

Canvas predicts values by using the information in the rest of the dataset, depending on the model type:

- For categorical prediction, Canvas puts each row into one of the categories listed in the **Target column**.
- For numeric prediction, Canvas uses the information in the dataset to predict the numeric values in the **Target column**.
- For time series forecasting, Canvas uses historical data to predict values for the **Target column** in the future.
• For image prediction, Canvas uses images that have been assigned labels to predict labels for unlabeled images.

• For text prediction, Canvas analyzes text data that has been assigned labels to predict labels for passages of unlabeled text.

Additional features to help you build your model

Note

The following features are available for numeric and categorical prediction and time series forecasting models.

Before building your model, you can filter your data or prepare it using advanced transforms. For more information about preparing your data for model building, see [Prepare data with advanced transformations](#).

You can also use visualization and analytics to explore your data and determine which features are best to include in your model. For more information, see [Explore and analyze your data](#).

To learn more about additional features such as previewing your model, validating your dataset, and changing the size of the random sample used to build your model, see [Preview your model](#).

For tabular datasets with multiple columns (such as datasets for building categorical, numeric, or time series forecasting model types), you might have rows with missing data points. While Canvas builds the model, it automatically adds missing values. Canvas uses the values in your dataset to perform a mathematical approximation for the missing values. For the highest model accuracy, we recommend adding in the missing data if you can find it. Note that the missing data feature is not supported for text prediction or image prediction models.

Get started

To get started with building a custom model, see [Build a model](#) and follow the procedure for the type of model that you want to build.

Build a model

The following sections show you how to build a model for each of the main types of custom models.
To build numeric prediction, 2 category prediction, or 3+ category prediction models, see Build a custom numeric or categorical prediction model.

To build single-label image prediction models, see Build a custom image prediction model.

To build multi-category text prediction models, see Build a custom text prediction model.

To build time series forecasting models, see Build a time series forecasting model.

**Note**

If you encounter an error during post-building analysis that tells you to increase your quota for `ml.m5.2xlarge` instances, see Request a Quota Increase.

---

**Build a custom numeric or categorical prediction model**

Numeric and categorical prediction models support both Quick builds and Standard builds.

To build a numeric or categorical prediction model, use the following procedure:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **My models**.
3. Choose **New model**.
4. In the **Create new model** dialog box, do the following:
   a. Enter a name in the **Model name** field.
   b. Select the **Predictive analysis** problem type.
   c. Choose **Create**.
5. For **Select dataset**, select your dataset from the list of datasets. If you haven’t already imported your data, choose **Import** to be directed through the import data workflow.
6. When you’re ready to begin building your model, choose **Select dataset**.
7. On the **Build** tab, for the **Target column** dropdown list, select the target for your model that you would like to predict.
8. For **Model type**, Canvas automatically detects the problem type for you. If you want to change the type or configure advanced model settings, choose **Configure model**.

When the **Configure model** dialog box opens, do the following:
a. For **Model type**, choose the model type that you want to build.

b. After you choose the model type, there are additional **Advanced settings**. For more information about each of the advanced settings, see Advanced model building configurations. To configure the advanced settings, do the following:

i. (Optional) For the **Objective metric** dropdown menu, select the metric that you want Canvas to optimize while building your model. If you don't select a metric, Canvas chooses one for you by default. For descriptions of the available metrics, see Metrics reference.

ii. For **Training method**, choose **Auto**, **Ensemble**, or **Hyperparameter optimization (HPO) mode**.

iii. For **Algorithms**, select the algorithms that you want to include for building model candidates.

iv. For **Data split**, specify in percentages how you want to split your data between the **Training set** and the **Validation set**. The training set is used for building the model, while the validation set is used for testing accuracy of model candidates.

v. For **Max candidates and runtime**, do the following:

   A. Set the **Max candidates** value, or the maximum number of model candidates that Canvas can generate. Note that **Max candidates** is only available in HPO mode.

   B. Set the hour and minute values for **Max job runtime**, or the maximum amount of time that Canvas can spend building your model. After the maximum time, Canvas stops building and selects the best model candidate.

c. After configuring the advanced settings, choose **Save**.

9. Select or deselect columns in your data to include or drop them from your build.

10. (Optional) Use the visualization and analytics tools that Canvas provides to visualize your data and determine which features you might want to include in your model. For more information, see Explore and analyze your data.

---

**Note**

If you make batch predictions with your model after building, Canvas adds dropped columns to your prediction results. However, Canvas does not add the dropped columns to your batch predictions for time series models.
11. (Optional) Use data transformations to clean, transform, and prepare your data for model building. For more information, see [Prepare your data with advanced transformations](#). You can view and remove your transforms by choosing **Model recipe** to open the **Model recipe** side panel.

12. (Optional) For additional features such as previewing the accuracy of your model, validating your dataset, and changing the size of the random sample that Canvas takes from your dataset, see [Preview your model](#).

13. After reviewing your data and making any changes to your dataset, choose **Quick build** or **Standard build** to begin a build for your model. The following screenshot shows the **Build** page and the **Quick build** and **Standard build** options.

After your model begins building, you can leave the page. When the model shows as **Ready** on the **My models** page, it's ready for analysis and predictions.

**Build a custom image prediction model**

Single-label image prediction models support both **Quick builds** and **Standard builds**.

To build a single-label image prediction model, use the following procedure:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **My models**.
3. Choose **New model**.

4. In the **Create new model** dialog box, do the following:
   
   a. Enter a name in the **Model name** field.
   
   b. Select the **Image analysis** problem type.
   
   c. Choose **Create**.

5. For **Select dataset**, select your dataset from the list of datasets. If you haven’t already imported your data, choose **Import** to be directed through the import data workflow.

6. When you’re ready to begin building your model, choose **Select dataset**.

7. On the **Build** tab, you see the **Label distribution** for the images in your dataset. The **Model type** is set to **Single-label image prediction**.

8. On this page, you can preview your images and edit the dataset. If you have any unlabeled images, choose **Edit dataset** and **Assign labels to unlabeled images**. You can also perform other tasks when you **Edit an image dataset**, such as renaming labels and adding images to the dataset.

9. After reviewing your data and making any changes to your dataset, choose **Quick build** or **Standard build** to begin a build for your model. The following screenshot shows the **Build** page of an image prediction model that is ready to be built.
After your model begins building, you can leave the page. When the model shows as Ready on the My models page, it’s ready for analysis and predictions.

**Build a custom text prediction model**

Multi-category text prediction models support both Quick builds and Standard builds.

To build a text prediction model, use the following procedure:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose My models.
3. Choose New model.
4. In the Create new model dialog box, do the following:
   a. Enter a name in the Model name field.
   b. Select the Text analysis problem type.
   c. Choose Create.
5. For Select dataset, select your dataset from the list of datasets. If you haven’t already imported your data, choose Import to be directed through the import data workflow.
6. When you’re ready to begin building your model, choose Select dataset.
7. On the Build tab, for the Target column dropdown list, select the target for your model that you would like to predict. The target column must have a binary or categorical data type, and there must be at least 25 entries (or rows of data) for each unique label in the target column.
8. For Model type, confirm that the model type is automatically set to Multi-category text prediction.
9. For the training column, select your source column of text data. This should be the column containing the text that you want to analyze.
10. Choose Quick build or Standard build to begin building your model. The following screenshot shows the Build page of a text prediction model that is ready to be built.
After your model begins building, you can leave the page. When the model shows as Ready on the My models page, it's ready for analysis and predictions.

**Build a time series forecasting model**

Time series forecasting models support both **Quick builds** and **Standard builds**.

To build a time series forecasting model, use the following procedure:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **My models**.
3. Choose **New model**.
4. In the **Create new model** dialog box, do the following:
   a. Enter a name in the **Model name** field.
   b. Select the **Time series forecasting** problem type.
   c. Choose **Create**.
5. For **Select dataset**, select your dataset from the list of datasets. If you haven’t already imported your data, choose **Import** to be directed through the import data workflow.
6. When you're ready to begin building your model, choose **Select dataset**.
7. On the **Build** tab, for the **Target column** dropdown list, select the target for your model that you would like to predict.

8. For **Model type**, Canvas automatically detects the problem type for you. If you want to change the type or configure advanced model settings, choose **Configure model**.

When the **Configure model** dialog box opens, do the following:

a. For **Model type**, choose the model type that you want to build.

b. For **Time series configuration**, specify the following values:

   i. For the **Item ID column** dropdown, choose a column in your dataset that uniquely identifies each row.

   ii. (Optional) For the **Group column** dropdown, choose a categorical column that you want to use for grouping your forecasting values.

   iii. For the **Time stamp column** dropdown, select the column with timestamps (in datetime format). For more information about the accepted datetime formats, see *Time Series Forecasts in Amazon SageMaker Canvas*.

   iv. For the **Forecast length** field, enter the period of time for which you want to forecast values.

   v. (Optional) Turn on the **Use holiday schedule** toggle to select a holiday schedule from various countries.

   vi. Choose **Save**.

c. After configuring your time series settings, there are additional **Advanced settings**. For more information about each of the advanced settings, see *Advanced model building configurations*. To configure the advanced settings, do the following:

   i. (Optional) For the **Objective metric** dropdown menu, select the metric that you want Canvas to optimize while building your model. If you don’t select a metric, Canvas chooses one for you by default. For descriptions of the available metrics, see *Metrics reference*.

   ii. For **Forecast quantiles**, enter up to 5 comma-separated quantile values to specify the upper and lower bounds of your forecast.

   iii. For **Forecast frequency**, specify the following:

      A. For **Frequency**, enter a **Unit** and **Value** to specify the frequency for the forecasted values.
B. For **Aggregation**, select an aggregation method for how Canvas handles data that doesn’t fit the frequency.

   iv. For **Max job runtime**, set hour and minute values for the maximum amount of time that Canvas can spend building your model. After the maximum time, Canvas stops building and selects the best model candidate.

d. After configuring the advanced settings, choose **Save**.

9. Select or deselect columns in your data to include or drop them from your build.

   ![Note]
   
   If you make batch predictions with your model after building, Canvas adds dropped columns to your prediction results. However, Canvas does not add the dropped columns to your batch predictions for time series models.

10. (Optional) Use the visualization and analytics tools that Canvas provides to visualize your data and determine which features you might want to include in your model. For more information, see **Explore and analyze your data**.

11. (Optional) Use data transformations to clean, transform, and prepare your data for model building. For more information, see **Prepare your data with advanced transformations**. You can view and remove your transforms by choosing **Model recipe** to open the **Model recipe** side panel.

12. (Optional) For additional features such as previewing the accuracy of your model, validating your dataset, and changing the size of the random sample that Canvas takes from your dataset, see **Preview your model**.

13. After reviewing your data and making any changes to your dataset, choose **Quick build** or **Standard build** to begin a build for your model.

After your model begins building, you can leave the page. When the model shows as **Ready** on the **My models** page, it’s ready for analysis and predictions.

**Advanced model building configurations**

Amazon SageMaker Canvas supports various advanced settings that you can configure when building a model. The following page lists all of the advanced settings along with additional information about their options and configurations.
The following advanced settings are currently only supported for numeric, categorical, and time series forecasting model types.

Advanced numeric and categorical prediction model settings

Canvas supports the following advanced settings for numeric and categorical prediction model types.

Objective metric

The objective metric is the metric that you want Canvas to optimize while building your model. If you don’t select a metric, Canvas chooses one for you by default. For descriptions of the available metrics, see the Metrics reference.

Training method

Canvas can automatically select the training method based on the dataset size, or you can select it manually. The following training methods are available for you to choose from:

- **Ensembling** – SageMaker leverages the AutoGluon library to train several base models. To find the best combination for your dataset, ensemble mode runs 5–10 trials with different model and meta parameter settings. Then, these models are combined using a stacking ensemble method to create an optimal predictive model. For a list of algorithms supported by ensemble mode for tabular data, see the following Algorithms section.

- **Hyperparameter optimization (HPO)** – SageMaker finds the best version of a model by tuning hyperparameters using Bayesian optimization or multi-fidelity optimization while running training jobs on your dataset. HPO mode selects the algorithms that are most relevant to your dataset and selects the best range of hyperparameters to tune your models. To tune your models, HPO mode runs up to 100 trials (default) to find the optimal hyperparameters settings within the selected range. If your dataset size is less than 100 MB, SageMaker uses Bayesian optimization. SageMaker chooses multi-fidelity optimization if your dataset is larger than 100 MB.

  For a list of algorithms supported by HPO mode for tabular data, see the following Algorithms section.
Auto – SageMaker automatically chooses either ensembling mode or HPO mode based on your dataset size. If your dataset is larger than 100 MB, SageMaker chooses HPO mode. Otherwise, it chooses ensembling mode.

Algorithms

In **Ensembling** mode, Canvas supports the following machine learning algorithms:

- **LightGBM** – An optimized framework that uses tree-based algorithms with gradient boosting. This algorithm uses trees that grow in breadth, rather than depth, and is highly optimized for speed.
- **CatBoost** – A framework that uses tree-based algorithms with gradient boosting. Optimized for handling categorical variables.
- **XGBoost** – A framework that uses tree-based algorithms with gradient boosting that grows in depth, rather than breadth.
- **Random Forest** – A tree-based algorithm that uses several decision trees on random sub-samples of the data with replacement. The trees are split into optimal nodes at each level. The decisions of each tree are averaged together to prevent overfitting and improve predictions.
- **Extra Trees** – A tree-based algorithm that uses several decision trees on the entire dataset. The trees are split randomly at each level. The decisions of each tree are averaged to prevent overfitting and to improve predictions. Extra trees add a degree of randomization in comparison to the random forest algorithm.
- **Linear Models** – A framework that uses a linear equation to model the relationship between two variables in observed data.
- Neural network torch – A neural network model that's implemented using Pytorch.
- Neural network fast.ai – A neural network model that's implemented using fast.ai.

In **HPO mode**, Canvas supports the following machine learning algorithms:

- **XGBoost** – A supervised learning algorithm that attempts to accurately predict a target variable by combining an ensemble of estimates from a set of simpler and weaker models.
- Deep learning algorithm – A multilayer perceptron (MLP) and feedforward artificial neural network. This algorithm can handle data that is not linearly separable.
Data split

You have the option to specify how you want to split your dataset between the training set (the portion of your dataset used for building the model) and the validation set, (the portion of your dataset used for verifying the model's accuracy). For example, a common split ratio is 80% training and 20% validation, where 80% of your data is used to build the model while 20% is saved for measuring model performance. If you don't specify a custom ratio, then Canvas splits your dataset automatically.

Max candidates

Note

This feature is only available in the HPO training mode.

You can specify the maximum number of model candidates that Canvas generates while building your model. We recommend that you use the default number of candidates, which is 100, to build the most accurate models. The maximum number you can specify is 250. Decreasing the number of model candidates may impact your model's accuracy.

Max job runtime

You can specify the maximum job runtime, or the maximum amount of time that Canvas spends building your model. After the time limit, Canvas stops building and selects the best model candidate.

The maximum time that you can specify is 720 hours. We highly recommend that you keep the maximum job runtime greater than 30 minutes to ensure that Canvas has enough time to generate model candidates and finish building your model.

Advanced time series forecasting model settings

For time series forecasting models, Canvas supports the following metrics listed in the previous section:

- Objective metric
- Algorithms
- Max job runtime
Time series forecasting models also support the following advanced settings:

**Aggregation**

If you set the forecasting frequency lower than the frequency of your recorded data, Canvas aggregates any data points that don’t match the new frequency. For example, if you have daily data points but would like to make a weekly forecast, you can set the frequency to weekly, and then Canvas combines all of the daily data points for each week into a single record. Aggregation is only supported for the target column, and the column values must be in datetime format.

The default aggregation method is to sum the values of the aggregated data points, but you can also set the aggregation method to the following:

- **avg** – Canvas sets the record's value to the average of all the aggregated data points.
- **first** – Canvas sets the record's value to the first value of the aggregated data points.
- **min** – Canvas sets the record's value to the minimum value found in the aggregated data points.
- **max** – Canvas sets the record's value to the maximum value found in the aggregated data points.

**Forecast quantiles**

For time series forecasting, SageMaker trains 6 model candidates with your target time series. Then, SageMaker combines these models using a stacking ensemble method to create an optimal forecasting model for a given objective metric. Each forecasting model generates a probabilistic forecast by producing forecasts at quantiles between P1 and P99. These quantiles are used to account for forecast uncertainty. By default, forecasts are generated for 0.1 (p10), 0.5 (p50), and 0.9 (p90). You can choose to specify up to five of your own quantiles from 0.01 (p1) to 0.99 (p99), by increments of 0.01 or higher.

**Preview your model**

Tip

The following functionalities are only available for custom models built with tabular datasets. Multi-category text prediction models are also excluded.

SageMaker Canvas provides you with tools to preview your model and validate data before you begin building. The following functionalities include previewing the accuracy of your model,
validating your dataset to prevent issues while building the model, and changing the size of the random sample for your model.

**Preview a model**

With Amazon SageMaker Canvas, you can get insights from your data before you build a model by choosing **Preview model**. For example, you can see how the data in each column is distributed. For models built using categorical data, you can also choose **Preview model** to generate an **Estimated accuracy** prediction of how well the model might analyze your data. The accuracy of a **Quick build** or a **Standard build** represents how well the model can perform on real data and is generally higher than the **Estimated accuracy**.

Amazon SageMaker Canvas automatically handles missing values in your dataset while it builds the model. It infers the missing values by using adjacent values that are present in the dataset.

**Validate data**

Before you build your model, SageMaker Canvas checks your dataset for issues that might cause your build to fail. If SageMaker Canvas finds any issues, then it warns you on the **Build** page before you attempt to build a model.

You can choose **Validate data** to see a list of the issues with your dataset. You can then use the SageMaker Canvas **data preparation features**, or your own tools, to fix your dataset before starting a build. If you don’t fix the issues with your dataset, then your build fails.
If you make changes to your dataset to fix the issues, you have the option to re-validate your dataset before attempting a build. We recommend that you re-validate your dataset before building.

The following table shows the issues that SageMaker Canvas checks for in your dataset and how to resolve them.

<table>
<thead>
<tr>
<th>Issue</th>
<th>Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wrong model type for your data</td>
<td>Try another model type or use a different dataset.</td>
</tr>
<tr>
<td>Missing values in your target column</td>
<td>Replace the missing values, drop rows with missing values, or use a different dataset.</td>
</tr>
<tr>
<td>Too many unique labels in your target column</td>
<td>Verify that you've used the correct column for your target column, or use a different dataset.</td>
</tr>
<tr>
<td>Too many non-numeric values in your target column</td>
<td>Choose a different target column, select another model type, or use a different dataset.</td>
</tr>
<tr>
<td>One or more column names contain double underscores</td>
<td>Rename the columns to remove any double underscores, and try again.</td>
</tr>
<tr>
<td>None of the rows in your dataset are complete</td>
<td>Replace the missing values, or use a different dataset.</td>
</tr>
<tr>
<td>Too many unique labels for the number of rows in your data</td>
<td>Check that you're using the right target column, increase the number of rows in your dataset, consolidate similar labels, or use a different dataset.</td>
</tr>
</tbody>
</table>

**Random sample**

SageMaker Canvas uses the random sampling method to sample your dataset. The random sample method means that each row has an equal chance of being picked for the sample. You can choose a column in the preview to get summary statistics for the random sample, such as the mean and the mode.
By default, SageMaker Canvas uses a random sample size of 20,000 rows from your dataset for datasets with more than 20,000 rows. For datasets smaller than 20,000 rows, the default sample size is the number of rows in your dataset. You can increase or decrease the sample size by choosing **Random sample** in the **Build** tab of the SageMaker Canvas application. You can use the slider to select your desired sample size, and then choose **Update** to change the sample size. The maximum sample size you can choose for a dataset is 40,000 rows, and the minimum sample size is 500 rows. If you choose a large sample size, the dataset preview and summary statistics might take a few moments to reload.

The **Build** page shows a preview of 100 rows from your dataset. If the sample size is the same size as your dataset, then the preview uses the first 100 rows of your dataset. Otherwise, the preview uses the first 100 rows of the random sample.

**Edit an image dataset**

In Amazon SageMaker Canvas, you can edit your image datasets and review your labels before building a model. You might want to perform tasks such as assigning labels to unlabeled images or adding more images to the dataset. These tasks can all be done in the Canvas application, providing you with one place to modify your dataset and build a model.

**Note**

Before building a model, you must assign labels to all images in your dataset. Also, you must have at least 25 images per label and a minimum of two labels. For more information about assigning labels, see the section on this page called **Assign labels to unlabeled images**. If you can’t determine a label for an image, you should delete it from your dataset. For more information about deleting images, see the section on this page **Add or delete images from the dataset**.

To begin editing your image dataset, you should be on the **Build** tab while building your single-label image prediction model.

A new page opens that shows the images in your dataset along with their labels. This page categorizes your image dataset into **Total images**, **Labeled images**, and **Unlabeled images**. You can also review the **Dataset preparation guide** for best practices on building a more accurate image prediction model.

The following screenshot shows the page for editing your image dataset.
From this page, you can do the following actions.

**View the properties for each image (label, size, dimensions)**

To view an individual image, you can search for it by file name in the search bar. Then, choose the image to open the full view. You can view the image properties and reassign the image’s label. Choose **Save** when you’re doing viewing the image.

**Add, rename, or delete labels in the dataset**

Canvas lists the labels for your dataset in the left navigation pane. You can add new labels to the dataset by entering a label in the **Add label** text field.

To rename or delete a label from your dataset, choose the **More options** icon ( ) next to the label and select either **Rename** or **Delete**. If you rename the label, you can enter the new label name and choose **Confirm**. If you delete the label, the label is removed from all images in your dataset that have that label. Any images with that label are left unlabeled.

**Assign labels to unlabeled images**

To view the unlabeled images in your dataset, choose **Unlabeled** in the left navigation pane. For each image, select it and open the label titled **Unlabeled** and select a label to assign to the image.
from the dropdown list. You can also select more than one image and perform this action, and all selected images are assigned the label you chose.

**Reassign labels to images**

You can reassign labels to images by selecting the image (or multiple images at a time) and opening the dropdown titled with the current label. Select your desired label, and the image or images are updated with the new label.

**Sort your images by label**

You can view all the images for a given label by choosing the label in the left navigation pane.

**Add or delete images from the dataset**

You can add more images to your dataset by choosing **Add images** in the top navigation pane. You’ll be taken through the workflow to import more images. The images you import are added to your existing dataset.

You can delete images from your dataset by selecting them and then choosing **Delete** in the top navigation pane.

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**Note**

After making any changes to your dataset, choose **Save dataset** to make sure that you don’t lose your changes.

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**Explore and analyze your data**

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**Note**

You can only use SageMaker Canvas visualizations and analytics for models built on tabular datasets. Multi-category text prediction models are also excluded.

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In Amazon SageMaker Canvas, you can explore the variables in your dataset using visualizations and analytics and create in-application visualizations and analytics. You can use these explorations to uncover relationships between your variables before building your model.
For more information about visualization techniques in Canvas, see [Explore your data using visualization techniques](#).

For more information about analytics in Canvas, see [Explore your data using analytics](#).

## Explore your data using visualization techniques

### Note
You can only use SageMaker Canvas visualizations for models built on tabular datasets. Multi-category text prediction models are also excluded.

With Amazon SageMaker Canvas, you can explore and visualize your data to gain advanced insights into your data before building your ML models. You can visualize using scatter plots, bar charts, and box plots, which can help you understand your data and discover the relationships between features that could affect the model accuracy.

In the **Build** tab of the SageMaker Canvas application, choose **Data visualizer** to begin creating your visualizations.

You can change the visualization sample size to adjust the size of the random sample taken from your dataset. A sample size that is too large might affect the performance of your data visualizations, so we recommend that you choose an appropriate sample size. To change the sample size, use the following procedure.

1. Choose **Visualization sample**.
2. Use the slider to select your desired sample size.
3. Choose **Update** to confirm the change to your sample size.

### Note
Certain visualization techniques require columns of a specific data type. For example, you can only use numeric columns for the x and y-axes of scatter plots.
Scatter plot

To create a scatter plot with your dataset, choose **Scatter plot** in the **Visualization** panel. Choose the features you want to plot on the x and y-axes from the **Columns** section. You can drag and drop the columns onto the axes or, once an axis has been dropped, you can choose a column from the list of supported columns.

You can use **Color by** to color the data points on the plot with a third feature. You can also use **Group by** to group the data into separate plots based on a fourth feature.

The following image shows a scatter plot that uses **Color by** and **Group by**. In this example, each data point is colored by the *MaritalStatus* feature, and grouping by the *Department* feature results in a scatter plot for the data points of each department.

Bar chart

To create a bar chart with your dataset, choose **Bar chart** in the **Visualization** panel. Choose the features you want to plot on the x and y-axes from the **Columns** section. You can drag and drop the columns onto the axes or, once an axis has been dropped, you can choose a column from the list of supported columns.

You can use **Group by** to group the bar chart by a third feature. You can use **Stack by** to vertically shade each bar based on the unique values of a fourth feature.
The following image shows a bar chart that uses **Group by** and **Stack by**. In this example, the bar chart is grouped by the `MaritalStatus` feature and stacked by the `JobLevel` feature. For each `JobRole` on the x axis, there is a separate bar for the unique categories in the `MaritalStatus` feature, and every bar is vertically stacked by the `JobLevel` feature.

**Box plot**

To create a box plot with your dataset, choose **Box plot** in the **Visualization** panel. Choose the features you want to plot on the x and y-axes from the **Columns** section. You can drag and drop the columns onto the axes or, once an axis has been dropped, you can choose a column from the list of supported columns.

You can use **Group by** to group the box plots by a third feature.

The following image shows a box plot that uses **Group by**. In this example, the x and y-axes show `JobLevel` and `JobSatisfaction`, respectively, and the colored box plots are grouped by the `Department` feature.
Explore your data using analytics

**Note**

You can only use SageMaker Canvas analytics for models built on tabular datasets. Multi-category text prediction models are also excluded.

With analytics in Amazon SageMaker Canvas, you can explore your dataset and gain insight on all of your variables before building a model. You can determine the relationships between features in your dataset using correlation matrices. You can use this technique to summarize your dataset into a matrix that shows the correlations between two or more values. This helps you identify and visualize patterns in a given dataset for advanced data analysis.

The matrix shows the correlation between each feature as positive, negative, or neutral. You might want to include features that have a high correlation with each other when building your model. Features that have little to no correlation might be irrelevant to your model, and you can drop those features when building your model.

To get started with correlation matrices in SageMaker Canvas, see the following section.
Create a correlation matrix

You can create a correlation matrix when you are preparing to build a model in the Build tab of the SageMaker Canvas application.

For instructions on how to begin creating a model, see Build a model.

After you've started preparing a model in the SageMaker Canvas application, do the following:

1. In the Build tab, choose Data visualizer.
2. Choose Analytics.
3. Choose Correlation matrix.

You should see a visualization similar to the following screenshot, which shows up to 15 columns of the dataset organized into a correlation matrix.

![Correlation Matrix Visualization](image)

After you've created the correlation matrix, you can customize it by doing the following:

1. Choose your columns

For Columns, you can select the columns that you want to include in the matrix. You can compare up to 15 columns from your dataset.
Note

You can use numeric, categorical, or binary column types for a correlation matrix. The correlation matrix doesn’t support datetime or text data column types.

To add or remove columns from the correlation matrix, select and deselect columns from the Columns panel. You can also drag and drop columns from the panel directly onto the matrix. If your dataset has a lot of columns, you can search for the columns you want in the Search columns bar.

To filter the columns by data type, choose the dropdown list and select All, Numeric, or Categorical. Selecting All shows you all of the columns from your dataset, whereas the Numeric and Categorical filters only show you the numeric or categorical columns in your dataset. Note that binary column types are included in the numeric or categorical filters.

For the best data insights, include your target column in the correlation matrix. When you include your target column in the correlation matrix, it appears as the last feature on the matrix with a target symbol.

2. Choose your correlation type

SageMaker Canvas supports different correlation types, or methods for calculating the correlation between your columns.

To change the correlation type, use the Columns filter mentioned in the preceding section to filter for your desired column type and columns. You should see the Correlation type in the side panel. For numeric comparisons, you have the option to select either Pearson or Spearman. For categorical comparisons, the correlation type is set as MI. For categorical and mixed comparisons, the correlation type is set as Spearman & MI.

For matrices that only compare numeric columns, the correlation type is either Pearson or Spearman. The Pearson measure evaluates the linear relationship between two continuous variables. The Spearman measure evaluates the monotonic relationship between two variables. For both Pearson and Spearman, the scale of correlation ranges from -1 to 1, with either end of the scale indicating a perfect correlation (a direct 1:1 relationship) and 0 indicating no correlation. You might want to select Pearson if your data has more linear relationships (as revealed by a scatter plot visualization). If your data is not linear, or contains a mixture of linear and monotonic relationships, then you might want to select Spearman.
For matrices that only compare categorical columns, the correlation type is set to Mutual Information Classification (MI). The MI value is a measure of the mutual dependence between two random variables. The MI measure is on a scale of 0 to 1, with 0 indicating no correlation and 1 indicating a perfect correlation.

For matrices that compare a mix of numeric and categorical columns, the correlation type **Spearman & MI** is a combination of the Spearman and MI correlation types. For correlations between two numeric columns, the matrix shows the Spearman value. For correlations between a numeric and categorical column or two categorical columns, the matrix shows the MI value.

Lastly, remember that correlation does not necessarily indicate causation. A strong correlation value only indicates that there is a relationship between two variables, but the variables might not have a causal relationship. Carefully review your columns of interest to avoid bias when building your model.

### 3. Filter your correlations

In the side panel, you can use the **Filter correlations** feature to filter for the range of correlation values that you want to include in the matrix. For example, if you want to filter for features that only have positive or neutral correlation, you can set the **Min** to 0 and the **Max** to 1 (valid values are -1 to 1).

For Spearman and Pearson comparisons, you can set the **Filter correlations** range anywhere from -1 to 1, with 0 meaning that there is no correlation. -1 and 1 mean that the variables have a strong negative or positive correlation, respectively.

For MI comparisons, the correlation range only goes from 0 to 1, with 0 meaning that there is no correlation and 1 meaning that the variables have a strong correlation, either positive or negative.

Each feature has a perfect correlation (1) with itself. Therefore, you might notice that the top row of the correlation matrix is always 1. If you want to exclude these values, you can use the filter to set the **Max** less than 1.

Keep in mind that if your matrix compares a mix of numeric and categorical columns and uses the **Spearman & MI** correlation type, then the *categorical x numeric* and *categorical x categorical* correlations (which use the MI measure) are on a scale of 0 to 1, whereas the *numeric x numeric* correlations (which use the Spearman measure) are on a scale of -1 to 1. Review your correlations of interest carefully to ensure that you know the correlation type being used to calculate each value.
4. Choose the visualization method

In the side panel, you can use **Visualize by** to change the visualization method of the matrix. Choose the **Numeric** visualization method to show the correlation (Pearson, Spearman, or MI) value, or choose the **Size** visualization method to visualize the correlation with differently sized and colored dots. If you choose **Size**, you can hover over a specific dot on the matrix to see the actual correlation value.

5. Choose a color palette

In the side panel, you can use **Color selection** to change the color palette used for the scale of negative to positive correlation in the matrix. Select one of the alternative color palettes to change the colors used in the matrix.

Prepare data with advanced transformations

**Note**

You can only use advanced transformations for models built on tabular datasets. Multi-category text prediction models are also excluded.

Your machine learning dataset might require data preparation before you build your model. You might want to clean your data due to various issues, which might include missing values or outliers, and perform feature engineering to improve the accuracy of your model. Amazon SageMaker Canvas provides ML data transforms with which you can clean, transform, and prepare your data for model building. You can use these transforms on your datasets without any code. SageMaker Canvas adds the transforms you use to the **Model recipe**, which is a record of the data preparation done on your data before building the model. Any data transforms you use only modify the input data for model building and do not modify your original data source.

The following transforms are available in SageMaker Canvas for you to prepare your data for building.

**Note**

The preview of your dataset shows the first 100 rows of the dataset. If your dataset has more than 20,000 rows, Canvas takes a random sample of 20,000 rows and previews the first 100 rows from that sample. You can only search for and specify values from the
previewed rows, and the filter functionality only filters the previewed rows and not the entire dataset.

**Drop columns**

You can exclude a column from your model build by dropping it in the **Build tab** of the SageMaker Canvas application. Deselect the column you want to drop, and it isn't included when building the model.

**Note**

If you drop columns and then make batch predictions with your model, SageMaker Canvas adds the dropped columns back to the output dataset available for you to download. However, SageMaker Canvas does not add the dropped columns back for time series models.

**Filter rows**

The filter functionality filters the previewed rows (the first 100 rows of your dataset) according to conditions that you specify. Filtering rows creates a temporary preview of the data and does not impact the model building. You can filter to preview rows that have missing values, contain outliers, or meet custom conditions in a column you choose.

**Filter rows by missing values**

Missing values are a common occurrence in machine learning datasets. If you have rows with null or empty values in certain columns, you might want to filter for and preview those rows.

To filter missing values from your previewed data, do the following.

1. In the **Build tab** of the SageMaker Canvas application, choose **Filter by rows**.
2. Choose the **Column** you want to check for missing values.
3. For the **Operation**, choose **Is missing**.

SageMaker Canvas filters for rows that contain missing values in the **Column** you selected and provides a preview of the filtered rows.
Filter rows by outliers

Outliers, or rare values in the distribution and range of your data, can negatively impact model accuracy and lead to longer building times. SageMaker Canvas enables you to detect and filter rows that contain outliers in numeric columns. You can choose to define outliers with either standard deviations or a custom range.

To filter for outliers in your data, do the following.

1. In the Build tab of the SageMaker Canvas application, choose Filter by rows.
2. Choose the Column you want to check for outliers.
3. For the Operation, choose Is outlier.
4. Set the Outlier range to either Standard deviation or Custom range.
5. If you choose Standard deviation, specify a SD (standard deviation) value from 1–3. If you choose Custom range, select either Percentile or Number, and then specify the Min and Max values.

The Standard deviation option detects and filters for outliers in numeric columns using the mean and standard deviation. You specify the number of standard deviations a value must vary from the mean to be considered an outlier. For example, if you specify 3 for SD, a value must fall more than 3 standard deviations from the mean to be considered an outlier.
The **Custom range** option detects and filters for outliers in numeric columns using minimum and maximum values. Use this method if you know your threshold values that delimit outliers. You can set the **Type** of the range to either **Percentile** or **Number**. If you choose **Percentile**, the **Min** and **Max** values should be the minimum and maximum of the percentile range (0-100) that you want to allow. If you choose **Number**, the **Min** and **Max** values should be the minimum and maximum numeric values that you want to filter in the data.

**Filter rows by custom values**

You can filter for rows with values that meet custom conditions. For example, you might want to preview rows that have a price value greater than 100 before removing them. With this functionality, you can filter rows that exceed the threshold you set and preview the filtered data.

To use the custom filter functionality, do the following.

1. In the **Build** tab of the SageMaker Canvas application, choose **Filter by rows**.
2. Choose the **Column** you want to check.
3. Select the type of **Operation** you want to use, and then specify the values for the selected condition.
For the **Operation**, you can choose one of the following options. Note that the available operations depend on the data type of the column you choose. For example, you cannot create a *is greater than* operation for a column containing text values.

<table>
<thead>
<tr>
<th>Operation</th>
<th>Supported data type</th>
<th>Supported feature type</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is equal to</td>
<td>Numeric, Text</td>
<td>Binary, Categorical</td>
<td>Filters rows where the value in <strong>Column</strong> equals the values you specify.</td>
</tr>
<tr>
<td>Is not equal to</td>
<td>Numeric, Text</td>
<td>Binary, Categorical</td>
<td>Filters rows where the value in <strong>Column</strong> doesn't equal the values you specify.</td>
</tr>
<tr>
<td>Is less than</td>
<td>Numeric</td>
<td>N/A</td>
<td>Filters rows where the value in <strong>Column</strong> is less than the value you specify.</td>
</tr>
<tr>
<td>Is less than or equal to</td>
<td>Numeric</td>
<td>N/A</td>
<td>Filters rows where the value in <strong>Column</strong> is less than or equal to the value you specify.</td>
</tr>
<tr>
<td>Is greater than</td>
<td>Numeric</td>
<td>N/A</td>
<td>Filters rows where the value in <strong>Column</strong> is greater than the value you specify.</td>
</tr>
<tr>
<td>Is greater than or equal to</td>
<td>Numeric</td>
<td>N/A</td>
<td>Filters rows where the value in <strong>Column</strong> is greater than or equal to the value you specify.</td>
</tr>
<tr>
<td>Is between</td>
<td>Numeric</td>
<td>N/A</td>
<td>Filters rows where the value in <strong>Column</strong> is between or equal to two values you specify.</td>
</tr>
<tr>
<td>Contains</td>
<td>Text</td>
<td>Categorical</td>
<td>Filters rows where the value in <strong>Column</strong> contains a values you specify.</td>
</tr>
</tbody>
</table>
### Supported data types

**Operation**

<table>
<thead>
<tr>
<th>Supported data type</th>
<th>Supported feature type</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starts with</td>
<td>Text</td>
<td>Categorical</td>
</tr>
<tr>
<td>Ends with</td>
<td>Categorical</td>
<td>Categorical</td>
</tr>
</tbody>
</table>

Filters rows where the value in **Column** begins with a value you specify.

Filters rows where the value in **Column** ends with a value you specify.

---

After you set the filter operation, SageMaker Canvas updates the preview of the dataset to show you the filtered data.

---

### Functions and operators

You can use mathematical functions and operators to explore and distribute your data. You can use the SageMaker Canvas supported functions or create your own formula with your existing data and create a new column with the result of the formula. For example, you can add the corresponding values of two columns and save the result to a new column.

You can nest statements to create more complex functions. The following are some examples of nested functions that you might use.
To calculate BMI, you could use the function `weight / (height ^ 2)`.

To classify ages, you could use the function `Case(age < 18, 'child', age < 65, 'adult', 'senior')`.

You can specify functions in the data preparation stage before you build your model. To use a function, do the following.

- In the Build tab of the SageMaker Canvas application, choose View all and then choose Custom formula to open the Custom formula panel.
- In the Custom formula panel, you can choose a Formula to add to your Model Recipe. Each formula is applied to all of the values in the columns you specify. For formulas that accept two or more columns as arguments, use columns with matching data types; otherwise, you get an error or null values in the new column.
- After you've specified a Formula, add a column name in the New Column Name field. SageMaker Canvas uses this name for the new column that is created.
- (Optional) Choose Preview to preview your transform.
- To add the function to your Model Recipe, choose Add.

SageMaker Canvas saves the result of your function to a new column using the name you specified in New Column Name. You can view or remove functions from the Model Recipe panel.

SageMaker Canvas supports the following operators for functions. You can use either the text format or the in-line format to specify your function.

<table>
<thead>
<tr>
<th>Operator</th>
<th>Description</th>
<th>Supported data types</th>
<th>Text format</th>
<th>In-line format</th>
</tr>
</thead>
<tbody>
<tr>
<td>Add</td>
<td>Returns the sum of the values</td>
<td>Numeric</td>
<td>Add(sales1, sales2)</td>
<td>sales1 + sales2</td>
</tr>
<tr>
<td>Subtract</td>
<td>Returns the difference between the values</td>
<td>Numeric</td>
<td>Subtract(sales1, sales2)</td>
<td>sales1 - sales2</td>
</tr>
<tr>
<td>Operator</td>
<td>Description</td>
<td>Supported data types</td>
<td>Text format</td>
<td>In-line format</td>
</tr>
<tr>
<td>----------</td>
<td>-------------</td>
<td>----------------------</td>
<td>-------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Multiply</td>
<td>Returns the product of the values</td>
<td>Numeric</td>
<td>Multiply(sales1, sales2)</td>
<td>sales1 * sales2</td>
</tr>
<tr>
<td>Divide</td>
<td>Returns the quotient of the values</td>
<td>Numeric</td>
<td>Divide(sales1, sales2)</td>
<td>sales1 / sales2</td>
</tr>
<tr>
<td>Mod</td>
<td>Returns the result of the modulo operator (the remainder after dividing the two values)</td>
<td>Numeric</td>
<td>Mod(sales1, sales2)</td>
<td>sales1 % sales2</td>
</tr>
<tr>
<td>Abs</td>
<td>Returns the absolute value of the value</td>
<td>Numeric</td>
<td>Abs(sales1)</td>
<td>N/A</td>
</tr>
<tr>
<td>Negate</td>
<td>Returns the negative of the value</td>
<td>Numeric</td>
<td>Negate(c1)</td>
<td>-c1</td>
</tr>
<tr>
<td>Exp</td>
<td>Returns $e$ (Euler's number) raised to the power of the value</td>
<td>Numeric</td>
<td>Exp(sales1)</td>
<td>N/A</td>
</tr>
<tr>
<td>Log</td>
<td>Returns the logarithm (base 10) of the value</td>
<td>Numeric</td>
<td>Log(sales1)</td>
<td>N/A</td>
</tr>
<tr>
<td>Ln</td>
<td>Returns the natural logarithm (base $e$) of the value</td>
<td>Numeric</td>
<td>Ln(sales1)</td>
<td>N/A</td>
</tr>
<tr>
<td>Pow</td>
<td>Returns the value raised to a power</td>
<td>Numeric</td>
<td>Pow(sales1, 2)</td>
<td>sales1 ^ 2</td>
</tr>
<tr>
<td>If</td>
<td>Returns a true or false label based on a condition you specify</td>
<td>Boolean, Numeric, Text</td>
<td>If(sales1 &gt; 7000, 'truelabel', 'falselabel')</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Use custom models
<table>
<thead>
<tr>
<th>Operator</th>
<th>Description</th>
<th>Supported data types</th>
<th>Text format</th>
<th>In-line format</th>
</tr>
</thead>
<tbody>
<tr>
<td>Or</td>
<td>Returns a Boolean value of whether one of the specified values or conditions is true or not</td>
<td>Boolean</td>
<td>Or(fullprice, discount)</td>
<td>fullprice</td>
</tr>
<tr>
<td>And</td>
<td>Returns a Boolean value of whether two of the specified values or conditions are true or not</td>
<td>Boolean</td>
<td>And(sales 1, sales2)</td>
<td>sales1 &amp;&amp; sales2</td>
</tr>
<tr>
<td>Not</td>
<td>Returns a Boolean value that is the opposite of the specified value or conditions</td>
<td>Boolean</td>
<td>Not(sales1)</td>
<td>!sales1</td>
</tr>
<tr>
<td>Case</td>
<td>Returns a Boolean value based on conditional statements (returns c1 if cond1 is true, returns c2 if cond2 is true, else returns c3)</td>
<td>Boolean, Numeric, Text</td>
<td>Case(cond1, c1, cond2, c2, c3)</td>
<td>N/A</td>
</tr>
<tr>
<td>Equal</td>
<td>Returns a Boolean value of whether two values are equal</td>
<td>Boolean, Numeric, Text</td>
<td>N/A</td>
<td>c1 = c2, c1 == c2</td>
</tr>
<tr>
<td>Not equal</td>
<td>Returns a Boolean value of whether two values are not equal</td>
<td>Boolean, Numeric, Text</td>
<td>N/A</td>
<td>c1 != c2</td>
</tr>
<tr>
<td>Less than</td>
<td>Returns a Boolean value of whether c1 is less than c2</td>
<td>Boolean, Numeric, Text</td>
<td>N/A</td>
<td>c1 &lt; c2</td>
</tr>
<tr>
<td>Greater than</td>
<td>Returns a Boolean value of whether c1 is greater than c2</td>
<td>Boolean, Numeric, Text</td>
<td>N/A</td>
<td>c1 &gt; c2</td>
</tr>
<tr>
<td>Operator</td>
<td>Description</td>
<td>Supported data types</td>
<td>Text format</td>
<td>In-line format</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>---------------------------</td>
<td>-------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Less than or equal</td>
<td>Returns a Boolean value of whether c1 is less than or equal to c2</td>
<td>Boolean, Numeric, Text</td>
<td>N/A</td>
<td>c1 &lt;= c2</td>
</tr>
<tr>
<td>Greater than or equal</td>
<td>Returns a Boolean value of whether c1 is greater than or equal to c2</td>
<td>Boolean, Numeric, Text</td>
<td>N/A</td>
<td>c1 &gt;= c2</td>
</tr>
</tbody>
</table>

SageMaker Canvas also supports aggregate operators, which can perform operations such as calculating the sum of all the values or finding the minimum value in a column. You can use aggregate operators in combination with standard operators in your functions. For example, to calculate the difference of values from the mean, you could use the function \( \text{Abs}(\text{height} - \text{avg}(\text{height})) \). SageMaker Canvas supports the following aggregate operators.

<table>
<thead>
<tr>
<th>Aggregate operator</th>
<th>Description</th>
<th>Format</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>sum</td>
<td>Returns the sum of all the values in a column</td>
<td>sum</td>
<td>sum(c1)</td>
</tr>
<tr>
<td>minimum</td>
<td>Returns the minimum value of a column</td>
<td>min</td>
<td>min(c2)</td>
</tr>
<tr>
<td>maximum</td>
<td>Returns the maximum value of a column</td>
<td>max</td>
<td>max(c3)</td>
</tr>
<tr>
<td>average</td>
<td>Returns the average value of a column</td>
<td>avg</td>
<td>avg(c4)</td>
</tr>
<tr>
<td>std</td>
<td>Returns the sample standard deviation of a column</td>
<td>std</td>
<td>std(c1)</td>
</tr>
<tr>
<td>stddev</td>
<td>Returns the standard deviation of the values in a column</td>
<td>stddev</td>
<td>stddev(c1)</td>
</tr>
<tr>
<td><strong>Aggregate operator</strong></td>
<td><strong>Description</strong></td>
<td><strong>Format</strong></td>
<td><strong>Example</strong></td>
</tr>
<tr>
<td>------------------------</td>
<td>----------------</td>
<td>------------</td>
<td>-------------</td>
</tr>
<tr>
<td>variance</td>
<td>Returns the unbiased variance of the values in a column</td>
<td>variance</td>
<td>variance(c1)</td>
</tr>
<tr>
<td>approx_count_distinct</td>
<td>Returns the approximate number of distinct items in a column</td>
<td>approx_count_distinct</td>
<td>approx_count_distinct(c1)</td>
</tr>
<tr>
<td>count</td>
<td>Returns the number of items in a column</td>
<td>count</td>
<td>count(c1)</td>
</tr>
<tr>
<td>first</td>
<td>Returns the first value of a column</td>
<td>first</td>
<td>first(c1)</td>
</tr>
<tr>
<td>last</td>
<td>Returns the last value of a column</td>
<td>last</td>
<td>last(c1)</td>
</tr>
<tr>
<td>stddev_pop</td>
<td>Returns the population standard deviation of a column</td>
<td>stddev_pop</td>
<td>stddev_pop(c1)</td>
</tr>
<tr>
<td>variance_pop</td>
<td>Returns the population variance of the values in a column</td>
<td>variance_pop</td>
<td>variance_pop(c1)</td>
</tr>
</tbody>
</table>

**Manage rows**

With the Manage rows transform, you can perform sort, random shuffle, and remove rows of data from the dataset.

**Sort rows**

To sort the rows in a dataset by a given column, do the following.

1. In the **Build** tab of the SageMaker Canvas application, choose **Manage rows** and then choose **Sort rows**.
2. For **Sort Column**, choose the column you want to sort by.
3. For **Sort Order**, choose either **Ascending** or **Descending**.
4. Choose **Add** to add the transform to the **Model recipe**.
Shuffle rows

To randomly shuffle the rows in a dataset, do the following.

1. In the Build tab of the SageMaker Canvas application, choose Manage rows and then choose Shuffle rows.
2. Choose Add to add the transform to the Model recipe.

Drop duplicate rows

To remove duplicate rows in a dataset, do the following.

1. In the Build tab of the SageMaker Canvas application, choose Manage rows and then choose Drop duplicate rows.
2. Choose Add to add the transform to the Model recipe.

Remove rows by missing values

Missing values are a common occurrence in machine learning datasets and can impact model accuracy. Use this transform if you want to drop rows with null or empty values in certain columns.

To remove rows that contain missing values in a specified column, do the following.

1. In the Build tab of the SageMaker Canvas application, choose Manage rows.
2. Choose Drop rows by missing values.
3. Choose Add to add the transform to the Model recipe.

SageMaker Canvas drops rows that contain missing values in the Column you selected. After removing the rows from the dataset, SageMaker Canvas adds the transform in the Model recipe section. If you remove the transform from the Model recipe section, the rows return to your dataset.
Remove rows by outliers

Outliers, or rare values in the distribution and range of your data, can negatively impact model accuracy and lead to longer building times. With SageMaker Canvas, you can detect and remove rows that contain outliers in numeric columns. You can choose to define outliers with either standard deviations or a custom range.

To remove outliers from your data, do the following.

1. In the **Build** tab of the SageMaker Canvas application, choose **Manage rows**.
2. Choose **Drop rows by outlier values**.
3. Choose the **Column** you want to check for outliers.
4. Set the **Operator** to **Standard deviation**, **Custom numeric range**, or **Custom quantile range**.
5. If you choose **Standard deviation**, specify a **Standard deviations** (standard deviation) value from 1–3. If you choose **Custom numeric range** or **Custom quantile range**, specify the **Min** and **Max** values (numbers for numeric ranges, or percentiles between 0–100% for quantile ranges).
6. Choose **Add** to add the transform to the **Model recipe**.

The **Standard deviation** option detects and removes outliers in numeric columns using the mean and standard deviation. You specify the number of standard deviations a value must vary from the
mean to be considered an outlier. For example, if you specify 3 for **Standard deviations**, a value must fall more than 3 standard deviations from the mean to be considered an outlier.

The **Custom numeric range** and **Custom quantile range** options detect and remove outliers in numeric columns using minimum and maximum values. Use this method if you know your threshold values that delimit outliers. If you choose a numeric range, the **Min** and **Max** values should be the minimum and maximum numeric values that you want to allow in the data. If you choose a quantile range, the **Min** and **Max** values should be the minimum and maximum of the percentile range (0–100) that you want to allow.

After removing the rows from the dataset, SageMaker Canvas adds the transform in the **Model recipe** section. If you remove the transform from the **Model recipe** section, the rows return to your dataset.

### Remove rows by custom values

You can remove rows with values that meet custom conditions. For example, you might want to exclude all of the rows with a price value greater than 100 when building your model. With this transform, you can create a rule that removes all rows that exceed the threshold you set.

To use the custom remove transform, do the following.

1. In the **Build** tab of the SageMaker Canvas application, choose **Manage rows**.
2. Choose **Drop rows by formula**.
3. Choose the **Column** you want to check.

4. Select the type of **Operation** you want to use, and then specify the values for the selected condition.

5. Choose **Add** to add the transform to the **Model recipe**.

For the **Operation**, you can choose one of the following options. Note that the available operations depend on the data type of the column you choose. For example, you cannot create a `is greater than` operation for a column containing text values.

<table>
<thead>
<tr>
<th>Operation</th>
<th>Supported data type</th>
<th>Supported feature type</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is equal to</td>
<td>Numeric, Text</td>
<td>Binary, Categorial</td>
<td>Removes rows where the value in <strong>Column</strong> equals the values you specify.</td>
</tr>
<tr>
<td>Is not equal to</td>
<td>Numeric, Text</td>
<td>Binary, Categorial</td>
<td>Removes rows where the value in <strong>Column</strong> doesn't equal the values you specify.</td>
</tr>
<tr>
<td>Is less than</td>
<td>Numeric</td>
<td>N/A</td>
<td>Removes rows where the value in <strong>Column</strong> is less than the value you specify.</td>
</tr>
<tr>
<td>Is less than or equal to</td>
<td>Numeric</td>
<td>N/A</td>
<td>Removes rows where the value in <strong>Column</strong> is less than or equal to the value you specify.</td>
</tr>
<tr>
<td>Is greater than</td>
<td>Numeric</td>
<td>N/A</td>
<td>Removes rows where the value in <strong>Column</strong> is greater than the value you specify.</td>
</tr>
<tr>
<td>Is greater than or equal to</td>
<td>Numeric</td>
<td>N/A</td>
<td>Removes rows where the value in <strong>Column</strong> is greater than or equal to the value you specify.</td>
</tr>
<tr>
<td>Operation</td>
<td>Supported data type</td>
<td>Supported feature type</td>
<td>Function</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------------</td>
<td>------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>Is between</td>
<td>Numeric</td>
<td>N/A</td>
<td>Removes rows where the value in <strong>Column</strong> is between or equal to two values you specify.</td>
</tr>
<tr>
<td>Contains</td>
<td>Text</td>
<td>Categorical</td>
<td>Removes rows where the value in <strong>Column</strong> contains a values you specify.</td>
</tr>
<tr>
<td>Starts with</td>
<td>Text</td>
<td>Categorical</td>
<td>Removes rows where the value in <strong>Column</strong> begins with a value you specify.</td>
</tr>
<tr>
<td>Ends with</td>
<td>Text</td>
<td>Categorical</td>
<td>Removes rows where the value in <strong>Column</strong> ends with a value you specify.</td>
</tr>
</tbody>
</table>

After removing the rows from the dataset, SageMaker Canvas adds the transform in the **Model recipe** section. If you remove the transform from the **Model recipe** section, the rows return to your dataset.
Rename columns

With the rename columns transform, you can rename columns in your data. When you rename a column, SageMaker Canvas changes the column name in the model input.

You can rename a column in your dataset by double-clicking on the column name in the **Build** tab of the SageMaker Canvas application and entering a new name. Pressing the **Enter** key submits the change, and clicking anywhere outside the input cancels the change. You can also rename a column by clicking the **More options** icon (⋯), located at the end of the row in list view or at the end of the header cell in grid view, and choosing **Rename**.

Your column name can't be longer than 32 characters or have double underscores (__), and you can't rename a column to the same name as another column. You also can't rename a dropped column.

The following screenshot shows how to rename a column by double-clicking the column name.

When you rename a column, SageMaker Canvas adds the transform in the **Model recipe** section. If you remove the transform from the **Model recipe** section, the column reverts to its original name.
Manage columns

With the following transforms, you can change the data type of columns and replace missing values or outliers for specific columns. SageMaker Canvas uses the updated data types or values when building your model but doesn’t change your original dataset. Note that if you’ve dropped a column from your dataset using the `Drop columns` transform, you can’t replace values in that column.

Replace missing values

Missing values are a common occurrence in machine learning datasets and can impact model accuracy. You can choose to drop rows that have missing values, but your model is more accurate if you choose to replace the missing values instead. With this transform, you can replace missing values in numeric columns with the mean or median of the data in a column, or you can also specify a custom value with which to replace missing values. For non-numeric columns, you can replace missing values with the mode (most common value) of the column or a custom value.

Use this transform if you want to replace the null or empty values in certain columns. To replace missing values in a specified column, do the following.

1. In the **Build** tab of the SageMaker Canvas application, choose **Manage columns**.
2. Choose **Replace missing values**.
3. Choose the **Column** in which you want to replace missing values.
4. Set **Mode** to **Manual** to replace missing values with values that you specify. With the **Automatic (default)** setting, SageMaker Canvas replaces missing values with imputed values that best fit your data. This imputation method is done automatically for each model build, unless you specify the **Manual** mode.
5. Set the **Replace with** value:
   - If your column is numeric, then select **Mean**, **Median**, or **Custom**. **Mean** replaces missing values with the mean for the column, and **Median** replaces missing values with the median for the column. If you choose **Custom**, then you must specify a custom value that you want to use to replace missing values.
   - If your column is non-numeric, then select **Mode** or **Custom**. **Mode** replaces missing values with the mode, or the most common value, for the column. For **Custom**, specify a custom value that you want to use to replace missing values.
6. Choose **Add** to add the transform to the **Model recipe**.
After replacing the missing values in the dataset, SageMaker Canvas adds the transform in the **Model recipe** section. If you remove the transform from the **Model recipe** section, the missing values return to the dataset.

Replace outliers

Outliers, or rare values in the distribution and range of your data, can negatively impact model accuracy and lead to longer building times. SageMaker Canvas enables you to detect outliers in numeric columns and replace the outliers with values that lie within an accepted range in your data. You can choose to define outliers with either standard deviations or a custom range, and you can replace outliers with the minimum and maximum values in the accepted range.

To replace outliers in your data, do the following.

1. In the **Build** tab of the SageMaker Canvas application, choose **Manage columns**.
2. Choose **Replace outlier values**.
3. Choose the **Column** in which you want to replace outliers.
4. For **Define outliers**, choose **Standard deviation**, **Custom numeric range**, or **Custom quantile range**.
5. If you choose **Standard deviation**, specify a **Standard deviations** (standard deviation) value from 1–3. If you choose **Custom numeric range** or **Custom quantile range**, specify the **Min** and **Max** values (numbers for numeric ranges, or percentiles between 0–100% for quantile ranges).
6. For **Replace with**, select **Min/max range**.

7. Choose **Add** to add the transform to the **Model recipe**.

The **Standard deviation** option detects outliers in numeric columns using the mean and standard deviation. You specify the number of standard deviations a value must vary from the mean to be considered an outlier. For example, if you specify 3 for **Standard deviations**, a value must fall more than 3 standard deviations from the mean to be considered an outlier. SageMaker Canvas replaces outliers with the minimum value or maximum value in the accepted range. For example, if you configure the standard deviations to only include values from 200–300, then SageMaker Canvas changes a value of 198 to 200 (the minimum).

The **Custom numeric range** and **Custom quantile range** options detect outliers in numeric columns using minimum and maximum values. Use this method if you know your threshold values that delimit outliers. If you choose a numeric range, the **Min** and **Max** values should be the minimum and maximum numeric values that you want to allow. SageMaker Canvas replaces any values that fall outside of the minimum and maximum to the minimum and maximum values. For example, if your range only allows values from 1–100, then SageMaker Canvas changes a value of 102 to 100 (the maximum). If you choose a quantile range, the **Min** and **Max** values should be the minimum and maximum of the percentile range (0–100) that you want to allow.

After replacing the values in the dataset, SageMaker Canvas adds the transform in the **Model recipe** section. If you remove the transform from the **Model recipe** section, the original values return to the dataset.
Change data type

SageMaker Canvas provides you with the ability to change the data type of your columns between numeric, text, and datetime, while also displaying the associated feature type for that data type. A data type refers to the format of the data and how it is stored, while the feature type refers to the characteristic of the data used in machine learning algorithms, such as binary or categorical. This gives you the flexibility to manually change the type of data in your columns based on the features. The ability to choose the right data type ensures data integrity and accuracy prior to building models. These data types are used when building models.

Note

Currently, changing the feature type (for example, from binary to categorical) is not supported.

The following table lists all of the supported data types in Canvas.
<table>
<thead>
<tr>
<th>Data type</th>
<th>Description</th>
<th>Example</th>
</tr>
</thead>
</table>
| Numeric   | Numeric data represents numerical values | 1, 2, 3  
1.1, 1.2, 1.3 |
| Text      | Text data represents sequences of characters, like names or descriptions | A, B, C, D  
apple, banana, orange  
1A!, 2A!, 3A! |
| Datetime  | Datetime data represents dates and times in timestamp format | 2019-07-01 01:00:00,  
2019-07-01 02:00:00,  
2019-07-01 03:00:00 |

The following table lists all of the supported feature types in Canvas.

<table>
<thead>
<tr>
<th>Feature type</th>
<th>Description</th>
<th>Example</th>
</tr>
</thead>
</table>
| Binary       | Binary features represent two possible values | 0, 1, 0, 1, 0 (2 distinct values)  
true, false, true (2 distinct values) |
| Categorical  | Categorical features represent distinct categories or groups | apple, banana, orange, apple  
(3 distinct values)  
A, B, C, D, E, A, D, C (5 distinct values) |

To modify data type of a column in a dataset, do the following.

1. In the **Build** tab of the SageMaker Canvas application, go to the **Column view** or **Grid view** and select the **Data type** dropdown for the specific column.
2. In the **Data type** dropdown, choose the data type to convert to. The following screenshot shows the dropdown menu.
3. For **Column**, choose or verify the column you want to change the data type for.

4. For **New data type**, choose or verify the new data type you want to convert to.

5. If the **New data type** is Datetime or Numeric, choose one of the following options under **Handle invalid values**:
   
   a. **Replace with empty value** – Invalid values are substituted with an empty value
   
   b. **Delete rows** – Rows with an invalid value are removed from the dataset
   
   c. **Replace with custom value** – Invalid values are substituted with the **Custom Value** that you specify.

6. Choose **Add** to add the transform to the **Model recipe**.

The data type for your column should now be updated.

### Prepare time series data

Use the following functionalities to prepare your time series data for building time series forecasting models.

### Resample time series data

By resampling time-series data, you can establish regular intervals for the observations in your time series dataset. This is particularly useful when working with time series data containing irregularly spaced observations. For instance, you can use resampling to transform a dataset with observations recorded every one hour, two hour and three hour intervals into a regular one hour
interval between observations. Forecasting algorithms require the observations to be taken at regular intervals.

To resample time series data, do the following.

1. In the Build tab of the SageMaker Canvas application, choose Time series.
2. Choose Resample.
3. For Timestamp column, choose the column you want to apply the transform to. You can only select columns of the Datetime type.
4. In the Frequency settings section, choose a Frequency and Rate. Frequency is the unit of frequency and Rate is the interval of the unit of frequency to be applied to the column. For example, choosing Calendar Day for Frequency value and 1 for Rate sets the interval to increase every 1 calendar day, such as 2023-03-26 00:00:00, 2023-03-27 00:00:00, 2023-03-28 00:00:00. See the table after this procedure for a complete list of Frequency value.
5. Choose Add to add the transform to the Model recipe.

The following table lists all of the Frequency types you can select when resampling time series data.

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Description</th>
<th>Example values (assuming Rate is 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Business Day</td>
<td>Resample observations in the datetime column to 5 business days of the week (Monday, Tuesday, Wednesday, Thursday, Friday)</td>
<td>2023-03-24 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-27 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-28 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-29 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-30 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-31 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-04-03 00:00:00</td>
</tr>
<tr>
<td>Calendar Day</td>
<td>Resample observations in the datetime column</td>
<td>2023-03-26 00:00:00</td>
</tr>
<tr>
<td>Frequency</td>
<td>Description</td>
<td>Example values (assuming Rate is 1)</td>
</tr>
<tr>
<td>--------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>--------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>to all 7 days of the week (Monday, Tuesday, Wednesday, Thursday, Friday, Saturday, Sunday)</td>
<td>2023-03-27 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-28 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-29 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-30 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-31 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-04-01 00:00:00</td>
</tr>
<tr>
<td>Week</td>
<td>Resample observations in the datetime column to the first day of each week</td>
<td>2023-03-13 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-20 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-03-27 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-04-03 00:00:00</td>
</tr>
<tr>
<td>Month</td>
<td>Resample observations in the datetime column to the first day of each month</td>
<td>2023-03-01 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-04-01 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-05-01 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-06-01 00:00:00</td>
</tr>
<tr>
<td>Annual Quarter</td>
<td>Resample observations in the datetime column to the last day of each quarter</td>
<td>2023-03-31 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-06-30 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-09-30 00:00:00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2023-12-31 00:00:00</td>
</tr>
<tr>
<td>Frequency</td>
<td>Description</td>
<td>Example values (assuming Rate is 1)</td>
</tr>
<tr>
<td>-----------</td>
<td>-------------</td>
<td>-------------------------------------</td>
</tr>
</tbody>
</table>
| Year      | Resample observations in the datetime column to the last day of each year | 2022-12-31 00:00:00  
2023-12-31 00:00:00  
2024-12-31 00:00:00 |
| Hour      | Resample observations in the datetime column to each hour of each day | 2023-03-24 00:00:00  
2023-03-24 01:00:00  
2023-03-24 02:00:00  
2023-03-24 03:00:00 |
| Minute    | Resample observations in the datetime column to each minute of each hour | 2023-03-24 00:00:00  
2023-03-24 00:01:00  
2023-03-24 00:02:00  
2023-03-24 00:03:00 |
| Second    | Resample observations in the datetime column to each second of each minute | 2023-03-24 00:00:00  
2023-03-24 00:00:01  
2023-03-24 00:00:02  
2023-03-24 00:00:03 |

When applying the resampling transform, you can use the Advanced option to specify how the resulting values of the rest of the columns (other than the timestamp column) in your dataset are modified. This can be achieved by specifying the resampling methodology, which can either be downsampling or upsampling for both numeric and non-numeric columns.

**Downsampling** increases the interval between observations in the dataset. For example, if you downsample observations that are taken either every hour or every two hours, each observation in your dataset is taken every two hours. The values of other columns of the hourly observations
are aggregated into a single value using a combination method. The following tables show an example of downsampling time series data by using mean as the combination method. The data is downsampled from every two hours to every hour.

The following table shows the hourly temperature readings over a day before downsampling.

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Temperature (Celsius)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:00 pm</td>
<td>30</td>
</tr>
<tr>
<td>1:00 am</td>
<td>32</td>
</tr>
<tr>
<td>2:00 am</td>
<td>35</td>
</tr>
<tr>
<td>3:00 am</td>
<td>32</td>
</tr>
<tr>
<td>4:00 am</td>
<td>30</td>
</tr>
</tbody>
</table>

The following table shows the temperature readings after downsampling to every two hours.

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Temperature (Celsius)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:00 pm</td>
<td>30</td>
</tr>
<tr>
<td>2:00 am</td>
<td>33.5</td>
</tr>
<tr>
<td>2:00 am</td>
<td>35</td>
</tr>
<tr>
<td>4:00 am</td>
<td>32.5</td>
</tr>
</tbody>
</table>

To downsample time series data, do the following:

1. Expand the Advanced section under the Resample transform.
2. Choose Non-numeric combination to specify the combination method for non-numeric columns. See the table below for a complete list of combination methods.
3. Choose Numeric combination to specify the combination method for numeric columns. See the table below for a complete list of combination methods.
If you don’t specify combination methods, the default values are Most Common for **Non-numeric combination** and Mean for **Numeric combination**. The following table lists the methods for numeric and non-numeric combination.

<table>
<thead>
<tr>
<th>Downsampling methodology</th>
<th>Combination method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-numeric combination</td>
<td>Most Common</td>
<td>Aggregate values in the non-numeric column by the most commonly occurring value</td>
</tr>
<tr>
<td>Non-numeric combination</td>
<td>Last</td>
<td>Aggregate values in the non-numeric column by the last value in the column</td>
</tr>
<tr>
<td>Non-numeric combination</td>
<td>First</td>
<td>Aggregate values in the non-numeric column by the first value in the column</td>
</tr>
<tr>
<td>Numeric combination</td>
<td>Mean</td>
<td>Aggregate values in the numeric column by the taking the mean of all the values in the column</td>
</tr>
<tr>
<td>Numeric combination</td>
<td>Median</td>
<td>Aggregate values in the numeric column by the taking the median of all the values in the column</td>
</tr>
<tr>
<td>Numeric combination</td>
<td>Min</td>
<td>Aggregate values in the numeric column by the taking the minimum of all the values in the column</td>
</tr>
<tr>
<td>Numeric combination</td>
<td>Max</td>
<td>Aggregate values in the numeric column by the taking the maximum of all the values in the column</td>
</tr>
<tr>
<td>Downsampling methodology</td>
<td>Combination method</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Numeric combination</td>
<td>Sum</td>
<td>Aggregate values in the numeric column by adding all the values in the column</td>
</tr>
<tr>
<td>Numeric combination</td>
<td>Quantile</td>
<td>Aggregate values in the numeric column by the taking the quantile of all the values in the column</td>
</tr>
</tbody>
</table>

*Upsampling* reduces the interval between observations in the dataset. For example, if you upsample observations that are taken every two hours into hourly observations, the values of other columns of the hourly observations are interpolated from the ones that have been taken every two hours.

To upsample time series data, do the following:

1. Expand the **Advanced** section under the **Resample** transform.
2. Choose **Non-numeric estimation** to specify the estimation method for non-numeric columns. See the table after this procedure for a complete list of methods.
3. Choose **Numeric estimation** to specify the estimation method for numeric columns. See the table below for a complete list of methods.
4. (Optional) Choose **ID Column** to specify the column that has the IDs of the observations of the time series. Specify this option if your dataset has two time series. If you have a column representing only one time series, don't specify a value for this field. For example, you can have a dataset that has the columns id and purchase. The id column has the following values: [1, 2, 2, 1]. The purchase column has the following values [$2, $3, $4, $1]. Therefore, the dataset has two time series—one time series is: 1: [$2, $1], and the other time series is 2: [$3, $4].

If you don't specify estimation methods, the default values are **Forward Fill** for **Non-numeric estimation** and **Linear** for **Numeric estimation**. The following table lists the methods for estimation.
<table>
<thead>
<tr>
<th>Upsampling methodology</th>
<th>Estimation method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-numeric estimation</td>
<td>Forward Fill</td>
<td>Interpolate values in the non-numeric column by taking the consecutive values after all the values in the column</td>
</tr>
<tr>
<td>Non-numeric estimation</td>
<td>Backward Fill</td>
<td>Interpolate values in the non-numeric column by taking the consecutive values before all the values in the column</td>
</tr>
<tr>
<td>Non-numeric estimation</td>
<td>Keep Missing</td>
<td>Interpolate values in the non-numeric column by showing empty values</td>
</tr>
<tr>
<td>Numeric estimation</td>
<td>Linear, Time, Index, Zero, S-Linear, Nearest, Quadratic, Cubic, Barycentric, Polynomial, Krogh, Piecewise Polynomial, Spline, P-chip, Akima, Cubic Spline, From Derivatives</td>
<td>Interpolate values in the numeric column by using the specified interpolator. For information on interpolation methods, see <code>pandas.DataFrame.interpolate</code> in the pandas documentation.</td>
</tr>
</tbody>
</table>

The following screenshot shows the **Advanced** settings with the fields for downsampling and upsampling filled out.
Use datetime extraction

With the datetime extraction transform, you can extract values from a datetime column to a separate column. For example, if you have a column containing dates of purchases, you can extract the month value to a separate column and use the new column when building your model. You can also extract multiple values to separate columns with a single transform.

Your datetime column must use a supported timestamp format. For a list of the formats that SageMaker Canvas supports, see Time Series Forecasts in Amazon SageMaker Canvas. If your dataset does not use one of the supported formats, update your dataset to use a supported timestamp format and re-import it to Amazon SageMaker Canvas before building your model.

To perform a datetime extraction, do the following.

1. In the **Build** tab of the SageMaker Canvas application, on the transforms bar, choose **View all**.
2. Choose **Extract features**.
3. Choose the **Timestamp column** from which you want to extract values.
4. For **Values**, select one or more values to extract from the column. The values you can extract from a timestamp column are **Year, Month, Day, Hour, Week of year, Day of year, and Quarter**.

5. (Optional) Choose **Preview** to preview the transform results.

6. Choose **Add** to add the transform to the **Model recipe**.

SageMaker Canvas creates a new column in the dataset for each of the values you extract. Except for **Year** values, SageMaker Canvas uses a 0-based encoding for the extracted values. For example, if you extract the **Month** value, January is extracted as 0, and February is extracted as 1.

You can see the transform listed in the **Model recipe** section. If you remove the transform from the **Model recipe** section, the new columns are removed from the dataset.

**Evaluate Your Model’s Performance in Amazon SageMaker Canvas**

After you’ve built your model, you can evaluate how well your model performed on your data before using it to make predictions. You can use information, such as the model’s accuracy when predicting labels and advanced metrics, to determine whether your model can make sufficiently accurate predictions for your data.

On the **Analyze** page for your model, Amazon SageMaker Canvas provides the following three tabs:
- **Overview** – Gives you a general overview of the model's performance, depending on the model type.
- **Scoring** – Shows visualizations that you can use to get more insights into your model's performance beyond the overall accuracy metrics.
- **Advanced metrics** – Contains your model's scores for advanced metrics and additional information that can give you a deeper understanding of your model's performance. You can also view information such as the column impacts.

The section [Evaluate your model's performance](#) describes how to view and interpret your model's Overview and Scoring tabs. The section [Use advanced metrics in your analyses](#) contains more detailed information about the Advanced metrics used to quantify your model's accuracy.

You can also view more advanced information for specific model candidates, which are all of the model iterations that Canvas runs through while building your model. Based on the advanced metrics for a given model candidate, you can select a different candidate to be the default, or the version that is used for making predictions and deploying. For each model candidate, you can view the Advanced metrics information to help you decide which model candidate you’d like to select as the default. You can view this information by selecting the model candidate from the **Model leaderboard**. For more information, see [View model candidates in the model leaderboard](#).

Canvas also provides the option to download a Jupyter notebook so that you can view and run the code used to build your model. This is useful if you’d like to make adjustments to the code or learn more about how your model was built. For more information, see [Download a model notebook](#).

### Evaluate your model's performance

Amazon SageMaker Canvas provides overview and scoring information for the different types of model. Your model's score can help you determine how accurate your model is when it makes predictions. The additional scoring insights can help you quantify the differences between the actual and predicted values.

To view the analysis of your model, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **My models**.
3. Choose the model that you built.
4. In the top navigation pane, choose the **Analyze** tab.
5. Within the **Analyze** tab, you can view the overview and scoring information for your model.
The following sections describe how to interpret the scoring for each model type.

**Evaluate categorical prediction models**

The **Overview** tab shows you the column impact for each column. **Column impact** is a percentage score that indicates how much weight a column has in making predictions in relation to the other columns. For a column impact of 25%, Canvas weighs the prediction as 25% for the column and 75% for the other columns.

The following screenshot shows the **Accuracy** score for the model, along with the **Optimization metric**, which is the metric that you choose to optimize when building the model. In this case, the **Optimization metric** is **Accuracy**. You can specify a different optimization metric if you build a new version of your model.

The **Scoring** tab for a categorical prediction model gives you the ability to visualize all the predictions. Line segments extend from the left of the page, indicating all the predictions the model has made. In the middle of the page, the line segments converge on a perpendicular segment to indicate the proportion of each prediction to a single category. From the predicted category, the segments branch out to the actual category. You can get a visual sense of how accurate the predictions were by following each line segment from the predicted category to the actual category.

The following image gives you an example **Scoring** section for a **3+ category prediction** model.
You can also view the **Advanced metrics** tab for more detailed information about your model's performance, such as the advanced metrics, error density plots, or confusion matrices. To learn more about the **Advanced metrics** tab, see [Use advanced metrics in your analyses](#).

### Evaluate numeric prediction models

The **Overview** tab shows you the column impact for each column. **Column impact** is a percentage score that indicates how much weight a column has in making predictions in relation to the other columns. For a column impact of 25%, Canvas weighs the prediction as 25% for the column and 75% for the other columns.

The following screenshot shows the **RMSE** score for the model on the **Overview** tab, which in this case is the **Optimization metric**. The **Optimization metric** is the metric that you choose to optimize when building the model. You can specify a different optimization metric if you build a new version of your model.
The **Scoring** tab for numeric prediction shows a line to indicate the model's predicted value in relation to the data used to make predictions. The values of the numeric prediction are often +/- the RMSE (root mean squared error) value. The value that the model predicts is often within the range of the RMSE. The width of the purple band around the line indicates the RMSE range. The predicted values often fall within the range.

The following image shows the **Scoring** section for numeric prediction.

You can also view the **Advanced metrics** tab for more detailed information about your model's performance, such as the advanced metrics, error density plots, or confusion matrices. To learn more about the **Advanced metrics** tab, see [Use advanced metrics in your analyses](#).
Evaluate time series forecasting models

On the Analyze page for time series forecasting models, you can see an overview of the model's metrics. You can hover over each metric for more information, or you can see Use advanced metrics in your analyses.

In the Column impact section, you can see the score for each column. Column impact is a percentage score that indicates how much weight a column has in making predictions in relation to the other columns. For a column impact of 25%, Canvas weighs the prediction as 25% for the column and 75% for the other columns.

The following screenshot shows the time series metrics scores for the model, along with the Optimization metric, which is the metric that you choose to optimize when building the model. In this case, the Optimization metric is RMSE. You can specify a different optimization metric if you build a new version of your model.

![Time series metrics screenshot](image)

Evaluate image prediction models

The Overview tab shows you the Per label performance, which gives you an overall accuracy score for the images predicted for each label. You can choose a label to see more specific details, such as the Correctly predicted and Incorrectly predicted images for the label.

You can turn on the Heatmap toggle to see a heatmap for each image. The heatmap shows you the areas of interest that have the most impact when your model is making predictions. For more information about heatmaps and how to use them to improve your model, choose the More info icon next to the Heatmap toggle.

The Scoring tab for single-label image prediction models shows you a comparison of what the model predicted as the label versus what the actual label was. You can select up to 10 labels at a time. You can change the labels in the visualization by choosing the labels dropdown menu and selecting or deselected labels.
You can also view insights for individual labels or groups of labels, such as the three labels with the highest or lowest accuracy, by choosing the **View scores for** dropdown menu in the **Model accuracy insights** section.

The following screenshot shows the **Scoring** information for a single-label image prediction model.

The model predicts the correct labels 96% of the time.

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**Evaluate text prediction models**

The **Overview** tab shows you the **Per label performance**, which gives you an overall accuracy score for the passages of text predicted for each label. You can choose a label to see more specific details, such as the **Correctly predicted** and **Incorrectly predicted** passages for the label.

The **Scoring** tab for multi-category text prediction models shows you a comparison of what the model predicted as the label versus what the actual label was.

In the **Model accuracy insights** section, you can see the **Most frequent category**, which tells you the category that the model predicted most frequently and how accurate those predictions were. If you model predicts a label of **Positive** correctly 99% of the time, then you can be fairly confident that your model is good at predicting positive sentiment in text.

The following screenshot shows the **Scoring** information for a multi-category text prediction model.
Use advanced metrics in your analyses

The following section describes how to find and interpret the advanced metrics for your model in Amazon SageMaker Canvas.

Note

Advanced metrics are only currently available for numeric and categorical prediction models.

To find the Advanced metrics tab, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose My models.
3. Choose the model that you built.
4. In the top navigation pane, choose the Analyze tab.
5. Within the Analyze tab, choose the Advanced metrics tab.
In the **Advanced metrics** tab, you can find the **Performance** tab. The page looks like the following screenshot.

At the top, you can see an overview of the metrics scores, including the **Optimization metric**, which is the metric that you selected (or that Canvas selected by default) to optimize when building the model.

The following sections describe more detailed information for the **Performance** tab within the **Advanced metrics**.

**Performance**

In the **Performance** tab, you'll see a **Metrics table**, along with visualizations that Canvas creates based on your model type. For categorical prediction models, Canvas provides a **confusion matrix**, whereas for numeric prediction models, Canvas provides you with **residuals** and **error density** charts.

In the **Metrics table**, you are provided with a full list of your model's scores for each advanced metric, which is more comprehensive than the scores overview at the top of the page. The metrics shown here depend on your model type. For a reference to help you understand and interpret each metric, see **Metrics reference**.

To understand the visualizations that might appear based on your model type, see the following options:
- **Confusion matrix** – Canvas uses confusion matrices to help you visualize when a model makes predictions correctly. In a confusion matrix, your results are arranged to compare the predicted values against the actual values. The following example explains how a confusion matrix works for a 2 category prediction model that predicts positive and negative labels:
  - True positive – The model correctly predicted positive when the true label was positive.
  - True negative – The model correctly predicted negative when the true label was negative.
  - False positive – The model incorrectly predicted positive when the true label was negative.
  - False negative – The model incorrectly predicted negative when the true label was positive.

- **Precision recall curve** – The precision recall curve is a visualization of the model's precision score plotted against the model's recall score. Generally, a model that can make perfect predictions would have precision and recall scores that are both 1. The precision recall curve for a decently accurate model is fairly high in both precision and recall.

- **Residuals** – Residuals are the difference between the actual values and the values predicted by the model. A residuals chart plots the residuals against the corresponding values to visualize their distribution and any patterns or outliers. A normal distribution of residuals around zero indicates that the model is a good fit for the data. However, if the residuals are significantly skewed or have outliers, it may indicate that the model is overfitting the data or that there are other issues that need to be addressed.

- **Error density** – An error density plot is a representation of the distribution of errors made by a model. It shows the probability density of the errors at each point, helping you to identify any areas where the model may be overfitting or making systematic errors.

**View model candidates in the model leaderboard**

When you build a model in Amazon SageMaker Canvas, SageMaker trains multiple model candidates, or different iterations of the model, and selects the one with the highest value for the optimization metric by default. The default model candidate is the only version that you can use with the other functionality in Canvas like making predictions, registering to the model registry or deploying to an endpoint.

However, you might want to review all of the model candidates and select a different candidate to be the default model. You can view all of the model candidates and more details about each candidate on the **Model leaderboard** in Canvas.

To view the **Model leaderboard**, do the following:
1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose **My models**.
3. Choose the model that you built.
4. In the top navigation pane, choose the **Analyze** tab.
5. Within the **Analyze** tab, choose **Model leaderboard**.

The **Model leaderboard** page opens, which looks like the following screenshot.

You can see that the first model candidate listed is marked as the **Default model**. This is the model candidate with which you can make predictions or deploy to endpoints.

To view more detailed metrics information about the model candidates to compare them, you can choose the **More options** icon ( 
 and choose **View model details**.)
Important

Loading the model details for non-default model candidates may take a few minutes (typically less than 10 minutes), and SageMaker Hosting charges apply. For more information, see SageMaker Pricing.

The model candidate opens in the Analyze tab, and the metrics shown are specific to that model candidate. When you’re done reviewing the model candidate’s metrics, you can go back or exit the view to return to the Model leaderboard.

If you’d like to set the Default model to a different candidate, you can choose the More options icon and choose Change to the default model. Changing the default model for a model trained using HPO mode might take several minutes.

Note

If your model is already deployed in production, registered to the model registry, or has automations set up, you must delete your deployment, model registration, or automations before changing the default model.

Metrics reference

The following sections describe the metrics that are available in Amazon SageMaker Canvas for each model type.

Metrics for numeric prediction

The following list defines the metrics for numeric prediction in SageMaker Canvas and gives you information about how you can use them.

- **InferenceLatency** – The approximate amount of time between making a request for a model prediction to receiving it from a real-time endpoint to which the model is deployed. This metric is measured in seconds and is only available for models built with the Ensembling mode.
- **MAE** – Mean absolute error. On average, the prediction for the target column is +/- {MAE} from the actual value.
Measures how different the predicted and actual values are when they're averaged over all values. MAE is commonly used in numeric prediction to understand model prediction error. If the predictions are linear, MAE represents the average distance from a predicted line to the actual value. MAE is defined as the sum of absolute errors divided by the number of observations. Values range from 0 to infinity, with smaller numbers indicating a better model fit to the data.

- **MAPE** – Mean absolute percent error. On average, the prediction for the target column is +/- {MAPE} % from the actual value.

MAPE is the mean of the absolute differences between the actual values and the predicted or estimated values, divided by the actual values and expressed as a percentage. A lower MAPE indicates better performance, as it means that the predicted or estimated values are closer to the actual values.

- **MSE** – Mean squared error, or the average of the squared differences between the predicted and actual values.

MSE values are always positive. The better a model is at predicting the actual values, the smaller the MSE value is.

- **R2** – The percentage of the difference in the target column that can be explained by the input column.

Quantifies how much a model can explain the variance of a dependent variable. Values range from one (1) to negative one (-1). Higher numbers indicate a higher fraction of explained variability. Values close to zero (0) indicate that very little of the dependent variable can be explained by the model. Negative values indicate a poor fit and that the model is outperformed by a constant function (or a horizontal line).

- **RMSE** – Root mean squared error, or the standard deviation of the errors.

Measures the square root of the squared difference between predicted and actual values, and is averaged over all values. It is used to understand model prediction error, and it's an important metric to indicate the presence of large model errors and outliers. Values range from zero (0) to infinity, with smaller numbers indicating a better model fit to the data. RMSE is dependent on scale, and should not be used to compare datasets of different types.
Metrics for categorical prediction

This section defines the metrics for categorical prediction in SageMaker Canvas and gives you information about how you can use them.

The following is a list of available metrics for 2-category prediction:

- **Accuracy** – The percentage of correct predictions. Or, the ratio of the number of correctly predicted items to the total number of predictions. Accuracy measures how close the predicted class values are to the actual values. Values for accuracy metrics vary between zero (0) and one (1). A value of 1 indicates perfect accuracy, and 0 indicates complete inaccuracy.

- **AUC** – A value between 0 and 1 that indicates how well your model is able to separate the categories in your dataset. A value of 1 indicates that it was able to separate the categories perfectly.

- **BalancedAccuracy** – Measures the ratio of accurate predictions to all predictions. This ratio is calculated after normalizing true positives (TP) and true negatives (TN) by the total number of positive (P) and negative (N) values. It is defined as follows: \( 0.5 \times \frac{(TP/P) + (TN/N)} \), with values ranging from 0 to 1. The balanced accuracy metric gives a better measure of accuracy when the number of positives or negatives differ greatly from each other in an imbalanced dataset, such as when only 1% of email is spam.

- **F1** – A balanced measure of accuracy that takes class balance into account. It is the harmonic mean of the precision and recall scores, defined as follows: \( F1 = \frac{2 \times \text{precision} \times \text{recall}}{\text{precision} + \text{recall}} \). F1 scores vary between 0 and 1. A score of 1 indicates the best possible performance, and 0 indicates the worst.

- **InferenceLatency** – The approximate amount of time between making a request for a model prediction to receiving it from a real-time endpoint to which the model is deployed. This metric is measured in seconds and is only available for models built with the **Ensembling** mode.

- **LogLoss** – Log loss, also known as cross-entropy loss, is a metric used to evaluate the quality of the probability outputs, rather than the outputs themselves. Log loss is an important metric to indicate when a model makes incorrect predictions with high probabilities. Values range from 0 to infinity. A value of 0 represents a model that perfectly predicts the data.

- **Precision** – Of all the times that {category x} was predicted, the prediction was correct {precision}% of the time.
Precision measures how well an algorithm predicts the true positives (TP) out of all of the positives that it identifies. It is defined as follows: \( \text{Precision} = \frac{TP}{TP+FP} \), with values ranging from zero (0) to one (1). Precision is an important metric when the cost of a false positive is high. For example, the cost of a false positive is very high if an airplane safety system is falsely deemed safe to fly. A false positive (FP) reflects a positive prediction that is actually negative in the data.

- **Recall** – The model correctly predicted \{recall\}% to be \{category x\} when \{target_column\} was actually \{category x\}.

Recall measures how well an algorithm correctly predicts all of the true positives (TP) in a dataset. A true positive is a positive prediction that is also an actual positive value in the data. Recall is defined as follows: \( \text{Recall} = \frac{TP}{TP+FN} \), with values ranging from 0 to 1. Higher scores reflect a better ability of the model to predict true positives (TP) in the data. Note that it is often insufficient to measure only recall, because predicting every output as a true positive yields a perfect recall score.

The following is a list of available metrics for 3+ category prediction:

- **Accuracy** – The percentage of correct predictions.

  Or, the ratio of the number of correctly predicted items to the total number of predictions. Accuracy measures how close the predicted class values are to the actual values. Values for accuracy metrics vary between zero (0) and one (1). A value of 1 indicates perfect accuracy, and 0 indicates complete inaccuracy.

- **BalancedAccuracy** – Measures the ratio of accurate predictions to all predictions.

  This ratio is calculated after normalizing true positives (TP) and true negatives (TN) by the total number of positive (P) and negative (N) values. It is defined as follows: \( 0.5 \times \left( \frac{TP}{P} + \frac{TN}{N} \right) \), with values ranging from 0 to 1. The balanced accuracy metric gives a better measure of accuracy when the number of positives or negatives differ greatly from each other in an imbalanced dataset, such as when only 1% of email is spam.

- **F1macro** – The F1macro score applies F1 scoring by calculating the precision and recall, and then taking their harmonic mean to calculate the F1 score for each class. Then, the F1macro averages the individual scores to obtain the F1macro score. F1macro scores vary between 0 and 1. A score of 1 indicates the best possible performance, and 0 indicates the worst.
- **InferenceLatency** – The approximate amount of time between making a request for a model prediction to receiving it from a real-time endpoint to which the model is deployed. This metric is measured in seconds and is only available for models built with the **Ensembling** mode.

- **LogLoss** – Log loss, also known as cross-entropy loss, is a metric used to evaluate the quality of the probability outputs, rather than the outputs themselves. Log loss is an important metric to indicate when a model makes incorrect predictions with high probabilities. Values range from 0 to infinity. A value of 0 represents a model that perfectly predicts the data.

- **PrecisionMacro** – Measures precision by calculating precision for each class and averaging scores to obtain precision for several classes. Scores range from zero (0) to one (1). Higher scores reflect the model's ability to predict true positives (TP) out of all of the positives that it identifies, averaged across multiple classes.

- **RecallMacro** – Measures recall by calculating recall for each class and averaging scores to obtain recall for several classes. Scores range from 0 to 1. Higher scores reflect the model's ability to predict true positives (TP) in a dataset, whereas a true positive reflects a positive prediction that is also an actual positive value in the data. It is often insufficient to measure only recall, because predicting every output as a true positive will yield a perfect recall score.

Note that for 3+ category prediction, you also receive the average F1, Accuracy, Precision, and Recall metrics. The scores for these metrics are just the metric scores averaged for all categories.

**Metrics for image and text prediction**

The following is a list of available metrics for image prediction and text prediction.

- **Accuracy** – The percentage of correct predictions.

  Or, the ratio of the number of correctly predicted items to the total number of predictions. Accuracy measures how close the predicted class values are to the actual values. Values for accuracy metrics vary between zero (0) and one (1). A value of 1 indicates perfect accuracy, and 0 indicates complete inaccuracy.

- **F1** – A balanced measure of accuracy that takes class balance into account.

  It is the harmonic mean of the precision and recall scores, defined as follows: $F_1 = \frac{2 \times \text{precision} \times \text{recall}}{\text{precision} + \text{recall}}$. F1 scores vary between 0 and 1. A score of 1 indicates the best possible performance, and 0 indicates the worst.

- **Precision** – Of all the times that {category x} was predicted, the prediction was correct {precision}% of the time.
Precision measures how well an algorithm predicts the true positives (TP) out of all of the positives that it identifies. It is defined as follows: \( \text{Precision} = \frac{TP}{TP+FP} \), with values ranging from zero (0) to one (1). Precision is an important metric when the cost of a false positive is high. For example, the cost of a false positive is very high if an airplane safety system is falsely deemed safe to fly. A false positive (FP) reflects a positive prediction that is actually negative in the data.

- Recall – The model correctly predicted \{\text{recall}\}% to be \{\text{category x}\} when \{\text{target_column}\} was actually \{\text{category x}\}.

Recall measures how well an algorithm correctly predicts all of the true positives (TP) in a dataset. A true positive is a positive prediction that is also an actual positive value in the data. Recall is defined as follows: \( \text{Recall} = \frac{TP}{TP+FN} \), with values ranging from 0 to 1. Higher scores reflect a better ability of the model to predict true positives (TP) in the data. Note that it is often insufficient to measure only recall, because predicting every output as a true positive yields a perfect recall score.

Note that for image and text prediction models where you are predicting 3 or more categories, you also receive the \textit{average} F1, Accuracy, Precision, and Recall metrics. The scores for these metrics are just the metric scores average for all categories.

**Metrics for time series forecasts**

The following defines the advanced metrics for time series forecasts in Amazon SageMaker Canvas and gives you information about how you can use them.

- Average Weighted Quantile Loss (wQL) – Evaluates the forecast by averaging the accuracy at the P10, P50, and P90 quantiles. A lower value indicates a more accurate model.
- Weighted Absolute Percent Error (WAPE) – The sum of the absolute error normalized by the sum of the absolute target, which measures the overall deviation of forecasted values from observed values. A lower value indicates a more accurate model, where WAPE = 0 is a model with no errors.
- Root Mean Square Error (RMSE) – The square root of the average squared errors. A lower RMSE indicates a more accurate model, where RMSE = 0 is a model with no errors.
- Mean Absolute Percent Error (MAPE) – The percentage error (percent difference of the mean forecasted value versus the actual value) averaged over all time points. A lower value indicates a more accurate model, where MAPE = 0 is a model with no errors.
Mean Absolute Scaled Error (MASE) – The mean absolute error of the forecast normalized by the mean absolute error of a simple baseline forecasting method. A lower value indicates a more accurate model, where MASE < 1 is estimated to be better than the baseline and MASE > 1 is estimated to be worse than the baseline.

Make predictions for your data

Use the custom model that you've built in SageMaker Canvas to make predictions for your data. The following sections show you how to make predictions for numeric and categorical prediction models, image prediction models, and text prediction models. For information about how to make predictions with a time series forecast model, see Make a time series forecast.

Numeric and categorical prediction, image prediction, and text prediction custom models support making the following types of predictions for your data:

- **Single predictions** — A Single prediction is when you only need to make one prediction. For example, you have one image or passage of text that you want to classify.
- **Batch predictions** — A Batch prediction is when you’d like to make predictions for an entire dataset. For example, you have a CSV file of customer reviews for which you'd like to predict the customer sentiment, or you have a folder of image files that you'd like to classify. You should make predictions with a dataset that matches your input dataset. Canvas provides you with the ability to do manual batch predictions, or you can configure automatic batch predictions that initiate whenever a specified dataset is updated in Canvas.

For each prediction or set of predictions, SageMaker Canvas returns the following:

- The predicted values
- The probability of the predicted value being correct

Get started

Choose one of the following workflows to make predictions with your custom model:

- Make batch predictions
- Make single predictions

After generating predictions with your model, you can also do the following:
• **Update your model by creating a new version.** If you want to try to improve the prediction accuracy of your model, you can build new versions of your model. You can update your data or change any advanced transformations you used, and then you can review and compare the versions of your model to choose the best one.

• **Register a model version in the SageMaker model registry.** You can register versions of your model to the SageMaker model registry, which is a feature for tracking and managing the status of model versions and machine learning pipelines. A data scientist or MLOps team user with access to the SageMaker model registry can review your model versions and approve or reject them before deploying them to production.

• **Send your batch predictions to Amazon QuickSight.** In Amazon QuickSight, you can build and publish dashboards with your batch prediction datasets. This can help you analyze and share results generated by your custom model.

## Make single predictions

### Note

This section describes how to get single predictions from your model inside the Canvas application. For information about making real-time invocations in a production environment by deploying your model to an endpoint, see [Deploy your models to an endpoint](#).

Make single predictions if you want to get a prediction for a single data point. You can use this feature to get real-time predictions or to experiment with changing individual values to see how they impact the prediction outcome.

Choose one of the following procedures based on your model type.

### Make single predictions with numeric and categorical prediction models

To make a single prediction for a numeric or categorical prediction model, do the following:

1. In the left navigation pane of the Canvas application, choose **My models**.
2. On the **My models** page, choose your model.
3. After opening your model, choose the **Predict** tab.
4. On the **Run predictions** page, choose **Single prediction**.
5. For each **Column** field, which represents the columns of your input data, you can change the **Value**. Select the dropdown menu for the **Value** you want to change. For numeric fields, you can enter a new number. For fields with labels, you can select a different label.

6. When you’re ready to generate the prediction, in the right **Prediction** pane, choose **Update**.

In the right **Prediction** pane, you’ll see the prediction result. You can **Copy** the prediction result chart, or you can also choose **Download** to either download the prediction result chart as an image or to download the values and prediction as a CSV file.

**Make single predictions with image prediction models**

To make a single prediction for a single-label image prediction model, do the following:

1. In the left navigation pane of the Canvas application, choose **My models**.
2. On the **My models** page, choose your model.
3. After opening your model, choose the **Predict** tab.
4. On the **Run predictions** page, choose **Single prediction**.
5. Choose **Import image**.
6. You’ll be prompted to upload an image. You can upload an image from your local computer or from an Amazon S3 bucket.
7. Choose **Import** to import your image and generate the prediction.

In the right **Prediction results** pane, the model lists the possible labels for the image along with a **Confidence** score for each label. For example, the model might predict the label **Sea** for an image, with a confidence score of 96%. The model may have predicted the image as a **Glacier** with only a confidence score of 4%. Therefore, you can determine that your model is fairly confident in predicting images of the sea.

**Make single predictions with text prediction models**

To make a single prediction for a multi-category text prediction model, do the following:

1. In the left navigation pane of the Canvas application, choose **My models**.
2. On the **My models** page, choose your model.
3. After opening your model, choose the **Predict** tab.
4. On the **Run predictions** page, choose **Single prediction**.
5. For the **Text field**, enter the text for which you’d like to get a prediction.
6. Choose **Generate prediction results** to get your prediction.

In the right **Prediction results** pane, you receive an analysis of your text in addition to a **Confidence** score for each possible label. For example, if you entered a good review for a product, you might get **Positive** with a confidence score of 85%, while the confidence score for **Neutral** might be 10% and the confidence score for **Negative** only 5%.

### Make batch predictions

Make batch predictions when you have an entire dataset for which you’d like to generate predictions.

There are two types of batch predictions you can make:

- **Manual** batch predictions are when you have a dataset for which you want to make one-time predictions.

- **Automatic** batch predictions are when you set up a configuration that runs a batch prediction whenever a specific dataset is updated. For example, if you’ve configured weekly updates to a SageMaker Canvas dataset of inventory data, you can set up automatic batch predictions that run whenever you update the dataset. After setting up an automated batch predictions workflow, see [Manage automations](#) for more information about viewing and editing the details of your configuration. For more information about setting up automatic dataset updates, see [Configure automatic updates for a dataset](#).

**Note**

You can only set up automatic batch predictions for datasets imported through local upload or Amazon S3. Additionally, automatic batch predictions can only run while you’re logged in to the Canvas application. If you log out of Canvas, automatic batch prediction jobs resume when you log back in.

To get started, reviewing the following section for batch prediction dataset requirements, and then choose one of the following manual or automatic batch prediction workflows.

### Batch prediction dataset requirements
For batch predictions, make sure that your datasets meet the requirements outlined in [Create a dataset](#).

You might not be able to make predictions on some datasets because they have incompatible schemas. A *schema* is an organizational structure. For a tabular dataset, the schema is the names of the columns and the data type of the data in the columns. An incompatible schema might happen for one of the following reasons:

- The dataset that you're using to make predictions has fewer columns than the dataset that you're using to build the model.
- The data types in the columns you used to build the dataset might be different from the data types in dataset that you're using to make predictions.
- The dataset that you're using to make predictions and the dataset that you've used to build the model have column names that don't match. The column names are case sensitive. Column1 is not the same as column1.

To ensure that you can successfully generate batch predictions, match the schema of your batch predictions dataset to the dataset you used to train the model.

**Note**

For batch predictions, if you dropped any columns when building your model, Canvas adds the dropped columns back to the prediction results. However, Canvas does not add the dropped columns to your batch predictions for time series models.

### Make manual batch predictions

Choose one of the following procedures to make manual batch predictions based on your model type.

#### Make manual batch predictions with numeric and categorical prediction models

To make manual batch predictions for a numeric or categorical prediction model, do the following:

1. In the left navigation pane of the Canvas application, choose My models.
2. On the My models page, choose your model.
3. After opening your model, choose the Predict tab.
4. On the Run predictions page, choose Batch prediction.

5. Choose Select dataset if you’ve already imported your dataset. If not, choose Import new dataset, and then you’ll be directed through the import data workflow.

6. From the list of available datasets, select your dataset and choose Generate predictions to get your predictions.

After the prediction job finishes running, on the Run predictions page, you see an output dataset listed under Predictions. This dataset contains your results, and if you select the More options icon ( ), you can choose Preview to preview the output data. You can see the input data matched to the prediction and the probability that the prediction is correct. Then, you can choose Download prediction to download the results as a file.

Make manual batch predictions with image prediction models

To make manual batch predictions for a single-label image prediction model, do the following:

1. In the left navigation pane of the Canvas application, choose My models.

2. On the My models page, choose your model.

3. After opening your model, choose the Predict tab.

4. On the Run predictions page, choose Batch prediction.

5. Choose Select dataset if you’ve already imported your dataset. If not, choose Import new dataset, and then you’ll be directed through the import data workflow.

6. From the list of available datasets, select your dataset and choose Generate predictions to get your predictions.

After the prediction job finishes running, on the Run predictions page, you see an output dataset listed under Predictions. This dataset contains your results, and if you select the More options icon ( ), you can choose View prediction results to see the output data. You can see the images along with their predicted labels and confidence scores. Then, you can choose Download prediction to download the results as a CSV or a ZIP file.

Make manual batch predictions with text prediction models

To make manual batch predictions for a multi-category text prediction model, do the following:
1. In the left navigation pane of the Canvas application, choose **My models**.
2. On the **My models** page, choose your model.
3. After opening your model, choose the **Predict** tab.
4. On the **Run predictions** page, choose **Batch prediction**.
5. Choose **Select dataset** if you’ve already imported your dataset. If not, choose **Import new dataset**, and then you’ll be directed through the import data workflow. The dataset you choose must have the same source column as the dataset with which you built the model.
6. From the list of available datasets, select your dataset and choose **Generate predictions** to get your predictions.

After the prediction job finishes running, on the **Run predictions** page, you see an output dataset listed under **Predictions**. This dataset contains your results, and if you select the **More options** icon, you can choose **Preview** to see the output data. You can see the images along with their predicted labels and confidence scores. Then, you can choose **Download prediction** to download the results.

**Make automatic batch predictions**

To set up a schedule for automatic batch predictions, do the following:

1. In the left navigation pane of Canvas, choose **My models**.
2. Choose your model.
3. Choose the **Predict** tab.
4. Choose **Batch prediction**.
5. For **Generate predictions**, choose **Automatic**.
6. The **Automate batch predictions** dialog box pops up. Choose **Select dataset** and choose the dataset for which you want to automate predictions. Note that you can only select a dataset that was imported through local upload or Amazon S3.
7. After selecting a dataset, choose **Set up**.

Canvas runs a batch predictions job for the dataset after you set up the configuration. Then, every time you **Update a dataset**, either manually or automatically, another batch predictions job runs.

After the prediction job finishes running, on the **Run predictions** page, you see an output dataset listed under **Predictions**. This dataset contains your results, and if you select the **More options** icon.
you can choose Preview to preview the output data. You can see the input data matched to the prediction and the probability that the prediction is correct. Then, you can choose Download to download the results.

The following sections describe how to view, update, and delete your automatic batch prediction configuration through the Datasets page in the Canvas application. You can only set up a maximum of 20 automatic configurations in Canvas. For more information about viewing your automated batch predictions job history or making changes to your automatic configuration through the Automations page, see Manage automations.

View your automatic batch prediction jobs

To view your job history for your automatic batch predictions, go to the Predict tab of your model.

Each automatic batch prediction job shows up in the Predict tab of your model. Under Predictions, you can see the All jobs tab and the Configuration tabs:

- **All jobs** – In this tab, you can see all of the batch prediction jobs for this model. You can filter the jobs by configuration name. For each job, you can see fields such as the Input dataset, which includes the version of the dataset, and the Prediction type, such as whether the predictions were automatic or manual. If you choose the More options icon, you can choose View prediction or Download prediction.

- **Configuration** – In this tab, you can see all of the automatic batch prediction configurations you’ve created for this model. For each configuration, you can see fields such as the timestamp for when it was Created, the Input dataset it tracks for updates, and the Next job scheduled. If you choose the More options icon, you can choose View all jobs to see the job history and in progress jobs for the configuration.

Edit your automatic batch prediction configuration

You might want to make changes to your auto update configuration for a dataset, such as changing the frequency of the updates. You might also want to turn off your automatic update configuration to pause the updates to your dataset.

When you edit a batch prediction configuration, you can change the target dataset but not the frequency (since automatic batch predictions occur whenever the dataset is updated).
To edit your auto update configuration, do the following:

1. Go to the **Predict** tab of your model.
2. Under **Predictions**, choose the **Configuration** tab.
3. Find your configuration and choose the **More options** icon.
4. From the dropdown menu, choose **Update configuration**.
5. The **Automate batch prediction** dialog box opens. You can select another dataset and choose **Set up** to save your changes.

Your automatic batch predictions configuration is now updated.

To pause your automatic batch predictions, turn off your automatic configuration by doing the following:

1. Go to the **Predict** tab of your model.
2. Under **Predictions**, choose the **Configuration** tab.
3. Find your configuration from the list and turn off the **Auto update** toggle.

Automatic batch predictions are now paused. You can turn the toggle back on at any time to resume the update schedule.

**Delete your automatic batch prediction configuration**

To learn how to delete your automatic batch prediction configuration, see [Delete an automatic configuration](#).

You can also delete your configuration by doing the following:

1. Go to the **Predict** tab of your model.
2. Under **Predictions**, choose the **Configuration** tab.
3. Find your configuration from the list and choose the **More options** icon.
4. From the dropdown menu, choose **Delete configuration**.

Your configuration should now be deleted.
Send predictions to Amazon QuickSight

⚠️ **Note**

You can send batch predictions to Amazon QuickSight for numeric and categorical prediction and time series forecasting models. You can also send predictions generated with BYOM models. Single-label image prediction and multi-category text prediction models are excluded.

Once you generate batch predictions with custom tabular models in SageMaker Canvas, you can send those predictions as CSV files to Amazon QuickSight, which is a business intelligence (BI) service to build and publish predictive dashboards.

For example, if you built a 2 category prediction model to determine whether a customer will churn, you can create a visual, predictive dashboard in Amazon QuickSight to show the percentage of customers that are expected to churn. To learn more about Amazon QuickSight, see the Amazon QuickSight User Guide.

The following sections show you how to send your batch predictions to Amazon QuickSight for analysis.

**Before you begin**

Your user must have the necessary AWS Identity and Access Management (IAM) permissions to send your predictions to Amazon QuickSight. Your administrator can set up the IAM permissions for your user. For more information, see Grant Your Users Permissions to Send Predictions to Amazon QuickSight.

Your Amazon QuickSight account must contain the default namespace, which is set up when you first create your Amazon QuickSight account. Contact your administrator to help you get access to Amazon QuickSight. For more information, see Setting up for Amazon QuickSight in the Amazon QuickSight User Guide.

Your Amazon QuickSight account must be created in the same Region as your Canvas application. If your Amazon QuickSight account’s home Region differs from your Canvas application’s Region, you must either close and recreate your Amazon QuickSight account, or set up a Canvas application in the same Region as your Amazon QuickSight account. You can check your Amazon QuickSight home Region by doing the following (assuming you already have an Amazon QuickSight account):
1. Open your Amazon QuickSight console.

2. When the page loads, your Amazon QuickSight home Region is appended to the URL in the following format: https://<your-home-region>.quicksight.aws.amazon.com/.

You must know the usernames of the Amazon QuickSight users to whom you want to send your predictions. You can send predictions to yourself or other users who have the right permissions. Any users to whom you send predictions must be in the default namespace of your Amazon QuickSight account and have the Author or Admin role in Amazon QuickSight.

Additionally, Amazon QuickSight must have access to the SageMaker default Amazon S3 bucket for your domain, which is named with the following format: sagemaker-{REGION}-{ACCOUNT_ID}. The Region should be the same as your Amazon QuickSight account's home Region and your Canvas application's Region. To learn how to give Amazon QuickSight access to the batch predictions stored in your Amazon S3 bucket, see the topic I can't connect to Amazon S3 in the Amazon QuickSight User Guide.

**Supported data formats**

Before sending your predictions, check that the data format of your batch predictions is compatible with Amazon QuickSight.

- To learn more about the accepted data formats for timeseries data, see Supported date formats in the Amazon QuickSight User Guide.

- To learn more about data values that might prevent you from sending to Amazon QuickSight, see Unsupported values in data in the Amazon QuickSight User Guide.

Also note that Amazon QuickSight uses the character " as a text qualifier, so if your Canvas data contains any " characters, make sure that you close all matching quotes. Any mismatching quotes can cause issues with sending your dataset to Amazon QuickSight.

**Send your batch predictions to Amazon QuickSight**

Use the following procedure to send your predictions to Amazon QuickSight:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose My models.
3. On the My models page, choose your model.
4. Choose the **Predict** tab.

5. Under **Predictions**, select the dataset (or datasets) of batch predictions that you’d like to share. You can share up to 5 datasets of batch predictions at a time.

6. After you select your dataset, choose **Send to Amazon QuickSight**.

   **Note**

   The **Send to Amazon QuickSight** button doesn’t activate unless you select one or more datasets.

Alternatively, you can preview your predictions by choosing the **More options** icon ( ) and then **View prediction results**. From the dataset preview, you can choose **Send to Amazon QuickSight**. The following screenshot shows you the **Send to Amazon QuickSight** button in a dataset preview.

7. In the **Send to Amazon QuickSight** dialog box, do the following:
a. For **QuickSight users**, enter the name of the Amazon QuickSight users to whom you want to send your predictions. If you want to send them to yourself, enter your own username. You can only send predictions to users in the default namespace of the Amazon QuickSight account, and the user must have the Author or Admin role in Amazon QuickSight.

b. Choose **Send**.

The following screenshot shows the **Send to Amazon QuickSight** dialog box:

```
Send to Amazon QuickSight

Gain insights into your batch predictions by creating visualizations in Amazon QuickSight. You can publish your QuickSight analyses as a dashboard to share with others. Learn more

Name
Canvas_batchinfer-Titanic_test_4.csv
Canvas_batchinfer-Titanic_test_3.csv

QuickSight users
Add QuickSight users
Reach out to a QuickSight peer or admin for usernames.

Cancel Send
```

After you send your batch predictions, the **QuickSight** field for the datasets you sent shows as Sent. In the confirmation box that confirms your predictions were sent, you can choose **Open Amazon QuickSight** to open your Amazon QuickSight application. If you're done using Canvas, you should **log out** of the Canvas application.

The Amazon QuickSight users that you've sent datasets to can open their Amazon QuickSight application and view the Canvas datasets that have been shared with them. Then, they can create predictive dashboards with the data. For more information, see **Getting started with Amazon QuickSight data analysis** in the **Amazon QuickSight User Guide**.

By default, all of the users to whom you send predictions have owner permissions for the dataset in Amazon QuickSight. Owners are able to create analyses, refresh, edit, delete, and re-share datasets. The changes that owners make to a dataset change the dataset for all users with access. To change
the permissions, go to the dataset in Amazon QuickSight and manage its permissions. For more information, see Viewing and editing the permissions users that a dataset is shared with in the Amazon QuickSight User Guide.

Download a model notebook

**Note**

The model notebook feature is only available for tabular models and fine-tuned foundation models. Model notebooks aren't supported for image prediction, text prediction, or time series forecasting models.

If you'd like to generate a model notebook for a tabular model built before this feature was launched, you must rebuild the model to generate a notebook.

For eligible models that you successfully build in Amazon SageMaker Canvas, a Jupyter notebook containing a report of all the model building steps is generated. This Jupyter notebook contains Python code that you can run locally or run in an environment like Amazon SageMaker Studio Classic to replicate the steps necessary to build your model. The notebook can be useful if you’d like to experiment with the code or see the backend details of how Canvas builds models.

To access the model notebook, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose My models.
3. Choose the model and version that you built.
4. On the model version’s page, choose the More options icon in the header.
5. From the dropdown menu, choose View notebook.
6. A popup appears with the notebook content. You can choose Download and then do one of the following:
   a. Choose Download to save the notebook content to your local device.
   b. Choose Copy S3 URI to copy the Amazon S3 location where the notebook is stored.
   The notebook is stored in the Amazon S3 bucket specified in your Canvas storage.
configuration, which is configured in the Prerequisites for setting up Amazon SageMaker Canvas section.

You should now be able to view the notebook either locally or as an object in Amazon S3. You can upload the notebook to an IDE to edit and run the code, or you can share the notebook with others in your organization to review.

Send your model to Amazon QuickSight

If you use Amazon QuickSight and want to leverage SageMaker Canvas in your Amazon QuickSight visualizations, you can build an Amazon SageMaker Canvas model and use it as a predictive field in your Amazon QuickSight dataset. A predictive field is a field in your Amazon QuickSight dataset that can make predictions for a given column in your dataset, similar to how Canvas users make single or batch predictions with a model. To learn more about how to integrate Canvas predictive abilities into your Amazon QuickSight datasets, see SageMaker Canvas integration in the Amazon QuickSight User Guide.

The following steps explain how you can add a predictive field to your Amazon QuickSight dataset using a Canvas model:

1. Open the Canvas application and build a model with your dataset.
2. After building the model in Canvas, send the model to Amazon QuickSight. A schema file automatically downloads to your local machine when you send the model to Amazon QuickSight. You upload this schema file to Amazon QuickSight in the next step.
3. Open Amazon QuickSight and choose a dataset with the same schema as the dataset you used to build your model. Add a predictive field to the dataset and do the following:
   a. Specify the model sent from Canvas.
   b. Upload the schema file that was downloaded in Step 2.
4. Save and publish your changes, and then generate predictions for the new dataset. Amazon QuickSight uses the model to fill in the target column with predictions.

In order to send a model from Canvas to Amazon QuickSight, you must meet the following prerequisites:

• You must have both Canvas and Amazon QuickSight set up. Your Amazon QuickSight account must be created in the same AWS Region as your Canvas application. If your Amazon QuickSight
account’s home Region differs from your Canvas application’s Region, you must either close and recreate your Amazon QuickSight account, or set up a Canvas application in the same Region as your Amazon QuickSight account. Your Amazon QuickSight account must also contain the default namespace, which you set up when you first create your Amazon QuickSight account. Contact your administrator to help you get access to Amazon QuickSight. For more information, see Setting up for Amazon QuickSight in the Amazon QuickSight User Guide.

- Your user must have the necessary AWS Identity and Access Management (IAM) permissions to send your predictions to Amazon QuickSight. Your administrator can set up the IAM permissions for your user. For more information, see Grant Your Users Permissions to Send Predictions to Amazon QuickSight.
- Amazon QuickSight must have access to the Amazon S3 bucket that you’ve specified for Canvas application storage. For more information, see Configure your Amazon S3 storage.

Time Series Forecasts in Amazon SageMaker Canvas

**Note**

Time series forecasting models are only supported for tabular datasets.

Amazon SageMaker Canvas gives you the ability to use machine learning time series forecasts. Time series forecasts give you the ability to make predictions that can vary with time.

You can make a time series forecast for the following examples:

- Forecasting your inventory in the coming months.
- The number of items sold in the next four months.
- The effect of reducing the price on sales during the holiday season.
- Item inventory in the next 12 months.
- The number of customers entering a store in the next several hours.
- Forecasting how a 10% reduction in the price of a product affects sales over a time period.

To make a time series forecast, your dataset must have the following:

- A timestamp column with all values having the datetime type.
• A target column that has the values that you’re using to forecast future values.
• An item ID column that contains unique identifiers for each item in your dataset, such as SKU numbers.

The datetime values in the timestamp column must use one of the following formats:

• YYYY-MM-DD MM:SS
• YYYY-MM-DDTHH:MM:SS
• YYYY-MM-DD
• MM/DD/YY
• MM/DD/YY HH:MM
• MM/DD/YYYY
• YYYY/MM/DD HH:MM:SS
• YYYY/MM/DD
• DD/MM/YYYY
• DD/MM/YY
• DD-MM-YY
• DD-MM-YYYY

You can make forecasts for the following intervals:

• 1 min
• 5 min
• 15 min
• 30 min
• 1 hour
• 1 day
• 1 week
• 1 month
• 1 year
Future values in your input dataset

Canvas automatically detects columns in your dataset that might potentially contain future values. If present, these values can enhance the accuracy of predictions. Canvas marks these specific columns with a Future values label. Canvas infers the relationship between the data in these columns and the target column that you are trying to predict, and utilizes that relationship to generate more accurate forecasts.

For example, you can forecast the amount of ice cream sold by a grocery store. To make a forecast, you must have a timestamp column and a column that indicates how much ice cream the grocery store sold. For a more accurate forecast, your dataset can also include the price, the ambient temperature, the flavor of the ice cream, or a unique identifier for the ice cream.

Ice cream sales might increase when the weather is warmer. A decrease in the price of the ice cream might result in more units sold. Having a column with ambient temperature data and a column with pricing data can improve your ability to forecast the number of units of ice cream the grocery store sells.

While providing future values is optional, it helps you to perform what-if analyses directly in the Canvas application, showing you how changes in future values could alter your predictions.

Handling missing values

You might have missing data for different reasons. The reason for your missing data might inform how you want Canvas to impute it. For example, your organization might use an automatic system that only tracks when a sale happens. If you're using a dataset that comes from this type of automatic system, you have missing values in the target column.

⚠️ Important
If you have missing values in the target column, we recommend using a dataset that doesn't have them. SageMaker Canvas uses the target column to forecast future values. Missing values in the target column can greatly reduce the accuracy of the forecast.

For missing values in the dataset, Canvas automatically imputes the missing values for you by filling the target column with 0 and other numeric columns with the median value of the column.

However, you can select your own filling logic for the target column and other numeric columns in your datasets. Target columns have different filling guidelines and restrictions than the rest of the
numeric columns. Target columns are filled up to the end of the historical period, whereas numeric columns are filled across both historical and future periods all the way to the end of the forecast horizon. Canvas only fills future values in a numeric column if your data has at least one record with a future timestamp and a value for that specific column.

You can choose one of the following filling logic options to impute missing values in your data:

- **zero** – Fill with 0.
- **NaN** – Fill with NaN, or not a number. This is only supported for the target column.
- **mean** – Fill with the mean value from the data series.
- **median** – Fill with the median value from the data series.
- **min** – Fill with the minimum value from the data series.
- **max** – Fill with the maximum value from the data series.

When choosing a filling logic, you should consider how your model interprets the logic. For example, in a retail scenario, recording zero sales of an available item is different from recording zero sales of an unavailable item, as the latter scenario doesn't necessarily imply a lack of customer interest in the unavailable item. In this case, filling with 0 in the target column of the dataset might cause the model to be under-biased in its predictions and infer a lack of customer interest in unavailable items. Conversely, filling with NaN might cause the model to ignore true occurrences of zero items being sold of available items.

**Types of forecasts**

You can make one of the following types of forecasts:

- **Single item**
- **All items**

For a forecast on all the items in your dataset, SageMaker Canvas returns a forecast for the future values for each item in your dataset.

For a single item forecast, you specify the item and SageMaker Canvas returns a forecast for the future values. The forecast includes a line graph that plots the predicted values over time.

**Topics**

- [Gain additional insights from your forecast](#)
• **Make a time series forecast**

**Gain additional insights from your forecast**

In Amazon SageMaker Canvas, you can use the following optional methods to get more insights from your forecast:

- Group column
- Holiday schedule
- What-if scenario

You can specify a column in your dataset as a **Group column**. Amazon SageMaker Canvas groups the forecast by each value in the column. For example, you can group the forecast on columns containing price data or unique item identifiers. Grouping a forecast by a column lets you make more specific forecasts. For example, if you group a forecast on a column containing item identifiers, you can see the forecast for each item.

Overall sales of items might be impacted by the presence of holidays. For example, in the United States, the number of items sold in both November and December might differ greatly from the number of items sold in January. If you use the data from November and December to forecast the sales in January, your results might be inaccurate. Using a holiday schedule prevents you getting inaccurate results. You can use a holiday schedule for 251 countries.

For a forecast on a single item in your dataset, you can use what-if scenarios. A what-if scenario gives you the ability to change values in your data and change the forecast. For example, you can answer the following questions by using a what-if scenario, "What if I lowered prices? How would that affect the number of items sold?"

**Make a time series forecast**

To make a time series forecast, you choose a target column. The target column contains the data that you want to predict. For example, your target column might have data on the number of items sold. After you select the target column, Amazon SageMaker Canvas selects a **Model type**. SageMaker Canvas uses the time-series data to automatically choose a time series model that you can use to make predictions on your data. After you build the model, you can evaluate its performance and use it to make predictions on new data.

Use the following procedure to make a time series forecast.
To make a time-series forecast, do the following.

1. Import the data.
2. Choose a target column in your dataset.
3. SageMaker Canvas automatically chooses **Time series forecasting** as the model type. Choose **Set configuration** to confirm that you're performing a time series forecast.
4. Specify the following fields:
   - **Item ID column** – The column that contains unique identifiers for each item in your dataset. For example, an SKU number uniquely identifies an item.
   - Optional: **Group column** – Groups the time series forecast by values in the column. For example, you can group your forecast for an item by store.
   - **Time stamp column** – The column containing the time stamps in your dataset. For a list of the supported date time formats for this column, see [Time Series Forecasts in Amazon SageMaker Canvas](#).
   - **Future timestamp** – A timestamp that indicates a future forecast time. SageMaker Canvas forecasts values up to the point in time specified by the timestamp.
   - Optional: **Holiday schedule** – Activate the holiday schedule to use a country's holiday schedule. Use it to make your forecasts with holiday data more accurate.

You can have one of the following types of missing values:

- Missing future values
- Missing values

Missing future values are missing values in the target column. SageMaker Canvas uses the values in the target column to forecast the values in the future. If you have missing values in the target column, your forecast might be less accurate. We highly recommend updating the dataset.

Missing values are values that are missing in any column other than the target column. With missing values that aren't in the target column, it's helpful to note the following:

- They generally don't reduce the accuracy of your forecast as much as missing future values.
- SageMaker Canvas automatically imputes the missing values.
You can evaluate the model by seeing how close the predictions are within the actual value. You can also use the **Column Impact** metric to determine the direction and magnitude of the column's impact on the model's predictions. For example, in the following image, holidays had the largest positive impact on the forecast for demand. Price had the largest negative impact on demand.

After you've built a model, you can make the following types of forecasts:

- **Single item** – Make a forecast for a single item in a dataset and a line graph of the values that SageMaker Canvas forecasts. For example, you can see how sales of an item vary over time.
- **All items** – Make a forecast for all items in a dataset.
- **What-if scenario** – See how changing values in the dataset can affect the overall forecast for a single item.
The following image shows a single item forecast with a what-if scenario. In a what-if scenario, you have the ability to change values that can vary with time. You can see how changing the values affects the forecast.

The points connected by the solid blue line are the values that the model forecasts. The points connected by the dashed lines show the what-if scenario.

### Updating a Model in Amazon SageMaker Canvas

Amazon SageMaker Canvas gives you the ability to update the models that you've built using new data. SageMaker Canvas shows you a model history, so that you can compare the models that you've built recently to those that you've generated in the past.

Each model that you build has a *version* number. The first model is Version 1, or V1. You can use model versions to see changes in prediction accuracy when you update your data or use advanced transformations.
Text prediction and image prediction models only support one model version.

For new versions of a model, you can only choose datasets that have the same target column as the target column in Version 1. You must build at least one version of a model to add a new version, and you can delete versions that aren’t useful to you anymore.

You can also see Register a model version in the SageMaker model registry to help you track your versions over time and collaborate with Studio Classic users who can approve or reject your model versions.

Use the following procedure to add a new model version or to view all of the versions for your model.

To add a new model version, do the following:

1. Open your SageMaker Canvas application.
2. In the left navigation pane, choose My models.
3. On the My models page, choose your model. You can Filter by problem type to find your model more easily.
4. After choosing your model, the Versions page opens, listing all of the versions of your model.
5. Choose Add version.

The following image shows the Versions page for a model, on which you can view your model versions and add new versions.
On the **Versions** page, you can view the following information for each of your model versions:

- **Status** – This field tells you whether your model is currently building (In building), done building (Ready), failed to build (Failed), or still being edited (In draft).

- **Model score, F1, Precision, Recall, and AUC** – If you turn on the **Show advanced metrics** toggle on this page, you can see these model metrics. These metrics indicate the accuracy and performance of your model. For more information, see [Evaluate your model](#).

- **Shared** – This field tells you whether or not you've shared the model version with SageMaker Studio Classic users.

- **Model registry** – This field tells you whether or not you've registered the version to a model registry. For more information, see [Register a model version in the SageMaker model registry](#).

After you choose a new version, you start the process of building another model. The process for building a new version of a model is almost the same as the process for building a model for the first time. For new versions of a model, you can only choose datasets that have the same target column as the target column in Version 1. For more information about building a model, see [Build a custom model](#).
Operationalize your models

After building a model in SageMaker Canvas that you feel confident about, you might want to integrate your model with the machine learning operations (MLOps) processes in your organization. MLOps includes common tasks such as deploying a model for use in production or setting up continuous integration and continuous deployment (CI/CD) pipelines.

The following topics describe how you can use features within Canvas to use a Canvas-built model in production.

Topics

- Register a model version in the SageMaker model registry
- Deploy your models to an endpoint

Register a model version in the SageMaker model registry

With SageMaker Canvas, you can build multiple iterations, or versions, of your model to improve it over time. You might want to build a new version of your model if you acquire better training data or if you want to attempt to improve the model's accuracy. For more information about adding versions to your model, see Update a model.

After you've built a model that you feel confident about, you might want to evaluate its performance and have it reviewed by a data scientist or MLOps engineer in your organization before using it in production. To do this, you can register your model versions to the SageMaker model registry. The SageMaker model registry is a repository that data scientists or engineers can use to catalog machine learning (ML) models and manage model versions and their associated metadata, such as training metrics. They can also manage and log the approval status of a model.

After you register your model versions to the SageMaker model registry, a data scientist or your MLOps team can access the SageMaker model registry through SageMaker Studio Classic, which is a web-based integrated development environment (IDE) for working with machine learning models. In the SageMaker model registry interface in Studio Classic, the data scientist or MLOps team can evaluate your model and update its approval status. If the model doesn't perform to their requirements, the data scientist or MLOps team can update the status to Rejected. If the model does perform to their requirements, then the data scientist or MLOps team can update the status to Approved. Then, they can deploy your model to an endpoint or automate model deployment with CI/CD pipelines. You can use the SageMaker model registry feature to seamlessly integrate models built in Canvas with the MLOps processes in your organization.
The following diagram summarizes an example of registering a model version built in Canvas to the SageMaker model registry for integration into an MLOps workflow.

You can register tabular, image, and text model versions to the SageMaker model registry. This includes time series forecasting models and SageMaker JumpStart based fine-tuned foundation models.

**Note**

Currently, you can't register BYOM model versions or Amazon Bedrock based fine-tuned foundation models built in Canvas to the SageMaker model registry.

The following sections show you how to register a model version to the SageMaker model registry from Canvas.

**Permissions management**

By default, you have permissions to register model versions to the SageMaker model registry. SageMaker grants these permissions for all new and existing Canvas user profiles through the AmazonSageMakerCanvasFullAccess policy, which is attached to the AWS IAM execution role for the SageMaker domain that hosts your Canvas application.

If your Canvas administrator is setting up a new domain or user profile, when they're setting up the domain and following the prerequisite instructions in the Getting started guide, SageMaker turns on the model registration permissions through the ML Ops permissions configuration option, which is enabled by default.
The Canvas administrator can manage model registration permissions at the user profile level as well. For example, if the administrator wants to grant model registration permissions to some user profiles but remove permissions for others, they can edit the permissions for a specific user. The following procedure shows how to turn off model registration permissions for a specific user profile:

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the user profile's domain.
5. On the **domain details** page, choose the **User profile** whose permissions you want to edit.
6. On the **User Details** page, choose **Edit**.
7. In the left navigation pane, choose **Canvas settings**.
8. In the **ML Ops permissions configuration** section, turn off the **Enable Model Registry registration permissions** toggle.
9. Choose **Submit** to save the changes to your domain settings.

The user profile should no longer have model registration permissions.

**Register a model version to the SageMaker model registry**

SageMaker model registry tracks all of the model versions that you build to solve a particular problem in a *model group*. When you build a SageMaker Canvas model and register it to SageMaker model registry, it gets added to a model group as a new model version. For example, if you build and register four versions of your model, then a data scientist or MLOps team working in the SageMaker model registry interface can view the model group and review all four versions of the model in one place.

When registering a Canvas model to the SageMaker model registry, a model group is automatically created and named after your Canvas model. Optionally, you can rename it to a name of your choice, or use an existing model group in the SageMaker model registry. For more information about creating a model group, see [Create a Model Group](#).

---

**Use custom models**
To register a model version to the SageMaker model registry from the Canvas application, use the following procedure:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose My models.
3. On the My models page, choose your model. You can Filter by problem type to find your model more easily.
4. After choosing your model, the Versions page opens, listing all of the versions of your model. You can turn on the Show advanced metrics toggle to view the advanced metrics, such as Recall and Precision, to compare your model versions and determine which one you’d like to register.
5. From the list of model versions, for the the version that you want to register, choose the More options icon.
   Alternatively, you can double click on the version that you need to register, and then on the version details page, choose the More options icon.
6. In the dropdown list, choose Add to Model Registry. The Add to Model Registry dialog box opens.
7. In the Add to Model Registry dialog box, do the following:
   a. (Optional) In the SageMaker Studio Classic model group section, for the Model group name field, enter the name of the model group to which you want to register your version. You can specify the name for a new model group that SageMaker creates for you, or you can specify an existing model group. If you don’t specify this field, Canvas registers your version to a default model group with the same name as your model.
   b. Choose Add.

Your model version should now be registered to the model group in the SageMaker model registry. When you register a model version to a model group in the SageMaker model registry, all
subsequent versions of the Canvas model are registered to the same model group (if you choose to register them). If you register your versions to a different model group, you need to go to the SageMaker model registry and delete the model group. Then, you can re-register your model versions to the new model group.

To view the status of your models, you can return to the Versions page for your model in the Canvas application. This page shows you the Model Registry status of each version. If the status is Registered, then the model has been successfully registered.

If you want to view the details of your registered model version, for the Model Registry status, you can hover over the Registered field to see the Model registry details pop-up box. These details contain more info, such as the following:

- The Model package group name is the model group that your version is registered to in the SageMaker model registry.
- The Approval status, which can be Pending Approval, Approved, or Rejected. If a Studio Classic user approves or rejects your version in the SageMaker model registry, then this status is updated on your model versions page when you refresh the page.

The following screenshot shows the Model registry details box, along with an Approval status of Approved for this particular model version.

Model Registry details

- Model package group name: canvas-test-cv-v1
- Model Registry version: Version 1
- Model Registry account ID: [redacted]
- Approval status: Approved

Deploy your models to an endpoint

In Amazon SageMaker Canvas, you can deploy your models to an endpoint to make predictions. SageMaker provides the ML infrastructure for you to host your model on an endpoint with the
compute instances that you choose. Then, you can *invoke* the endpoint (send a prediction request) and get a real-time prediction from your model. With this functionality, you can use your model in production to respond to incoming requests, and you can integrate your model with existing applications and workflows.

To get started, you should have a model version that is ready to deploy. For more information about building a model in Canvas, see [Build a custom model](#).

⚠️ **Important**

You can deploy any model type in SageMaker Canvas except for time series forecasting models.

Review the following **Permissions management** section, and then begin creating new deployments in the **Deploy a model** section.

**Permissions management**

By default, you have permissions to deploy model versions to the SageMaker Hosting endpoints. SageMaker grants these permissions for all new and existing Canvas user profiles through the `AmazonSageMakerCanvasFullAccess` policy, which is attached to the AWS IAM execution role for the SageMaker domain that hosts your Canvas application.

If your Canvas administrator is setting up a new domain or user profile, when they're setting up the domain and following the prerequisite instructions in the [Prerequisites for setting up Amazon SageMaker Canvas](#), SageMaker turns on the model deployment permissions through the **Enable direct deployment of Canvas models** option, which is enabled by default.

The Canvas administrator can manage model deployment permissions at the user profile level as well. For example, if the administrator wants to grant model deployment permissions to some user profiles but remove permissions for others, they can edit the permissions for a specific user.

The following procedure shows how to turn off model deployment permissions for a specific user profile:

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. From the list of domains, select the user profile’s domain.

5. On the **domain details** page, choose the **User profile** whose permissions you want to edit.

6. On the **User Details** page, choose **Edit**.

7. In the left navigation pane, choose **Canvas settings**.

8. In the **ML Ops permissions configuration** section, turn off the **Enable direct deployment of Canvas models** toggle.

9. Choose **Submit** to save the changes to your domain settings.

The user profile should no longer have model deployment permissions.

**Deploy a model**

To get started with deploying your model, you create a new deployment in Canvas and specify the model version that you want to deploy along with the ML infrastructure, such as the type and number of compute instances that you would like to use for hosting the model.

Canvas suggests a default type and number of instances based on your model type, or you can learn more about the various SageMaker instance types on the [Amazon SageMaker pricing page](https://aws.amazon.com/sagemaker/pricing/). You are charged based on the SageMaker instance pricing while your endpoint is active.

After your model is deployed to a SageMaker Hosting **real-time inference endpoint**, you can begin making predictions by *invoking* the endpoint.

There are several different ways for you to deploy a model version from the Canvas application. You can access the model deployment option through any of the following methods:

- On the **My models** page of the Canvas application, you can choose the model that you want to deploy. Then, from the model's **Versions** page, you can choose the **More options** icon ([1](#)) next to a model version and select **Deploy**.

- When on the details page for a model version, on the **Analyze** tab, you can choose the **Deploy** option.

- When on the details page for a model version, on the **Predict** tab, you can choose the **More options** icon ([1](#)) at the top of the page and select **Deploy**.
On the **Operations** page of the Canvas application, you can choose the **Deployments** tab and then choose **Create deployment**.

All of these methods open the **Deploy model** side panel, where you specify the deployment configuration for your model. To deploy the model from this panel, do the following:

1. (Optional) If you're creating a deployment from the **Operations** page, you'll have the option to **Select model and version**. Use the dropdown menus to select the model and model version that you want to deploy.

2. Enter a name in the **Deployment name** field.

3. For **Instance type**, SageMaker detects a default instance type and number that is suitable for your model. However, you can change the instance type that you would like to use for hosting your model.

4. For **Instance count**, you can set the number of active instances that are used for your endpoint. SageMaker detects a default number that is suitable for your model, but you can change this number.

5. When you're ready to deploy your model, choose **Deploy**.

Your model should now be deployed to an endpoint. For information about how to view your deployment details or perform various actions, see the following sections.

**View your deployments**

You might want to check the status or details of a model deployment in Canvas. For example, if your deployment failed, you might want to check the details to troubleshoot.

You can view your Canvas model deployments from the Canvas application or from the Amazon SageMaker console.

To view deployment details from Canvas, choose one of the following procedures:
To view your deployment details from the Operations page, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation pane, choose Operations.
3. Choose the Deployments tab.
4. Choose your deployment by name from the list.

To view your deployment details from a model version’s page, do the following:

1. In the SageMaker Canvas application, go to your model version’s details page.
2. Choose the Deploy tab.
3. On the Deployments section that lists all of the deployment configurations associated with that model version, find your deployment.
4. Choose the More options icon
   and then select View details to open the details page.

The details page for your deployment opens, and you can view information such as the time of the most recent prediction, the endpoint’s status and configuration, and the model version that is currently deployed to the endpoint.

You can also view your currently active Canvas workspace instances and active endpoints from the SageMaker dashboard in the SageMaker console. Your Canvas endpoints are listed alongside any other SageMaker Hosting endpoints that you’ve created, and you can filter them by searching for endpoints with the Canvas tag.

The following screenshot shows the SageMaker dashboard. In the Canvas section, you can see that one workspace instance is in service and four endpoints are active.
Update a deployment configuration

You can also update your deployment configuration. For example, you can deploy an updated model version to the endpoint, or you can update the instance type or number of instances behind the endpoint based on your capacity needs.

There are several different ways for you to update your deployment from the Canvas application. You can use any of the following methods:

- On the **Operations** page of the Canvas application, you can choose the **Deployments** tab and select the deployment that you want to update. Then, choose **Update configuration**.
- When on the details page for a model version, on the **Deploy** tab, you can view the deployments for that version. Next to the deployment, choose the **More options** icon.
and then choose **Update configuration**.

Both of the preceding methods open the **Update configuration** side panel, where you can make changes to your deployment configuration. To update the configuration, do the following:

1. For the **Select version** dropdown menu, you can select a different model version to deploy to the endpoint.

   **Note**
   When updating a deployment configuration, you can only choose a different model version to deploy. To deploy a different model, create a new deployment.

2. For **Instance type**, you can select a different instance type for hosting your model.
3. For **Instance count**, you can change the number of active instances that are used for your endpoint.
4. Choose **Save**.

Your deployment configuration should now be updated.

**Test your deployment**

You can test your deployment by invoking the endpoint, or making single prediction requests, through the Canvas application. The endpoint returns a response with the prediction along with the probabilities of that prediction being correct. You can use this functionality to confirm that your endpoint responds to requests before invoking your endpoint programmatically in a production environment.

**Note**
Execution length is an estimate of the time taken to invoke and get a response from the endpoint in Canvas. For detailed latency metrics, see [SageMaker Endpoint Invocation Metrics](#).

To test your endpoint through the Canvas application, do the following:
1. Open the SageMaker Canvas application.
2. In the left navigation panel, choose **Operations**.
3. Choose the **Deployments** tab.
4. From the list of deployments, choose the one with the endpoint that you want to invoke.
5. On the deployment’s details page, choose the **Test deployment** tab.
6. On the deployment testing page, you can modify the **Value** fields to specify a new data point.
7. After modifying the values, choose **Update** to get the prediction result.

The prediction loads, along with the **Invocation result** fields which indicate whether or not the invocation was successful and how long the request took to process.

The following screenshot shows a prediction performed in the Canvas application on the **Test deployment** tab.

![Canvas Application Screenshot]

For all model types except numeric prediction, the prediction returns the following fields:

- **predicted_label** – the predicted output
- **probability** – the probability that the predicted label is correct
• **labels** – the list of all the possible labels

• **probabilities** – the probabilities corresponding to each label (the order of this list matches the order of the labels)

For numeric prediction models, the prediction only contains the **score** field, which is the predicted output of the model, such as the predicted price of a house.

You can continue making single predictions through the deployment testing page, or you can see the following section **Invoke your endpoint** to learn how to invoke your endpoint programmatically from applications.

**Invoke your endpoint**

After testing your deployment, you can use your endpoint in production with your applications by invoking the endpoint programmatically the same way that you can invoke any other SageMaker real-time endpoint. Invoking an endpoint programmatically returns a response object which contains the same fields as mentioned in the preceding section **Test your deployment**.

For more detailed information about how to programmatically invoke endpoints, see **Invoke models for real-time inference**.

The following Python examples show you how to invoke your endpoint based on the model type.

**Numeric and categorical prediction models**

The following example shows you how to invoke numeric or categorical prediction models.

```python
import boto3
import pandas as pd

client = boto3.client("runtime.sagemaker")
body = pd.DataFrame(["feature_column1", "feature_column2"], ["feature_column1", "feature_column2"]).to_csv(header=False, index=False).encode("utf-8")

response = client.invoke_endpoint(
    EndpointName="endpoint_name",
    ContentType="text/csv",
    Body=body,
    Accept="application/json"
)
```
**Image prediction models**

The following example shows you how to invoke image prediction models.

```python
import boto3
client = boto3.client("runtime.sagemaker")
with open("example_image.jpg", "rb") as file:
    body = file.read()
    response = client.invoke_endpoint(
        EndpointName="endpoint_name",
        ContentType="application/x-image",
        Body=body,
        Accept="application/json"
    )
```

**Text prediction models**

The following example shows you how to invoke text prediction models.

```python
import boto3
import pandas as pd
client = boto3.client("runtime.sagemaker")
body = pd.DataFrame(["Example text 1"], ["Example text 2"])
  .to_csv(header=False, index=False).encode("utf-8")
response = client.invoke_endpoint(
    EndpointName="endpoint_name",
    ContentType="text/csv",
    Body=body,
    Accept="application/json"
)
```

**Delete a model deployment**

You can delete your model deployment from the Canvas application. This action also deletes the endpoint from the SageMaker console and shuts down any endpoint-related resources.

ℹ️ **Note**

Optionally, you can delete your endpoint through the [SageMaker console](https://aws.amazon.com/sagemaker/) or using the SageMaker DeleteEndpoint API. For more information, see [Delete Endpoints](https://docs.aws.amazon.com/sagemaker/latest/dg/delete-endpoint.html) and [Endpoints](https://docs.aws.amazon.com/sagemaker/latest/dg/endpoints.html).
To delete a deployment in Canvas, do the following:

1. Open the SageMaker Canvas application.
2. In the left navigation panel, choose Operations.
3. Choose the Deployments tab.
4. From the list of deployments, choose the one that you want to delete.
5. At the top of the deployment details page, choose the More options icon.
6. Choose Delete deployment.
7. In the Delete deployment dialog box, choose Delete.

Your deployment and SageMaker Hosting endpoint should now be deleted from both Canvas and the SageMaker console.

Manage automations

In SageMaker Canvas, you can create automations that update your dataset or generate predictions from your model on a schedule. For example, you might receive new shipping data on a daily basis. You can set up an automatic update for your dataset and automatic batch predictions that run whenever the dataset is updated. Using these features, you can set up an automated workflow and reduce the amount of time you spend manually updating datasets and making predictions.

Note

You can only set up a maximum of 20 automatic configurations in your Canvas application. Automations are only active while you’re logged in to the Canvas application. If you log out of Canvas, your automatic jobs pause until you log back in.

The following sections describe how to view, edit, and delete configurations for existing automations. To learn how to set up automations, see the following topics:
• To set up automatic dataset updates, see Update a dataset.

• To set up automatic batch predictions, see Make batch predictions.

View your automations

You can also view all of your auto update jobs by going to the left navigation pane of Canvas and choosing Automations. The Automations page combines automations for both automatic dataset updates and automatic batch predictions. From the Automations page, you can see the following tabs:

• All jobs – You can see every instance of a Dataset update or Batch prediction job that Canvas has done. For each job, you can see fields such as the associated Input dataset, the Configuration name of the associated auto update configuration, and the Status showing whether the job was successful or not. You can filter the jobs by configuration name:
  • For dataset update jobs, you can choose the latest version of the dataset, or the most recent job, to preview the dataset.
  • For batch prediction jobs, you can choose the More options icon to view or download the predictions for that job.

• Configuration – You can see all of the Dataset update and Batch prediction configurations you've created. For each configuration, you can see fields such as the associated Input dataset and the Frequency of the jobs. You can also turn off or turn on the Auto update toggle to pause or resume automatic updates. If you choose the More options icon for a specific configuration, you can choose to View all jobs for the configuration, Update configuration, or Delete configuration.

Edit your automatic configurations

After setting up a configuration, you might want to make changes to it. For automatic dataset updates, you can update the Amazon S3 location for Canvas to import data, the frequency of the updates, and the starting time. For automatic batch predictions, you can change the dataset that the configuration tracks for updates. You can also turn off the automation to temporarily pause updates until you choose to resume them.

The following sections show you how to update each type of configuration.
**Note**

You can't change the frequency for automatic batch predictions because automatic batch predictions run every time the target dataset is updated.

**Edit your automatic dataset update configuration**

You might want to make changes to your auto update configuration for a dataset, such as changing the frequency of the updates. You might also want to turn off your automatic update configuration to pause the updates to your dataset.

To make changes to your auto update configuration for a dataset, do the following:

1. In the left navigation pane of Canvas, choose **Automations**.
2. Choose the **Configuration** tab.
3. For your auto update configuration, choose the **More options** icon.
4. In the dropdown menu, choose **Update configuration**. You are taken to the **Auto updates** tab of the dataset.
5. Make your changes to the configuration. When you're done making changes, choose **Save**.

To pause your dataset updates, turn off your automatic configuration. One way to turn off auto updates is by doing the following:

1. In the left navigation pane of Canvas, choose **Automations**.
2. Choose the **Configuration** tab.
3. Find your configuration from the list and turn off the **Auto update** toggle.

Automatic updates for your dataset are now paused. You can turn this toggle back on at any time to resume the update schedule.

**Edit your automatic batch prediction configuration**

When you edit a batch prediction configuration, you can change the target dataset but not the frequency (since automatic batch predictions occur whenever the dataset is updated).
To make changes to your automatic batch predictions configuration, do the following:

1. In the left navigation pane of Canvas, choose **Automations**.
2. Choose the **Configuration** tab.
3. For your auto update configuration, choose the **More options** icon
   
   ).
4. In the dropdown menu, choose **Update configuration**. You are taken to the **Auto updates** tab of the dataset.
5. The **Automate batch prediction** dialog box opens. You can select another dataset and choose **Set up** to save your changes.

Your automatic batch predictions configuration is now updated.

To pause your automatic batch predictions, turn off your automatic configuration. Use the following procedure to turn off your configuration:

1. In the left navigation pane of Canvas, choose **Automations**.
2. Choose the **Configuration** tab.
3. Find your configuration from the list and turn off the **Auto update** toggle.

Automatic batch predictions for your dataset are now paused. You can turn this toggle back on at any time to resume the update schedule.

**Delete an automatic configuration**

You might want to delete a configuration to stop your automated workflow in SageMaker Canvas.

To delete a configuration for automatic dataset updates or automatic batch predictions, do the following:

1. In the left navigation pane of Canvas, choose **Automations**.
2. Choose the **Configuration** tab.
3. Find your auto update configuration, and choose the **More options** icon

   ).
4. Choose **Delete configuration**.
5. In the dialog box that pops up, choose **Delete**.
Collaborate with data scientists

Note

The functionality described on this page only applies to Amazon SageMaker Studio Classic. Currently, you can only share models to Canvas (or view shared Canvas models) in Studio Classic. If you’re currently using the latest version of Studio, you must run Studio Classic from within the latest version of Studio to share models to Canvas or view models shared from Canvas. For more information about accessing Studio Classic, see the Studio Classic documentation.

With Amazon SageMaker Canvas, business analysts using Canvas and data scientists using Amazon SageMaker Studio Classic can share ML models and collaborate with each other while working in their own environments to share domain knowledge and provide expert inputs towards improving models.

Using SageMaker Canvas collaboration, you can share Standard build models from Canvas with data scientists in Studio Classic to review, update, and share back with Canvas users. Users in Canvas can share one version of a model with up to 23 Studio Classic users.

Note

Collaboration on models with Studio Classic users isn’t supported for single-label image prediction, multi-category text prediction, or time series forecasting model types. Additionally, SageMaker Canvas doesn’t support sharing your model to the same user profile as the one that created the model. You must have two separate user profiles to share a model.

The following sections describe the steps for collaboration:

• In the Canvas application, a business analyst shares their model with a Studio Classic user.
• The Studio Classic user receives the shared model in the Studio Classic application. They can choose to share feedback with the analyst, make updates to the model, or share an alternate model version.
• The business analyst receives the feedback or updated model in Canvas and can generate predictions in view-only mode.

To collaborate, the Canvas user and Studio Classic user must be in the same Amazon SageMaker domain. For more information about setting up your domain and users, see the SageMaker Canvas Prerequisites.

Note
Model collaboration is different from Bring your own model to SageMaker Canvas, where you can bring a model that you’ve trained anywhere and import it into Canvas for generating predictions.

Prerequisites

Before a Canvas user and Studio Classic user can collaborate on models, the users' IAM role must have AWS Identity and Access Management (IAM) permissions to share models. If you haven’t already set up permissions, see Grant Users Permissions to Collaborate with Studio Classic.

The Canvas user must also have a Standard build model trained in Canvas and ready to share.

Note
Collaboration does not support Quick build models.

You should also have the user profile name of the Studio Classic user with whom you want to collaborate. The Studio Classic user must be in the same Amazon SageMaker domain as your Canvas user. You can find a user’s profile name by using the following procedure:

2. In the navigation panel, choose Domains.
3. From the list of Domains, choose your domain. This opens the domain details page, where you can find all of the User profiles for the domain.

Keep the user profile name ready for the first step of the following tutorial.
Canvas users: Share a model with Studio Classic users

Within the Canvas application, share your model version with Studio Classic users or request feedback from them. You should use a model version that has been built; you can’t share a model version that is a draft or currently building. You can only share one version per model.

To share your Canvas model with Studio Classic users, use the following procedure.

1. Open the SageMaker Canvas application.
2. From the Models page, select the model that you want to share. You can only share Standard build models.
3. In the header, choose Share.
4. In the Share Model dialog box, do the following:
   a. From the Choose a model version to share dropdown list, select the model version for which you want feedback.
   b. From the SageMaker Studio users dropdown list, select Studio Classic users by their profile names. You can add up to 23 Studio Classic users.
   c. For the Add a note field, you can enter a quick note that accompanies your model when you send it to the Studio Classic users.
   d. Choose Share.
   e. In the Share Model confirmation box that appears, choose Share.

You have now shared your model with the Studio Classic users, and the users receive a notification in Studio Classic that a model has been shared with them.

Studio Classic users: Receive a model in Studio Classic from Canvas users

In Studio Classic, if a model has been shared with you, you receive a notification similar to the following when you open the Studio Classic application.

Choose View shared models to open the Shared models and notebooks page in Studio Classic.
If you miss the notification, you can find the Shared models and notebooks page by doing the following:

1. Open your Amazon SageMaker Studio Classic application.
2. In the side navigation pane, choose the **Home** icon.

3. In the side navigation bar that opens, choose **Models**.

4. In the dropdown list, choose **Shared models** to open the **Shared models and notebooks** page.

On the **Shared models and notebooks** page, select the filter **Shared with me**. You should see the Canvas model that has been shared with you in the list of shared models. Choose **View model** on the shared model, which opens the model details page in Autopilot. The opened model should have a banner at the top that looks similar to the following screenshot.

From this page, you can see the model details, as well as any notes about the model shared with you by the Canvas user. In the Canvas banner at the top, you can choose the following actions:

- Share feedback with the Canvas user.
- Make updates to the shared model and share the updates with the Canvas user.
- Share an alternate version of the model with the Canvas user. Canvas uses **Autopilot** to train multiple versions of the model and select the best version. You can select a different version if you decide that it’s better for your use case.

For more information on the preceding actions, see the following sections.

**Share feedback**

You might want to send a comment or feedback to the Canvas user without making any changes to the model.

To share feedback on the shared model, use the following procedure:

1. On the model details page, choose **Share feedback**.
2. In the **Share feedback** dialog box, add a note in the **Add feedback** field.
3. Choose **Share** to send the feedback to the Canvas user.
After giving feedback, you can view the feedback you sent in the Canvas banner at the top of the model details page. The Canvas user receives the feedback in the Canvas application and can make changes based on your feedback.

**Share an updated model with the Canvas user**

You might want to make changes to the model that the Canvas user shared with you. For example, you might want to use advanced data transformations such as one-hot encoding to improve the accuracy of the model. You can update the model with Amazon SageMaker Data Wrangler and Amazon SageMaker Autopilot in Studio Classic, which are features that help you make data transformations and train your model.

⚠ **Warning**

If you exit the following workflow at any time, your model updates are not saved, and you must restart the workflow.

To update the model and send the updated model to the Canvas user, use the following procedure:

1. **On the model details page, in the Canvas banner, choose **Update model**.**
2. **In the banner’s dropdown list, choose **Update data transformations**.**
3. **The workflow opens your model in Amazon SageMaker Data Wrangler, where you can choose to edit the data transformations used for the model. Make your data transformations in the Data Wrangler interface. For more information about Data Wrangler and the data transformations you can use, see the Data Wrangler documentation.**
4. **After you’ve finished your data transformations, choose **Retrain model** on the Canvas banner to open the Export data and train a model with SageMaker Autopilot page in the Data Wrangler interface.**
5. Verify the fields on the **Export data and train a model with SageMaker Autopilot** page, and then choose **Export and train** to export your data transformations to Amazon SageMaker Autopilot.

6. The workflow opens the **Create an Autopilot experiment** page in Autopilot, where you can create an Autopilot experiment and retrain the model with the updated data transformations. Fill out the fields for each of the **Create an Autopilot experiment** pages.

   For more information about Autopilot and Autopilot experiments, see [Create an experiment](#) in the Autopilot documentation.

7. After you've finished configuring your Autopilot experiment and reviewed the final settings, choose **Create experiment** in the Autopilot interface to begin training the model. The model trains, during which you can choose **Stop training** in the Autopilot interface at any time.

8. After the model has trained, the Canvas banner at the top of the page compares the metrics of the old model with the updated model. The **Best model summary** lists the metrics, such as Recall and Precision, and whether the new model improves the metrics or not. Review the metrics and decide whether you would like to share the updated model or not. For more information about Autopilot metrics, see [Metrics and validation](#).

9. If you decide that you want to share the updated model with the Canvas user, choose **Share** in the banner.

10. In the **Share** dialog box, do the following:

    a. For the **Select a model to share** dropdown list, the best model from your Autopilot experiment should already be selected and marked with a label **Best Candidate**. If the model version that you want to share is not selected, open the dropdown and select the correct version.

    b. For the **Add feedback** field, you can enter a note for the Canvas user.

    c. Choose **Share** to share the updated model and note with the Canvas user.

   After sharing the model, you receive a notification that your model was shared successfully similar to the following screenshot.
You can choose **View shared models** in the banner to return to the **Shared models and notebooks** page. From this page, you can see the updated model that you shared with the Canvas user under the **Shared by me** label.

### Share an alternate model with the Canvas user

When SageMaker Canvas builds a model, Amazon SageMaker Autopilot trains multiple versions of the model and selects the best one. You might decide that an alternate version of the model is better according to your needs. You can share an alternate Autopilot version of the model with the Canvas user instead of making changes to the one they sent. For more information about Autopilot, see the [Autopilot documentation](#).

To share an alternate model, use the following procedure:

1. On the model details page, in the Canvas banner, choose **Update model**.
2. In the banner’s dropdown list, choose **Recommend an alternate Auto ML candidate**.
3. The page for the Autopilot job opens where you can review all of the trained model versions. When you're ready to share an alternate version, in the Canvas banner at the top of the page, choose **Share**.
4. In the **Share** dialog box, do the following:
   a. For the **Select a model to share** dropdown list, the best model from the Autopilot experiment is selected and marked with the label **Best Candidate**. Open the dropdown and select the alternate model version that you want to share.
   b. For the **Add feedback** field, you can enter a note for the Canvas user.
   c. Choose **Share** to share the alternate model version and note with the Canvas user.

After sharing the model, you receive a notification that your alternate model was shared successfully similar to the following screenshot.

![Notification of successfully shared alternate model](image)

You can choose **View shared models** in the banner to return to the **Shared models and notebooks** page. From this page, you can see the updated model that you shared with the Canvas user under the **Shared by me** label.
Canvas users: Receive model updates from a Studio Classic user

When a Studio Classic user shares an updated or alternate model with the Canvas user, the Canvas user receives a notification.

In the Canvas app, the notification looks like the following screenshot.

![Notification Screenshot]

You can choose **View update** to see the updated model, or you can go to the **Models** page in the Canvas application and select the shared model to view it.

**Note**

Canvas users can't edit a model that has been shared with them by a Studio Classic user. Models imported from Studio Classic are view and predict only.

A model on which a Studio Classic user has collaborated looks like the following card on the **Models** page.
Customer Churn Model

- Accuracy: --
- Dataset: --
- Target: Plan
- Problem type: Multiclass
- Received: 7/3/2021 18:11

The model import from Studio Classic can take up to 20 minutes, during which the model shows as Importing.
After importing the model, you can view its metrics and generate predictions with it.

The following screenshot shows the **Analyze** tab, where you can evaluate the model accuracy and metrics. For more information, see [Evaluate Your Model's Performance in Amazon SageMaker Canvas](#).

The following screenshot shows the **Predict** tab, where you can generate predictions with the model. For more information on generating predictions in Canvas, see [Make predictions for your data](#).

Use custom models
On both the **Analyze** and **Predict** tabs, you can see the **Shared History** panel, which shows you the model versions and comments shared with you by Studio Classic users.

### Bring your own model to SageMaker Canvas

**Note**

The functionality described on this page only applies to Amazon SageMaker Studio Classic. Currently, you can only share models to Canvas (or view shared Canvas models) in Studio Classic. If you’re currently using the latest version of Studio, you must run Studio Classic from within the latest version of Studio to share models to Canvas or view models shared from Canvas. For more information about accessing Studio Classic, see the [Studio Classic documentation](#).

Business analysts can benefit from ML models already built by data scientists to solve business problems instead of creating a new model in Amazon SageMaker Canvas. However, it might be difficult to use these models outside the environments in which they are built due to technical requirements, rigidity of tools, and manual processes to import models. This often forces users to rebuild ML models, resulting in the duplication of effort and additional time and resources.
SageMaker Canvas removes these limitations so you can generate predictions in Canvas with models that you've trained anywhere. You can register ML models in SageMaker Model Registry, which is a metadata store for ML models, and import them into SageMaker Canvas. Additionally, you can generate predictions with models that data scientists have trained in Amazon SageMaker Autopilot or SageMaker JumpStart. Canvas users can then analyze and generate predictions from any model that has been shared with them.

After you’ve satisfied the Prerequisites, see the following sections for instructions on how to bring your own models into Canvas and generate predictions. The workflow begins in Studio Classic, where a Studio Classic user shares a model with a Canvas user. Then, the Canvas user signs in to their Canvas app to receive the shared model and generate predictions with it.

### Note

You can share models trained with tabular, text, and image data to Canvas. You can't share time series models. Also, Canvas bring your own model (BYOM) only supports CPU-based models (or models that use CPU instances to make predictions).

### Prerequisites

To bring your model into SageMaker Canvas, complete the following prerequisites:

- You must have a Amazon SageMaker Studio Classic user who has onboarded to Amazon SageMaker domain. The Studio Classic user must be in the same domain as the Canvas user. Model sharing occurs when a Studio Classic user shares a model with a Canvas user from within Studio Classic. If you don’t already have a Studio Classic user set up, see the Studio Classic documentation and Onboard to Amazon SageMaker domain.

- You must have a trained model from SageMaker Autopilot, SageMaker JumpStart, or SageMaker Model Registry. For any model that you've built outside of SageMaker, you must register your model in Model Registry before importing it into Canvas. For more information, see the Model Registry documentation.

- The Canvas user with whom you want to share your model must have permission to access the Amazon S3 bucket in which you store your datasets and model artifacts. For instructions on how admins can give Canvas users the permissions they need, see Grant Users Permissions to Collaborate with Studio Classic.
You should also have the user profile name of the Canvas user with whom you want to collaborate. The Canvas user must be in the same Amazon SageMaker domain as your Studio Classic user. You can find a user's profile name by using the following procedure:

2. In the navigation panel, choose Domains.
3. From the list of Domains, choose your domain. This opens the domain details page, where you can find all of the User profiles for the domain.

Keep the user profile name ready for the first step of the following tutorial.

If your SageMaker Canvas app is running in a private customer VPC, any Autopilot models shared from Studio Classic must use Autopilot HPO mode to support generating predictions in Canvas. For more information about HPO mode, see Training modes and algorithm support in the Autopilot documentation.

**Note**

If you want feedback from data scientists on a model built inside Canvas, see Collaborate with data scientists, where a Canvas user shares a model with a Studio Classic user, and the Studio Classic user shares feedback or model updates.

**Studio Classic users: Share a model to SageMaker Canvas**

You should have a model trained with tabular data that you're ready to share with Canvas users. See the following sections for information on how to share your models from features within Studio Classic.

**Autopilot**

You can share a model to Canvas from Amazon SageMaker Autopilot in Studio Classic. Autopilot is a feature that enables you to train and and deploy your models in SageMaker.

You need to have a Studio Classic user and a trained model ready to share from Autopilot. For more information on how to set up Studio Classic, see the Studio Classic documentation. For more information about Autopilot, see the Autopilot documentation.

To share a model from Autopilot to Canvas, use the following procedure.
1. Open your Amazon SageMaker Studio Classic application.
2. In the side navigation pane, choose the **Home** icon
   ![Home icon]
3. In the side navigation bar of Studio Classic, choose **AutoML** to open Autopilot.
4. On the **Autopilot** page, select the Autopilot model that you want to share with the Canvas user. You can only share one model at a time.
5. From the Autopilot job details page, in the **Models** tab, select the model version that you want to share.
6. Choose **Share**.
7. In the **Share model** dialog box, do the following:
   a. For the **Add Canvas users** field, enter the Canvas user’s profile name. You can enter up to 23 Canvas users. If a user profile you specify doesn’t have a Canvas app associated with it, you can’t enter the profile name.
   b. For the **Add a note** field, add a description or note for the Canvas user when they receive the model.
   c. Choose **Share** to share the model.

You have now shared the model with the Canvas user.

**JumpStart**

You can share a model to Canvas from SageMaker JumpStart in Studio Classic. With JumpStart, you can access and tune pretrained models before deploying them.

You need to have a Studio Classic user and a successfully completed training job in JumpStart. For more information about how to set up Studio Classic, see the [Studio Classic documentation](#). For more information about JumpStart, see the [JumpStart documentation](#).

To share a model from JumpStart to Canvas, use the following procedure.

1. Open your Amazon SageMaker Studio Classic application.
2. In the side navigation pane, choose the **Home** icon
   ![Home icon]
3. In the side navigation bar that opens, choose **SageMaker JumpStart**.
4. Choose **Launched JumpStart assets** to open the page that lists your JumpStart training jobs, models, and endpoints.

5. Choose the **Training jobs** tab to view the list of your model training jobs.

6. From the **Training jobs** list, select the training job that you want to share with the Canvas user. You can only share one job at a time. This opens the training job details page.

7. In the header for the training job, choose **Share**, and select **Share to Canvas**.

8. In the **Share to Canvas** dialog box, do the following:

   a. For the **Add Canvas users to share** field, enter the Canvas user’s profile name. You can enter up to 23 Canvas users. If a user profile you specify doesn’t have a Canvas app associated with it, you can't enter the profile name.

   b. For the **Add a note** field, add a description or note for the Canvas user when they receive the model.

   c. Choose **Share** to share the model.

You have now shared the model with the Canvas user.

**Model Registry**

You can share a model to Canvas from SageMaker Model Registry in Studio Classic. With Model Registry, you can register models that you bring from outside of SageMaker and integrate them with your ML pipelines.

You need to have a Studio Classic user and a model version saved in the Model Registry. For more information about how to set up Studio Classic, see the [Studio Classic documentation](#). If you don’t have a model version in the Model Registry, create a model group and register a version to it. For more information about Model Registry, see the [Model Registry documentation](#).

To share a model version from Model Registry to Canvas, use the following procedure.

1. Open your Amazon SageMaker Studio Classic application.
2. In the side navigation pane, choose the **Home** icon.

3. In the side navigation bar that opens, choose **Models**.

4. Select **Model Registry** from the dropdown list to open the Model Registry page and show all of the model groups registered in your account.

5. Choose the model group that has the model version that you want to share.

6. You can share a model version either from the model group page or the model version page.

   - To share a model version from the model group page, complete the following steps:
     1. Choose **Versions**, and check the box next to the model version you want to share with the Canvas user. You can only share one model version at a time.
     2. In the **Actions** dropdown menu, choose **Share model artifacts**.

   - To share a model version from the model version page, complete the following steps:
     1. Choose **Versions**, and select the name of the model version you want to share with the Canvas user. You can only share one model version at a time.
     2. In the **Actions** dropdown menu, choose **Share model artifacts**.

7. In the **Share model** dialog box, do the following:

   a. For the **Add Canvas users to share** field, enter the Canvas user’s profile name. You can enter up to 23 Canvas users. If a user profile you specify doesn’t have a Canvas app associated with it, you can't enter the profile name.

   b. For **Add model details**, do the following:

      i. For the **Training dataset** field, enter the Amazon S3 path for your training dataset.

      ii. For the **Validation dataset** field, enter the Amazon S3 path for your validation dataset.

      iii. For **Target column**, either select **Use the first column** if the first column in your dataset is the target, or select **Specify the target column name** to set the target as a different column in your dataset.

      iv. For **Column headers**, select one of the following options:

         A. Select **Use the first row** if the first row of your dataset contains the column headers.
B. Select **Specify a different dataset in S3 for column headers** if you have a file stored in Amazon S3 containing headers that can be mapped to your dataset. The headers file must have the same number of columns as your dataset.

C. Select **Automatically generate** if you don’t already have column headers and would like SageMaker to generate generic column names for your dataset.

v. From the **Problem type** dropdown list, select your model type.

vi. If you selected the **Binary classification** or **Multi-class** problem types, the **Configure model outputs** option appears.

If you already have a file stored in Amazon S3 that maps default target column class names to your desired class names, then turn on **Model output names** and enter the Amazon S3 path to the mapping file. If you don’t have a mapping file, then turn off **Model output names** and manually enter the **Number of model outputs** (the number of target column classes in your data). Then, enter your desired class names to replace the default class names.

c. (Optional) For the **Add a note** field, add a description or note for the Canvas user when they receive the model.

d. Choose **Share** to share the model version.

You have now shared the model with the Canvas user.

**Shared models and notebooks**

On the **Shared models and notebooks** page in Amazon SageMaker Studio Classic, you can view the models that you've shared and that have been shared with you. This page gives you a central place to view and manage all of your models in Studio Classic.

You need to have a Studio Classic user and a model ready to share from Autopilot, JumpStart, or Model Registry. For more information on how to set up Studio Classic, see the [Studio Classic documentation](#). For more information about the **Shared models and notebooks** page, see the [Shared models and notebooks documentation](#).

The following example walks you through sharing an Amazon SageMaker Autopilot model, but you can use the sharing feature on the **Shared models and notebooks** page to share models from any of the other features in the previous sections, such as JumpStart and Model Registry.

To share an Autopilot model from the **Shared models and notebooks** page, use the following procedure.
1. Open your Amazon SageMaker Studio Classic application.
2. In the side navigation pane, choose the **Home** icon.
3. In the side navigation bar of Studio Classic, choose **Models**.
4. In the dropdown list, choose **Shared models** to open the **Shared models and notebooks** page.
5. Choose the filter icon, and in the **Shared from** dropdown list, choose **Autopilot**.
6. Select the Autopilot model from the list that you want to share with the Canvas user. You can only share one model at a time. Alternatively, you can select the model to open the model details page.
7. From either the Autopilot jobs page or the model details page, choose **Share**.
8. In the **Share model** dialog box, do the following:
   a. For the **Add Canvas users to share** field, enter the Canvas user's profile name. You can enter up to 23 Canvas users. If a user profile you specify doesn't have a Canvas app associated with it, you can't enter the profile name.
   b. For the **Add a note** field, add a description or note for the Canvas user when they receive the model.
   c. Choose **Share** to share the model.

You have now shared the model with the Canvas user.

After you share the model, you receive a notification popup in Studio Classic similar to the following screenshot.

---

You shared **pyt-cpu-models-16** successfully!

---

You can choose **View model** to open the **Shared models and notebooks** page in Studio Classic. You can also view your shared models at any time from the **Shared models and notebooks** page.

From this page, you can see the models that you've shared with the Canvas user under the **Shared by me** label, as shown in the following screenshot.
Models that you've shared to Canvas have text on the card similar to the following example:
Shared to: 12 Canvas users.

**Canvas users: Receive a shared model in SageMaker Canvas**

When a Studio Classic user shares a model with a Canvas user, you receive a notification within the Canvas application that a Studio Classic user has shared a model with you.
In the Canvas application, the notification is similar to the following screenshot.

You can choose **View update** to see the shared model, or you can go to the **Models** page in the Canvas application to discover all of the models that have been shared with you.

**Note**

Canvas users can’t edit a model that has been shared with them by a Studio Classic user. Models imported from Studio Classic are view and predict only.

A model that has been shared by a Studio Classic user looks like the following card on the **Models** page. This is different from **Collaborate with data scientists**, where a Canvas user shares a model and a Studio Classic user shares updates or feedback with the Canvas user.
Customer Churn Model

Accuracy — —

Dataset --

Target Plan

Problem type Multiclass

Received 7/3/2021 18:11

View
The model import from Studio Classic can take up to 20 minutes, during which the model shows as **Importing**.

After importing the model, you can view its metrics and generate predictions with it. SageMaker Canvas uses Amazon SageMaker Serverless Inference resources to generate model analysis and predictions for shared models. You might see costs associated with Serverless Inference in your AWS account.

The following screenshot shows the **Analyze** tab in the Canvas application for a shared model, where you can evaluate the model accuracy and metrics. For more information, see [Evaluate Your Model's Performance in Amazon SageMaker Canvas](#).

![Canvas Analyze Tab](image)

**Model status**

43052.733

The model often predicts a value that is within +/- 43052.733 of the actual value for median_house_value.

The following screenshot shows the **Predict** tab, where you can generate predictions with the model. For more information on generating predictions in Canvas, see [Make predictions for your data](#).
On both the **Analyze** and **Predict** tabs, you can see the **Shared History** panel, which shows you the model versions and comments shared with you by Studio Classic users.

### Logging out of Amazon SageMaker Canvas

After completing your work in Amazon SageMaker Canvas, you can log out or configure your application to automatically terminate the *workspace instance*. A workspace instance is dedicated for your use every time you launch a Canvas application, and you are billed for as long as the instance runs. Logging out or terminating the workspace instance stops the workspace instance billing. For more information, see [SageMaker Pricing](https://aws.amazon.com/sagemaker/pricing).

The following sections describe how to log out of your Canvas application and how to configure your application to automatically shut down on a schedule.

### Log out of Canvas

When you log out of Canvas, your models and datasets aren't affected, but SageMaker Canvas cancels any **Quick build** tasks. If you log out of SageMaker Canvas while running a **Quick build**, your build might be interrupted until you relaunch the application. When you relaunch, SageMaker Canvas automatically restarts the build. **Standard builds** continue even if you log out.
To log out, choose the Log out button ( ) on the left panel of the SageMaker Canvas application.

You can also log out from the SageMaker Canvas application by closing your browser tab and then deleting the application in the console.

After you log out, SageMaker Canvas tells you to relaunch in a different tab. Logging in takes between 3 minutes and 8 minutes. If you have an administrator who set up SageMaker Canvas for you, use the instructions they gave you to log back in. If don’t have an administrator, see the procedure for accessing SageMaker Canvas in Prerequisites for setting up Amazon SageMaker Canvas.

Automatically shut down Canvas

If you’re a Canvas administrator, you might want to regularly shut down applications to reduce costs. You can either create a schedule to shut down active Canvas applications, or you can create an automation to shut down Canvas applications as soon as they’re idle (meaning the user hasn’t been active for 2 hours).

You can create these solutions using AWS Lambda functions that call the DeleteApp API and delete Canvas applications given certain conditions. For more information about these solutions and access to AWS CloudFormation templates that you can use, see the blog Optimizing costs for Amazon SageMaker Canvas with automatic shutdown of idle apps.

Note

You might experience missing Amazon CloudWatch metrics if there was an error when setting up your idle shut down schedule or a CloudWatch error. We recommend that you add a CloudWatch alarm that monitors for missing metrics. If you encounter this issue, reach out to AWS Support for help.

Limitations and troubleshooting

The following section outlines troubleshooting help and limitations that apply when using Amazon SageMaker Canvas. You can use these this topic to help troubleshoot any issues you encounter.
Troubleshooting issues with granting permissions through the SageMaker console

If you’re having trouble granting Canvas base permissions or Ready-to-use models permissions to your user, your user might have an AWS IAM execution role with more than one trust relationship to other AWS services. A trust relationship is a policy attached to your role that defines which principals (users, roles, accounts, or services) can assume the role. For example, you might encounter an issue granting additional Canvas permissions to your user if their execution role has a trust relationship to both Amazon SageMaker and Amazon Forecast.

You can fix this problem by choosing one of the following options.

1. **Remove all but one trusted service from the role.**

   This solution requires you to edit the trust relationship for your user profile’s IAM role and remove all AWS services except SageMaker.

   To edit the trust relationship for your IAM execution role, do the following:

   2. In the navigation pane of the IAM console, choose Roles. The console displays the roles for your account.
   3. Choose the name of the role that you want to modify, and select the Trust relationships tab on the details page.
   4. Choose Edit trust policy.
   5. In the **Edit trust policy editor**, paste the following, and then choose Update Policy.

   ```json
   {
     "Version": "2012-10-17",
     "Statement": [
       {
         "Effect": "Allow",
         "Principal": {
           "Service": [
             "sagemaker.amazonaws.com"
           ]
         },
         "Action": "sts:AssumeRole"
       }
     ]
   }
   ```
You can also update this policy document using the IAM CLI. For more information, see `update-trust` in the IAM Command Line Reference.

You can now retry granting the Canvas base permissions or the Ready-to-use models permissions to your user.

2. **Use a different role with one or fewer trusted services.**

This solution requires you to specify a different IAM role for your user profile. Use this option if you already have an IAM role that you can substitute.

To specify a different execution role for your user, do the following:

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain that you want to view a list of user profiles for.
5. On the domain details page, choose the User profiles tab.
6. Choose the user whose permissions you want to edit. On the User details page, choose Edit.
7. On the General settings page, choose the Execution role dropdown list and select the role that you want to use.
8. Choose Submit to save your changes to the user profile.

Your user should now be using an execution role with only one trusted service (SageMaker).

You can retry granting the Canvas base permissions or the Ready-to-use models permissions to your user.

3. **Manually attach the AWS managed policy to the execution role instead of using the toggle in the SageMaker domain settings.**

Instead of using the toggle in the domain or user profile settings, you can manually attach the AWS managed policies that grant a user the correct permissions.

To grant a user Canvas base permissions, attach the AmazonSageMakerCanvasFullAccess policy. To grant a user Ready-to-use models permissions, attach the AmazonSageMakerCanvasAIServicesAccess policy.

Use the following procedure to attach an AWS managed policy to your role:
2. Choose **Roles**.
3. In the search box, search for the user's IAM role by name and select it.
4. On the page for the user's role, under **Permissions**, choose **Add permissions**.
5. From the dropdown menu, choose **Attach policies**.
6. Search for and select the policy or policies that you want to attach to the user’s execution role:
   a. To grant the Canvas base permissions, search for and select the **AmazonSageMakerCanvasFullAccess** policy.
   b. To grant the Ready-to-use models permissions, search for and select the **AmazonSageMakerCanvasAIServicesAccess** policy.
7. Choose **Add permissions** to attach the policy to the role.

After attaching an AWS managed policy to the user’s role through the IAM console, your user should now have the Canvas base permissions or Ready-to-use models permissions.

**Limitations for collaboration**

The following general limitations apply when you are collaborating with data scientists in Amazon SageMaker Studio Classic.

- You can only share successfully trained models from Canvas to Studio Classic. Similarly, you can only share models that have been successfully trained in Studio Classic back to Canvas.
- You can't share **Quick build** models from Canvas to Studio Classic. You can only share **Standard build** models.
- You can only share one version of a **Standard build** model trained in Canvas. You can train additional versions of your model within Canvas, but you can't share them to Studio Classic.
- From Studio Classic, you can only share feedback or share an updated model with Canvas. You can't perform both actions at the same time.
- The length limitation for comments shared from Studio Classic to Canvas and Canvas to Studio Classic is 1024 characters.
- You can only share your Canvas or Studio Classic models with a different user profile. You can't share models between Canvas and Studio Classic within your own user profile.
- You can't share from a Canvas user to a Canvas user, or from a Studio Classic user to a Studio Classic user.
There are also limitations that apply depending on the type of model you want to share. See the following sections for limitations on time series forecasting models and numeric and categorical prediction models.

**Limitations for collaborating on time series forecasting models**

The following limitations apply when you are collaborating on time series forecasting models between Canvas and Studio Classic.

- You can’t make predictions with time series forecasting models in Studio Classic through an automated Share button. However, you can create a Jupyter notebook and write your own code.
- For time series forecasting models, you can’t change the model recipe or data transformations in Studio Classic. You can only make the following updates to time series forecasting models in Studio Classic:
  - You can update the length of the forecast horizon.
  - You can update the item’s metadata field, which groups your data by a certain column.
  - You can update other dimension fields, such as specifying a holiday schedule.

**Limitations for collaborating on numeric and categorical prediction models**

The following limitations apply when you are collaborating on numeric and categorical prediction model types between Canvas and Studio Classic.

- When updating or training models in Studio Classic, if you close the tab with the collaboration banner at the top, it ends the share model workflow and you lose your progress. In that case, you must restart the share model workflow from the Shared With Me section on the Shared Models page. For more information, see Collaborate with data scientists.
- When updating models in Studio Classic, you can’t change the target column if you want to share the model updates back to Canvas. If you want to change the target column and re-train the model, train the model and then use the Share button to share to Canvas. For more information about sharing a new model to Canvas, see Bring your own model to SageMaker Canvas.
- When updating models in the Amazon SageMaker Data Wrangler Recipe interface in Studio Classic, there are limits to which changes a Studio Classic user can apply that Canvas supports:
  - You can only share a model to Canvas that has been trained from the last node in a Data Wrangler linear data flow.
  - Only transformation nodes are supported.
• You can’t perform operations on the **Target** column.

• You can’t update the data type of columns.

• You can’t update the data source or add a new data source.

• When sharing an alternative candidate to Canvas from the Studio Classic Autopilot page, you can’t select the model from the leaderboard. You must choose the shared model from the banner and then select an alternative from the list. For more information, see [Share an alternate model with the Canvas user](#) in the Canvas documentation.

• Only models that are compatible with [SageMaker Neo](#) can be shared back to Canvas successfully. Compatible models are Autopilot models that use XGBoost or MLP algorithms. Incompatible models include Autopilot models that use the linear learner algorithm.

• For custom formula transforms using Spark SQL, Canvas only supports Unary operations, Aggregate functions, the String concatenation operation and the Power operation. Other operations are not supported.

## Limitations for bring your own model (BYOM)

The following general limitations apply when you want to [bring your own model](#) to SageMaker Canvas.

• When a model is shared from Studio Classic to Canvas, the Canvas user cannot update or view details on the dataset that was used to build the model.

• When a Canvas user wants to run a single prediction on an imported model, there are no data type restrictions when updating column values. You must manually make sure that when you update values for single predictions, you match the data type of the existing values.

• When a Canvas user wants to run batch predictions on an imported model, Canvas assumes that you (the Canvas user) know what the expected input dataset should look like. You should have a dataset with columns and data types that match the dataset that was used to train the model. If not, consult with the user who shared the model with you and import a dataset that you can use for running batch predictions.

• The Canvas application internally uses a [serverless endpoint](#) to run predictions and generate model metrics. The model shared to Canvas must be compatible with serverless endpoints:

  • The maximum memory size is 6144 MB.

  • When configuring the inference input response keys in your container, use the following configuration:
You can choose either a SageMaker-provided inference container or bring your own image inference container to be used for endpoint. SageMaker provides containers for its built-in algorithms and prebuilt Docker images for some of the most common machine learning frameworks. If you are bringing your own container, you must modify it to work with SageMaker. For more information about bringing your own container, see Adapting Your Own Inference Container.

The Feature exclusions for serverless endpoints also apply.

To share a model from Studio Classic to Canvas successfully, Canvas accepts model inference outputs in the format below:

TEXT/CSV

Regression: The model inference response should be a byte string where each of the output predictions are separated by \n:

```
b'-0.0007884334772825241\n0.015136942267417908\n0.050063662230968475\n0.02891816757619381'
```

Classification: The model inference response should be a byte string where each of predicted_label, predicted_probability, probabilities, and labels are separated by \n. The following example is for binary classification:

```
b'no,0.9967488050460815,[0.9967488050460815, 0.003251201706007123],['no', 'yes']
no,0.9999420642857283,[0.9999420642857283, 5.793538366560824e-05],['no', 'yes']
no,0.9999846816062927,[0.9999846816062927, 1.5326571883633733e-05],['no', 'yes']
no,0.9999727606773376,[0.9999727606773376, 2.7267418772680685e-05],['no', 'yes']
```

The following example is for multi-class classification:

```
b'Iris-setosa,1.0,[1.0, 0.0, 0.0],['Iris-setosa', 'Iris-versicolor', 'Iris-virginica']
'Iris-setosa,1.0,[1.0, 0.0, 0.0],['Iris-setosa', 'Iris-versicolor', 'Iris-virginica']
'Iris-setosa,1.0,[1.0, 0.0, 0.0],['Iris-setosa', 'Iris-versicolor', 'Iris-virginica']
'Iris-setosa,1.0,[1.0, 0.0, 0.0],['Iris-setosa', 'Iris-versicolor', 'Iris-virginica']
```
APPLICATION/JSON

- Regression: The model inference response should be a JSON string which contains the prediction key, and its value should be the list of output predictions:

```javascript
let response = {
    "predictions": [
        // First instance prediction.
        1.75
        // Second instance prediction.
        3.25
    ]
}
```

- Classification: The model inference response should be a JSON string which contains the probabilities key, and its value should be the list of probabilities.

The following example is for binary classification:

```javascript
let response = {
    "probabilities": [
        // First instance prediction.
        [0.9, 0.1]
        // Second instance prediction.
        [0.2, 0.8]
    ]
}
```

The following example is for multi-class classification:

```javascript
let response = {
    "probabilities": [
        // First instance prediction.
        [0.7, 0.2, 0.1]
        // Second instance prediction.
        [0.2, 0.5, 0.3]
    ]
}
```
There are also limitations that apply depending on the type of model you want to bring:

**Bring your own model from SageMaker JumpStart**

Review the following information and limits when sharing a SageMaker JumpStart model with Canvas.

- The following are the supported algorithms for which you can import models into Canvas. For more details, see the [SageMaker JumpStart documentation](SageMaker JumpStart documentation).
  - Tabular classification: LightGBM, CatBoost, XGBoost, AutoGluon-Tabular, TabTransformer, Linear Learner
  - Tabular regression: LightGBM, CatBoost, XGBoost, AutoGluon-Tabular, TabTransformer, Linear Learner
- In SageMaker JumpStart, the **Share** button is only turned on if the model is ready to share to Canvas. If your trained model does not have a **Share** to SageMaker Canvas button, your model is not supported for BYOM.
- You must provide training and validation datasets when training the SageMaker JumpStart model. The datasets should be stored in Amazon S3, and your Studio Classic and Canvas users' execution role must have access to the Amazon S3 location. You can use the same Amazon S3 URIs to share the training and validation datasets with Canvas, or you can share different datasets with the same data schema.

Your training or validation data file should look like the following (in CSV format). You should index your files with the first column as the target.

<table>
<thead>
<tr>
<th>3</th>
<th>1</th>
<th>22</th>
<th>1</th>
<th>1</th>
<th>0</th>
<th>4</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>38</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>67</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>67</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>40</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>56</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>6</td>
</tr>
</tbody>
</table>

- By default, SageMaker JumpStart uses the first column of the training and validation datasets as the target when training a model. The target column (or by default, the first column) of the datasets is shared to Canvas.
- You must provide the column headers of the training and validation datasets when training the SageMaker JumpStart model. By default, SageMaker JumpStart only accepts datasets without column headers, so you must add the column headers as a file while training your model. The
Amazon S3 URI for the column headers file is shared to Canvas as well. Your column headers file should look like the following example (in CSV format). The first column should be the target.

```
Segmentation EverMarried Age Graduated WorkExperience SpendingScore FamilySize Var1
```

- The training job in SageMaker JumpStart must be Complete before you can share with Canvas.
- For classification problems (or categorical prediction in Canvas), original class names need to be provided in the **Configure model output** section when sharing to Canvas. The order of the class names must match the indexing used in the model. Your mapping relation file should look like the following example in CSV format, where index 0 (the first index) is mapped to the class name A:

```
A B C D
```

When the Canvas user views the model metrics in the Canvas application, they can only see the index of each class (0, 1, 2). However, the user can see the class names when viewing the results for a single prediction.

**Bring your own model from Autopilot**

Review the following information and limits when sharing a model from Autopilot to Canvas.

- You can only share models to Canvas that you’ve successfully trained from an AutoML job with **Ensembling**, **HPO**, or **Auto** mode (for **Auto** mode, Autopilot chooses **Ensembling** or **HPO** mode based on the training dataset size). The currently supported Autopilot problem types are Regression, Multi-class classification, Binary classification.
- For each Autopilot job, you can choose any model (the **Best model** or any other candidates) to share to Canvas one at a time. You only need to choose the **Share model** button and then specify the Canvas users with whom you’d like to share the model and a note.
- AutoGluon-Tabular models that use Data Wrangler transformers for inference cannot be shared to Canvas. This is because Data Wrangler transformers cause the model to use more than one container.
- HPO models that aren’t **compatible with SageMaker Neo** can’t be shared to Canvas successfully. Compatible models are Autopilot models that use XGBoost or MLP algorithms. Incompatible models include Autopilot models that use the linear learner algorithm.
Bring your own model from Model Registry

Review the following information and limits when sharing a model from Model Registry to Canvas.

- Unlike the **Share** button provided by SageMaker JumpStart, Model Registry doesn't provide model validation, so it's possible that a registered model shared successfully from Studio Classic can fail while importing to Canvas due to model incompatibility. Review the following tips before sharing to Canvas from Model Registry:

  - Use a single inference container for your model. You can register models with **multiple containers** within the **AdditionalInference Specifications** field, but Canvas is only optimized for one inference container per model. For example, when you use a inference pipeline and register multiple containers in the **AdditionalInference Specifications** field with multiple data preprocessing containers and an inference container, by default the first container is selected for model inference in Canvas. Evaluate if this works for your use case if you're using machine learning pipelines.

  - Use a SageMaker **built-in tabular algorithm** with compatible inference formats. Tested sample algorithms with compatible inference outputs are Autogluon-Tabular, CatBoost, LightGBM, TabTransformer and XGBoost. Algorithms like Factorization Machines don't accept CSV as file input, and the inference output formats for algorithms like Linear Learner and K-NN are not supported by Canvas.

  - You can also bring your own image container and share to Canvas, or modify pre-built SageMaker containers.

    - If you are bringing your own container, you must modify it to work with SageMaker. For more information about bringing your own container, see **Adapting Your Own Inference Container**.

    - For detailed formatting for your inference output formats, see **Limitations for bring your own model (BYOM)**.

- When registering your model in a **model package group**, remember to provide the following attributes with your inference container:

  - **Environment**:

    ```json
    "{"SAGEMAKER_CONTAINER_LOG_LEVEL": "20", "SAGEMAKER_PROGRAM": "inference.py ", "SAGEMAKER_REGION": "us-west-2", "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/ model/code"}"
    ```

  - **Image**:
Amazon SageMaker

ModelDataUrl

```
"s3://sagemaker-us-west-2-<account-id>/model-regression-abalone-2022-10-14-23-02-45/model.tar.gz"
```

- **ModelDataUrl**

```
"<account-id>.dkr.ecr.us-west-2.amazonaws.com/sagemaker-xgboost:1.3-1"
```

- You must provide training and validation datasets when sharing the model from Model Registry to Canvas. The datasets should be stored in Amazon S3, and the Studio Classic and Canvas users' execution role must have access to the Amazon S3 location. You can use the same Amazon S3 URIs to share the training and validation datasets with Canvas, or you can share different datasets with the same data schema. The datasets must have the exact input formatting that feeds your model's inference container.

- You must provide the target column to Canvas, or the first column of your training/validation dataset is used by default.

- In the **Add model details** section when sharing to Canvas, you can provide the first row your training and validation datasets as the headers, or you can specify the headers as a different file.

- For classification problems (or categorical prediction in Canvas), original class names need to be provided when sharing to SageMaker Canvas through the **Configure model outputs** option. The order of the class names must match the indexing used with the shared model. The mapping can be either a CSV file in Amazon S3, or you can manually input the class names.

### Manage billing and cost in SageMaker Canvas

To track the costs associated with your SageMaker Canvas application, you can use the AWS Billing and Cost Management service. Billing and Cost Management provides tools to help you gather information related to your cost and usage, analyze your cost drivers and usage trends, and take action to budget your spending. For more information, see [What is AWS Billing and Cost Management?](#)

Billing in SageMaker Canvas consists of the following components:

- **Workspace instance charges** – You are charged for the number of hours that you are logged in to or using SageMaker Canvas. We recommend that you log out or create a schedule to shut down any Canvas applications that you're not actively using to reduce costs. For more information, see [Logging out of Amazon SageMaker Canvas](#).
• AWS service charges – You are charged for building and making predictions with custom models, or for making predictions with Ready-to-use models:

• Training charges – For all model types, you are charged based on your resource usage while the model builds. These resources include any compute instances that Canvas spins up. You may see these charges on your account as Hosting, Training, Processing, or Batch Transform jobs.

• Prediction charges – You are charged for the resources used to generate predictions, depending on the type of custom model that you built or the type of Ready-to-use model you used.

The Ready-to-use models in Canvas leverage other AWS services to generate predictions. When you use a Ready-to-use model, you are charged by the respective service, and their pricing conditions apply:

• For sentiment analysis, entities extraction, language detection, and personal information detection, you’re charged with Amazon Comprehend pricing.

• For object detection in images and text detection in images, you’re charged with Amazon Rekognition pricing.

• For expense analysis, identity document analysis, and document analysis, you’re charged with Amazon Textract pricing.

For more information, see SageMaker Canvas pricing.

To help you track your costs in Billing and Cost Management, you can assign custom tags to your SageMaker Canvas app and users. You can track the costs your apps incur, and by tagging individual user profiles, you can track costs based on the user profile. For more information about tags, see Using Cost Allocation Tags.

You can add tags to your SageMaker Canvas app and users by doing the following:

• If you are setting up your Amazon SageMaker domain and SageMaker Canvas for the first time, follow the Getting Started instructions and add tags when creating your domain or users. You can add tags either through the General settings in the domain console setup, or through the APIs (CreateDomain or CreateUserProfile). SageMaker adds the tags specified in your domain or UserProfile to any SageMaker Canvas apps or users you create after you create the domain.

• If you want to add tags to apps in an existing domain, you must add tags to either the domain or the UserProfile. You can adds tags through either the console or the AddTags API. If you add tags
through the console, then you must delete and relaunch your SageMaker Canvas app in order for the tags to propagate to the app. If you use the API, the tags are added directly to the app. For more information about deleting and relaunching a SageMaker Canvas app, see Manage apps.

After you add tags to your domain, it might take up to 24 hours for the tags to appear in the AWS Billing and Cost Management console for activation. After they appear in the console, it takes another 24 hours for the tags to activate.

On the Cost explorer page, you can group and filter your costs by tags and usage types to separate your Workspace instance charges from your Training charges. The charges for each are listed as the following:

- **Workspace instance charges**: Charges show up under the usage type REGION-Canvas:Session-Hrs (Hrs).
- **Training charges**: Charges show up under the usage types for SageMaker Hosting, Training, Processing, or Batch Transform jobs.

**Amazon SageMaker geospatial capabilities**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. If prior to November 30, 2023 you created a Amazon SageMaker domain, Studio Classic remains the default experience. domains created after November 30, 2023 default to the new Studio experience. Amazon SageMaker geospatial features and resources are only available in Studio Classic. To learn more about setting up a domain and getting started with Studio, see Getting started with Amazon SageMaker geospatial.

Amazon SageMaker geospatial capabilities makes it easier for data scientists and machine learning (ML) engineers to build, train, and deploy ML models faster using geospatial data. You have access to open-source and third-party data, processing, and visualization tools to make it more efficient to prepare geospatial data for ML. You can increase your productivity by using purpose-built algorithms and pre-trained ML models to speed up model building and training, and use built-in
visualization tools to explore prediction outputs on an interactive map and then collaborate across teams on insights and results.

Note

Currently, SageMaker geospatial capabilities are only supported in the US West (Oregon) Region.
If you don't see the SageMaker geospatial UI available in your current Studio Classic instance check to make sure you are currently in the US West (Oregon) Region.

Why use SageMaker geospatial capabilities?

You can use SageMaker geospatial capabilities to make predictions on geospatial data faster than do-it-yourself solutions. SageMaker geospatial capabilities make it easier to access geospatial data from your existing customer data lakes, open-source datasets, and other SageMaker geospatial data providers. SageMaker geospatial capabilities minimize the need for building custom infrastructure and data preprocessing functions by offering purpose-built algorithms for efficient data preparation, model training, and inference. You can also create and share custom visualizations and data with your company from Amazon SageMaker Studio Classic. SageMaker geospatial capabilities offer pre-trained models for common uses in agriculture, real estate, insurance, and financial services.

How can I use SageMaker geospatial capabilities?

You can use SageMaker geospatial capabilities in two ways.

- Through the SageMaker geospatial UI, as a part of Amazon SageMaker Studio Classic UI.
- Through a Studio Classic notebook instance that uses the Geospatial 1.0 image.

SageMaker has the following geospatial capabilities

- Use a purpose built SageMaker geospatial image that supports both CPU and GPU-based notebook instances, and also includes commonly used open-source libraries found in geospatial machine learning workflows.
- Use the Amazon SageMaker Processing and the SageMaker geospatial container to run large-scale workloads with your own datasets, including soil, weather, climate, LiDAR, and commercial aerial and satellite imagery.
• Run an Earth Observation job for raster data processing.
• Run a Vector Enrichment job to convert latitude and longitude into human readable addresses, and match noisy GPS traces to specific roads.
• Use built-in visualization tools right in Studio Classic to interactively view geospatial data or model predictions on a map.

You can also use data from a collection of geospatial data providers. Currently, the data collections available include:

• USGS Landsat
• Sentinel-1
• Sentinel-2
• Copernicus DEM
• National Agriculture Imagery Program

Are you a first-time user of SageMaker geospatial?

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. New domains created after November 30, 2023 default to the Studio experience. Access to SageMaker geospatial is limited to Studio Classic, to learn more see Accessing SageMaker geospatial.

If you’re a first-time user of AWS or Amazon SageMaker, we recommend that you do the following:

1. Create an AWS account.

   To learn about setting up an AWS account and getting started with SageMaker, see Set Up Amazon SageMaker Prerequisites.

2. Create a user role and execution role that work with SageMaker geospatial.

   As a managed service, Amazon SageMaker geospatial capabilities performs operations on your behalf on the AWS hardware that SageMaker manages. A SageMaker execution role can perform only the operations that users grant. To work with SageMaker geospatial capabilities, you must set up a user role and an execution role. For more information, see SageMaker geospatial capabilities roles.

3. Update your trust policy to include SageMaker geospatial.
SageMaker geospatial defines an additional service principal. To learn how to create or update your SageMaker execution role’s trust policy, see Adding the SageMaker geospatial service principal to an existing SageMaker execution role.

4. **Set up an Amazon SageMaker domain to access Amazon SageMaker Studio Classic.**

To use SageMaker geospatial, a domain is required. For domains created before November 30, 2023 the default experience is Studio Classic. domains created after November 30, 2023 default to the Studio experience. To learn more about accessing Studio Classic from Studio, see Accessing SageMaker geospatial.

5. **Remember, shut down resources.**

When you have finished using SageMaker geospatial capabilities, shut down the instance it runs on to avoid incurring additional charges. For more information, see Shut Down Resources.

**Topics**
- Getting started with Amazon SageMaker geospatial
- Using a processing jobs for custom geospatial workloads
- Earth Observation Jobs
- Vector Enrichment Jobs
- Visualization Using SageMaker geospatial capabilities
- Amazon SageMaker geospatial Map SDK
- SageMaker geospatial capabilities FAQ
- SageMaker geospatial Security and Permissions
- Types of compute instances
- Data collections

**Getting started with Amazon SageMaker geospatial**

SageMaker geospatial provides a purpose built **Image** and **Instance type** for Amazon SageMaker Studio Classic notebooks. You can use either CPU or GPU enabled notebooks with the SageMaker geospatial **Image**. You can also visualize your geospatial data using a purpose built visualizer. Furthermore, SageMaker geospatial also provides APIs that allow you to query raster data collections. You can also use pre-trained models to analyze geospatial data, reverse geocoding, and map matching.
Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. If prior to November 30, 2023 you created a Amazon SageMaker domain, Studio Classic remains the default experience. domains created after November 30, 2023 default to the new Studio experience.

To access and get started using Amazon SageMaker geospatial, do the following:

Topics

• Accessing SageMaker geospatial
• Create an Amazon SageMaker Studio Classic notebook using the geospatial image
• Access the Sentinel-2 raster data collection and create an earth observation job to perform land segmentation

Accessing SageMaker geospatial

Note

Currently, SageMaker geospatial capabilities are only supported in the US West (Oregon) Region and in Studio Classic.

If you don't see the SageMaker geospatial UI available in your current Studio Classic instance check to make sure you are currently in the US West (Oregon) Region.

A domain is required to access SageMaker geospatial. If you created a domain prior to November 30, 2023 the default experience is Studio Classic.

If you created a domain after November 30, 2023 or if you have migrated to Studio, then you can use the following procedure to activate Studio Classic from within Studio to use SageMaker geospatial features.

To learn more about creating a domain, see Onboard to Amazon SageMaker domain.

To access Studio Classic from Studio

1. Launch Amazon SageMaker Studio.
2. Under **Applications**, choose **Studio Classic**.
3. Then, choose **Create Studio Classic space**.
4. On the **Create Studio Classic space** page, enter a **Name**.
5. Disable the **Share with my domain** option. SageMaker geospatial is not available in shared domains.
6. Then choose **Create space**.

When successful the **Status** changes to **Updating**. When your Studio Classic application is ready to be used the status changes to **Stopped**.

To start your Studio Classic application, choose **Run**.

**Create an Amazon SageMaker Studio Classic notebook using the geospatial image**

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](#).

⚠️ **Note**

Currently, SageMaker geospatial is only supported in the US West (Oregon) Region. If you don't see SageMaker geospatial available in your current domain or notebook instance, make sure that you're currently in the US West (Oregon) Region.

Use the following procedure to create Studio Classic notebook with the SageMaker geospatial image. If your default studio experience is Studio, see [Accessing SageMaker geospatial](#) to learn about starting a Studio Classic application.

**To create a Studio Classic notebook with the SageMaker geospatial image**

1. Launch Studio Classic
2. Choose **Home** in the menu bar.
3. Under **Quick actions**, choose **Open Launcher**.

4. When the **Launcher** dialog box opens, choose **Change environment** under **Notebooks and compute resources**.

5. When the **Change environment** dialog box opens, choose the **Image** dropdown and choose or type **Geospatial 1.0**.

![Change environment dialog box](image)

6. Next, choose an **Instance type** from the dropdown.

   SageMaker geospatial supports two types of notebook instances: CPU and GPU. The supported CPU instance is called **ml.geospatial.interactive**. Any of the G5-family of GPU instances can be used with the Geospatial 1.0 image.

   **Note**

   If you receive a ResourceLimitExceeded error when attempting to start a GPU based instance, you need to request a quota increase. To get started on a Service Quotas quota increase request, see [Requesting a quota increase](#) in the *Service Quotas User Guide*.

7. Choose **Select**.
8. Choose **Create notebook**.

After creating a notebook, to learn more about SageMaker geospatial, try the [SageMaker geospatial tutorial](#). It shows you how to process Sentinel-2 image data and perform land segmentation on it using the earth observation jobs API.

**Access the Sentinel-2 raster data collection and create an earth observation job to perform land segmentation**

This Python-based tutorial uses the SDK for Python (Boto3) and an Amazon SageMaker Studio Classic notebook. To complete this demo successfully, make sure that you have the required AWS Identity and Access Management (IAM) permissions to use SageMaker geospatial and Studio Classic. SageMaker geospatial requires that you have a user, group, or role which can access Studio Classic. You must also have a SageMaker execution role that specifies the SageMaker geospatial service principal, sagemaker-geospatial.amazonaws.com in its trust policy.

To learn more about these requirements, see [SageMaker geospatial IAM roles](#).

This tutorial shows you how to use SageMaker geospatial API to complete the following tasks:

- Find the available raster data collections with `list_raster_data_collections`.
- Search a specified raster data collection by using `search_raster_data_collection`.
- Create an earth observation job (EOJ) by using `start_earth_observation_job`.

**Using `list_raster_data_collections` to find available data collections**

SageMaker geospatial supports multiple raster data collections. To learn more about the available data collections, see [Data collections](#).

This demo uses satellite data that's collected from [Sentinel-2 Cloud-Optimized GeoTIFF](#) satellites. These satellites provide global coverage of Earth's land surface every five days. In addition to collecting surface images of Earth, the Sentinel-2 satellites also collect data across a variety of spectral bands.

To search an area of interest (AOI), you need the ARN that's associated with the Sentinel-2 satellite data. To find the available data collections and their associated ARNs in your AWS Region, use the `list_raster_data_collections` API operation.
Because the response can be paginated, you must use the `get_paginator` operation to return all of the relevant data:

```python
import boto3
import sagemaker
import sagemaker_geospatial_map
import json

## SageMaker Geospatial is currently only available in US-WEST-2
session = boto3.Session(region_name='us-west-2')
execution_role = sagemaker.get_execution_role()

## Creates a SageMaker Geospatial client instance
geospatial_client = session.client(service_name="sagemaker-geospatial")

# Creates a resusable Paginator for the list_raster_data_collections API operation
paginator = geospatial_client.get_paginator("list_raster_data_collections")

# Create a PageIterator from the paginator class
page_iterator = paginator.paginate()

# Use the iterator to iterate through the results of list_raster_data_collections
results = []
for page in page_iterator:
    results.append(page['RasterDataCollectionSummaries'])

print(results)
```

This is a sample JSON response from the `list_raster_data_collections` API operation. It's truncated to include only the data collection (Sentinel-2) that's used in this code example. For more details about a specific raster data collection, use `get_raster_data_collection`:

```json
{
    "Description": "Sentinel-2a and Sentinel-2b imagery, processed to Level 2A (Surface Reflectance) and converted to Cloud-Optimized GeoTIFFs",
    "DescriptionPageUrl": "https://registry.opendata.aws/sentinel-2-l2a-cogs",
    "Name": "Sentinel 2 L2A COGs",
    "SupportedFilters": [
        {
            "Maximum": 100,
        }
    ]
}
```
After running the previous code sample, you get the ARN of the Sentinel-2 raster data collection, 

**Searching the Sentinel-2 raster data collection using search_raster_data_collection**

In the preceding section, you used list_raster_data_collections to get the ARN for the Sentinel-2 data collection. Now you can use that ARN to search the data collection over a given area of interest (AOI), time range, properties, and the available UV bands.

To call the search_raster_data_collection API you must pass in a Python dictionary to the RasterDataCollectionQuery parameter. This example uses AreaOfInterest, TimeRangeFilter, PropertyFilters, and BandFilter. For ease, you can specify the Python dictionary using the variable `search_rdc_query` to hold the search query parameters:

```python
search_rdc_query = {
    "AreaOfInterest": {
        "AreaOfInterestGeometry": {
            "PolygonGeometry": {
                "Coordinates": [
                    # coordinates are input as longitude followed by latitude
                ]
            }
        }
    },
    "Maximum": 90,
    "Minimum": 0,
    "Name": "ViewOffNadir",
    "Type": "number"
},
    "Name": "Platform",
    "Type": "string"
],
"Tags": {},
"Type": "PUBLIC"
}
```
In this example, you query an AreaOfInterest that includes Lake Mead in Utah. Furthermore, Sentinel-2 supports multiple types of image bands. To measure the change in the surface of the water, you only need the visual band.

After you create the query parameters, you can use the search_raster_data_collection API to make the request.

The following code sample implements a search_raster_data_collection API request. This API does not support pagination using the get_paginator API. To make sure that the full API response has been gathered the code sample uses a while loop to check that NextToken exists.
The code sample then uses `extend()` to append the satellite image URLs and other response metadata to the `items_list`.


```python
search_rdc_response = sm_geo_client.search_raster_data_collection(
    RasterDataCollectionQuery=search_rdc_query
)

## items_list is the response from the API request.
items_list = []

## Use the python .get() method to check that the 'NextToken' exists, if null returns None breaking the while loop
while search_rdc_response.get('NextToken'):
    items_list.extend(search_rdc_response['Items'])
    search_rdc_response = sm_geo_client.search_raster_data_collection(
        RasterDataCollectionQuery=search_rdc_query,
        NextToken=search_rdc_response['NextToken']
    )

## Print the number of observation return based on the query
print (len(items_list))
```

The following is a JSON response from your query. It has been truncated for clarity. Only the "BandFilter": ["visual"] specified in the request is returned in the Assets key-value pair:

```json
{
    'Assets': {
        'visual': {
            'Href': 'https://sentinel-cogs.s3.us-west-2.amazonaws.com/sentinel-s2-l2a-cogs/15/T/UH/2022/6/S2A_15TUH_20220623_0_L2A/TCI.tif'
        }
    },
    'DateTime': datetime.datetime(2022, 6, 23, 17, 22, 5, 926000, tzinfo = tzlocal()),
    'Geometry': {
        'Coordinates': [
```
Now that you have your query results, in the next section you can visualize the results by using matplotlib. This is to verify that results are from the correct geographical region.

**Visualizing your search_raster_data_collection using matplotlib**

Before you start the earth observation job (EOJ), you can visualize a result from our query with matplotlib. The following code sample takes the first item, `items_list[0]['Assets']["visual"]['Href']`, from the `items_list` variable created in the previous code sample and prints an image using matplotlib.

```python
# Visualize an example image.
import os
from urllib import request
import tiffiff
import matplotlib.pyplot as plt

image_dir = './images/lake_mead'
os.makedirs(image_dir, exist_ok=True)

image_url = items_list[0]['Assets']['visual']['Href']
img_id = image_url.split('/')[-2]
path_to_image = image_dir + '/' + img_id + '_TCI.tif'
response = request.urlretrieve(image_url, path_to_image)
```
print("Downloaded image: " + img_id)

tci = tiff.imread(path_to_image)
plt.figure(figsize=(6, 6))
plt.imshow(tci)
plt.show()

After checking that the results are in the correct geographical region, you can start the Earth Observation Job (EOJ) in the next step. You use the EOJ to identify the water bodies from the satellite images by using a process called land segmentation.

**Starting an earth observation job (EOJ) that performs land segmentation on a series of Satellite images**

SageMaker geospatial provides multiple pre-trained models that you can use to process geospatial data from raster data collections. To learn more about the available pre-trained models and custom operations, see [Types of Operations](#).

To calculate the change in the water surface area, you need to identify which pixels in the images correspond to water. Land cover segmentation is a semantic segmentation model supported by the `start_earth_observation_job` API. Semantic segmentation models associate a label with every pixel in each image. In the results, each pixel is assigned a label that's based on the class map for the model. The following is the class map for the land segmentation model:

```json
{
    0: "No_data",
    1: "Saturated_or_defective",
    2: "Dark_area_pixels",
    3: "Cloud_shadows",
    4: "Vegetation",
    5: "Not_vegetated",
    6: "Water",
    7: "Unclassified",
    8: "Cloud_medium_probability",
    9: "Cloud_high_probability",
    10: "Thin_cirrus",
    11: "Snow_ice"
}
```

To start an earth observation job, use the `start_earth_observation_job` API. When you submit your request, you must specify the following:
• **InputConfig** (*dict*) – Used to specify the coordinates of the area that you want to search, and other metadata that's associated with your search.

• **JobConfig** (*dict*) – Used to specify the type of EOJ operation that you performed on the data. This example uses **LandCoverSegmentationConfig**.

• **ExecutionRoleArn** (*string*) – The ARN of the SageMaker execution role with the necessary permissions to run the job.

• **Name** (*string*) – A name for the earth observation job.

The **InputConfig** is a Python dictionary. Use the following variable `eoj_input_config` to hold the search query parameters. Use this variable when you make the `start_earth_observation_job` API request.

```python
# Perform land cover segmentation on images returned from the Sentinel-2 dataset.

eoj_input_config = {
    "RasterDataCollectionQuery": {
        "AreaOfInterest": {
            "AreaOfInterestGeometry": {
                "PolygonGeometry": {
                    "Coordinates": [
                        [-114.529, 36.142],
                        [-114.373, 36.142],
                        [-114.373, 36.411],
                        [-114.529, 36.411],
                        [-114.529, 36.142],
                    ]
                }
            }
        },
        "TimeRangeFilter": {
            "StartTime": "2021-01-01T00:00:00Z",
            "EndTime": "2022-07-10T23:59:59Z",
        },
        "PropertyFilters": {
            "Properties": [{'Property': {'EoCloudCover': {'LowerBound': 0, "UpperBound": 1}}},
                "LogicalOperator": "AND",
            ]
        }
    }
}
```
The JobConfig is a Python dictionary that is used to specify the EOJ operation that you want performed on your data:

```python
eoj_config = {"LandCoverSegmentationConfig": {}}
```

With the dictionary elements now specified, you can submit your `start_earth_observation_job` API request using the following code sample:

```python
# Gets the execution role arn associated with current notebook instance
execution_role_arn = sagemaker.get_execution_role()

# Starts an earth observation job
response = sm_geo_client.start_earth_observation_job(
    Name="lake-mead-landcover",
    InputConfig=eoj_input_config,
    JobConfig=eoj_config,
    ExecutionRoleArn=execution_role_arn,
)

print(response)
```

The start an earth observation job returns an ARN along with other metadata.

To get a list of all ongoing and current earth observation jobs use the `list_earth_observation_jobs` API. To monitor the status of a single earth observation job use the `get_earth_observation_job` API. To make this request, use the ARN created after submitting your EOJ request. To learn more, see `GetEarthObservationJob` in the Amazon SageMaker API Reference.

To find the ARNs associated with your EOJs use the `list_earth_observation_jobs` API operation. To learn more, see `ListEarthObservationJobs` in the Amazon SageMaker API Reference.

```python
# List all jobs in the account
sg_client.list_earth_observation_jobs()[:"EarthObservationJobSummaries"]
```

The following is an example JSON response:
After the status of your EOJ job changes to COMPLETED, proceed to the next section to calculate the change in Lake Mead's surface area.

**Calculating the change in the Lake Mead surface area**

To calculate the change in Lake Mead's surface area, first export the results of the EOJ to Amazon S3 by using `export_earth_observation_job`:

```python
sagemaker_session = sagemaker.Session()
s3_bucket_name = sagemaker_session.default_bucket()  # Replace with your own bucket if needed
s3_bucket = session.resource("s3").Bucket(s3_bucket_name)
prefix = "export-lake-mead-eoj"  # Replace with the S3 prefix desired
export_bucket_and_key = f"s3://{s3_bucket_name}/{prefix}/"

eoj_output_config = {"S3Data": {"S3Uri": export_bucket_and_key}}
export_response = sm_geo_client.export_earth_observation_job(
    ExecutionRoleArn=execution_role_arn,
    OutputConfig=eoj_output_config,
)
```
To see the status of your export job, use `get_earth_observation_job`:

```python
export_job_details =
    sm_geo_client.get_earth_observation_job(Arn=export_response["Arn"])
```

To calculate the changes in Lake Mead's water level, download the land cover masks to the local SageMaker notebook instance and download the source images from our previous query. In the class map for the land segmentation model, the water's class index is 6.

To extract the water mask from a Sentinel-2 image, follow these steps. First, count the number of pixels marked as water (class index 6) in the image. Second, multiply the count by the area that each pixel covers. Bands can differ in their spatial resolution. For the land cover segmentation model all bands are down sampled to a spatial resolution equal to 60 meters.

```python
import os
from glob import glob
import cv2
import numpy as np
import tifffile
import matplotlib.pyplot as plt
from urllib.parse import urlparse
from botocore import UNSIGNED
from botocore.config import Config

# Download land cover masks
mask_dir = "./masks/lake_mead"
os.makedirs(mask_dir, exist_ok=True)
image_paths = []
for s3_object in s3_bucket.objects.filter(Prefix=prefix).all():
    path, filename = os.path.split(s3_object.key)
    if "output" in path:
        mask_name = mask_dir + "/" + filename
        s3_bucket.download_file(s3_object.key, mask_name)
        print("Downloaded mask: " + mask_name)

# Download source images for visualization
for tci_url in tci_urls:
    url_parts = urlparse(tci_url)
    img_id = url_parts.path.split("/")[2]
    # Rest of the code...
```
tci_download_path = image_dir + "/" + img_id + "_TCI.tif"
cogs_bucket = session.resource("s3", config=Config(signature_version=UNSIGNED, region_name="us-west-2"))
cogs_bucket.download_file(url_parts.hostname.split("."))[0], tci_download_path)
print("Downloaded image: " + img_id)

print("Downloads complete.")

image_files = glob("images/lake_mead/*.tif")
mask_files = glob("masks/lake_mead/*.tif")
image_files.sort(key=lambda x: x.split("SQA_"))[1]
mask_files.sort(key=lambda x: x.split("SQA_"))[1]
overlay_dir = "/masks/lake_mead_overlay"
mask_dates = []

for image_file, mask_file in zip(image_files, mask_files):
    image_id = image_file.split('/')[1].split("_TCI"."00]
    mask_id = mask_file.split('/')[1].split(".tif"."0"
    mask_date = mask_id.split("_"."2"
    mask_dates.append(mask_date)
    assert image_id == mask_id
    image = tifffile.imread(image_file)
    image_ds = cv2.resize(image, (1830, 1830), interpolation=cv2.INTER_LINEAR)
    mask = tifffile.imread(mask_file)
    water_mask = np.isin(mask, [6]).astype(np.uint8) # water has a class index 6
    lake_mask = water_mask[1000:, :1100]
    lake_area = lake_mask.sum() * 60 * 60 / (1000 * 1000)  # calculate the surface area
    lake_areas.append(lake_area)
    contour, _ = cv2.findContours(water_mask, cv2.RETR_TREE, cv2.CHAIN_APPROX_SIMPLE)
    combined = cv2.drawContours(image_ds, contour, -1, (255, 0, 0), 4)
    lake_crop = combined[1000:, :1100]
    cv2.putText(lake_crop, f"{mask_date}", (10,50), cv2.FONT_HERSHEY_SIMPLEX, 1.5, (0, 0, 0), 3, cv2.LINE_AA)
    cv2.putText(lake_crop, f"{lake_area} [sq km]", (10,100), cv2.FONT_HERSHEY_SIMPLEX, 1.5, (0, 0, 0), 3, cv2.LINE_AA)
    overlay_file = overlay_dir + '/'+mask_date + '.png'
    cv2.imwrite(overlay_file, cv2.cvtColor(lake_crop, cv2.COLOR_RGB2BGR))

# Plot water surface area vs. time.
plt.figure(figsize=(20,10))
plt.title('Lake Mead surface area for the 2021.02 - 2022.07 period.', fontsize=20)
Using matplotlib, you can visualize the results with a graph. The graph shows that the surface area of Lake Mead decreased from January 2021–July 2022.

Using a processing jobs for custom geospatial workloads

With Amazon SageMaker Processing, you can use a simplified, managed experience on SageMaker to run your data processing workloads with the purpose-built geospatial container.

The underlying infrastructure for a Amazon SageMaker Processing job is fully managed by SageMaker. During a processing job, cluster resources are provisioned for the duration of your job, and cleaned up when a job completes.
The preceding diagram shows how SageMaker spins up a geospatial processing job. SageMaker takes your geospatial workload script, copies your geospatial data from Amazon Simple Storage Service (Amazon S3), and then pulls the specified geospatial container. The underlying infrastructure for the processing job is fully managed by SageMaker. Cluster resources are provisioned for the duration of your job, and cleaned up when a job completes. The output of the processing job is stored in the bucket you specified.

⚠️ Path naming constraints
The local paths inside a Processing jobs container must begin with /opt/ml/processing/.

SageMaker geospatial provides a purpose-built container, 081189585635.dkr.ecr.us-west-2.amazonaws.com/sagemaker-geospatial-v1-0:latest that can be specified when running a processing job.

Topics
- **Overview**: Run processing jobs using ScriptProcessor and a SageMaker geospatial container
- **Using ScriptProcessor to calculate the Normalized Difference Vegetation Index (NDVI) using Sentinel-2 satellite data**
Overview: Run processing jobs using ScriptProcessor and a SageMaker geospatial container

SageMaker geospatial provides a purpose-built processing container, 081189585635.dkr.ecr.us-west-2.amazonaws.com/sagemaker-geospatial-v1-0:latest. You can use this container when running a job with Amazon SageMaker Processing. When you create an instance of the ScriptProcessor class that is available through the Amazon SageMaker Python SDK for Processing, specify this image_uri.

Note
If you receive a ResourceLimitExceeded error when attempting to start a processing job, you need to request a quota increase. To get started on a Service Quotas quota increase request, see Requesting a quota increase in the Service Quotas User Guide

Prerequisites for using ScriptProcessor

1. You have created a Python script that specifies your geospatial ML workload.
2. You have granted the SageMaker execution role access to any Amazon S3 buckets that are needed.
3. Prepare your data for import into the container. Amazon SageMaker Processing jobs support either setting the s3_data_type equal to "ManifestFile" or to "S3Prefix".

The following procedure show you how to create an instance of ScriptProcessor and submit a Amazon SageMaker Processing job using the SageMaker geospatial container.

To create a ScriptProcessor instance and submit a Amazon SageMaker Processing job using a SageMaker geospatial container

1. Instantiate an instance of the ScriptProcessor class using the SageMaker geospatial image:

```python
from sagemaker.processing import ScriptProcessor, ProcessingInput, ProcessingOutput
sm_session = sagemaker.session.Session()
execution_role_arn = sagemaker.get_execution_role()

# purpose-built geospatial container
```
```
image_uri = '081189585635.dkr.ecr.us-west-2.amazonaws.com/sagemaker-geospatial-v1-0:latest'

script_processor = ScriptProcessor(
    command=['python3'],
    image_uri=image_uri,
    role=execution_role_arn,
    instance_count=4,
    instance_type='ml.m5.4xlarge',
    sagemaker_session=sm_session
)
```

Replace `execution_role_arn` with the ARN of the SageMaker execution role that has access to the input data stored in Amazon S3 and any other AWS services that you want to call in your processing job. You can update the `instance_count` and the `instance_type` to match the requirements of your processing job.

2. To start a processing job, use the `.run()` method:

```
# Can be replaced with any S3 compliant string for the name of the folder.
s3_folder = geospatial-data-analysis

# Use .default_bucket() to get the name of the S3 bucket associated with your current SageMaker session
s3_bucket = sm_session.default_bucket()

s3_manifest_uri = f's3://{s3_bucket}/{s3_folder}/manifest.json'
s3_prefix_uri =  f's3://{s3_bucket}/{s3_folder}/image-prefix

script_processor.run(
    code='preprocessing.py',
    inputs=[
        ProcessingInput(
            source=s3_manifest_uri | s3_prefix_uri ,
            destination='/opt/ml/processing/input_data/',
            s3_data_type = "ManifestFile" | "S3Prefix",
            s3_data_distribution_type= "ShardedByS3Key" | "FullyReplicated"
        )
    ],
    outputs=[
        ProcessingOutput(
            source='/opt/ml/processing/output_data/',
            destination=s3_output_prefix_url
        )
    ]
)`
• Replace `preprocessing.py` with the name of your own Python data processing script.

• A processing job supports two methods for formatting your input data. You can either create a manifest file that points to all of the input data for your processing job, or you can use a common prefix on each individual data input. If you created a manifest file set `s3_manifest_uri` equal to "ManifestFile". If you used a file prefix set `s3_manifest_uri` equal to "S3Prefix". You specify the path to your data using `source`.

• You can distribute your processing job data two ways:

  • Distribute your data to all processing instances by setting `s3_data_distribution_type` equal to `FullyReplicated`.

  • Distribute your data in shards based on the Amazon S3 key by setting `s3_data_distribution_type` equal to `ShardedByS3Key`. When you use `ShardedByS3Key` one shard of data is sent to each processing instance.

You can use a script to process SageMaker geospatial data. That script can be found in Step 3: Writing a script that can calculate the NDVI. To learn more about the `.run()` API operation, see `run` in the Amazon SageMaker Python SDK for Processing.

To monitor the progress of your processing job, the `ProcessingJobs` class supports a `describe` method. This method returns a response from the `DescribeProcessingJob` API call. To learn more, see `DescribeProcessingJob` in the Amazon SageMaker API Reference.

The next topic show you how to create an instance of the `ScriptProcessor` class using the SageMaker geospatial container, and then how to use it to calculate the Normalized Difference Vegetation Index (NDVI) with Sentinel-2 images.

### Using ScriptProcessor to calculate the Normalized Difference Vegetation Index (NDVI) using Sentinel-2 satellite data

The following code samples show you how to calculate the normalized difference vegetation index of a specific geographical area using the purpose-built geospatial image within a Studio Classic notebook and run a large-scale workload with Amazon SageMaker Processing using `ScriptProcessor` from the SageMaker Python SDK.
This demo also uses an Amazon SageMaker Studio Classic notebook instance that uses the geospatial kernel and instance type. To learn how to create a Studio Classic geospatial notebook instance, see Create an Amazon SageMaker Studio Classic notebook using the geospatial image.

You can follow along with this demo in your own notebook instance by copying and pasting the following code snippets:

1. Use `search_raster_data_collection` to query a specific area of interest (AOI) over a given time range using a specific raster data collection, Sentinel-2.
2. Create a manifest file that specifies what data will be processed during the processing job.
3. Write a data processing Python script calculating the NDVI.
4. Create a `ScriptProcessor` instance and start the Amazon SageMaker Processing job.
5. Visualizing the results of your processing job.

**Query the Sentinel-2 raster data collection using SearchRasterDataCollection**

With `search_raster_data_collection` you can query supported raster data collections. This example uses data that's pulled from Sentinel-2 satellites. The area of interest (AreaOfInterest) specified is rural northern Iowa, and the time range (TimeRangeFilter) is January 1, 2022 to December 30, 2022. To see the available raster data collections in your AWS Region use `list_raster_data_collections`. To see a code example using this API, see `ListRasterDataCollections` in the Amazon SageMaker Developer Guide.


A `search_raster_data_collection` API request requires two parameters:

- You need to specify an Arn parameter that corresponds to the raster data collection that you want to query.
- You also need to specify a `RasterDataCollectionQuery` parameter, which takes in a Python dictionary.

The following code example contains the necessary key-value pairs needed for the `RasterDataCollectionQuery` parameter saved to the `search_rdc_query` variable.
To make the `search_raster_data_collection` request, you must specify the ARN of the Sentinel-2 raster data collection: `arn:aws:sagemaker-geospatial:us-west-2:378778860802:raster-data-collection/public/nmqj48dcu3g7ayw8`. You also must need to pass in the Python dictionary that was defined previously, which specifies query parameters.

```python
search_rdc_query = {
    "AreaOfInterest": {
        "AreaOfInterestGeometry": {
            "PolygonGeometry": {
                "Coordinates": [
                    [-94.50938680498298, 43.22487436936203],
                    [-94.50938680498298, 42.843474642037194],
                    [-93.86520004156142, 42.843474642037194],
                    [-93.86520004156142, 43.22487436936203],
                    [-94.50938680498298, 43.22487436936203]
                ]
            }
        },
        "TimeRangeFilter": {
            "StartTime": "2022-01-01T00:00:00Z",
            "EndTime": "2022-12-30T23:59:59Z"
        }
    }
}
```

```python
# Creates a SageMaker Geospatial client instance
sm_geo_client = session.create_client(service_name="sagemaker-geospatial")
```
The results of this API can not be paginated. To collect all the satellite images returned by the `search_raster_data_collection` operation, you can implement a while loop. This checks for `NextToken` in the API response:

```python
## Holds the list of API responses from search_raster_data_collection
items_list = []
while search_rdc_response1.get('NextToken') and search_rdc_response1['NextToken'] != None:
    items_list.extend(search_rdc_response1['Items'])

search_rdc_response1 = sm_geo_client.search_raster_data_collection(
    RasterDataCollectionQuery=search_rdc_query,
    NextToken=search_rdc_response1['NextToken']
)
```

The API response returns a list of URLs under the `Assets` key corresponding to specific image bands. The following is a truncated version of the API response. Some of the image bands were removed for clarity.

```json
{
    'Assets': {
        'aot': {
            'Href': 'https://sentinel-cogs.s3.us-west-2.amazonaws.com/sentinel-s2-l2a-cogs/15/T/UH/2022/12/S2A_15TUH_20221230_0_L2A/AOT.tif'
        },
        'blue': {
            'Href': 'https://sentinel-cogs.s3.us-west-2.amazonaws.com/sentinel-s2-l2a-cogs/15/T/UH/2022/12/S2A_15TUH_20221230_0_L2A/B02.tif'
        },
        'swir22-jp2': {
            'Href': 's3://sentinel-s2-l2a/tiles/15/T/UH/2022/12/30/0/B12.jp2'
        },
        'visual-jp2': {
            'Href': 's3://sentinel-s2-l2a/tiles/15/T/UH/2022/12/30/0/TCI.jp2'
        }
    }
}
```
In the next section, you create a manifest file using the 'Id' key from the API response.

**Create an input manifest file using the Id key from the search_raster_data_collection API response**

When you run a processing job, you must specify a data input from Amazon S3. The input data type can either be a manifest file, which then points to the individual data files. You can also add a prefix to each file that you want processed. The following code example defines the folder where your manifest files will be generated.

Use SDK for Python (Boto3) to get the default bucket and the ARN of the execution role that is associated with your Studio Classic notebook instance:

```python
sm_session = sagemaker.session.Session()
s3 = boto3.resource('s3')
# Gets the default excution role associated with the notebook
```
Next, you create a manifest file. It will hold the URLs of the satellite images that you wanted processed when you run your processing job later in step 4.

```python
# Format of a manifest file
manifest_prefix = {}
manifest_prefix['prefix'] = 's3://' + s3_bucket + '/' + s3_folder + '/
manifest = [manifest_prefix]
print(manifest)
```

The following code sample returns the S3 URI where your manifest files will be created.

```python
[{'prefix': 's3://sagemaker-us-west-2-111122223333/script-processor-input-manifest/'}]
```

All the response elements from the search_raster_data_collection response are not needed to run the processing job.

The following code snippet removes the unnecessary elements 'Properties', 'Geometry', and 'DateTime'. The 'Id' key-value pair, 'Id': 'S2A_15TUH_20221230_0_L2A', contains the year and the month. The following code example parses that data to create new keys in the Python dictionary `dict_month_items`. The values are the assets that are returned from the SearchRasterDataCollection query.

```python
# For each response get the month and year, and then remove the metadata not related to the satellite images.
dict_month_items = {}
for item in items_list:
    # Example ID being split: 'S2A_15TUH_20221230_0_L2A'
    yyyymm = item['Id'].split('_')[2][:6]
    if yyyymm not in dict_month_items:
        dict_month_items[yyyymm] = []
```

Geospatial processing job
# Removes unneeded metadata elements for this demo
```
item.pop('Properties', None)
item.pop('Geometry', None)
item.pop('DateTime', None)
```

# Appends the response from search_raster_data_collection to newly created key above
```
dict_month_items[yyyymm].append(item)
```

This code example uploads the `dict_month_items` to Amazon S3 as a JSON object using the `.upload_file()` API operation:

```
## key_ is the yyyymm timestamp formatted above
## value_ is the reference to all the satellite images collected via our searchRDC query
for key_, value_ in dict_month_items.items():
    filename = f'manifest_{key_}.json'
    with open(filename, 'w') as fp:
        json.dump(value_, fp)
        s3.meta.client.upload_file(filename, s3_bucket, s3_folder + '/' + filename)
    manifest.append(filename)
    os.remove(filename)
```

This code example uploads a parent `manifest.json` file that points to all the other manifests uploaded to Amazon S3. It also saves the path to a local variable: `s3_manifest_uri`. You'll use that variable again to specify the source of the input data when you run the processing job in step 4.

```
with open('manifest.json', 'w') as fp:
    json.dump(manifest, fp)
    s3.meta.client.upload_file('manifest.json', s3_bucket, s3_folder + '/manifest.json')
    os.remove('manifest.json')
```

```
s3_manifest_uri = f's3://{s3_bucket}/{s3_folder}/manifest.json'
```

Now that you created the input manifest files and uploaded them, you can write a script that processes your data in the processing job. It processes the data from the satellite images, calculates the NDVI, and then returns the results to a different Amazon S3 location.
Write a script that calculates the NDVI

Amazon SageMaker Studio Classic supports the use of the %%writefile cell magic command. After running a cell with this command, its contents will be saved to your local Studio Classic directory. This is code specific to calculating NDVI. However, the following can be useful when you write your own script for a processing job:

- In your processing job container, the local paths inside the container must begin with `/opt/ml/processing/`. In this example, `input_data_path = '/opt/ml/processing/input_data/'` and `processed_data_path = '/opt/ml/processing/output_data/'` are specified in that way.

- With Amazon SageMaker Processing, a script that a processing job runs can upload your processed data directly to Amazon S3. To do so, make sure that the execution role associated with your ScriptProcessor instance has the necessary requirements to access the S3 bucket. You can also specify an outputs parameter when you run your processing job. To learn more, see the `.run()` API operation in the Amazon SageMaker Python SDK. In this code example, the results of the data processing are uploaded directly to Amazon S3.

- To manage the size of the Amazon EBS container attached to your processing job use the `volume_size_in_gb` parameter. The container's default size is 30 GB. You can also optionally use the Python library Garbage Collector to manage storage in your Amazon EBS container.

The following code example loads the arrays into the processing job container. When arrays build up and fill in the memory, the processing job crashes. To prevent this crash, the following example contains commands that remove the arrays from the processing job's container.

```python
%%writefile compute_ndvi.py

import os
import pickle
import sys
import subprocess
import json
import rioxarray

if __name__ == "__main__":
    print("Starting processing")

    input_data_path = '/opt/ml/processing/input_data/
    input_files = []
```
for current_path, sub_dirs, files in os.walk(input_data_path):
    for file in files:
        if file.endswith(".json"):
            input_files.append(os.path.join(current_path, file))

print("Received {} input_files: ").format(len(input_files), input_files))

items = []
for input_file in input_files:
    full_file_path = os.path.join(input_data_path, input_file)
    print(full_file_path)
    with open(full_file_path, 'r') as f:
        items.append(json.load(f))

items = [item for sub_items in items for item in sub_items]

for item in items:
    red_uri = item["Assets"]['red']["Href"]
    nir_uri = item["Assets"]['nir']["Href"]

    red = rioxarray.open_rasterio(red_uri, masked=True)
    nir = rioxarray.open_rasterio(nir_uri, masked=True)

    ndvi = (nir - red) / (nir + red)

    file_name = 'ndvi_' + item["Id"] + '.tif'
    output_path = '/opt/ml/processing/output_data'
    output_file_path = f'{output_path}/{file_name}'

    ndvi.rio.to_raster(output_file_path)
    print("Written output:", output_file_path)

You now have a script that can calculate the NDVI. Next, you can create an instance of the ScriptProcessor and run your Processing job.

Creating an instance of the ScriptProcessor class

This demo uses the ScriptProcessor class that is available via the Amazon SageMaker Python SDK. First, you need to create an instance of the class, and then you can start your Processing job by using the .run() method.

from sagemaker.processing import ScriptProcessor, ProcessingInput, ProcessingOutput
image_uri = '081189585635.dkr.ecr.us-west-2.amazonaws.com/sagemaker-geospatial-v1-0:latest'

processor = ScriptProcessor(
    command=['python3'],
    image_uri=image_uri,
    role=execution_role_arn,
    instance_count=4,
    instance_type='ml.m5.4xlarge',
    sagemaker_session=sm_session)

print('Starting processing job.')

When you start your Processing job, you need to specify a ProcessingInput object. In that object, you specify the following:

- The path to the manifest file that you created in step 2, s3_manifest_uri. This is the source of the input data to the container.
- The path to where you want the input data to be saved in the container. This must match the path that you specified in your script.
- Use the s3_data_type parameter to specify the input as "ManifestFile".

s3_output_prefix_url = f's3://{s3_bucket}/{s3_folder}/output'

processor.run(
    code='compute_ndvi.py',
    inputs=[
        ProcessingInput(
            source=s3_manifest_uri,
            destination='/opt/ml/processing/input_data/',
            s3_data_type="ManifestFile",
            s3_data_distribution_type="ShardedByS3Key",
        ),
    ],
    outputs=[
        ProcessingOutput(
            source='/opt/ml/processing/output_data/',
            destination=s3_output_prefix_url,
            s3_upload_mode="Continuous"
The following code example uses the `.describe()` method to get details of your Processing job.

```python
preprocessing_job_descriptor = processor.jobs[-1].describe()
s3_output_uri = preprocessing_job_descriptor['ProcessingOutputConfig']['Outputs'][0]["S3Output"]["S3Uri"]
print(s3_output_uri)
```

**Visualizing your results using matplotlib**

With the Matplotlib Python library, you can plot raster data. Before you plot the data, you need to calculate the NDVI using sample images from the Sentinel-2 satellites. The following code example opens the image arrays using the `.open_rasterio()` API operation, and then calculates the NDVI using the nir and red image bands from the Sentinel-2 satellite data.

```python
# Opens the python arrays
import rioxarray

red_uri = items[25]["Assets"]["red"]["Href"]
nir_uri = items[25]["Assets"]["nir"]["Href"]

red = rioxarray.open_rasterio(red_uri, masked=True)
nir = rioxarray.open_rasterio(nir_uri, masked=True)

# Calculates the NDVI
ndvi = (nir - red) / (nir + red)

# Common plotting library in Python
import matplotlib.pyplot as plt

f, ax = plt.subplots(figsize=(18, 18))
ndvi.plot(cmap='viridis', ax=ax)
aset_title("NDVI for {}".format(items[25]["Id"]))
aset_axis_off()
plt.show()
```

The output of the preceding code example is a satellite image with the NDVI values overlaid on it. An NDVI value near 1 indicates lots of vegetation is present, and values near 0 indicate no vegetation is present.
This completes the demo of using ScriptProcessor.
Earth Observation Jobs

Using an Earth Observation job (EOJ), you can acquire, transform, and visualize geospatial data to make predictions. You can choose an operation based on your use case from a wide range of operations and models. You get the flexibility of choosing your area of interest, selecting the data providers, and setting time-range based and cloud-cover-percentage-based filters. After SageMaker creates an EOJ for you, you can visualize the inputs and outputs of the job using the visualization functionality. An EOJ has various use cases that include comparing deforestation over time and diagnosing plant health. You can create an EOJ by using a SageMaker notebook with a SageMaker geospatial image. You can also access the SageMaker geospatial UI as a part of Amazon SageMaker Studio Classic UI to view the list of all your jobs. You can also use the UI to pause or stop an ongoing job. You can choose a job from the list of available EOJ to view the Job summary, the Job details as well as visualize the Job output.

Topics

• Create an Earth Observation Job Using a Amazon SageMaker Studio Classic Notebook with a SageMaker geospatial Image
• Types of Operations

Create an Earth Observation Job Using a Amazon SageMaker Studio Classic Notebook with a SageMaker geospatial Image

To use a SageMaker Studio Classic notebook with a SageMaker geospatial image:

1. From the Launcher, choose Change environment under Notebooks and compute resources.
2. Next, the Change environment dialog opens.
3. Select the Image dropdown and choose Geospatial 1.0. The Instance type should be ml.geospatial.interactive. Do not change the default values for other settings.
4. Choose Select.
5. Choose Create notebook.

You can initiate an EOJ using a Amazon SageMaker Studio Classic notebook with a SageMaker geospatial image using the code provided below.

```python
import boto3
import sagemaker
```
import sagemaker_geospatial_map

session = boto3.Session()
execution_role = sagemaker.get_execution_role()
sg_client = session.client(service_name="sagemaker-geospatial")

The following is an example showing how to create an EOJ in the US West (Oregon) Region.

```python
# Query and Access Data
search_rdc_args = {
    "RasterDataCollectionQuery": {
        "AreaOfInterest": {
            "AreaOfInterestGeometry": {
                "PolygonGeometry": {
                    "Coordinates": [
                        [-114.529, 36.142],
                        [-114.373, 36.142],
                        [-114.373, 36.411],
                        [-114.529, 36.411],
                        [-114.529, 36.142],
                        [-114.529, 36.142],
                    ]
                }
            }
        },
        "TimeRangeFilter": {
            "StartTime": "2021-01-01T00:00:00Z",
            "EndTime": "2022-07-10T23:59:59Z",
        },
        "PropertyFilters": {
            "Properties": [{"Property": {"EoCloudCover": {"LowerBound": 0,
            "UpperBound": 1}}}],
            "LogicalOperator": "AND",
        },
        "BandFilter": ["visual"],
    }
}

tci_urls = []
data_manifests = []
```
while search_rdc_args.get("NextToken", True):
    search_result = sg_client.search_raster_data_collection(**search_rdc_args)
    if search_result.get("NextToken"):
        data_manifests.append(search_result)
    for item in search_result["Items"]:  
        tci_url = item["Assets"]['visual']['Href']
        print(tci_url)
        tci_urls.append(tci_url)

    search_rdc_args["NextToken"] = search_result.get("NextToken")

# Perform land cover segmentation on images returned from the sentinel dataset.
eoj_input_config = {
    "RasterDataCollectionQuery": {
        "AreaOfInterest": {
            "AreaOfInterestGeometry": {
                "PolygonGeometry": {
                    "Coordinates": [
                        [-114.529, 36.142],
                        [-114.373, 36.142],
                        [-114.373, 36.411],
                        [-114.529, 36.411],
                        [-114.529, 36.142],
                    ]
                }
            }
        },
        "TimeRangeFilter": {
            "StartTime": "2021-01-01T00:00:00Z",
            "EndTime": "2022-07-10T23:59:59Z",
        },
        "PropertyFilters": {
            "Properties": [{"Property": {"EoCloudCover": {"LowerBound": 0,
            "UpperBound": 1}}}],
            "LogicalOperator": "AND",
        }
    }
}
eoj_config = {"LandCoverSegmentationConfig": {}}
response = sg_client.start_earth_observation_job(
    Name="lake-mead-landcover",
    InputConfig=eoj_input_config,
    JobConfig=eoj_config,
    ExecutionRoleArn=execution_role,
)

After your EOJ is created, the Arn is returned to you. You use the Arn to identify a job and perform further operations. To get the status of a job, you can run

```
sg_client.get_earth_observation_job(Arn = response['Arn'])
```

The following example shows how to query the status of an EOJ until it is completed.

```
eoj_arn = response['Arn']
job_details = sg_client.get_earth_observation_job(Arn=eoj_arn)
{k: v for k, v in job_details.items() if k in ['Arn', 'Status', 'DurationInSeconds']}
# List all jobs in the account
sg_client.list_earth_observation_jobs()['EarthObservationJobSummaries']
```

After the EOJ is completed, you can visualize the EOJ outputs directly in the notebook. The following example shows you how an interactive map can be rendered.

```
map = sagemaker_geospatial_map.create_map({
    'is_raster': True
})
map.set_sagemaker_geospatial_client(sg_client)
# render the map
map.render()
```

The following example shows how the map can be centered on an area of interest and the input and output of the EOJ can be rendered as separate layers within the map.

```
# visualize the area of interest
config = {"label": "Lake Mead A0I"}
aoi_layer = map.visualize_eoj_aoi(Arn=eoj_arn, config=config)

# Visualize input.
time_range_filter = {
    "start_date": "2022-07-01T00:00:00Z",
    "end_date": "2022-07-10T23:59:59Z",
}
```
config = {"label": "Input"}

input_layer = map.visualize_eoj_input(
    Arn=eoj_arn, config=config, time_range_filter=time_range_filter
)

# Visualize output, EOJ needs to be in completed status.
time_range_filter = {
    "start_date": "2022-07-01T00:00:00Z",
    "end_date": "2022-07-10T23:59:59Z",
}
config = {"preset": "singleBand", "band_name": "mask"}
output_layer = map.visualize_eoj_output(
    Arn=eoj_arn, config=config, time_range_filter=time_range_filter
)

You can use the `export_earth_observation_job` function to export the EOJ results to your Amazon S3 bucket. The export function makes it convenient to share results across teams. SageMaker also simplifies dataset management. We can simply share the EOJ results using the job ARN, instead of crawling thousands of files in the S3 bucket. Each EOJ becomes an asset in the data catalog, as results can be grouped by the job ARN. The following example shows how you can export the results of an EOJ.

```python
sagemaker_session = sagemaker.Session()
s3_bucket_name = sagemaker_session.default_bucket()  # Replace with your own bucket if needed
s3_bucket = session.resource("s3").Bucket(s3_bucket_name)
prefix = "eoj_lakemead"  # Replace with the S3 prefix desired
export_bucket_and_key = f"s3://{s3_bucket_name}/{prefix}/"

eoj_output_config = {"S3Data": {"S3Uri": export_bucket_and_key}}
export_response = sg_client.export_earth_observation_job(
    Arn=eoj_arn,
    ExecutionRoleArn=execution_role,
    OutputConfig=eoj_output_config,
    ExportSourceImages=False,
)
```

You can monitor the status of your export job using the following snippet.

```python
# Monitor the export job status
export_job_details = sg_client.get_earth_observation_job(Arn=export_response["Arn"])
```
You are not charged the storage fees after you delete the EOJ.

For an example that showcases how to run an EOJ, see this blog post.

For more example notebooks on SageMaker geospatial capabilities, see this GitHub repository.

Types of Operations

When you create an EOJ, you select an operation based on your use case. Amazon SageMaker geospatial capabilities provide a combination of purpose-built operations and pre-trained models. You can use these operations to understand the impact of environmental changes and human activities over time or identify cloud and cloud-free pixels.

Cloud Masking

Identify clouds in satellite images is an essential pre-processing step in producing high-quality geospatial data. Ignoring cloud pixels can lead to errors in analysis, and over-detection of cloud pixels can decrease the number of valid observations. Cloud masking has the ability to identify cloudy and cloud-free pixels in satellite images. An accurate cloud mask helps get satellite images for processing and improves data generation. The following is the class map for cloud masking.

```json
{
    0: "No_cloud",
    1: "cloud"
}
```

Cloud Removal

Cloud removal for Sentinel-2 data uses an ML-based semantic segmentation model to identify clouds in the image. Cloudy pixels can be replaced by with pixels from other timestamps. USGS Landsat data contains landsat metadata that is used for cloud removal.

Temporal Statistics

Temporal statistics calculate statistics for geospatial data through time. The temporal statistics currently supported include mean, median, and standard deviation. You can calculate these
statistics by using GROUPBY and set it to either all or yearly. You can also mention the TargetBands.

**Zonal Statistics**

Zonal statistics performs statistical operations over a specified area on the image.

**Resampling**

Resampling is used to upscale and downscale the resolution of a geospatial image. The value attribute in resampling represents the length of a side of the pixel.

**Geomosaic**

Geomosaic allows you to stitch smaller images into a large image.

**Band Stacking**

Band stacking takes more than one image band as input and stacks them into a single GeoTIFF. The OutputResolution attribute determines the resolution of the output image. Based on the resolutions of the input images, you can set it to lowest, highest or average.

**Band Math**

Band Math, also known as Spectral Index, is a process of transforming the observations from multiple spectral bands to a single band, indicating the relative abundance of features of interests. For instance, Normalized Difference Vegetation Index (NDVI) and Enhanced Vegetation Index (EVI) are helpful for observing the presence of green vegetation features.

**Land Cover Segmentation**

Land Cover segmentation is a semantic segmentation model that has the capability to identify the physical material, such as vegetation, water, and bare ground, at the earth surface. Having an accurate way to map the land cover patterns helps you understand the impact of environmental change and human activities over time. Land Cover segmentation is often used for region planning, disaster response, ecological management, and environmental impact assessment. The following is the class map for Land Cover segmentation.

```json
{
  0: "No_data",
  1: "Saturated_or_defective",
  ...
}
```
Availability of EOJ Operations

The availability of operations depends on whether you are using the SageMaker geospatial UI or the Amazon SageMaker Studio Classic notebooks with a SageMaker geospatial image. Currently, notebooks support all functionalities. To summarize, the following geospatial operations are supported by SageMaker:

<table>
<thead>
<tr>
<th>Operations</th>
<th>Description</th>
<th>Availability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cloud Masking</td>
<td>Identify cloud and cloud-free pixels to get improved and accurate satellite imagery.</td>
<td>UI, Notebook</td>
</tr>
<tr>
<td>Cloud Removal</td>
<td>Remove pixels containing parts of a cloud from satellite imagery.</td>
<td>Notebook</td>
</tr>
<tr>
<td>Temporal Statistics</td>
<td>Calculate statistics through time for a given GeoTIFF.</td>
<td>Notebook</td>
</tr>
<tr>
<td>Zonal Statistics</td>
<td>Calculate statistics on user-defined regions.</td>
<td>Notebook</td>
</tr>
<tr>
<td>Resampling</td>
<td>Scale images to different resolutions.</td>
<td>Notebook</td>
</tr>
<tr>
<td>Geomosaic</td>
<td>Combine multiple images for greater fidelity.</td>
<td>Notebook</td>
</tr>
<tr>
<td>Operations</td>
<td>Description</td>
<td>Availability</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Band Stacking</td>
<td>Combine multiple spectral bands to create a single image.</td>
<td>Notebook</td>
</tr>
<tr>
<td>Band Math / Spectral Index</td>
<td>Obtain a combination of spectral bands that indicate the abundance of features of interest.</td>
<td>UI, Notebook</td>
</tr>
<tr>
<td>Land Cover Segmentation</td>
<td>Identify land cover types such as vegetation and water in satellite imagery.</td>
<td>UI, Notebook</td>
</tr>
</tbody>
</table>

**Vector Enrichment Jobs**

A Vector Enrichment Job (VEJ) performs operations on your vector data. Currently, you can use a VEJ to do reverse geocoding or map matching.

**Reverse Geocoding**

With a reverse geocoding VEJ, you can convert geographic coordinates (latitude, longitude) to human-readable addresses powered by Amazon Location Service. When you upload a CSV file containing the longitude and latitude coordinates, it returns the address number, country, label, municipality, neighborhood, postal code and region of that location. The output file consists of your input data along with columns containing these the values appended at the end. These jobs are optimized to accept tens of thousands of GPS traces.

**Map Matching**

Map matching allows you to snap GPS coordinates to road segments. The input should be a CSV file containing the trace ID (route), longitude, latitude and the timestamp attributes. There can be multiple GPS co-ordinates per route. The input can contain multiple routes too. The output is a GeoJSON file that contains links of the predicted route. It also has the snap points provided in the input. These jobs are optimized to accept tens of thousands of drives in one request. Map matching is supported by OpenStreetMap. Map matching fails if the names in the input source field don't match the ones in MapMatchingConfig. The error message you receive contains
The field names present in the input file and the expected field name that is not found in `MapMatchingConfig`.

The input CSV file for a VEJ must contain the following:

- A header row
- Latitude and longitude in separate columns
- The ID and Timestamp columns can be in numeric or string format. All other column data must be in numeric format only
- No miss matching quotes

For the timestamp column, SageMaker geospatial capabilities supports epoch time in seconds and milliseconds (long integer). The string formats supported are as follows:

- "dd.MM.yyyy HH:mm:ss z"
- "yyyy-MM-dd'T'HH:mm:ss.SSS'Z'"
- "yyyy-MM-dd'T'HH:mm:ss"
- "yyyy-MM-dd hh:mm:ss a"
- "yyyy-MM-dd HH:mm:ss"
- "yyyyMMddHHmmss"

While you need to use an Amazon SageMaker Studio Classic notebook to execute a VEJ, you can view all the jobs you create using the UI. To use the visualization in the notebook, you first need to export your output to your S3 bucket. The VEJ actions you can perform are as follows.

- `StartVectorEnrichmentJob`
- `GetVectorEnrichmentJob`
- `ListVectorEnrichmentJobs`
- `StopVectorEnrichmentJob`
- `DeleteVectorEnrichmentJob`

**Visualization Using SageMaker geospatial capabilities**

Using the visualization functionalities provided by Amazon SageMaker geospatial you can visualize geospatial data, the inputs to your EOJ or VEJ jobs as well as the outputs exported from your
Amazon S3 bucket. The visualization tool is powered by Foursquare Studio. The following image depicts the visualization tool supported by SageMaker geospatial capabilities.

![Visualization tool](image)

You can use the left navigation panel to add data, layers, filters, and columns. You can also make modifications to how you interact with the map.

**Dataset**

The source of data used for visualization is called a Dataset. To add data for visualization, choose Add Data in the left navigation panel. You can either upload the data from your Amazon S3 bucket or your local machine. The data formats supported are CSV, JSON and GeoJSON. You can add multiple datasets to your map. After you upload the dataset, you can see it loaded on the map screen.

**Layers**

In the layer panel, a layer is created and populated automatically when you add a dataset. If your map consists of more than one dataset, you can select which dataset belongs to a layer. You can create new layers and group them. SageMaker geospatial capabilities support various layer types, including point, arc, icon, and polygon.

You can choose any data point in a layer to have an Outline. You can also further customize the data points. For example, you can choose the layer type as Point and then Fill Color based on any column of your dataset. You can also change the radius of the points.
The following image shows the layers panel supported by SageMaker geospatial capabilities.

**Columns**

You can view the columns present in your dataset by using the **Columns** tab in the left navigation panel.

**Filters**

You can use filters to limit the data points that display on the map.

**Interactions**

In the **Interactions** panel, you can customize how you interact with the map. For example, you can choose what metrics to display when you hover the tooltip over a data point.

**Base map**

Currently, SageMaker only supports the Amazon Dark base map.

**Split Map Modes**
You can have a **Single Map**, **Dual Maps** or **Swipe Maps**. With **Dual Maps**, you can compare the same map side-by-side using different layers. Use **Swipe Maps** to overlay two maps on each other and use the sliding separator to compare them. You can choose the split map mode by choosing the **Split Mode** button on the top right corner of your map.

**Legends for EOJ in the SageMaker geospatial UI**

The output visualization of an EOJ depends on the operation you choose to create it. The legend is based on the default color scale. You can view the legend by choosing the **Show legend** button on the top right corner of your map.

**Spectral Index**

When you visualize the output for an EOJ that uses the spectral index operation, you can map the category based on the color from the legend as shown.

![Spectral Index Legend](image)

**Cloud Masking**

When you visualize the output for an EOJ that uses the cloud masking operation, you can map the category based on the color from the legend as shown.

![Cloud Masking Legend](image)
Land Cover Segmentation

When you visualize the output for an EOJ that uses the Land Cover Segmentation operation, you can map the category based on the color from the legend as shown.

Amazon SageMaker geospatial Map SDK

You can use Amazon SageMaker geospatial capabilities to visualize maps within the SageMaker geospatial UI as well as SageMaker notebooks with a geospatial image. These visualizations are supported by the map visualization library called Foursquare Studio.

You can use the APIs provided by the SageMaker geospatial map SDK to visualize your geospatial data, including the input, output, and AoI for EOJ.

Topics
• add_dataset API
• update_dataset API
• add_layer API
• update_layer API
• visualize_eoj_aoi API
• visualize_eoj_input API
• visualize_eoj_output API

add_dataset API

Adds a raster or vector dataset object to the map.

Request syntax

```python
Request =
    add_dataset(  
        self,  
        dataset: Union[Dataset, Dict, None] = None,  
        *,  
        auto_create_layers: bool = True,  
        center_map: bool = True,  
        **kwargs: Any,  
    ) -> Optional[Dataset]
```

Request parameters

The request accepts the following parameters.

Positional arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dataset</td>
<td>Union[Dataset, Dict, None]</td>
<td>Data used to create a dataset, in CSV, JSON, or GeoJSON format (for local datasets) or a UUID string.</td>
</tr>
</tbody>
</table>
Keyword arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>auto_create_layers</td>
<td>Boolean</td>
<td>Whether to attempt to create new layers when adding a dataset. Default value is False.</td>
</tr>
<tr>
<td>center_map</td>
<td>Boolean</td>
<td>Whether to center the map on the created dataset. Default value is True.</td>
</tr>
<tr>
<td>id</td>
<td>String</td>
<td>Unique identifier of the dataset. If you do not provide it, a random ID is generated.</td>
</tr>
<tr>
<td>label</td>
<td>String</td>
<td>Dataset label which is displayed.</td>
</tr>
<tr>
<td>color</td>
<td>Tuple[float, float, float]</td>
<td>Color label of the dataset.</td>
</tr>
<tr>
<td>metadata</td>
<td>Dictionary</td>
<td>Object containing tileset metadata (for tiled datasets).</td>
</tr>
</tbody>
</table>

Response

This API returns the Dataset object that was added to the map.

update_dataset API

Updates an existing dataset's settings.

Request syntax

```python
Request = 
    update_dataset(
        self,
        dataset_id: str,  
        values: Union[_DatasetUpdateProps, dict, None] = None,
)```
**kwargs: Any,
) -> Dataset

**Request parameters**

The request accepts the following parameters.

Positional arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dataset_id</td>
<td>String</td>
<td>The identifier of the dataset to be updated.</td>
</tr>
<tr>
<td>values</td>
<td>Union[_DatasetUpdateProps, dict, None]</td>
<td>The values to update.</td>
</tr>
</tbody>
</table>

Keyword arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>label</td>
<td>String</td>
<td>Dataset label which is displayed.</td>
</tr>
<tr>
<td>color</td>
<td>RGBColor</td>
<td>Color label of the dataset.</td>
</tr>
</tbody>
</table>

**Response**

This API returns the updated dataset object for interactive maps, or None for non-interactive HTML environments.

**add_layer API**

Adds a new layer to the map. This function requires at least one valid layer configuration.

**Request syntax**

```
Request =
```
add_layer(
    self,
    layer: Union[LayerCreationProps, dict, None] = None,
    **kwargs: Any
) -> Layer

Request parameters

The request accepts the following parameters.

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>layer</td>
<td>Union[LayerCreationProps, dict, None]</td>
<td>A set of properties used to create a layer.</td>
</tr>
</tbody>
</table>

Response

The layer object that was added to the map.

update_layer API

Update an existing layer with given values.

Request syntax

Request =
    update_layer(
        self,
        layer_id: str,
        values: Union[LayerUpdateProps, dict, None],
        **kwargs: Any
    ) -> Layer

Request parameters

The request accepts the following parameters.

Arguments
<table>
<thead>
<tr>
<th>Positional argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>layer_id</td>
<td>String</td>
<td>The ID of the layer to be updated.</td>
</tr>
<tr>
<td>values</td>
<td>Union[LayerUpdateProps, dict, None]</td>
<td>The values to update.</td>
</tr>
</tbody>
</table>

**Keyword arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>type</td>
<td>LayerType</td>
<td>The type of layer.</td>
</tr>
<tr>
<td>data_id</td>
<td>String</td>
<td>Unique identifier of the dataset this layer visualizes.</td>
</tr>
<tr>
<td>fields</td>
<td>Dict [string, Optional[string]]</td>
<td>Dictionary that maps fields that the layer requires for visualization to appropriate dataset fields.</td>
</tr>
<tr>
<td>label</td>
<td>String</td>
<td>Canonical label of this layer.</td>
</tr>
<tr>
<td>is_visible</td>
<td>Boolean</td>
<td>Whether the layer is visible or not.</td>
</tr>
<tr>
<td>config</td>
<td>LayerConfig</td>
<td>Layer configuration specific to its type.</td>
</tr>
</tbody>
</table>

**Response**

Returns the updated layer object.

**visualize_eoj_aoi API**

Visualize the AoI of the given job ARN.
Request parameters

The request accepts the following parameters.

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arn</td>
<td>String</td>
<td>The ARN of the job.</td>
</tr>
<tr>
<td>config</td>
<td>Dictionary</td>
<td>An option to pass layer properties.</td>
</tr>
</tbody>
</table>

config = { label: <string> custom label of the added AoI layer, default AoI }

Response

Reference of the added input layer object.

visualize_eoj_input API

Visualize the input of the given EOJ ARN.

Request parameters

The request accepts the following parameters.

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arn</td>
<td>String</td>
<td>The ARN of the job.</td>
</tr>
<tr>
<td>time_range_filter</td>
<td>Dictionary</td>
<td>An option to provide the start and end time. Defaults to the raster data collection search start and end date.</td>
</tr>
</tbody>
</table>

time_range_filter = {
  start_date: <string> date in ISO format
}
### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>end_date: &lt;string&gt;</td>
<td>date in ISO format</td>
<td></td>
</tr>
</tbody>
</table>

#### config

```
config = { label: <string>  
          custom label of the added output layer, default Input }
```

- **Type**: Dictionary
- **Description**: An option to pass layer properties.

### Response

Reference of the added input layer object.

#### visualize_eoj_output API

Visualize the output of the given EOJ ARN.

### Request parameters

The request accepts the following parameters.

#### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arn</td>
<td>String</td>
<td>The ARN of the job.</td>
</tr>
<tr>
<td>time_range_filter</td>
<td>Dictionary</td>
<td>An option to provide the start and end time. Defaults to the raster data collection search start and end date.</td>
</tr>
</tbody>
</table>

```
time_range_filter = {  
                      start_date: <string> date in ISO format  
                      end_date: <string> date in ISO format  
```

- **Type**: Dictionary
- **Description**: An option to provide the start and end time. Defaults to the raster data collection search start and end date.
<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>config</td>
<td>Dictionary</td>
<td>An option to pass layer properties.</td>
</tr>
</tbody>
</table>

```python
config = {
    'label': <string> custom label of the added output layer, default Output
    'preset': <string> singleBand or trueColor,
    'band_name': <string>, only required for 'singleBand' preset. Allowed bands for a EOJ
}
```

**Response**

Reference of the added output Layer object.

To learn more about visualizing your geospatial data, refer to [Visualization Using Amazon SageMaker geospatial](#).

**SageMaker geospatial capabilities FAQ**

Use the following FAQ items to find answers to commonly asked questions about SageMaker geospatial capabilities.

1. **What regions are Amazon SageMaker geospatial capabilities available in?**

   Currently, SageMaker geospatial capabilities are only supported in the US West (Oregon) Region. To view SageMaker geospatial, choose the name of the currently displayed Region in the navigation bar of the console. Then choose the US West (Oregon) Region.
2. **What AWS Identity and Access Management permissions and policies are required to use SageMaker geospatial?**

To use SageMaker geospatial you need a user, group, or role that can access SageMaker. You also need to create a SageMaker execution role so that SageMaker geospatial can perform operations on your behalf. To learn more, see [SageMaker geospatial capabilities roles](#).

3. **I have an existing SageMaker execution role. Do I need to update it?**

Yes. To use SageMaker geospatial you must specify an additional service principal in your IAM trust policy: sagemaker-geospatial.amazonaws.com. To learn about specifying a service principal in a trust relationship, see [Adding the SageMaker geospatial service principal to an existing SageMaker execution role](#) in the *Amazon SageMaker Developer Guide*.

4. **Can I use SageMaker geospatial capabilities through my VPC environment?**

Yes, you can use SageMaker geospatial through a VPN. To learn more, see [Use Amazon SageMaker geospatial capabilities in Your Amazon Virtual Private Cloud](#).

5. **Why can't I see the SageMaker geospatial map visualizer, image or instance type when I navigate to Amazon SageMaker Studio Classic?**

Verify that you are launching Amazon SageMaker Studio Classic in the US West (Oregon) Region and that you are not using a shared space.

6. **Why can't I see the SageMaker geospatial image or instance type when I try to create a notebook instance in Studio Classic?**

Verify that you are launching Amazon SageMaker Studio Classic in the US West (Oregon) Region and that you are not using a shared space. To learn more, see [Create an Amazon SageMaker Studio Classic notebook using the geospatial image](#).

7. **What bands supported for various raster data collections?**

Use the `GetRasterDataCollection` API response and refer to the `ImageSourceBands` field to find the bands supported for that particular data collection.

---

**SageMaker geospatial Security and Permissions**

Use the topics on this page to learn about SageMaker geospatial capabilities security features. Additionally, learn how to use SageMaker geospatial capabilities in an Amazon Virtual Private Cloud as well as protect your data at rest using encryption.
For more information about IAM users and roles, see Identities (Users, Groups, and Roles) in the IAM User Guide.

To learn more about using IAM with SageMaker, see Identity and Access Management for Amazon SageMaker.

Topics

- Configuration and Vulnerability Analysis in SageMaker geospatial
- Use Amazon SageMaker geospatial capabilities in Your Amazon Virtual Private Cloud
- Use AWS KMS Permissions for Amazon SageMaker geospatial capabilities

Configuration and Vulnerability Analysis in SageMaker geospatial

Configuration and IT controls are a shared responsibility between AWS and you, our customer. AWS handles basic security tasks like guest operating system (OS) and database patching, firewall configuration, and disaster recovery. These procedures have been reviewed and certified by the appropriate third parties. For more details, see the following resources:

- Shared Responsibility Model.
- Amazon Web Services: Overview of Security Processes.


Amazon SageMaker geospatial capabilities provide a number of security features to consider as you develop and implement your own security policies. The following best practices are general guidelines and don't represent a complete security solution. Because these best practices might not be appropriate or sufficient for your environment, treat them as helpful considerations rather than prescriptions.

Apply principle of least privilege

Amazon SageMaker geospatial capabilities provide granular access policy for applications using IAM roles. We recommend that the roles be granted only the minimum set of privileges required by the job. We also recommend auditing the jobs for permissions on a regular basis and upon any change to your application.

Role-based access control (RBAC) permissions
Administrators should strictly control Role-based access control (RBAC) permissions for Amazon SageMaker geospatial capabilities.

**Use temporary credentials whenever possible**

Where possible, use temporary credentials instead of long-term credentials, such as access keys. For scenarios in which you need IAM users with programmatic access and long-term credentials, we recommend that you rotate access keys. Regularly rotating long-term credentials helps you familiarize yourself with the process. This is useful in case you are ever in a situation where you must rotate credentials, such as when an employee leaves your company. We recommend that you use IAM access last used information to rotate and remove access keys safely. For more information, see [Rotating access keys](#) and [Security best practices in IAM](#).

**Use AWS CloudTrail to view and log API calls**

AWS CloudTrail tracks anyone making API calls in your AWS account. API calls are logged whenever anyone uses the Amazon SageMaker geospatial capabilities API, the Amazon SageMaker geospatial capabilities console or Amazon SageMaker geospatial capabilities AWS CLI commands. Enable logging and specify an Amazon S3 bucket to store the logs.

Your trust, privacy, and the security of your content are our highest priorities. We implement responsible and sophisticated technical and physical controls designed to prevent unauthorized access to, or disclosure of, your content and ensure that our use complies with our commitments to you. For more information, see [AWS Data Privacy FAQ](#).

**Use Amazon SageMaker geospatial capabilities in Your Amazon Virtual Private Cloud**

The following topic gives information on how to use SageMaker notebooks with a SageMaker geospatial image in a Amazon SageMaker domain with VPC only mode. For more information on VPCs in Amazon SageMaker Studio Classic see [Choose an Amazon VPC](#).

**VPC only communication with the internet**

By default, SageMaker domain uses two Amazon VPC. One of the Amazon VPC is managed by Amazon SageMaker and provides direct internet access. You specify the other Amazon VPC, which provides encrypted traffic between the domain and your Amazon Elastic File System (Amazon EFS) volume.

You can change this behavior so that SageMaker sends all traffic over your specified Amazon VPC. If VPC only has been choosen as the network access mode during the SageMaker domain
creation, the following requirements need to be considered to still allow usage of SageMaker Studio Classic notebooks within the created SageMaker domain.

**Requirements to use VPC only mode**

1. **Note**
   
   In order to use the visualization components of SageMaker geospatial capabilities, the browser you use to access the SageMaker Studio Classic UI needs to be connected to the internet.

When you choose VpcOnly, follow these steps:

1. You must use private subnets only. You cannot use public subnets in VpcOnly mode.
2. Ensure your subnets have the required number of IP addresses needed. The expected number of IP addresses needed per user can vary based on use case. We recommend between 2 and 4 IP addresses per user. The total IP address capacity for a Studio Classic domain is the sum of available IP addresses for each subnet provided when the domain is created. Ensure that your estimated IP address usage does not exceed the capacity supported by the number of subnets you provide. Additionally, using subnets distributed across many availability zones can aid in IP address availability. For more information, see [VPC and subnet sizing for IPv4](#).

3. **Note**
   
   You can configure only subnets with a default tenancy VPC in which your instance runs on shared hardware. For more information on the tenancy attribute for VPCs, see [Dedicated Instances](#).

3. Set up one or more security groups with inbound and outbound rules that together allow the following traffic:

   - **NFS traffic over TCP on port 2049** between the domain and the Amazon EFS volume.
   - **TCP traffic within the security group.** This is required for connectivity between the JupyterServer app and the KernelGateway apps. You must allow access to at least ports in the range 8192-65535.

4. If you want to allow internet access, you must use a [NAT gateway](#) with access to the internet, for example through an [internet gateway](#).
5. If you don't want to allow internet access, create interface VPC endpoints (AWS PrivateLink) to allow Studio Classic to access the following services with the corresponding service names. You must also associate the security groups for your VPC with these endpoints.

**Note**
Currently, SageMaker geospatial capabilities are only supported in the US West (Oregon) Region.

- SageMaker API: com.amazonaws.us-west-2.sagemaker.api
- SageMaker runtime: com.amazonaws.us-west-2.sagemaker.runtime. This is required to run Studio Classic notebooks with a SageMaker geospatial image.
- Amazon S3: com.amazonaws.us-west-2.s3.
- SageMaker geospatial capabilities: com.amazonaws.us-west-2.sagemaker-geospatial

If you use the SageMaker Python SDK to run remote training jobs, you must also create the following Amazon VPC endpoints.

- AWS Security Token Service: com.amazonaws.region.sts
- Amazon CloudWatch: com.amazonaws.region.logs. This is required to allow SageMaker Python SDK to get the remote training job status from Amazon CloudWatch.

**Note**
For a customer working within VPC mode, company firewalls can cause connection issues with SageMaker Studio Classic or between JupyterServer and the KernelGateway. Make the following checks if you encounter one of these issues when using SageMaker Studio Classic from behind a firewall.

- Check that the Studio Classic URL is in your networks allowlist.
- Check that the websocket connections are not blocked. Jupyter uses websocket under the hood. If the KernelGateway application is InService, JupyterServer may not be able
Use AWS KMS Permissions for Amazon SageMaker geospatial capabilities

You can protect your data at rest using encryption for SageMaker geospatial capabilities. By default, it uses server-side encryption with an Amazon SageMaker geospatial owned key. SageMaker geospatial capabilities also supports an option for server-side encryption with a customer managed KMS key.

Server-Side Encryption with Amazon SageMaker geospatial managed key (Default)

SageMaker geospatial capabilities encrypts all your data, including computational results from your Earth Observation jobs (EOJ) and Vector Enrichment jobs (VEJ) along with all your service metadata. There is no data that is stored within SageMaker geospatial capabilities unencrypted. It uses a default AWS owned key to encrypt all your data.

Server-Side Encryption with customer managed KMS key (Optional)

SageMaker geospatial capabilities supports the use of a symmetric customer managed key that you create, own, and manage to add a second layer of encryption over the existing AWS owned encryption. Because you have full control of this layer of encryption, you can perform such tasks as:

- Establishing and maintaining key policies
- Establishing and maintaining IAM policies and grants
- Enabling and disabling key policies
- Rotating key cryptographic material
- Adding tags
- Creating key aliases
- Scheduling keys for deletion

For more information, see Customer managed keys in the AWS Key Management Service Developer Guide.
How SageMaker geospatial capabilities uses grants in AWS KMS

SageMaker geospatial capabilities requires a grant to use your customer managed key. When you create an EOJ or an VEJ encrypted with a customer managed key, SageMaker geospatial capabilities creates a grant on your behalf by sending a CreateGrant request to AWS KMS. Grants in AWS KMS are used to give SageMaker geospatial capabilities access to a KMS key in a customer account. You can revoke access to the grant, or remove the service's access to the customer managed key at any time. If you do, SageMaker geospatial capabilities won't be able to access any of the data encrypted by the customer managed key, which affects operations that are dependent on that data.

Create a customer managed key

You can create a symmetric customer managed key by using the AWS Management Console, or the AWS KMS APIs.

To create a symmetric customer managed key

Follow the steps for Creating symmetric encryption KMS keys in the AWS Key Management Service Developer Guide.

Key policy

Key policies control access to your customer managed key. Every customer managed key must have exactly one key policy, which contains statements that determine who can use the key and how they can use it. When you create your customer managed key, you can specify a key policy. For more information, see Determining access to AWS KMS keys in the AWS Key Management Service Developer Guide.

To use your customer managed key with your SageMaker geospatial capabilities resources, the following API operations must be permitted in the key policy. The principal for these operations should be the Execution Role you provide in the SageMaker geospatial capabilities request. SageMaker geospatial capabilities assumes the provided Execution Role in the request to perform these KMS operations.

- `kms:CreateGrant`
- `kms:GenerateDataKey`
- `kms:Decrypt`
- `kms:GenerateDataKeyWithoutPlaintext`
The following are policy statement examples you can add for SageMaker geospatial capabilities:

CreateGrant

```
"Statement" : [
    {
        "Sid" : "Allow access to Amazon SageMaker geospatial capabilities",
        "Effect" : "Allow",
        "Principal" : {
            "AWS" : "<Customer provided Execution Role ARN>"
        },
        "Action" : [
            "kms:CreateGrant",
            "kms:Decrypt",
            "kms:GenerateDataKey",
            "kms:GenerateDataKeyWithoutPlaintext"
        ],
        "Resource" : "*",
    },
]
```

For more information about specifying permissions in a policy, see AWS KMS permissions in the AWS Key Management Service Developer Guide. For more information about troubleshooting, see Troubleshooting key access in the AWS Key Management Service Developer Guide.

If your key policy does not have your account root as key administrator, you need to add the same KMS permissions on your execution role ARN. Here is a sample policy you can add to the execution role:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Action": [
                "kms:CreateGrant",
                "kms:Decrypt",
                "kms:GenerateDataKey",
                "kms:GenerateDataKeyWithoutPlaintext"
            ],
            "Resource": ["<KMS key Arn>"
```


Monitoring your encryption keys for SageMaker geospatial capabilities

When you use an AWS KMS customer managed key with your SageMaker geospatial capabilities resources, you can use AWS CloudTrail or Amazon CloudWatch Logs to track requests that SageMaker geospatial sends to AWS KMS.

Select a tab in the following table to see examples of AWS CloudTrail events to monitor KMS operations called by SageMaker geospatial capabilities to access data encrypted by your customer managed key.

CreateGrant

```json
{
   "eventVersion": "1.08",
   "userIdentity": {
      "type": "AssumedRole",
      "principalId": "AROAIGDTESTANDEXAMPLE:SageMaker-Geospatial-StartEOJ-KMSAccess",
      "accountId": "111122223333",
      "accessKeyId": "AKIAIOSFODNN7EXAMPLE3",
      "sessionContext": {
         "sessionIssuer": {
            "type": "Role",
            "principalId": "AKIAIOSFODNN7EXAMPLE3",
            "arn": "arn:aws:sts::111122223333:assumed-role/SageMakerGeospatialCustomerRole",
            "accountId": "111122223333",
            "userName": "SageMakerGeospatialCustomerRole"
         },
         "webIdFederationData": {},
         "attributes": {
            "creationDate": "2023-03-17T18:02:06Z",
            "mfaAuthenticated": "false"
         }
      },
      "invokedBy": "arn:aws:iam::111122223333:root"
   }
}
```
GenerateDataKey

```json
{
    "eventVersion": "1.0",
    "eventSource": "aws:kms.amazonaws.com",
    "eventTime": "2023-03-17T18:02:06Z",
    "eventSourceARN": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE",
    "eventName": "GenerateDataKey",
    "awsRegion": "us-west-2",
    "sourceIPAddress": "172.12.34.56",
    "userAgent": "ExampleDesktop/1.0 (V1; OS)",
    "requestParameters": {
        "KeyId": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE",
        "GrantTokens": null,
        "operations": [
            "Encrypt"
        ],
        "granteePrincipal": "sagemaker-geospatial.us-west-2.amazonaws.com"
    },
    "responseElements": {
        "GrantId": "0abc0ac0d0b000f00ea00cc0a0e00fc00bce000c000f0000000c0bc0000aaafSAMPLE",
        "KeyId": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE"
    },
    "requestID": "ff000af-00eb-00ce-0e00-ea000fb0fba0SAMPLE",
    "eventID": "ff000af-00eb-00ce-0e00-ea000fb0fba0SAMPLE",
    "readOnly": false,
    "resources": [
        {
            "accountId": "111122223333",
            "type": "AWS::KMS::Key",
            "ARN": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE"
        }
    ],
    "eventType": "AwsApiCall",
    "managementEvent": true,
    "recipientAccountId": "111122223333",
    "eventCategory": "Management"
}
```
"userIdentity": {
    "type": "AWSService",
    "invokedBy": "sagemaker-geospatial.amazonaws.com"
},
"eventTime": "2023-03-24T00:29:45Z",
"eventSource": "kms.amazonaws.com",
"eventName": "GenerateDataKey",
"awsRegion": "us-west-2",
"sourceIPAddress": "sagemaker-geospatial.amazonaws.com",
"userAgent": "sagemaker-geospatial.amazonaws.com",
"requestParameters": {
    "encryptionContext": {
        "aws:s3:arn": "arn:aws:s3:::axis-earth-observation-
job-378778860802/111122223333/napy9eintp64/output/
consolidated/32PPR/2022-01-04T09:58:03Z/S2B_32PPR_20220104_0_L2A_msavi.tif"
    },
    "keyId": "arn:aws:kms:us-
west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE",
    "keySpec": "AES_256"
},
"responseElements": null,
"requestID": "ff000af-00eb-00ce-0e00-ea000fb0fba0SAMPLE",
"eventID": "ff000af-00eb-00ce-0e00-ea000fb0fba0SAMPLE",
"readOnly": true,
"resources": [
    {
        "accountId": "111122223333",
        "type": "AWS::KMS::Key",
        "ARN": "arn:aws:kms:us-
west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE"
    }
],
"eventType": "AwsApiCall",
"managementEvent": true,
"recipientAccountId": "111122223333",
"eventCategory": "Management"
}

Decrypt

{
    "eventVersion": "1.08",
    "userIdentity": {

GenerateDataKeyWithoutPlainText

{
    "eventVersion": "1.08",
    "userIdentity": {
        "type": "AssumedRole",
        "principalId": "AROAIGDTESTANDEXAMPLE:SageMaker-Geospatial-StartEOJ-KMSAccess",
        "arn": "arn:aws:sts::111122223333:assumed-role/SageMaker-Geospatial-StartEOJ-KMSAccess/ARNofAssumedRole",
        "accountId": "111122223333",
        "callerAccountIds": null,
        "callerType": "Role",
        "servicePrincipal": "sts.amazonaws.com",
        "sessionContext": {
            "sessionIssuer": {
                "arn": "arn:aws:iam::111122223333:invitation/ARNofInvitation",
                "principalId": "PARTIAL_ArnOfPrincipal",
                "accountId": "111122223333",
                "awsRegion": "us-west-2"
            }
        }
    },
    "eventTime": "2023-03-28T22:04:24Z",
    "eventSource": "kms.amazonaws.com",
    "eventName": "Encrypt",
    "awsRegion": "us-west-2",
    "sourceIPAddress": "sagemaker-geospatial.amazonaws.com",
    "userAgent": "sagemaker-geospatial.amazonaws.com",
    "requestParameters": {
        "KeyType": "_CUSTOMERMASTERKEY",
        "Description": "GenerateDataKeyWithoutPlainText",
        "EncryptionContext": {
            "aws:s3:KMSKeyId": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-1234567890 SAMPLE"
        }
    },
    "responseElements": null,
    "requestID": "ff000af-00eb-00ce-0e00-ea000fb0fba0SAMPLE",
    "eventID": "ff000af-00eb-00ce-0e00-ea000fb0fba0SAMPLE",
    "readOnly": true,
    "resources": [
        {
            "accountId": "111122223333",
            "type": "AWS::KMS::Key",
            "ARN": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-1234567890 SAMPLE"
        }
    ],
    "eventType": "AwsApiCall",
    "managementEvent": true,
    "recipientAccountId": "111122223333",
    "eventCategory": "Management"
}
"accountId": "111122223333",
"accessKeyId": "AKIAIOSFODNN7EXAMPLE3",
"sessionContext": {
    "sessionIssuer": {
        "type": "Role",
        "principalId": "AKIAIOSFODNN7EXAMPLE3",
        "arn": "arn:aws:sts::111122223333:assumed-role/SageMakerGeospatialCustomerRole",
        "accountId": "111122223333",
        "userName": "SageMakerGeospatialCustomerRole"
    },
    "webIdFederationData": {},
    "attributes": {
        "creationDate": "2023-03-17T18:02:06Z",
        "mfaAuthenticated": "false"
    }
},
"invokedBy": "arn:aws:iam::111122223333:root",
"eventTime": "2023-03-28T22:09:16Z",
"eventSource": "kms.amazonaws.com",
"eventName": "GenerateDataKeyWithoutPlaintext",
"awsRegion": "us-west-2",
"sourceIPAddress": "172.12.34.56",
"userAgent": "ExampleDesktop/1.0 (V1; OS)",
"requestParameters": {
    "keySpec": "AES_256",
    "keyId": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE"
},
"responseElements": null,
"requestID": "ff000af-00eb-00ce-0e00-ea000f0fba0SAMPLE",
"eventID": "ff000af-00eb-00ce-0e00-ea000f0fba0SAMPLE",
"readOnly": true,
"resources": [
    {
        "accountId": "111122223333",
        "type": "AWS::KMS::Key",
        "ARN": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-123456SAMPLE"
    }
],
Types of compute instances

SageMaker geospatial capabilities offer three types of compute instances.

- **SageMaker Studio Classic geospatial notebook instances** – SageMaker geospatial supports both CPU and GPU-based notebook instances in Studio Classic. Notebook instances are used to build, train, and deploy ML models. For a list of available notebook instance types that work with the geospatial image, see [Supported notebook instance types](#).

- **SageMaker geospatial jobs instances** – Run processing jobs to transform satellite image data.

- **SageMaker geospatial model inference types** – Make predictions by using pre-trained ML models on satellite imagery.

The instance type is determined by the operations that you run.

The following table shows the available SageMaker geospatial specific operations and instance types that you can use.

<table>
<thead>
<tr>
<th>Operations</th>
<th>Instance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temporal Statistics</td>
<td>ml.geospatial.jobs</td>
</tr>
<tr>
<td>Zonal Statistics</td>
<td>ml.geospatial.jobs</td>
</tr>
<tr>
<td>Resampling</td>
<td>ml.geospatial.jobs</td>
</tr>
<tr>
<td>Geomosaic</td>
<td>ml.geospatial.jobs</td>
</tr>
<tr>
<td>Band Stacking</td>
<td>ml.geospatial.jobs</td>
</tr>
<tr>
<td>Band Math</td>
<td>ml.geospatial.jobs</td>
</tr>
<tr>
<td>Cloud Removal with Landsat8</td>
<td>ml.geospatial.jobs</td>
</tr>
</tbody>
</table>
### SageMaker geospatial supported notebook instance types

SageMaker geospatial supports both CPU and GPU-based notebook instances in Studio Classic. If when starting a GPU enabled notebook instance you receive a ResourceLimitExceeded error, you need to request a quota increase. To get started on a Service Quotas quota increase request, see Requesting a quota increase in the Service Quotas User Guide.

### Supported Studio Classic notebook instance types

<table>
<thead>
<tr>
<th>Name</th>
<th>Instance type</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.geospatial.interactive</td>
<td>CPU</td>
</tr>
<tr>
<td>ml.g5.xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.2xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.4xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.8xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.16xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.12xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.24xlarge</td>
<td>GPU</td>
</tr>
<tr>
<td>ml.g5.48xlarge</td>
<td>GPU</td>
</tr>
</tbody>
</table>

You are charged different rates for each type of compute instance that you use. For more information about pricing, see Geospatial ML with Amazon SageMaker.
### SageMaker geospatial libraries

The SageMaker geospatial specific **Instance type, ml.geospatial.interactive** contains the following Python libraries.

#### Geospatial libraries available on the geospatial instance type

<table>
<thead>
<tr>
<th>Library name</th>
<th>Version available</th>
</tr>
</thead>
<tbody>
<tr>
<td>numpy</td>
<td>1.23.4</td>
</tr>
<tr>
<td>scipy</td>
<td>1.11.2</td>
</tr>
<tr>
<td>pandas</td>
<td>1.4.4</td>
</tr>
<tr>
<td>gdal</td>
<td>3.2.2</td>
</tr>
<tr>
<td>fiona</td>
<td>1.8.22</td>
</tr>
<tr>
<td>geopandas</td>
<td>0.11.1</td>
</tr>
<tr>
<td>shapely</td>
<td>1.8.4</td>
</tr>
<tr>
<td>seaborn</td>
<td>0.11.2</td>
</tr>
<tr>
<td>notebook</td>
<td>1.8.22</td>
</tr>
<tr>
<td>scikit-image</td>
<td>0.11.2</td>
</tr>
<tr>
<td>rasterio</td>
<td>6.4.12</td>
</tr>
<tr>
<td>scikit-learn</td>
<td>0.19.2</td>
</tr>
<tr>
<td>ipyleaflet</td>
<td>1.0.1</td>
</tr>
<tr>
<td>rtree</td>
<td>0.17.2</td>
</tr>
<tr>
<td>opencv</td>
<td>4.6.0.66</td>
</tr>
<tr>
<td>supy</td>
<td>2022.4.7</td>
</tr>
<tr>
<td>SNAP toolbox</td>
<td>9.0</td>
</tr>
<tr>
<td>Library name</td>
<td>Version available</td>
</tr>
<tr>
<td>---------------</td>
<td>------------------</td>
</tr>
<tr>
<td>cdsapi</td>
<td>0.6.1</td>
</tr>
<tr>
<td>arosics</td>
<td>1.8.1</td>
</tr>
<tr>
<td>rasterstats</td>
<td>0.18.0</td>
</tr>
<tr>
<td>rioxarray</td>
<td>0.14.1</td>
</tr>
<tr>
<td>pyroSAR</td>
<td>0.20.0</td>
</tr>
<tr>
<td>eo-learn</td>
<td>1.4.1</td>
</tr>
<tr>
<td>deepforest</td>
<td>1.2.7</td>
</tr>
<tr>
<td>scrapy</td>
<td>2.8.0</td>
</tr>
<tr>
<td>netCDF4</td>
<td>1.6.3</td>
</tr>
<tr>
<td>xarray[complete]</td>
<td>0.20.1</td>
</tr>
<tr>
<td>Orfeotoolbox</td>
<td>OTB-8.1.1</td>
</tr>
<tr>
<td>pytorch</td>
<td>2.0.1</td>
</tr>
<tr>
<td>pytorch-cuda</td>
<td>11.8</td>
</tr>
<tr>
<td>torchvision</td>
<td>0.15.2</td>
</tr>
<tr>
<td>torchaudio</td>
<td>2.0.2</td>
</tr>
<tr>
<td>pytorch-lightning</td>
<td>2.0.6</td>
</tr>
<tr>
<td>tensorflow</td>
<td>2.13.0</td>
</tr>
</tbody>
</table>

**Data collections**

Amazon SageMaker geospatial supports the following raster data collections. Of the following data collections, you can use the USGS Landsat and the Sentinel-2 Cloud-Optimized GeoTIFF data
collections when starting an Earth Observation Job (EOJ). To learn more about the EOJs, see Earth Observation Jobs.

- Copernicus Digital Elevation Model (DEM) – GLO-30
- Copernicus Digital Elevation Model (DEM) – GLO-90
- Sentinel-2 Cloud-Optimized GeoTIFFs
- Sentinel-1
- National Agriculture Imagery Program (NAIP) on AWS
- USGS Landsat 8

To find the list of available raster data collections in your AWS Regions, use `ListRasterDataCollections`. In the `ListRasterDataCollections response`, you get a `RasterDataCollectionMetadata` object that contains details about the available raster data collections.

Example Example – Calling the `ListRasterDataCollections` API using the AWS SDK for Python (Boto3)

When you use the SDK for Python (Boto3) and SageMaker geospatial, you must create a geospatial client, `geospatial_client`. Use the following Python snippet to make a call to the `list_raster_data_collections` API:

```python
import boto3
import sagemaker
import sagemaker_geospatial_map
import json

## SageMaker Geospatial Capabilities is currently only available in US-WEST-2
session = boto3.Session(region_name='us-west-2')
execution_role = sagemaker.get_execution_role()

## Creates a SageMaker Geospatial client instance
geospatial_client = session.client(service_name="sagemaker-geospatial")

# Creates a reusable Paginator for the list_raster_data_collections API operation
paginator = geospatial_client.get_paginator("list_raster_data_collections")

# Create a PageIterator from the Paginator
page_iterator = paginator.paginate()
```
# Use the iterator to iterate through the results of list_raster_data_collections
results = []
for page in page_iterator:
    results.append(page['RasterDataCollectionSummaries'])

print(results)

In the JSON response, you will receive the following, which has been truncated for clarity:

```json
{
    "Description": "Copernicus DEM is a Digital Surface Model which represents the surface of the Earth including buildings, infrastructure, and vegetation. GLO-30 is instance of Copernicus DEM that provides limited worldwide coverage at 30 meters.",
    "DescriptionPageUrl": "https://registry.opendata.aws/copernicus-dem/",
    "Name": "Copernicus DEM GLO-30",
    "Tags": {},
    "Type": "PUBLIC"
}
```

**Image band information from the USGS Landsat and Sentinel-2 data collections**

Image band information from the USGS Landsat 8 and Sentinel-2 data collections are provided in the following table.

**USGS Landsat**

<table>
<thead>
<tr>
<th>Band name</th>
<th>Wave length range (nm)</th>
<th>Units</th>
<th>Valid range</th>
<th>Fill value</th>
<th>Spatial resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>coastal</td>
<td>435 - 451</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>blue</td>
<td>452 - 512</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>green</td>
<td>533 - 590</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>red</td>
<td>636 - 673</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>nir</td>
<td>851 - 879</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>Band name</td>
<td>Wave length range (nm)</td>
<td>Units</td>
<td>Valid range</td>
<td>Fill value</td>
<td>Spatial resolution</td>
</tr>
<tr>
<td>---------------</td>
<td>------------------------</td>
<td>-------------</td>
<td>-------------</td>
<td>---------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>swir16</td>
<td>1566 - 1651</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>swir22</td>
<td>2107 - 2294</td>
<td>Unitless</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>qa_aerosol</td>
<td>NA</td>
<td>Bit Index</td>
<td>0 - 255</td>
<td>1</td>
<td>30m</td>
</tr>
<tr>
<td>qa_pixel</td>
<td>NA</td>
<td>Bit Index</td>
<td>1 - 65455</td>
<td>1 (bit 0)</td>
<td>30m</td>
</tr>
<tr>
<td>qa_radsat</td>
<td>NA</td>
<td>Bit Index</td>
<td>1 - 65455</td>
<td>NA</td>
<td>30m</td>
</tr>
<tr>
<td>t</td>
<td>10600 - 11190</td>
<td>Scaled Kelvin</td>
<td>1 - 65455</td>
<td>0 (No Data)</td>
<td>30m (scaled from 100m)</td>
</tr>
<tr>
<td>atran</td>
<td>NA</td>
<td>Unitless</td>
<td>0 - 10000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>cdist</td>
<td>NA</td>
<td>Kilometers</td>
<td>0 - 24000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>drad</td>
<td>NA</td>
<td>W/(m^2 sr (\mu)m)/DN</td>
<td>0 - 28000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>urad</td>
<td>NA</td>
<td>W/(m^2 sr (\mu)m)/DN</td>
<td>0 - 28000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>trad</td>
<td>NA</td>
<td>W/(m^2 sr (\mu)m)/DN</td>
<td>0 - 28000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>emis</td>
<td>NA</td>
<td>Emissivity coefficient</td>
<td>1 - 10000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
<tr>
<td>emsd</td>
<td>NA</td>
<td>Emissivity coefficient</td>
<td>1 - 10000</td>
<td>-9999 (No Data)</td>
<td>30m</td>
</tr>
</tbody>
</table>
### Sentinel-2

<table>
<thead>
<tr>
<th>Band name</th>
<th>Wave length range (nm)</th>
<th>Scale</th>
<th>Valid range</th>
<th>Fill value</th>
<th>Spatial resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>coastal</td>
<td>443</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>60m</td>
</tr>
<tr>
<td>blue</td>
<td>490</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>10m</td>
</tr>
<tr>
<td>green</td>
<td>560</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>10m</td>
</tr>
<tr>
<td>red</td>
<td>665</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>10m</td>
</tr>
<tr>
<td>rededge1</td>
<td>705</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>rededge2</td>
<td>740</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>rededge3</td>
<td>783</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>nir</td>
<td>842</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>10m</td>
</tr>
<tr>
<td>nir08</td>
<td>865</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>nir08</td>
<td>865</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>nir09</td>
<td>940</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>60m</td>
</tr>
<tr>
<td>swir16</td>
<td>1610</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>swir22</td>
<td>2190</td>
<td>0.0001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
<tr>
<td>aot</td>
<td>Aerosol optical thickness</td>
<td>0.001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>10m</td>
</tr>
<tr>
<td>wvp</td>
<td>Scene-average water vapor</td>
<td>0.001</td>
<td>NA</td>
<td>0 (No Data)</td>
<td>10m</td>
</tr>
<tr>
<td>Band name</td>
<td>Wave length range (nm)</td>
<td>Scale</td>
<td>Valid range</td>
<td>Fill value</td>
<td>Spatial resolution</td>
</tr>
<tr>
<td>-----------</td>
<td>------------------------</td>
<td>-------</td>
<td>-------------</td>
<td>------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>scl</td>
<td>Scene classification data</td>
<td>NA</td>
<td>1 - 11</td>
<td>0 (No Data)</td>
<td>20m</td>
</tr>
</tbody>
</table>

**RStudio on Amazon SageMaker**

RStudio is an integrated development environment for R, with a console, syntax-highlighting editor that supports direct code execution, and tools for plotting, history, debugging and workspace management. Amazon SageMaker supports RStudio as a fully-managed integrated development environment (IDE) integrated with Amazon SageMaker domain through Posit Workbench. For more information about Posit Workbench, see the [Posit website](#).

RStudio allows customers to create data science insights using an R environment. With RStudio integration, you can launch an RStudio environment in the domain to run your RStudio workflows on SageMaker resources.

SageMaker integrates RStudio through the creation of a RStudioServerPro app.

The following are supported by RStudio on SageMaker.

- R developers use the RStudio IDE interface with popular developer tools from the R ecosystem. Users can launch new RStudio sessions, write R code, install dependencies from RStudio Package Manager, and publish Shiny apps using RStudio Connect.
- R developers can quickly scale underlying compute resources to run large scale data processing and statistical analysis.
- Platform administrators can set up user identities, authorization, networking, storage, and security for their data science teams through AWS IAM Identity Center and AWS Identity and Access Management integration. This includes connection to private Amazon Virtual Private Cloud (Amazon VPC) resources and internet-free mode with AWS PrivateLink.
- Integration with AWS License Manager.

For information on the onboarding steps to create a domain with RStudio enabled, see [Amazon SageMaker domain overview](#).
## Region availability

The following table gives information about the AWS Regions that RStudio on SageMaker is supported in.

<table>
<thead>
<tr>
<th>Region name</th>
<th>Region</th>
</tr>
</thead>
<tbody>
<tr>
<td>US East (Ohio)</td>
<td>us-east-2</td>
</tr>
<tr>
<td>US East (N. Virginia)</td>
<td>us-east-1</td>
</tr>
<tr>
<td>US West (N. California)</td>
<td>us-west-1</td>
</tr>
<tr>
<td>US West (Oregon)</td>
<td>us-west-2</td>
</tr>
<tr>
<td>Asia Pacific (Mumbai)</td>
<td>ap-south-1</td>
</tr>
<tr>
<td>Asia Pacific (Seoul)</td>
<td>ap-northeast-2</td>
</tr>
<tr>
<td>Asia Pacific (Singapore)</td>
<td>ap-southeast-1</td>
</tr>
<tr>
<td>Asia Pacific (Sydney)</td>
<td>ap-southeast-2</td>
</tr>
<tr>
<td>Asia Pacific (Tokyo)</td>
<td>ap-northeast-1</td>
</tr>
<tr>
<td>Canada (Central)</td>
<td>ca-central-1</td>
</tr>
<tr>
<td>Europe (Frankfurt)</td>
<td>eu-central-1</td>
</tr>
<tr>
<td>Europe (Ireland)</td>
<td>eu-west-1</td>
</tr>
<tr>
<td>Europe (London)</td>
<td>eu-west-2</td>
</tr>
<tr>
<td>Europe (Paris)</td>
<td>eu-west-3</td>
</tr>
<tr>
<td>Europe (Stockholm)</td>
<td>eu-north-1</td>
</tr>
<tr>
<td>South America (São Paulo)</td>
<td>sa-east-1</td>
</tr>
</tbody>
</table>
## RStudio components

- **RStudioServerPro**: The RStudioServerPro app is a multiuser app that is a shared resource among all user profiles in the domain. Once an RStudio app is created in a domain, the admin can give permissions to users in the domain.
- **RStudio user**: RStudio users are users within the domain that are authorized to use the RStudio license.
- **RStudio admin**: An RStudio on Amazon SageMaker admin can access the RStudio administrative dashboard. RStudio on Amazon SageMaker admins differ from "stock" Posit Workbench admins because they do not have root access to the instance running the RStudioServerPro app and can't modify the RStudio configuration file.
- **RStudio Server**: The RStudio Server instance is responsible for serving the RStudio UI to all authorized Users. This instance is launched on an Amazon SageMaker instance.
- **RSession**: An RSession is a browser-based interface to the RStudio IDE running on an Amazon SageMaker instance. Users can create and interact with their RStudio projects through the RSession.
- **RSessionGateway**: The RSessionGateway app is used to support an RSession.
- **RStudio administrative dashboard**: This dashboard gives information on the RStudio users in the Amazon SageMaker domain and their sessions. This dashboard can only be accessed by users that have RStudio admin authorization.

### Differences from Posit Workbench

RStudio on Amazon SageMaker has some significant differences from [Posit Workbench](#).

- When using RStudio on SageMaker, users don’t have access to the RStudio configuration files. Amazon SageMaker manages the configuration file and sets defaults. You can modify the RStudio Connect and RStudio Package Manager URLs when creating your RStudio-enabled Amazon SageMaker domain.
- Project sharing, realtime collaboration, and Job Launcher are not currently supported when using RStudio on Amazon SageMaker.
- When using RStudio on SageMaker, the RStudio IDE runs on Amazon SageMaker instances for on-demand containerized compute resources.
- RStudio on SageMaker only supports the RStudio IDE and does not support other IDEs supported by a Posit Workbench installation.
- RStudio on SageMaker only supports the RStudio version specified in [Upgrade the RStudio Version](#).

## Manage RStudio on Amazon SageMaker

The following topics give information on managing RStudio on Amazon SageMaker. This includes information on your RStudio environment configuration, user sessions, and necessary resources. For information on how to use RStudio on SageMaker, see [Use RStudio on Amazon SageMaker](#).

For information about creating a Amazon SageMaker domain with RStudio enabled, see [Amazon SageMaker domain overview](#).

For information about the AWS Regions that RStudio on SageMaker is supported in, see [Supported Regions and Quotas](#).

### Topics

- [RStudio license](#)
- [Upgrade the RStudio Version](#)
- [Network and Storage](#)
- [RStudioServerPro instance type](#)
- [RStudio Connect URL](#)
- [RStudio Package Manager](#)
- [Create an Amazon SageMaker domain with RStudio using the AWS CLI](#)
- [Add RStudio support to an existing domain](#)
- [Bring your own image to RStudio on SageMaker](#)
- [Manage users](#)
- [RStudio administrative dashboard](#)
- [Shut down and restart RStudio](#)
- [Manage billing and cost](#)
- [Diagnose issues and get support](#)

### RStudio license

RStudio on Amazon SageMaker is a paid product and requires that each user is appropriately licensed. Licenses for RStudio on Amazon SageMaker may be obtained from RStudio PBC directly,
or by purchasing a subscription to Posit Workbench on AWS Marketplace. For existing customers of Posit Workbench Enterprise, licenses are issued at no additional cost.

To use an RStudio license with Amazon SageMaker, you must first have a valid RStudio license registered with AWS License Manager. Subscriptions to Posit Workbench on AWS Marketplace automatically trigger license creation with AWS License Manager. For licenses purchased directly through RStudio PBC, a licenses grant for your AWS Account must be created. Contact RStudio for direct license purchases or to enable existing licenses in AWS License Manager. For more information about registering a license with AWS License Manager, see Seller issued licenses in AWS License Manager.

The following topics show how to acquire and validate a license granted by RStudio PBC.

**Get an RStudio license**

1. If you don't have an RStudio license, you may purchase one from the AWS Marketplace or from RStudio PBC directly.
   - To purchase a subscription from the AWS Marketplace, complete the steps in Subscribing to an AMI product with contract pricing public offer by searching for Posit Workbench.
   - To purchase from RStudio PBC directly, navigate to RStudio Pricing or contact sales@rstudio.com. When buying or updating an RStudio license, you must provide the AWS Account that will host your Amazon SageMaker domain.

   If you have an existing RStudio license, contact your RStudio Sales representative or sales@rstudio.com to add RStudio on Amazon SageMaker to your existing Posit Workbench Enterprise license, or to convert your Posit Workbench Standard license. The RStudio Sales representative will send you the appropriate electronic order form.

2. RStudio grants a Posit Workbench license to your AWS Account through AWS License Manager in the US East (N. Virginia) Region. Although the RStudio license is granted in the US East (N. Virginia) Region, your license can be consumed in any AWS Region that RStudio on Amazon SageMaker is supported in. You can expect the license grant process to complete within three business days after you share your AWS account ID with RStudio.

3. When this license is granted, you receive an email from your RStudio Sales representative with instructions to accept your license grant.

**Validate your RStudio license to be used with Amazon SageMaker**
1. Log into the AWS License Manager console in the same region as your Amazon SageMaker domain. If you are using AWS License Manager for the first time, AWS License Manager prompts you to grant permission to use AWS License Manager.

2. Select **Start using AWS License manager**.

3. Select **I grant AWS License Manager the required permissions** and select **Grant Permissions**.

4. Navigate to **Granted Licenses** on the left panel.

5. Select the license grant with RSW-SageMaker as the **Product name** and select **View**.

6. From the license detail page, select **Accept & activate license**.

**RStudio administrative dashboard**

You can use the RStudio administrative dashboard to see the number of users on the license following the steps in **RStudio administrative dashboard**.

**Upgrade the RStudio Version**

This guide provides information about the 2023.03.2-547.pro5 version update for RStudio on SageMaker. Starting February 27, 2024, new domains with RStudio support are created with Posit Workbench version 2023.03.2-547.pro5. This applies to the RStudioServerPro applications and default RSessionGateway applications.

The following sections give information about the 2023.03.2-547.pro5 release.

**Latest version updates**

The patch version 2023.03.2-547.pro5 release includes the following change:

- Fixed intermittent RServer crash when joining an RSession that was started with the job launcher and is not immediately available.

The latest RStudio version is 2023.03.2-454.pro2. This version includes the following changes:

- Added RTools 4.3 support
- Added support for R 4.3
- Upgraded Quarto to 1.2.335
- Improved session management
For more information about the changes in this release, see https://docs.posit.co/ide/news/.

**Note**

If you see the following warning, there is a version mismatch between the RSession and the Posit Workbench version used in RStudio on SageMaker. To resolve this issue, update the RStudio version for the domain. For information about updating the RStudio version, see Upgrade to the new version. Despite this warning, versions 2023.03.2-547.pro5 and 2023.03.2-454.pro2 are compatible images.

Session version 2023.03.2+454.pro2 does not match server version 2023.03.3-547.pro5 - this is an unsupported configuration, and you may experience unexpected issues as a result.

**Versioning**

There are currently two versions of Posit Workbench supported by SageMaker.

- Latest version supported: 2023.03.2-547.pro5
- Previous version supported: 2022.02.2-485.pro2

The default Posit Workbench version that's selected by SageMaker depends on the creation date of the domain.

- For domains created after February 27, 2024, version 2023.03.2-547.pro5 is the default selected version.
- For domains created after June 27, 2023 and before February 27, 2024, version 2023.03.2-454.pro2 is the default selected version. You can update your domains to the latest version (2023.03.2-547.pro5) by setting it as the default version for the domain. For more information, see Upgrade to the new version.
- For domains created before June 27, 2023, version 2022.02.2-485.pro2 is the default selected version. You can update your domains to the latest version (2023.03.2-547.pro5) by setting it as the default version for the domain. For more information, see Upgrade to the new version.
Note

The default RSessionGateway application version matches the current version of the RStudioServerPro application.

The following table lists the image ARNs for both versions for each AWS Region. These ARNs are passed as part of an update-domain command to set the desired version.

<table>
<thead>
<tr>
<th>Region</th>
<th>2022.02.2-485.pro2 Image ARN</th>
<th>2023.03.2-547.pro5 Image ARN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region</td>
<td>ARN for RStudio 2021.08</td>
<td>ARN for RStudio 2023.03</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------------------</td>
<td>-------------------------</td>
</tr>
</tbody>
</table>
Upgrade to the new version

Existing domains using version `2022.02.2-485.pro2` or `2023.03.2-454.pro2` can upgrade to `2023.03.2-547.pro5` version in one of two ways:

- Create a new domain from the AWS CLI with RStudio enabled.
- Update an existing domain to use the `2023.03.2-547.pro5` version.

The following procedure shows how to delete the RStudio application for an existing domain, set the default version to `2023.03.2-547.pro5`, and then create an RStudio application.

1. Delete the RStudioServerPro application and all RSessionGateway applications associated with your existing domain. For information about how to find your domain ID, see View domains. For more information about deleting applications, see Shut down and restart RStudio.

   ```bash
   aws sagemaker delete-app
   --region region
   --domain-id domainId
   --user-profile-name domain-shared
   --app-type RStudioServerPro
   --app-name default
   ```

2. If your domain is using RStudio version `2022.02.2-485.pro2`, update the domain to set `2023.03.2-547.pro5` as the default Posit Workbench version. The `SageMakerImageArn` value in the following update-domain command specifies the RStudio
2023.03.2-547.pro5 version as the default. This ARN must match the Region that your domain is in. For a list of all available ARNs, see Versioning.

Pass an execution role ARN for the domain that provides permissions to update the domain.

```
aws sagemaker update-domain \
  --region region \
  --domain-id domainId \
  --domain-settings-for-update "{"RStudioServerProDomainSettingsForUpdate":"{"DefaultResourceSpec": {"SageMakerImageArn": \"arn-for-2023.03.2-547.pro5-version\", "InstanceType": \"system\"}, "DomainExecutionRoleArn": \"execution-role-arn\"}}"
```

3. Create a new RStudioServerPro application in the existing domain.

```
aws sagemaker create-app \
  --region region \
  --domain-id domainId \
  --user-profile-name domain-shared \
  --app-type RStudioServerPro \
  --app-name default
```

Your RStudioServerPro application is now updated to version 2023.03.2-547.pro5. You can now relaunch your RSessionGateway applications.

**Downgrade to the existing version**

You can manually downgrade the version of your existing RStudio application to the 2022.02.2-485.pro2 version.

**To downgrade to the existing version**

1. Delete the RStudioServerPro application that's associated with your existing domain. For information about how to find your domain ID, see View domains.

```
aws sagemaker delete-app \
  --domain-id domainId \
  --user-profile-name domain-shared \
  --app-type RStudioServerPro \
  --app-name default
```
2. Pass the corresponding `2022.02.2-485.pro2` ARN for your Region as part of the update-domain command. For a list of all available ARNs, see Versioning. You must also pass an execution role ARN for the domain that provides permissions to update the domain.

```bash
aws sagemaker update-domain \
  --region region \
  --domain-id domainId \
  --domain-settings-for-update "{"RStudioServerProDomainSettingsForUpdate": {"DefaultResourceSpec": {"SageMakerImageArn": "arn-for-2022.02.2+485.pro2-version", "InstanceType": "system"}, "DomainExecutionRoleArn": "execution-role-arn"}}"
```

3. Create a new RStudioServerPro application in the existing domain. The RStudio version defaults to `2022.02.2-485.pro2`.

```bash
aws sagemaker create-app \
  --domain-id domainId \
  --user-profile-name domain-shared \
  --app-type RStudioServerPro \
  --app-name default
```

Your RStudioServerPro application is now downgraded to version `2022.02.2-485.pro2`.

**Changes to BYOI Images**

If you use a BYOI image with RStudio and update your RStudioServerPro version to `2023.03.2-547.pro5`, you must upgrade your custom images to use the `2023.03.2-547.pro5` release and redeploy your existing RSessions. If you attempt to load a non-compatible image in an RSession of a domain using the `2023.03.2-547.pro5` version, the RSession fails because it cannot parse parameters that it receives. To prevent failure, update all of the deployed custom images in your existing RStudioServerPro application.

The RSW_VERSION in the Dockerfile must be consistent with the Posit Workbench version used in RStudio on SageMaker. You can validate the current version in Posit Workbench. To do so, use the version name that's located in the lower left corner of the Posit Workbench launcher page.

```bash
... 
ARG RSW_VERSION=2023.03.3-547.pro5 
ENV RSTUDIO_FORCE_NON_ZERO_EXIT_CODE="1" 
ARG RSW_NAME=rstudio-workbench
```
ARG OS_CODE_NAME=bionic
ARG RSW_DOWNLOAD_URL=https://s3.amazonaws.com/rstudio-ide-build/server/${OS_CODE_NAME}/amd64
RUN RSW_VERSION_URL=`echo -n "${RSW_VERSION}" | sed 's/+/-/g'` \
    && curl -o rstudio-workbench.deb ${RSW_DOWNLOAD_URL}/${RSW_NAME}-${RSW_VERSION_URL}-amd64.deb \
    && gdebi -n ./rstudio-workbench.deb

Note

If you see the following warning, there is a version mismatch between the RSW_VERSION and the Posit Workbench version used in RStudio on SageMaker. Despite this warning, versions 2023.03.2-547.pro5 and 2023.03.2-454.pro2 are compatible images.

Session version 2023.03.2+454.pro2 does not match server version 2023.03.3-547.pro5 - this is an unsupported configuration, and you may experience unexpected issues as a result.

Network and Storage

The following topic describes network access and data storage considerations for your RStudio instance. For general information about network access and data storage when using Amazon SageMaker, see Data Protection in Amazon SageMaker.

Amazon EFS volume

RStudio on Amazon SageMaker shares an Amazon EFS volume with the Amazon SageMaker Studio Classic application in the domain. When the RStudio application is added to a domain, SageMaker creates a folder named shared in the Amazon EFS directory. If this shared folder is deleted or changed manually, then the RStudio application may no longer function. For more information about the Amazon EFS volume, see Manage Your Amazon EFS Storage Volume in SageMaker Studio Classic.

Installed packages and scripts

Packages that you install from within RStudio are scoped to the user profile level. This means that the installed package persists through RSession shut down, restarts, and across RSessions for each user profile that they are installed in. R Scripts that are saved in RSessions behave the same way. Both packages and R Scripts are saved in the user's Amazon EFS volume.
Encryption

RStudio on Amazon SageMaker supports encryption at rest.

Use RStudio in VPC-only mode

RStudio on Amazon SageMaker supports AWS PrivateLink integration. With this integration, you can use RStudio on SageMaker in VPC-only mode without direct access to the internet. When you use RStudio in VPC-only mode, your security groups are automatically managed by the service. This includes connectivity between your RServer and your RSessions.

The following are required to use RStudio in VPC-only mode. For more information on selecting a VPC, see Choose an Amazon VPC.

- A private subnet with either access the internet to make a call to Amazon SageMaker & License Manager, or Amazon Virtual Private Cloud (Amazon VPC) endpoints for both Amazon SageMaker & License Manager.
- The domain cannot have any more than two associated Security Groups.
- A Security Group ID for use with the domain in domain Settings. This must allow all outbound access.
- A Security Group ID for use with the Amazon VPC endpoint. This security group must allow inbound traffic from the domain Security Group ID.
- Amazon VPC Endpoint for sagemaker.api and AWS License Manager. This must be in the same Amazon VPC as the private subnet.

RStudioServerPro instance type

When deciding which Amazon EC2 instance type to use for your RStudioServerPro app, the main factor to consider is bandwidth. Bandwidth is important because the RStudioServerPro instance is responsible for serving the RStudio UI to all users. This includes UI heavy workflows, such as generating figures, animations, and displaying many data rows. Therefore, there may be some UI performance degradation depending on the workload across all users. The following are the available instance types to use for your RStudioServerPro. For pricing information about these instances, see Amazon SageMaker Pricing.

- `ml.t3.medium`: This instance type is recommended for Domains with low UI use and is free to use.
Note

The system value is translated to ml.t3.medium.

- `ml.c5.4xlarge`: This instance type is recommended for Domains with moderate UI use.
- `ml.c5.9xlarge`: This instance type is recommended for Domains with heavy UI use.

**Changing RStudio instance type**

To change the instance type of your RStudioServerPro, pass the new instance type as part of a call to the `update-domain` CLI command. You then need to delete the existing RStudioServerPro app using the `delete-app` CLI command and create a new RStudioServerPro app using the `create-app` CLI command.

**RStudio Connect URL**

RStudio Connect is a publishing platform for Shiny applications, R Markdown reports, dashboards, plots, and more. RStudio Connect makes it easy to surface machine learning and data science insights by making hosting content simple and scalable. If you have an RStudio Connect server, then you can set the server as the default place where apps are published. For more information about RStudio Connect, see [RStudio Connect](#).

When you onboard to RStudio on Amazon SageMaker domain, an RStudio Connect server is not created. You can create an RStudio Connect server on an Amazon EC2 instance to use Connect with Amazon SageMaker domain. For information about how to set up your RStudio Connect server, see [Host RStudio Connect and Package Manager for ML development in RStudio on Amazon SageMaker](#).

**Add an RStudio Connect URL**

If you have an RStudio Connect URL, you can update the default URL so that your RStudio Users can publish to it.

1. Navigate to the **domains** page.
2. Select the desired domain.
3. Choose **domain Settings**.
4. Under **General Settings**, select **Edit**.
5. From the new page, select **RStudio Settings** on the left side.

6. Under **RStudio Connect URL**, enter the RStudio Connect URL to add.

7. Select **Submit**.

---

**CLI**

You can set a default RStudio Connect URL when you create your domain. The only way to update your RStudio Connect URL from the AWS CLI is to delete your domain and create a new one with the updated RStudio Connect URL.

**RStudio Package Manager**

RStudio Package Manager is a repository management server used to organize and centralize packages across your organization. For more information on RStudio Package Manager, see [RStudio Package Manager](https://rstudio.com/package-manager/). If you don't supply your own Package Manager URL, Amazon SageMaker domain uses the default Package Manager repository when you onboard RStudio following the steps in [Amazon SageMaker domain overview](https://docs.aws.amazon.com/sagemaker/latest/dg/domain-overview.html). For more information, see [Host RStudio Connect and Package Manager for ML development in RStudio on Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/host-rstudio-package-manager.html).

**Update Package Manager URL**

You can update the Package Manager URL used for your RStudio-enabled domain as follows.

1. Navigate to the **domains** page.
2. Select the desired domain.
3. Choose **domain Settings**.
4. Under **General Settings**, select **Edit**.
5. From the new page, select **RStudio Settings** on the left side.
6. Under **RStudio Package Manager**, enter your RStudio Package Manager URL.
7. Select **Submit**.

**CLI**

The only way to update your Package Manager URL from the AWS CLI is to delete your domain and create a new one with the updated Package Manager URL.
Create an Amazon SageMaker domain with RStudio using the AWS CLI

The following topic shows how to onboard to Amazon SageMaker domain with RStudio enabled using the AWS CLI. To onboard using the AWS Management Console, see Amazon SageMaker domain overview.

Prerequisites

- Install and configure AWS CLI version 2
- Configure the AWS CLI with IAM credentials

Create DomainExecution role

To launch the RStudio App, you must provide a DomainExecution role. This role is used to determine whether RStudio needs to be launched as part of Amazon SageMaker domain creation. This role is also used by Amazon SageMaker to access the RStudio License and push RStudio logs.

Note

The DomainExecution role should have at least AWS License Manager permissions to access RStudio License, and CloudWatch permissions to push logs in your account.

The following procedure shows how to create the DomainExecution role with the AWS CLI.

1. Create a file named assume-role-policy.json with the following content.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Action": "sts:AssumeRole",
            "Effect": "Allow",
            "Principal": {
                "Service": ["sagemaker.amazonaws.com"]
            }
        }
    ]
}
```
2. Create the DomainExecution role. `<REGION>` should be the AWS Region to launch your domain in.

```bash
aws iam create-role --region <REGION> --role-name DomainExecution --assume-role-policy-document file://assume-role-policy.json
```

3. Create a file named `domain-setting-policy.json` with the following content. This policy allows the RStudioServerPro app to access necessary resources and allows Amazon SageMaker to automatically launch an RStudioServerPro app when the existing RStudioServerPro app is in a Deleted or Failed status.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "VisualEditor0",
            "Effect": "Allow",
            "Action": [
                "license-manager:ExtendLicenseConsumption",
                "license-manager:ListReceivedLicenses",
                "license-manager:GetLicense",
                "license-manager:CheckoutLicense",
                "license-manager:CheckInLicense",
                "logs:CreateLogDelivery",
                "logs:CreateLogGroup",
                "logs:CreateLogStream",
                "logs:DeleteLogDelivery",
                "logs:Describe*",
                "logs:GetLogDelivery",
                "logs:GetLogEvents",
                "logs:ListLogDeliveries",
                "logs:PutLogEvents",
                "logs:PutResourcePolicy",
                "logs:UpdateLogDelivery",
                "sagemaker:CreateApp"
            ],
            "Resource": "*"
        }
    ]
}
```
4. Create the domain setting policy that is attached to the DomainExecution role. Be aware of the PolicyArn from the response, you will need to enter that ARN in the following steps.

```bash
aws iam create-policy --region <REGION> --policy-name domain-setting-policy --policy-document file://domain-setting-policy.json
```

5. Attach `domain-setting-policy` to the DomainExecution role. Use the PolicyArn returned in the previous step.

```bash
aws iam attach-role-policy --role-name DomainExecution --policy-arn <POLICY_ARN>
```

---

**Create Amazon SageMaker domain with RStudio App**

The RStudioServerPro app is launched automatically when you create a Amazon SageMaker domain using the `create-domain` CLI command with the `RStudioServerProDomainSettings` parameter specified. When launching the RStudioServerPro App, Amazon SageMaker checks for a valid RStudio license in the account and fails domain creation if the license is not found.

The creation of a Amazon SageMaker domain differs based on the authentication method and the network type. These options must be used together, with one authentication method and one network connection type selected. For more information about the requirements to create a new domain, see [CreateDomain](#).

The following authentication methods are supported.

- IAM Auth
- SSO Auth

The following network connection types are supported:

- PublicInternet
- VPCOnly

**Authentication methods**

**IAM Auth Mode**
The following shows how to create an Amazon SageMaker domain with RStudio enabled and an IAM Auth Network Type. For more information about AWS Identity and Access Management, see What is IAM?

- DomainExecutionRoleArn should be the ARN for the role created in the previous step.
- ExecutionRole is the ARN of the role given to users in the Amazon SageMaker domain.
- vpc-id should be the ID of your Amazon Virtual Private Cloud. subnet-ids should be a space-separated list of subnet IDs. For information about vpc-id and subnet-ids, see VPCs and subnets.
- RStudioPackageManagerUrl and RStudioConnectUrl are optional and should be set to the URLs of your RStudio Package Manager and RStudio Connect server, respectively.
- app-network-access-type should be either PublicInternetOnly or VPCOnly.

```bash
aws sagemaker create-domain --region <REGION> --domain-name <DOMAIN_NAME> \
  --auth-mode IAM \
  --default-user-settings ExecutionRole=<DEFAULT_USER_EXECUTIONROLE> \
  --domain-settings
RStudioServerProDomainSettings={RStudioPackageManagerUrl=<PACKAGE_MANAGER_URL>,RStudioConnectUrl=<CONNECT_URL>,DomainExecutionRoleArn=<DOMAINEXECUTION_ROLE_ARN} \
  --vpc-id <VPC_ID> \
  --subnet-ids <SUBNET_IDS> \
  --app-network-access-type <NETWORK_ACCESS_TYPE>
```

Authentication using IAM Identity Center

The following shows how to create an Amazon SageMaker domain with RStudio enabled and an SSO Auth Network Type. AWS IAM Identity Center must be enabled for the region that the domain is launched on. For more information about IAM Identity Center, see What is AWS IAM Identity Center?

- DomainExecutionRoleArn should be the ARN for the role created in the previous step.
- ExecutionRole is the ARN of the role given to users in the Amazon SageMaker domain.
- vpc-id should be the ID of your Amazon Virtual Private Cloud. subnet-ids should be a space-separated list of subnet IDs. For information about vpc-id and subnet-ids, see VPCs and subnets.
- RStudioPackageManagerUrl and RStudioConnectUrl are optional and should be set to the URLs of your RStudio Package Manager and RStudio Connect server, respectively.
• app-network-access-type should be either PublicInternetOnly or VPCOnly.

```
aws sagemaker create-domain --region <REGION> --domain-name <DOMAIN_NAME> \
   --auth-mode SSO \ 
   --default-user-settings ExecutionRole=<DEFAULT_USER_EXECUTIONROLE> \ 
   --domain-settings
RStudioServerProDomainSettings={RStudioPackageManagerUrl=<PACKAGE_MANAGER_URL>,RStudioConnectUrl=<CONNECT_URL>,DomainExecutionRoleArn=<DOMAINEXECUTION_ROLE_ARN},
\ 
   --vpc-id <VPC_ID> \
   --subnet-ids <SUBNET_IDS> \
   --app-network-access-type <NETWORK_ACCESS_TYPE>
```

Connection types

PublicInternet/Direct Internet network type

The following shows how to create a Amazon SageMaker domain with RStudio enabled and a PublicInternet Network Type.

• DomainExecutionRoleArn should be the ARN for the role created in the previous step.
• ExecutionRole is the ARN of the role given to users in the Amazon SageMaker domain.
• vpc-id should be the ID of your Amazon Virtual Private Cloud. subnet-ids should be a space-separated list of subnet IDs. For information about vpc-id and subnet-ids, see VPCs and subnets.
• RStudioPackageManagerUrl and RStudioConnectUrl are optional and should be set to the URLs of your RStudio Package Manager and RStudio Connect server, respectively.
• auth-mode should be either SSO or IAM.

```
aws sagemaker create-domain --region <REGION> --domain-name <DOMAIN_NAME> \
   --auth-mode <AUTH_MODE> \ 
   --default-user-settings ExecutionRole=<DEFAULT_USER_EXECUTIONROLE> \ 
   --domain-settings
RStudioServerProDomainSettings={RStudioPackageManagerUrl=<PACKAGE_MANAGER_URL>,RStudioConnectUrl=<CONNECT_URL>,DomainExecutionRoleArn=<DOMAINEXECUTION_ROLE_ARN},
\ 
   --vpc-id <VPC_ID> \
   --subnet-ids <SUBNET_IDS> \
   --app-network-access-type PublicInternetOnly
```
VPCOnly mode

The following shows how to launch an Amazon SageMaker domain with RStudio enabled and a VPCOnly Network Type. For more information about using the VPCOnly network access type, see [Connect SageMaker Studio Notebooks in a VPC to External Resources](#).

- `DomainExecutionRoleArn` should be the ARN for the role created in the previous step.
- `ExecutionRole` is the ARN of the role given to users in the Amazon SageMaker domain.
- `vpc-id` should be the ID of your Amazon Virtual Private Cloud. `subnet-ids` should be a space-separated list of subnet IDs. Your private subnet must be able to either access the internet to make a call to Amazon SageMaker, and AWS License Manager or have Amazon VPC endpoints for both Amazon SageMaker and AWS License Manager. For information about Amazon VPC endpoints, see [Interface Amazon VPC endpoints](#). For information about `vpc-id` and `subnet-ids`, see [VPCs and subnets](#).
- `SecurityGroups` must allow outbound access to the Amazon SageMaker and AWS License Manager endpoints.
- `auth-mode` should be either SSO or IAM.

### Note

When using Amazon Virtual Private Cloud endpoints, the security group attached to your Amazon Virtual Private Cloud endpoints must allow inbound traffic from the security group you pass as part of the domain-setting parameter of the create-domain CLI call.

With RStudio, Amazon SageMaker manages security groups for you. This means that Amazon SageMaker manages security group rules to ensure RSessions can access RStudioServerPro Apps. Amazon SageMaker creates one security group rule per user profile.

```bash
aws sagemaker create-domain --region <REGION> --domain-name <DOMAIN_NAME> \
    --auth-mode <AUTH_MODE> \
    --default-user-settings
    SecurityGroups=<USER_SECURITY_GROUP>,ExecutionRole=<DEFAULT_USER_EXECUTIONROLE> \
    --domain-settings
    SecurityGroupId=<DOMAIN_SECURITY_GROUP>,RStudioServerProDomainSettings={DomainExecutionRoleArn=\ 
        --vpc-id <VPC_ID> \n```
Note: The RStudioServerPro app is launched by a special user profile named domain-shared. As a result, this app is not returned as part of list-app API calls by any other user profiles.

You may have to increase the Amazon VPC quota in your account to increase the number of users. For more information, see Amazon VPC quotas.

Verify domain creation

Use the following command to verify that your domain has been created with a Status of InService. Your domain-id is appended to the domains ARN. For example, arn:aws:sagemaker:<REGION>:<ACCOUNT_ID>:domain/<DOMAIN_ID>.

```
aws sagemaker describe-domain --domain-id <DOMAIN_ID> --region <REGION>
```

Add RStudio support to an existing domain

If you have added an RStudio License through AWS License Manager, you can create a new Amazon SageMaker domain with support for RStudio on SageMaker. If you have an existing domain that does not support RStudio, you can add RStudio support to that domain without having to delete and recreate the domain.

The following topic outlines how to add this support.

Prerequisites

You must complete the following steps before you update your current domain to add support for RStudio on SageMaker.

- Install and configure AWS CLI version 2
- Configure the AWS CLI with IAM credentials
- Create a domain execution role following the steps in Create a SageMaker Domain with RStudio using the AWS CLI. This domain-level IAM role is required by the RStudioServerPro app. The role requires access to AWS License Manager for verifying a valid Posit Workbench license and Amazon CloudWatch Logs for publishing server logs.
- Bring your RStudio license to AWS License Manager following the steps in RStudio license.
• (Optional) If you want to use RStudio in VPCOnly mode, complete the steps in [RStudio in VPC-Only](#).

• Ensure that the security groups you have configured for each [UserProfile](#) in your domain meet the account-level quotas. When configuring the default user profile during domain creation, you can use the `DefaultUserSettings` parameter of the [CreateDomain](#) API to add `SecurityGroups` that are inherited by all the user profiles created in the domain. You can also provide additional security groups for a specific user as part of the `UserSettings` parameter of the [CreateUserProfile](#) API. If you have added security groups this way, you must ensure that the total number of security groups per user profile doesn't exceed the maximum quota of 2 in VPCOnly mode and 4 in PublicInternetOnly mode. If the resulting total number of security groups for any user profile exceeds the quota, you can combine multiple security groups' rules into one security group.

Add RStudio support to an existing domain

After you have completed the prerequisites, you can add RStudio support to your existing domain. The following steps outline how to update your existing domain to add support for RStudio.

**Step 1: Delete all apps in the domain**

To add support for RStudio in your domain, SageMaker must update the underlying security groups for all existing user profiles. To complete this, you must delete and recreate all existing apps in the domain. The following procedure shows how to delete all of the apps.

1. List all of the apps in the domain.

   ```bash
   aws sagemaker list-apps --domain-id-equals <DOMAIN_ID>
   ```

2. Delete each app for each user profile in the domain.

   ```bash
   // JupyterServer apps
   aws sagemaker delete-app
   --domain-id <DOMAIN_ID> \
   --user-profile-name <USER_PROFILE> \
   --app-type JupyterServer \
   --app-name <APP_NAME>
   ```
// KernelGateway apps
aws sagemaker \
  delete-app \
  --domain-id <DOMAIN_ID> \
  --user-profile-name <USER_PROFILE> \
  --app-type KernelGateway \
  --app-name <APP_NAME>

**Step 2 - Update all user profiles with the new list of security groups**

This is a one-time action that you must complete for all of the existing user profiles in your domain when you have refactored your existing security groups. This prevents you from hitting the quota for the maximum number of security groups. The UpdateUserProfile API call fails if the user has any apps that are in **InService** status. Delete all apps, then call UpdateUserProfile API to update the security groups.

⚠️ **Note**

The following requirement for VPCOnly mode outlined in [Connect Amazon SageMaker Studio Classic Notebooks in a VPC to External Resources](https://docs.aws.amazon.com/sagemaker/latest/dg/connect-vpc.html) is no longer needed when adding RStudio support because AppSecurityGroupManagement is managed by the SageMaker service:

"**TCP traffic within the security group.** This is required for connectivity between the JupyterServer app and the KernelGateway apps. You must allow access to at least ports in the range 8192–65535."

```
aws sagemaker \
  update-user-profile \
  --domain-id <DOMAIN_ID> \
  --user-profile-name <USER_PROFILE> \
  --user-settings "{"SecurityGroups": ["<SECURITY_GROUP>", 
"<SECURITY_GROUP>"]}"
```

**Step 3 - Activate RStudio by calling the UpdateDomain API**

1. Call the [UpdateDomain](https://docs.aws.amazon.com/sagemaker/latest/dg/API_UpdateDomain.html) API to add support for RStudio on SageMaker. The `defaultusersettings` parameter is only needed if you have refactored the default security groups for your user profiles.
• For VPCOnly mode:

```bash
aws sagemaker \
  update-domain \
  --domain-id <DOMAIN_ID> \
  --app-security-group-management Service \
  --domain-settings-for-update
RStudioServerProDomainSettingsForUpdate={DomainExecutionRoleArn=<DOMAIN_EXECUTION_ROLE_ARN> \
  --default-user-settings "{"SecurityGroups": ["<SECURITY_GROUP>", 
  "<SECURITY_GROUP>"}}"
```

• For PublicInternetOnly mode:

```bash
aws sagemaker \
  update-domain \
  --domain-id <DOMAIN_ID> \
  --domain-settings-for-update
RStudioServerProDomainSettingsForUpdate={DomainExecutionRoleArn=<DOMAIN_EXECUTION_ROLE_ARN> \
  --default-user-settings "{"SecurityGroups": ["<SECURITY_GROUP>", 
  "<SECURITY_GROUP>"}}"
```

2. Verify that the domain status is InService. After the domain status is InService, support for RStudio on SageMaker is added.

```bash
aws sagemaker \
  describe-domain \
  --domain-id <DOMAIN_ID>
```

3. Verify that the RStudioServerPro app’s status is InService using the following command.

```bash
aws sagemaker list-apps --user-profile-name domain-shared
```

**Step 4 - Add RStudio access for existing users**

As part of the update in Step 3, SageMaker marks the RStudio AccessStatus of all existing user profiles in the domain as DISABLED by default. This prevents exceeding the number of users allowed by your current license. To add access for existing users, there is a one-time opt-in step. Perform the opt-in by calling the UpdateUserProfile API with the following RStudioServerProAppSettings:
• **AccessStatus** = ENABLED

• *Optional* - **UserGroup** = R_STUDIO_USER or R_STUDIO_ADMIN

```
aws sagemaker \
  update-user-profile \
  --domain-id <DOMAIN_ID> \
  --user-profile-name <USER_PROFILE> \
  --user-settings "{"RStudioServerProAppSettings": {"AccessStatus": "ENABLED"}}"
```

⚠️ **Note**

By default, the number of users that can have access to RStudio is 60.

**Step 5 – Deactivate RStudio access for new users**

Unless otherwise specified when calling `UpdateDomain`, RStudio support is added by default for all new user profiles created after you have added support for RStudio on SageMaker. To deactivate access for a new user profile, you must explicitly set the **AccessStatus** parameter to **DISABLED** as part of the `CreateUserProfile` API call. If the **AccessStatus** parameter is not specified as part of the `CreateUserProfile` API, the default access status is **ENABLED**.

```
aws sagemaker \
  create-user-profile \
  --domain-id <DOMAIN_ID> \
  --user-profile-name <USER_PROFILE> \
  --user-settings "{"RStudioServerProAppSettings": {"AccessStatus": "DISABLED"}}"
```

**Bring your own image to RStudio on SageMaker**

A SageMaker image is a file that identifies language packages and other dependencies that are required to run RStudio on Amazon SageMaker. SageMaker uses these images to create an environment where you run RStudio. Amazon SageMaker provides a built-in RStudio image for you to use. If you need different functionality, you can bring your own custom images.

The process to bring your own image to use with RStudio on SageMaker takes three steps:
1. Build a custom image from a Dockerfile and push it to a repository in Amazon Elastic Container Registry (Amazon ECR).

2. Create a SageMaker image that points to a container image in Amazon ECR and attach it to your Amazon SageMaker domain.

3. Launch a new session in RStudio with your custom image.

You can create images and image versions, and attach image versions to your domain, using the SageMaker control panel, the AWS SDK for Python (Boto3), and the AWS Command Line Interface (AWS CLI). You can also create images and image versions using the SageMaker console, even if you haven't onboarded to a domain.

The following topics show how to bring your own image to RStudio on SageMaker by creating, attaching, and launching a custom image.

**Key terminology**

The following section defines key terms for bringing your own image to use with RStudio on SageMaker.

- **Dockerfile**: A Dockerfile is a file that identifies the language packages and other dependencies for your Docker image.

- **Docker image**: The Docker image is a built Dockerfile. This image is checked into Amazon ECR and serves as the basis of the SageMaker image.

- **SageMaker image**: A SageMaker image is a holder for a set of SageMaker image versions based on Docker images.

- **Image version**: An image version of a SageMaker image represents a Docker image that is compatible with RStudio and stored in an Amazon ECR repository. Each image version is immutable. These image versions can be attached to a domain and used with RStudio on SageMaker.

**Prerequisites**

You must complete the following prerequisites before bringing your own image to use with RStudio on Amazon SageMaker.
• If you have an existing domain with RStudio that was created before April 7, 2022, you must delete your RStudioServerPro application and recreate it. For information about how to delete an application, see Shut down and Update SageMaker Studio Classic.

• Install the Docker application. For information about setting up Docker, see Orientation and setup.

• Create a local copy of an RStudio-compatible Dockerfile that works with SageMaker. For information about creating a sample RStudio dockerfile, see Use a custom image to bring your own development environment to RStudio on Amazon SageMaker.

• Use an AWS Identity and Access Management execution role that has the AmazonSageMakerFullAccess policy attached. If you have onboarded to domain, you can get the role from the domain Summary section of the SageMaker control panel.

Add the following permissions to access the Amazon Elastic Container Registry (Amazon ECR) service to your execution role.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "VisualEditor0",
         "Effect": "Allow",
         "Action": [
            "ecr:CreateRepository",
            "ecr:BatchGetImage",
            "ecr:CompleteLayerUpload",
            "ecr:DescribeImages",
            "ecr:DescribeRepositories",
            "ecr:UploadLayerPart",
            "ecr:ListImages",
            "ecr:InitiateLayerUpload",
            "ecr:BatchCheckLayerAvailability",
            "ecr:PutImage"
         ],
         "Resource": "*"
      }
   ]
}
```

• Install and configure AWS CLI with the following (or higher) version. For information about installing the AWS CLI, see Installing or updating the latest version of the AWS CLI.
Custom RStudio image specifications

In this guide, you'll learn custom RStudio image specifications to use when you bring your own image. There are two sets of requirements that you must satisfy with your custom RStudio image to use it with Amazon SageMaker. These requirements are imposed by RStudio PBC and the Amazon SageMaker Studio Classic platform. If either of these sets of requirements aren't satisfied, then your custom image won't function properly.

RStudio PBC requirements

RStudio PBC requirements are laid out in the Using Docker images with RStudio Workbench / RStudio Server Pro, Launcher, and Kubernetes article. Follow the instructions in this article to create the base of your custom RStudio image.

For instructions about how to install multiple R versions in your custom image, see Installing multiple versions of R on Linux.

Amazon SageMaker Studio Classic requirements

Amazon SageMaker Studio Classic imposes the following set of installation requirements for your RStudio image.

- You must use an RStudio base image of at least 2023.03.2-454.pro2. For more information, see Upgrade the RStudio Version.
- You must install the following packages:

```bash
yum install -y sudo 
openjdk-11-jdk 
libpng-dev 
&& yum clean all 
&& /opt/R/\${R_VERSION}/bin/R -e "install.packages('reticulate', repos='https://packagemanager.rstudio.com/cran/__linux__/centos7/latest')" 
&& /opt/python/\${PYTHON_VERSION}/bin/pip install --upgrade 
  'boto3>1.0<2.0' 
  'awscli>1.0<2.0' 
  'sagemaker[local]<3'
```
• You must provide default values for the RSTUDIO_CONNECT_URL and RSTUDIO_PACKAGE_MANAGER_URL environment values.

```bash
ENV RSTUDIO_CONNECT_URL "YOUR_CONNECT_URL"
ENV RSTUDIO_PACKAGE_MANAGER_URL "YOUR_PACKAGE_MANAGER_URL"
ENV RSTUDIO_FORCE_NON_ZERO_EXIT_CODE 1
```

The following general specifications apply to the image that is represented by an RStudio image version.

**Running the image**

ENTRYPOINT and CMD instructions are overridden so that the image is run as an RSession application.

**Stopping the image**

The DeleteApp API issues the equivalent of a docker stop command. Other processes in the container won’t get the SIGKILL/SIGTERM signals.

**File system**

The /opt/.sagemakerinternal and /opt/ml directories are reserved. Any data in these directories might not be visible at runtime.

**User data**

Each user in a SageMaker domain gets a user directory on a shared Amazon Elastic File System volume in the image. The location of the current user’s directory on the Amazon Elastic File System volume is /home/sagemaker-user.

**Metadata**

A metadata file is located at /opt/ml/metadata/resource-metadata.json. No additional environment variables are added to the variables defined in the image. For more information, see [Get App Metadata](#).

**GPU**

On a GPU instance, the image is run with the --gpus option. Only the CUDA toolkit should be included in the image, not the NVIDIA drivers. For more information, see [NVIDIA User Guide](#).
Metrics and logging

Logs from the RSession process are sent to Amazon CloudWatch in the customer's account. The name of the log group is /aws/sagemaker/studio. The name of the log stream is $domainID/$userProfileName/RSession/$appName.

Image size

Image size is limited to 25 GB. To view the size of your image, run docker image ls.

Create a custom RStudio image

This topic describes how you can create a custom RStudio image using the SageMaker console and the AWS CLI. If you use the AWS CLI, you must run the steps from your local machine. The following steps do not work from within Amazon SageMaker Studio Classic.

When you create an image, SageMaker also creates an initial image version. The image version represents a container image in Amazon Elastic Container Registry (ECR). The container image must satisfy the requirements to be used in RStudio. For more information, see Custom RStudio image specifications.

For information about testing your image locally and resolving common issues, see the SageMaker Studio Custom Image Samples repo.

Topics

- Add a SageMaker-compatible RStudio Docker container image to Amazon ECR
- Create a SageMaker image from the console
- Create an image from the AWS CLI

Add a SageMaker-compatible RStudio Docker container image to Amazon ECR

Use the following steps to add a Docker container image to Amazon ECR:

- Create an Amazon ECR repository.
- Authenticate to Amazon ECR.
- Build a SageMaker-compatible RStudio Docker image.
- Push the image to the Amazon ECR repository.
Note

The Amazon ECR repository must be in the same AWS Region as your domain.

To build and add a Docker image to Amazon ECR

1. Create an Amazon ECR repository using the AWS CLI. To create the repository using the Amazon ECR console, see Creating a repository.

   ```bash
   aws ecr create-repository \
   --repository-name rstudio-custom \
   --image-scanning-configuration scanOnPush=true
   ```

   Response:

   ```json
   {
     "repository": {
       "repositoryArn": "arn:aws:ecr:us-east-2:acct-id:repository/rstudio-custom",
       "registryId": "acct-id",
       "repositoryName": "rstudio-custom",
       "repositoryUri": "acct-id.dkr.ecr.us-east-2.amazonaws.com/rstudio-custom",
       ...
     }
   }
   ```

2. Authenticate to Amazon ECR using the repository URI returned as a response from the create-repository command. Make sure that the Docker application is running. For more information, see Registry Authentication.

   ```bash
   aws ecr get-login-password | \n   docker login --username AWS --password-stdin <repository-uri>
   ```

   Response:

   Login Succeeded

3. Build the Docker image. Run the following command from the directory that includes your Dockerfile.
Create a SageMaker image from the console

To create an image

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
4. On the Custom images page, choose Create image.
5. For Image source, enter the registry path to the container image in Amazon ECR. The path is in the following format:

   `acct-id.dkr.ecr.region.amazonaws.com/repo-name[:tag] or [@digest]`

6. Choose Next.
7. Under Image properties, enter the following:
   - Image name – The name must be unique to your account in the current AWS Region.
   - (Optional) Image display name – The name displayed in the domain user interface. When not provided, Image name is displayed.
   - (Optional) Description – A description of the image.
• IAM role – The role must have the AmazonSageMakerFullAccess policy attached. Use the dropdown menu to choose one of the following options:
  • Create a new role – Specify any additional Amazon Simple Storage Service (Amazon S3) buckets that you want your notebooks users to access. If you don't want to allow access to additional buckets, choose None.
    SageMaker attaches the AmazonSageMakerFullAccess policy to the role. The role allows your notebook users to access the Amazon S3 buckets listed next to the check marks.
  • Enter a custom IAM role ARN – Enter the Amazon Resource Name (ARN) of your IAM role.
  • Use existing role – Choose one of your existing roles from the list.
  • (Optional) Image tags – Choose Add new tag. You can add up to 50 tags. Tags are searchable using the SageMaker console or the SageMaker Search API.

8. Under Image type, select RStudio image.
9. Choose Submit.

The new image is displayed in the Custom images list and briefly highlighted. After the image has been successfully created, you can choose the image name to view its properties or choose Create version to create another version.

To create another image version

1. Choose Create version on the same row as the image.
2. For Image source, enter the registry path to the Amazon ECR image. The image shouldn't be the same image as used in a previous version of the SageMaker image.

To use the custom image in RStudio, you must attach it to your domain. For more information, see Attach a custom SageMaker image.

Create an image from the AWS CLI

This section shows how to create a custom Amazon SageMaker image using the AWS CLI.

Use the following steps to create a SageMaker image:

• Create an Image.
• Create an ImageVersion.
• Create a configuration file.
• Create an AppImageConfig.

To create the SageMaker image entities

1. Create a SageMaker image. The role ARN must have at least the AmazonSageMakerFullAccessPolicy policy attached.

   ```
   aws sagemaker create-image \
     --image-name rstudio-custom-image \
     --role-arn arn:aws:iam::<acct-id>:role/service-role/<execution-role>
   ```

   Response:
   ```
   {
     "ImageArn": "arn:aws:sagemaker:us-east-2:<acct-id>:image/rstudio-custom-image"
   }
   ```

2. Create a SageMaker image version from the image. Pass the unique tag value that you chose when you pushed the image to Amazon ECR.

   ```
   aws sagemaker create-image-version \
     --image-name rstudio-custom-image \
     --base-image <repository-uri>::<tag>
   ```

   Response:
   ```
   {
     "ImageVersionArn": "arn:aws:sagemaker:us-east-2:<acct-id>:image-version/rstudio-image/1"
   }
   ```

3. Check that the image version was successfully created.

   ```
   aws sagemaker describe-image-version \
     --image-name rstudio-custom-image \
     --version 1
   ```

   Response:
4. Create a configuration file, named `app-image-config-input.json`. The app image config is used to configuration for running a SageMaker image as a Kernel Gateway application.

```json
{
   "AppImageConfigName": "rstudio-custom-config"
}
```

5. Create the AppImageConfig using the file that you created in the previous step.

```bash
aws sagemaker create-app-image-config \
   --cli-input-json file://app-image-config-input.json
```

Response:

```json
{
}
```

**Attach a custom SageMaker image**

This guide shows how to attach a custom RStudio image to your Amazon SageMaker domain using the SageMaker console or the AWS Command Line Interface (AWS CLI).
To use a custom SageMaker image, you must attach a custom RStudio image to your domain. When you attach an image version, it appears in the RStudio Launcher and is available in the **Select image** dropdown list. You use the dropdown to change the image used by RStudio.

There is a limit to the number of image versions that you can attach. After you reach the limit, you must first detach a version so that you can attach a different version of the image.

**Topics**

- [Attach an image version to your domain using the console](#)
- [Attach an existing image version to your domain using the AWS CLI](#)

**Attach an image version to your domain using the console**

You can attach a custom SageMaker image version to your domain using the SageMaker console's control panel. You can also create a custom SageMaker image, and an image version, and then attach that version to your domain.

**To attach an existing image**

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. Select the desired domain.
5. Choose **Environment**.
6. Under **Custom SageMaker Studio Classic images attached to domain**, choose **Attach image**.
7. For **Image source**, choose **Existing image** or **New image**.

   If you select **Existing image**, choose an image from the Amazon SageMaker image store.

   If you select **New image**, provide the Amazon ECR registry path for your Docker image. The path must be in the same AWS Region as the domain. The Amazon ECR repo must be in the same account as your domain, or cross-account permissions for SageMaker must be enabled.

8. Choose an existing image from the list.
9. Choose a version of the image from the list.
10. Choose **Next**.
11. Enter values for **Image name**, **Image display name**, and **Description**.
12. Choose the IAM role. For more information, see Create a custom RStudio image.

13. (Optional) Add tags for the image.

14. (Optional) Choose Add new tag, then add a configuration tag.

15. For Image type, select RStudio Image.

16. Choose Submit.

Wait for the image version to be attached to the domain. After the version is attached, it appears in the Custom images list and is briefly highlighted.

**Attach an existing image version to your domain using the AWS CLI**

Two methods are presented to attach the image version to your domain using the AWS CLI. In the first method, you create a new domain with the version attached. This method is simpler but you must specify the Amazon Virtual Private Cloud (Amazon VPC) information and execution role that's required to create the domain.

If you have already onboarded to the domain, you can use the second method to attach the image version to your current domain. In this case, you don't need to specify the Amazon VPC information and execution role. After you attach the version, delete all of the applications in your domain and relaunch RStudio.

**Attach the SageMaker image to a new domain**

To use this method, you must specify an execution role that has the AmazonSageMakerFullAccess policy attached.

Use the following steps to create the domain and attach the custom SageMaker image:

- Get your default VPC ID and subnet IDs.
- Create the configuration file for the domain, which specifies the image.
- Create the domain with the configuration file.

**To add the custom SageMaker image to your domain**

1. Get your default VPC ID.

   ```bash
   aws ec2 describe-vpcs --filters Name=isDefault,Values=true
   ```
2. Get your default subnet IDs using the VPC ID from the previous step.

```bash
aws ec2 describe-subnets \
   --filters Name=vpc-id,Values=<vpc-id> \
   --query "Subnets[*].SubnetId" --output json
```

Response:
```
[
  "subnet-b55171dd",
  "subnet-8a5f99c6",
  "subnet-e88d1392"
]
```

3. Create a configuration file named `create-domain-input.json`. Insert the VPC ID, subnet IDs, ImageName, and AppImageConfigName from the previous steps. Because ImageVersionNumber isn't specified, the latest version of the image is used, which is the only version in this case. Your execution role must satisfy the requirements in Prerequisites.

```json
{
   "DomainName": "domain-with-custom-r-image",
   "VpcId": "<vpc-id>",
   "SubnetIds": [
      "<subnet-ids>"
   ],
   "DomainSettings": {
      "RStudioServerProDomainSettings": {
         "DomainExecutionRoleArn": "<execution-role>"
      }
   },
   "DefaultUserSettings": {
      "ExecutionRole": "<execution-role>",
      "RSessionAppSettings": {
         "CustomImages": [
            {
```
"AppImageConfigName": "rstudio-custom-config",
"ImageName": "rstudio-custom-image"
]
]
}
}
,"AuthMode": "IAM"
}

4. Create the domain with the attached custom SageMaker image.

```bash
aws sagemaker create-domain \
  --cli-input-json file://create-domain-input.json
```

Response:

```
{
  "DomainArn": "arn:aws:sagemaker:region:acct-id:domain/domain-id",
  "Url": "https://domain-id.studio.region.sagemaker.aws/..."
}
```

**Attach the SageMaker image to an existing domain**

This method assumes that you've already onboarded to domain. For more information, see [Amazon SageMaker domain overview](#).

**Note**

You must delete all of the applications in your domain to update the domain with the new image version. For information about deleting these applications, see [Delete an Amazon SageMaker domain](#).

Use the following steps to add the SageMaker image to your current domain.

- Get your DomainID from the SageMaker console.
- Use the DomainID to get the DefaultUserSettings for the domain.
- Add the ImageName and AppImageConfig as a CustomImage to the DefaultUserSettings.
- Update your domain to include the custom image.
To add the custom SageMaker image to your domain

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Select the desired domain.
5. Choose domain settings.
6. Under General Settings, find the domain ID. The ID is in the following format: d-xxxxxxxxxxxx.
7. Use the domain ID to get the description of the domain.

   ```bash
   aws sagemaker describe-domain
   --domain-id <d-xxxxxxxxxxxx>
   ```

Response:

```json
{
   "DomainId": "d-xxxxxxxxxxxx",
   "DefaultUserSettings": {
      "KernelGatewayAppSettings": {
         "CustomImages": [
            ...
         ]
      }
   }
}
```

8. Save the DefaultUserSettings section of the response to a file named update-domain-input.json.
9. Insert the ImageName and AppImageConfigName from the previous steps as a custom image. Because ImageVersionNumber isn't specified, the latest version of the image is used, which is the only version in this case.

```json
{
   "DefaultUserSettings": {
      "RSessionAppSettings": {
         "CustomImages": [
            {
               "ImageName": "rstudio-custom-image",
            }
         ]
      }
   }
}
```
10. Use the domain ID and default user settings file to update your domain.

```bash
aws sagemaker update-domain \
   --domain-id <d-xxxxxxxxxxxx> \
   --cli-input-json file://update-domain-input.json
```

Response:

```json
{
   "DomainArn": "arn:aws:sagemaker:region:acct-id:domain/domain-id"
}
```

11. Delete the RStudioServerPro application. You must restart the RStudioServerPro domain-shared application for the RStudio Launcher UI to pick up the latest changes.

```bash
aws sagemaker delete-app \
   --domain-id <d-xxxxxxxxxxxx> --user-profile-name domain-shared \
   --app-type RStudioServerPro --app-name default
```

12. Create a new RStudioServerPro application. You must create this application using the AWS CLI.

```bash
aws sagemaker create-app \
   --domain-id <d-xxxxxxxxxxxx> --user-profile-name domain-shared \
   --app-type RStudioServerPro --app-name default
```

**Launch a custom SageMaker image in RStudio**

You can use your custom image when launching an RStudio application from the console. After you create your custom SageMaker image and attach it to your domain, the image appears in the image selector dialog box of the RStudio Launcher. To launch a new RStudio app, follow the steps in [Open RStudio Launcher and launch RSessions](#) and select your custom image as shown in the following image.
Clean up image resources

This guide shows how to clean up RStudio image resources that you created in the previous sections. To delete an image, complete the following steps using either the SageMaker console or the AWS CLI, as shown in this guide.

- Detach the image and image versions from your Amazon SageMaker domain.
- Delete the image, image version, and app image config.

After you've completed these steps, you can delete the container image and repository from Amazon ECR. For more information about how to delete the container image and repository, see Deleting a repository.
Clean up resources from the SageMaker console

When you detach an image from a domain, all versions of the image are detached. When an image is detached, all users of the domain lose access to the image versions.

To detach an image

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Select the desired domain.
5. Choose Environment.
6. Under Custom images attached to domain, choose the image and then choose Detach.
7. (Optional) To delete the image and all versions from SageMaker, select Also delete the selected images .... This does not delete the associated images from Amazon ECR.
8. Choose Detach.

Clean up resources from the AWS CLI

To clean up resources

1. Detach the image and image versions from your domain by passing an empty custom image list to the domain. Open the update-domain-input.json file that you created in Attach the SageMaker image to your current domain.
2. Delete the RSessionAppSettings custom images and then save the file. Do not modify the KernelGatewayAppSettings custom images.

```json
{
   "DomainId": "d-xxxxxxxxxxxxx",
   "DefaultUserSettings": {
      "KernelGatewayAppSettings": {
         "CustomImages": [
            ...
         ],
         "RSessionAppSettings": {
            "CustomImages": [
            ]
      }
   }
}```
3. Use the domain ID and default user settings file to update your domain.

```
aws sagemaker update-domain \
  --domain-id <d-xxxxxxxxxxxxx> \
  --cli-input-json file://update-domain-input.json
```

Response:

```
{
  "DomainArn": "arn:aws:sagemaker:us-east-2:acct-id:domain/d-xxxxxxxxxxxxx"
}
```

4. Delete the app image config.

```
aws sagemaker delete-app-image-config \
  --app-image-config-name rstudio-image-config
```

5. Delete the SageMaker image, which also deletes all image versions. The container images in Amazon ECR that are represented by the image versions are not deleted.

```
aws sagemaker delete-image \
  --image-name rstudio-image
```

**Manage users**

After your RStudio-enabled Amazon SageMaker domain is running, you can add user profiles (UserProfiles) to the domain. The following topics show how to create user profiles that are authorized to use RStudio, as well as update an existing user profile. For information on how to delete an RStudio App, UserProfile, or domain, follow the steps in [Delete an Amazon SageMaker domain](#).
The limit for the total number of UserProfiles in a Amazon SageMaker domain is 60.

There are two types of users:

- Unauthorized: This user cannot access the RStudio app. By default, a new user is Unauthorized if the domain is enabled for RStudio.
- Authorized: This user can access the RStudio app and use one of the RStudio license seats.

If a user is authorized, they can be given one of the following levels of access to RStudio.

- RStudio User: This is a standard RStudio user and can access RStudio.
- RStudio Admin: The admin of your Amazon SageMaker domain has the ability to create users, add existing users, and update the permissions of existing users. Admins can also access the RStudio Administrative dashboard. However, this admin is not able to update parameters that are managed by Amazon SageMaker.

**Methods to create a user**

The following topics show how to create a user in your RStudio-enabled Amazon SageMaker domain.

**Create user console**

To create a user in your RStudio-enabled Amazon SageMaker domain from the console, complete the steps in [Add user profiles](#).

**Create user CLI**

The following command shows how to add users to a Amazon SageMaker domain with IAM authentication. A User can belong to either the R_STUDIO_USER or R_STUDIO_ADMIN User group.

```
aws sagemaker create-user-profile --region <REGION> \
    --domain-id <DOMAIN-ID> \
    --user-profile-name <USER_PROFILE_NAME-ID> \
    --user-settings RStudioServerProAppSettings={UserGroup=<USER-GROUP>}
```
The following command shows how to add users to a Amazon SageMaker domain with authentication using IAM Identity Center. A user can belong to either the R_STUDIO_USER or R_STUDIO_ADMIN User group.

```
aws sagemaker create-user-profile --region <REGION> \
--domain-id <DOMAIN-ID> \
--user-profile-name <USER_PROFILE_NAME-ID> \
--user-settings RStudioServerProAppSettings={UserGroup=<USER-GROUP>} \
--single-sign-on-user-identifier UserName \
--single-sign-on-user-value <USER-NAME>
```

**Update existing user**

You cannot update the authorization of an existing user. You must delete the existing user and create a new one with the updated authorization.

**Log in to RStudio as another user**

2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **domains**.
4. Select the domain containing the user profile.
5. Select a user name from the list of users. This opens a new page with details about the user profile and the apps that are running.
6. Select **Launch**.
7. From the dropdown, select **RStudio** to launch an RStudio instance.

**Terminate sessions for another user**

1. From the list of running apps, identify the app you want to delete.
2. Click the respective **Delete app** button for the app you are deleting.

**Delete another user**

You cannot delete a user if the user is running any apps. Delete all apps before attempting to delete a user.

1. From the **User Profile** page, select **Edit**. This opens a new **General settings** page.
2. Under **Delete user**, select **Delete user**.

**RStudio administrative dashboard**

This topic shows how to access and use the RStudio administrative dashboard. With the RStudio administrative dashboard, admins can manage users and RSessions, as well as view information about RStudio Server instance utilization and Amazon CloudWatch Logs.

**Launch the RStudio administrative dashboard**

The R_STUDIO_ADMIN authorization allows the user to access the RStudio administrative dashboard. An R_STUDIO_ADMIN user can access the RStudio administrative dashboard by replacing `workspaces` with `admin` in their RStudio URL manually. The following shows how to modify the URL to access the RStudio administrative dashboard.

For example, the following RStudio URL:

```
https://<DOMAIN-ID>.studio.us-east-2.sagemaker.aws/rstudio/default/s/<SESSION-ID>/workspaces
```

Can be converted to:

```
https://<DOMAIN-ID>.studio.us-east-2.sagemaker.aws/rstudio/default/s/<SESSION-ID>/admin
```

**Dashboard tab**

This tab gives an overview of your RStudio Server instance utilization, as well as information on the number of active RSessions.

**Sessions tab**

This tab gives information on the active RSessions, such as the user that launched the RSessions, the time that the RSessions have been running, and their resource utilization.

**Users tab**

This tab gives information on the RStudio authorized users in the domain, such as the time that the last RSession was launched and their resource utilization.
Stats tab

This tab gives information on the utilization of your RStudio Server instance.

Logs tab

This tab displays Amazon CloudWatch Logs for the RStudio Server instance. For more information about logging events with Amazon CloudWatch Logs, see What is Amazon CloudWatch Logs?.

Shut down and restart RStudio

To shut down and restart your Posit Workbench and the associated RStudioServerPro app, you must first shut down all of your existing RSessions. You can shut down the RSessionGateway apps from within RStudio. You can then shut down the RStudioServerPro app using the AWS CLI. After the RStudioServerPro app is shut down, you must reopen RStudio through the SageMaker console.

Any unsaved notebook information is lost in the process. The user data in the Amazon EFS volume isn't impacted.

Note

If you are using a custom image with RStudio, ensure that your docker image is using an RStudio version that is compatible with the version of Posit Workbench being used by SageMaker after you restart your RStudioServerPro app.

The following topics show how to shut down the RSessionGateway and RStudioServerPro apps and restart them.

Suspend your RSessions

Complete the following procedure to suspend all of your RSessions.

1. From the RStudio Launcher, identify the RSession that you want to suspend.
2. Select Suspend for the session.
3. Repeat this for all RSessions.

Delete your RSessions

Complete the following procedure to shut down all of your RSessions.
1. From the RStudio Launcher, identify the RSession that you want to delete.

2. Select Quit for the session. This opens a new Quit Session window.

3. From the Quit Session window, select Force Quit, to end all child processes in the session.

4. Select Quit Session to confirm deletion of the session.

5. Repeat this for all RSessions.

Delete your RStudioServerPro app

Run the following commands from the AWS CLI to delete and restart your RStudioServerPro app.

1. Delete the RStudioServerPro application by using your current domain id.

   ```bash
   aws sagemaker delete-app \
   --domain-id <domainId> \
   --user-profile-name domain-shared \
   --app-type RStudioServerPro \n   --app-name default
   ```

2. Re-create the RStudioServerPro application.

   ```bash
   aws sagemaker create-app \
   --domain-id <domainId> \
   --user-profile-name domain-shared \
   --app-type RStudioServerPro \n   --app-name default
   ```

Manage billing and cost

To track the costs associated with your RStudio environment, you can use the AWS Billing and Cost Management service. AWS Billing and Cost Management provides useful tools to help you gather information related to your cost and usage, analyze your cost drivers and usage trends, and take action to budget your spending. For more information, see What is AWS Billing and Cost Management?

The following describes components required to run RStudio on Amazon SageMaker and how each component factors into billing for your RStudio instance.
• RStudio License – You must purchase an RStudio license. There is no additional charge for using your RStudio license with Amazon SageMaker. For more information about your RStudio license, see [RStudio license](#).

• RSession - These are RStudio working sessions launched by end users. You are charged while the RSession is running.

• RStudio Server - A multi-tenant server manages all the RSessions. You can choose the instance type to run RStudio Server on, and pay the related costs. The default instance, "system", is free, but you can choose to pay for higher tiers. For more information about the available instance types for your RStudio Server, see [RStudioServerPro instance type](#).

**Tracking billing at user level**

To track billing at the user level using Cost Allocation Tags, see [Using Cost Allocation Tags](#).

**Diagnose issues and get support**

The following sections describe how to diagnose issues with RStudio on Amazon SageMaker. To get support for RStudio on Amazon SageMaker, contact Amazon SageMaker support. For help with purchasing an RStudio license or modifying the number of license seats, contact sales@rstudio.com.

**Upgrade your version**

If you receive a warning that there is a version mismatch between your RSession and RStudioServerPro apps, then you must upgrade the version of your RStudioServerPro app. For more information, see [Upgrade the RStudio Version](#).

**View Metrics and Logs**

You can monitor your workflow performance while using RStudio on Amazon SageMaker. View data logs and information about metrics with the RStudio administrative dashboard or Amazon CloudWatch.

**View your RStudio logs from the RStudio administrative dashboard**

You can view metrics and logs directly from the RStudio administrative dashboard.

1. Log in to your [Amazon SageMaker domain](#).

2. Navigate to the RStudio administrative dashboard following the steps in [RStudio administrative dashboard](#).
3. Select the **Logs** tab.

**View your RStudio logs from Amazon CloudWatch Logs**

Amazon CloudWatch monitors your AWS resources and the applications that you run on AWS in real time. You can use Amazon CloudWatch to collect and track metrics, which are variables that you can measure for your resources and applications. To ensure that your RStudio apps have permissions for Amazon CloudWatch, you must include the permissions described in [Amazon SageMaker domain overview](#). You don’t need to do any setup to gather Amazon CloudWatch Logs.

The following steps show how to view Amazon CloudWatch Logs for your RSession.

These logs can be found in the /aws/sagemaker/studio log stream from the AWS CloudWatch console.

2. Select Logs from the left side. From the dropdown menu, select Log groups.
3. On the Log groups screen, search for aws/sagemaker/studio. Select the Log group.
4. On the aws/sagemaker/studio Log group screen, navigate to the Log streams tab.
5. To find the logs for your domain, search Log streams using the following format:

   `<DomainId>/domain-shared/rstudioserverpro/default`

**Use RStudio on Amazon SageMaker**

With RStudio support in Amazon SageMaker, you can put your production workflows in place and take advantage of SageMaker features. The following topics show how to launch an RStudio session and complete key workflows. For information about managing RStudio on SageMaker, see [Manage RStudio on Amazon SageMaker](#).

For information about the onboarding steps to create an Amazon SageMaker domain with RStudio enabled, see [Amazon SageMaker domain overview](#).

For information about the AWS Regions that RStudio on SageMaker is supported in, see [Supported Regions and Quotas](#).

**Topics**
Collaborate in RStudio

To share your RStudio project, you can connect RStudio to your Git repo. For information on setting this up, see [Version Control with Git and SVN](#).

Note: Project sharing and realtime collaboration are not currently supported when using RStudio on Amazon SageMaker.

Base R image

When launching your RStudio instance, the Base R image serves as the basis of your instance. This image extends the `r-session-complete` Docker image.

This Base R image includes the following:

- R v4.0 or higher
- `awscli`, `sagemaker`, and `boto3` Python packages
- [Reticulate](https://github.com/reticulate/reticulate) package for R SDK integration

RSession application colocation

Users can create multiple RSession applications on the same instance. Each instance type supports up to four colocated RSession applications. This applies to each user independently. For example, if two users create applications, then SageMaker allocates different underlying instances to each user. Each of these instances would support 4 RSession applications.

Customers only pay for the instance type used regardless of how many RSession applications are running on the instance. If a user creates an RSession with a different associated instance type, then a new underlying instance is created.
Open RStudio Launcher and launch RSessions

The following topics show how to use the RStudio Launcher to launch RSessions.

Open RStudio Launcher

Open the RStudio launcher using the following set of procedures that matches your environment.

Open RStudio Launcher from the Amazon SageMaker Console

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. From the left navigation, select RStudio.
3. Under Get Started, select the domain and user profile to launch.

Open RStudio Launcher from Amazon SageMaker Studio

1. Navigate to Studio following the steps in Launch Amazon SageMaker Studio.
3. From the RStudio landing page, choose Launch application.

Open RStudio Launcher from the AWS CLI

The procedure to open the RStudio Launcher using the AWS CLI differs depending on the method used to manage your users.

IAM Identity Center

1. Use the AWS access portal to open your Amazon SageMaker domain.
2. Modify the URL path to “/rstudio/default” as follows.

```
# Studio URL
https://<domain-id>.studio.<region>.sagemaker.aws/jupyter/default/lab

# Modified URL
https://<domain-id>.studio.<region>.sagemaker.aws/rstudio/default
```
To open the RStudio Launcher from the AWS CLI in IAM mode, complete the following procedure.

1. Create a presigned URL using the following command.

   ```bash
   aws sagemaker create-presigned-domain-url --region <REGION> \
   --domain-id <DOMAIN-ID> \
   --user-profile-name <USER-PROFILE-NAME>
   ```

2. Append `&redirect=RStudioServerPro` to the generated URL.

3. Navigate to the updated URL.

**Launch RSessions**

After you’ve launched the RStudio Launcher, you can create a new RSession.

1. Select **New Session**.

2. Enter a **Session Name**.

3. Select an instance type that your RSession runs on. This defaults to `ml.t3.medium`.

4. Select an Image that your RSession uses as the kernel.

5. Select Start Session.

6. After your session has been created, you can start it by selecting the name.

   **Note**

   If you receive a warning that there is a version mismatch between your RSession and RStudioServerPro apps, then you must upgrade the version of your RStudioServerPro app. For more information, see [Upgrade the RStudio Version](#).

**Suspend your RSessions**

1. From the RStudio Launcher, identify the RSsession that you want to suspend.

2. Select **Suspend** for the session.

**Delete your RSessions**

1. From the RStudio Launcher, identify the RSsession that you want to delete.
2. Select Quit for the session. This opens a new Quit Session window.
3. From the Quit Session window, select Force Quit, to end all child processes in the session.
4. Select Quit Session to confirm deletion of the session.

Publish to RStudio Connect

RStudio Connect enables data scientists to publish insights, dashboard and web applications from RStudio on Amazon SageMaker. For more information, see Host RStudio Connect and Package Manager for ML development in RStudio on Amazon SageMaker.

For more information on RStudio Connect, see the RStudio Connect User Guide.

Access Amazon SageMaker features with RStudio on Amazon SageMaker

One of the benefits of using RStudio on Amazon SageMaker is the integration of Amazon SageMaker features. This includes integration with Amazon SageMaker Studio Classic and Reticulate.

Use Amazon SageMaker Studio Classic and RStudio on Amazon SageMaker

Your Amazon SageMaker Studio Classic and RStudio instances share the same Amazon EFS file system. This means that files that you import and create using Studio Classic can be accessed using RStudio and vice versa. This allows you to work on the same files using both Studio Classic and RStudio without having to move your files between the two. For more information on this workflow, see the Announcing Fully Managed RStudio on Amazon SageMaker for Data Scientists blog.

Use Amazon SageMaker SDK with reticulate

The reticulate package is used as an R interface to Amazon SageMaker Python SDK to make API calls to Amazon SageMaker. The reticulate package translates between R and Python objects, and Amazon SageMaker provides a serverless data science environment to train and deploy Machine Learning (ML) models at scale. For general information about the reticulate package, see R Interface to Python.

For a blog that outlines how to use the reticulate package with Amazon SageMaker, see Using R with Amazon SageMaker.

The following examples show how to use reticulate for specific use cases.
• For a notebook that describes how to use reticulate to do batch transform to make predictions, see [Batch Transform Using R with Amazon SageMaker](https://aws.amazon.com/sagemaker/).  
• For a notebook that describes how to use reticulate to conduct hyperparameter tuning and generate predictions, see [Hyperparameter Optimization Using R with Amazon SageMaker](https://aws.amazon.com/sagemaker/).

**Get started with Code Editor in Amazon SageMaker Studio**


Code Editor has the [AWS Toolkit for VS Code](https://marketplace.visualstudio.com) extension pre-installed, which enables connections to AWS services such as [Amazon CodeWhisperer](https://aws.amazon.com/sagemaker/), a general purpose, machine learning-powered code generator that provides code recommendations in real time. For more information about extensions, see [Code Editor Connections and Extensions](https://aws.amazon.com/sagemaker/).

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](https://aws.amazon.com/sagemaker/).

To launch Code Editor, create a Code Editor private space. The Code Editor space uses a single Amazon Elastic Compute Cloud (Amazon EC2) instance for your compute and a single Amazon Elastic Block Store (Amazon EBS) volume for your storage. Everything in your space such as your code, Git profile, and environment variables are stored on the same Amazon EBS volume. The volume has 3000 IOPS and a throughput of 125 MBps. Your administrator has configured the default Amazon EBS storage settings for your space.

The default storage size is 5 GB, but your administrator can increase the amount of space you get. For more information, see [Change the default storage size](https://aws.amazon.com/sagemaker/).

You can scale your compute up or down by changing the Amazon EC2 instance type that runs your Code Editor application. Before you change the associated instance type, you must first stop your Code Editor space. For more information, see [Code Editor application instances and images](https://aws.amazon.com/sagemaker/).
Your administrator might provide you with a lifecycle configuration to customize your environment. You can specify the lifecycle configuration when you create the space. For more information, see [Code Editor lifecycle configurations](#).

You can also bring your own file storage system if you have an Amazon EFS volume.

## Topics

- [Code Editor user guide](#)
- [Code Editor administrator guide](#)

### Code Editor user guide

The topics in this section provide guides for using Code Editor, including how to launch, add connections to AWS services, shut down resources, and more. After creating a Code Editor space, you can access your Code Editor session directly through the browser.

Within your Code Editor environment, you can do the following:

- Access all artifacts persisted in your home directory
- Clone your GitHub repositories and commit changes
- Access the SageMaker Python SDK
You can return to Studio to review any assets created in your Code Editor environment such as experiments, pipelines, or training jobs.

**Topics**

- Check the version of Code Editor
- Code Editor application instances and images
- Launch a Code Editor application in Studio
- Launch a Code Editor application using the AWS CLI
- Clone a repository in Code Editor
- Code Editor Connections and Extensions
- Log out and shut down resources

**Check the version of Code Editor**

The following steps show how to check the version of your Code Editor application.

**To check the Code Editor application version**

1. Launch and run a Code Editor space and navigate to the Code Editor application UI. For more information, see [Launch a Code Editor application in Studio](#).
2. In the upper-left corner of the Code Editor UI, choose the menu button ( )

   Then, choose Help. Then, choose About.

**Note**


**Code Editor application instances and images**

Only some instances are compatible with Code Editor applications. You can choose the instance type that is compatible with your use case from the **Instance** dropdown menu.
The **Fast launch** instances start up much faster than the other instances. For more information about fast launch instance types in Studio, [Available Studio Classic Instance Types](#).

**Note**

If you use a GPU instance type when configuring your Code Editor application, you must also use a GPU-based image. The Code Editor space UI automatically selects a compatible image when you select your instance type.

Within a space, your data is stored in an Amazon EBS volume that persists independently from the life of an instance. You won't lose your data when you change instances. If your Code Editor space is Running, you must stop your space before changing instance types.

The following table lists the ARNs of the available Code Editor CPU and GPU images for each Region.

<table>
<thead>
<tr>
<th>Region</th>
<th>CPU</th>
<th>GPU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region</td>
<td>distribution CPU ARN</td>
<td>distribution GPU ARN</td>
</tr>
<tr>
<td>-----------------</td>
<td>----------------------------------------------------------</td>
<td>----------------------------------------------------------</td>
</tr>
<tr>
<td>Region</td>
<td>ARN for CPU</td>
<td>ARN for GPU</td>
</tr>
<tr>
<td>----------------</td>
<td>--------------</td>
<td>--------------</td>
</tr>
</tbody>
</table>
If you encounter instance limits, contact your administrator. To get more storage and compute for a user, administrators can request an increase to a user's AWS quotas. For more information about requesting a quota increase, see Amazon SageMaker endpoints and quotas.

**Launch a Code Editor application in Studio**

To configure and access your Code Editor integrated development environment through Studio, you must create a Code Editor space. For more information about spaces in Studio, see Amazon SageMaker Studio spaces.
The following procedure shows how to create and run a Code Editor space.

To create and run a Code Editor space

1. Launch the updated Studio experience. For more information, see Launch Amazon SageMaker Studio.
2. Do one of the following:
   - Within the updated Amazon SageMaker Studio UI, select Code Editor from the Applications menu.
   - Within the updated Amazon SageMaker Studio UI, choose View Code Editor spaces in the Overview section of the Studio homepage.
3. In the upper-right corner of the Code Editor landing page, choose Create Code Editor space.
4. Enter a name for your Code Editor space. The name must be 1–62 characters in length using letters, numbers, and dashes only.
5. Choose Create space.
6. After the space is created, you have some options before you choose to run the space:
   - You can edit the Storage (GB), Lifecycle Configuration, or Attach custom EFS filesystem settings. Options for these settings are available based on administrator specification.
• From the **Instance** dropdown menu, you can choose the instance type most compatible with your use case.

If you use a GPU instance type when configuring your Code Editor application, you must also use a GPU-based image. Within a space, your data is stored in an Amazon EBS volume that persists independently from the life of an instance. You won't lose your data when you change instances.

**Note**

To update space settings, you must first stop your space. If your Code Editor uses an instance with NVMe instance stores, any data stored on the NVMe store is deleted when the space is stopped.

7. After updating your settings, choose **Run Space** in the space detail page.
8. After the status of the space is **Running**, choose **Open Code Editor** to go to your Code Editor session.

Within the Code Editor Studio landing page, you can filter and manage existing spaces.
To manage your Code Editor spaces

1. Navigate to the Code Editor Studio landing page and filter your Code Editor spaces by Private to me or Running.

2. Do one of the following:
   - Within the Code Editor Studio landing page, in the row of the space name of your choice, you can Stop, Start, or Open that space in the Action column.
   - Choose the name of a space in the Code Editor Studio landing page. This takes you to the space detail page where you can also Stop, Start, or Open that space or update the space settings.

Launch a Code Editor application using the AWS CLI

To configure and access your Code Editor integrated development environment through the AWS Command Line Interface (AWS CLI), you must create a Code Editor space. Be sure to meet the Prerequisites before going through the following steps. Use the following procedure to create and run a Code Editor space.

To create and run a Code Editor space

1. Access a space using AWS Identity and Access Management (IAM) or AWS IAM Identity Center authentication. For more information about accessing spaces using the AWS CLI, see Accessing spaces using the AWS Command Line Interface in Amazon SageMaker Studio spaces.

2. Create an application and specify CodeEditor as the app-type using the following command.

   If you use a GPU instance type when creating your Code Editor application, you must also use a GPU-based image.

   ```bash
   aws sagemaker create-app
   --domain-id domain-id
   --space-name space-name
   --app-type CodeEditor
   --app-name default
   ```
For more information about available Code Editor image ARNs, see Code Editor application instances and images.

3. After the Code Editor application is in service, launch the application using a presigned URL. You can use the `describe-app` API to check if your application is in service. Use the `create-presigned-domain-url` API to create a presigned URL:

```bash
aws sagemaker create-presigned-domain-url --domain-id domain-id --space-name space-name --user-profile-name user-profile-name --session-expiration-duration-in-seconds 43200 --landing-uri app:CodeEditor:
```

4. Open the generated URL to start working in your Code Editor application.

### Clone a repository in Code Editor

You can navigate through folders and clone a repository in the Explorer window of the Code Editor application UI.

To clone a repository, go through the following steps:

#### To clone a repository

1. Open your Code Editor application in the browser, and choose the **Exploration** button in the left navigation pane.

2. Choose **Clone Repository** in the Explorer window. Then, provide a repository URL or pick a repository source in the prompt.

3. Choose a folder to clone your repository into. Note that the default Code Editor folder is `/home/sagemaker-user/`. Cloning your repository may take some time.

4. To open the cloned repository, choose either **Open in New Window** or **Open**.

5. To return to the Code Editor application UI homepage, choose **Cancel**.

6. Within the repository, a prompt asks if you trust the authors of the files in your new repository. You have two choices:
To trust the folder and enable all features, choose **Yes, I trust the authors**.

To browse the repository content in *restricted mode*, choose **No, I don't trust the authors**.

In restricted mode, tasks are not allowed to run, debugging is disabled, workspace settings are not applied, and extensions have limited functionality.

To exit restricted mode, trust the authors of all files in your current folder or its parent folder, and enable all features, choose **Manage** in the **Restricted Mode** banner.

**Code Editor Connections and Extensions**

Code Editor supports IDE connections to AWS services as well as extensions available in the [Open VSX Registry](#).

**Connections to AWS**

Code Editor environments are integrated with the [AWS Toolkit for VS Code](#) to add connections to AWS services. To get started with connections to AWS services, you must have valid AWS Identity and Access Management (IAM) credentials. For more information, see [Authentication and access for the AWS Toolkit for Visual Studio Code](#).

Within your Code Editor environment, you can add connections to:

- **AWS Explorer** – View, modify, and deploy AWS resources in Amazon S3, CloudWatch, and more.
  
  Accessing certain features within AWS Explorer requires certain AWS permissions. For more information, see [Authentication and access for the AWS Toolkit for Visual Studio Code](#).

- **Amazon CodeWhisperer** – Build applications faster with AI-powered code suggestions.

To use Amazon CodeWhisperer with Code Editor, you must add the following permissions to your SageMaker execution role.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "CodeWhispererPermissions",
            "Effect": "Allow",
            "Action": ["codewhisperer:GenerateRecommendations"],
            "Resource": "*"
        }
    ]
}
```
For more information, see Creating IAM policies and Adding and removing IAM identity permissions in the IAM User Guide.

Extensions

Code Editor supports IDE extensions available in the Open VSX Registry.

To get started with extensions in your Code Editor environment, choose the Extensions icon in the left navigation pane. Here, you can configure connections to AWS by installing the AWS Toolkit. For more information, see Installing the AWS Toolkit for Visual Studio Code.

In the search bar, you can search directly for additional extensions through the Open VSX Registry, such as the AWS Toolkit, Jupyter, Python, and more.

Log out and shut down resources

In the upper-left corner of the Code Editor environment, choose the menu icon. Then, choose SageMaker: Log out.

Stop your space through Studio

To stop your Code Editor space in Studio use the following steps:

To stop your Code Editor space in Studio

1. Return to the Code Editor landing page by doing one of the following:
   a. In the navigation bar in the upper-left corner, choose Code Editor.
   b. Alternatively, in the left navigation pane, choose Code Editor in the Applications menu.
2. Find the name of the Code Editor space you created. If the status of your space is Running, choose Stop in the Action column. You can also stop your space directly in the space detail page by choosing Stop space. The space may take some time to stop.
Additional resources such as SageMaker endpoints, Amazon EMR (Amazon EMR) clusters and Amazon Simple Storage Service (Amazon S3) buckets created from Studio are not automatically deleted when your space instance shuts down. To stop accruing charges from resources, delete any additional resources. For more information, see Delete unused resources.

**Shut down resources using the AWS CLI**

You can delete your Code Editor application and space using the AWS Command Line Interface (AWS CLI).

- [DeleteApp](#)
- [DeleteSpace](#)

**Code Editor administrator guide**

You can use Code Editor with an On-Demand Instance for faster startup time, and configurable storage. You can launch a Code Editor application through Amazon SageMaker Studio or through the AWS CLI. You can also edit Code Editor default settings within the domain console. For more information, see View and edit domains.

**Topics**

- [Prerequisites](#)
- [Give your users access to private spaces](#)
- [Change the default storage size](#)
- [Code Editor lifecycle configurations](#)
Prerequisites

To use Code Editor, based on Code-OSS, Visual Studio Code - Open Source, first onboard to Amazon SageMaker domain and create a user profile. For more information, see Amazon SageMaker domain overview.

If you are interacting with your Code Editor application using the AWS CLI, you must also complete the following prerequisites.

- Update the AWS CLI by following the steps in Installing the current AWS CLI Version.
- From your local machine, run `aws configure` and provide your AWS credentials. For information about AWS credentials, see Understanding and getting your AWS credentials.

To get more storage and compute for your application, you can request an increase to your AWS quotas. For more information about requesting a quota increase, see Amazon SageMaker endpoints and quotas.

Give your users access to private spaces

This section provides a policy that grants user access to private spaces. You can also use the policy to restrict private spaces and applications that are associated with them to the owner associated with the user profile.

You must provide your users with permissions to the following:

- Private spaces
- The user profile required for access to the private spaces

To provide permissions, attach the following policy to the IAM roles of your users.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "sagemaker:CreateApp",
            "sagemaker:DeleteApp"
         ]
      }
   ]
}
```
"Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:app/*",
"Condition": {
  "Null": {
    "sagemaker:OwnerUserProfileArn": "true"
  }
},
{
  "Sid": "SMStudioCreatePresignedDomainUrlForUserProfile",
  "Effect": "Allow",
  "Action": [
    "sagemaker:CreatePresignedDomainUrl"
  ],
  "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:user-profile/
${sagemaker:DomainId}/${sagemaker:UserProfileName}"
},
{
  "Sid": "SMStudioAppPermissionsListAndDescribe",
  "Effect": "Allow",
  "Action": [
    "sagemaker:ListApps",
    "sagemaker:ListDomains",
    "sagemaker:ListUserProfiles",
    "sagemaker:ListSpaces",
    "sagemaker:DescribeApp",
    "sagemaker:DescribeDomain",
    "sagemaker:DescribeUserProfile",
    "sagemaker:DescribeSpace"
  ],
  "Resource": "*"
},
{
  "Sid": "SMStudioAppPermissionsTagOnCreate",
  "Effect": "Allow",
  "Action": ["sagemaker:AddTags"],
  "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:/*/",
  "Condition": {
    "Null": {
      "sagemaker:TaggingAction": "false"
    }
  }
},
}
null

```json
{
  "Sid": "SMStudioRestrictSharedSpacesWithoutOwners",
  "Effect": "Allow",
  "Action": [
    "sagemaker:CreateSpace",
    "sagemaker:UpdateSpace",
    "sagemaker:DeleteSpace"
  ],
  "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:space/${sagemaker:DomainId}/*",
  "Condition": {
    "Null": {
      "sagemaker:OwnerUserProfileArn": "true"
    }
  }
},
{
  "Sid": "SMStudioRestrictSpacesToOwnerUserProfile",
  "Effect": "Allow",
  "Action": [
    "sagemaker:CreateSpace",
    "sagemaker:UpdateSpace",
    "sagemaker:DeleteSpace"
  ],
  "Resource": "arn:aws:sagemaker:{{Region}}:{{AccountId}}:space/${sagemaker:DomainId}/*",
  "Condition": {
    "ArnLike": {
      "sagemaker:OwnerUserProfileArn": "arn:aws:sagemaker:$$AWS Region:111122223333:user-profile/${sagemaker:DomainId}/${sagemaker:UserProfileName}"
    },
    "StringEquals": {
      "sagemaker:SpaceSharingType": [
        "Private",
        "Shared"
      ]
    }
  }
},
{
  "Sid": "SMStudioRestrictCreatePrivateSpaceAppsToOwnerUserProfile",
  "Effect": "Allow",
  "Action": [
    "sagemaker:CreateApp",
    "sagemaker:DeleteApp",
    "sagemaker:UpdateApp",
    "sagemaker:CreateImageVersion",
    "sagemaker:DeleteImageVersion",
    "sagemaker:UpdateImageVersion",
    "sagemaker:DescribeImageVersion",
    "sagemaker:DescribeImage",
    "sagemaker:ListImageVersions",
    "sagemaker:ListImages",
    "sagemaker:CreateRepository",
    "sagemaker:DeleteRepository",
    "sagemaker:UpdateRepository",
    "sagemaker:DescribeRepository",
    "sagemaker:ListRepositories",
    "sagemaker:ListFileSetDefinitions",
    "sagemaker:DescribeFileSetDefinition",
    "sagemaker:CreateFileSetDefinition",
    "sagemaker:DeleteFileSetDefinition",
    "sagemaker:CreateTrialInstance",
    "sagemaker:DeleteTrialInstance",
    "sagemaker:UpdateTrialInstance",
    "sagemaker:DescribeTrialInstance",
    "sagemaker:ListTrialInstances",
    "sagemaker:CreateTrialInstanceGroup",
    "sagemaker:DeleteTrialInstanceGroup",
    "sagemaker:UpdateTrialInstanceGroup",
    "sagemaker:DescribeTrialInstanceGroup",
    "sagemaker:ListTrialInstanceGroups",
    "sagemaker:CreateTrialJob",
    "sagemaker:DeleteTrialJob",
    "sagemaker:UpdateTrialJob",
    "sagemaker:DescribeTrialJob",
    "sagemaker:ListTrialJobs",
    "sagemaker:CreateModelPackage",
    "sagemaker:DeleteModelPackage",
    "sagemaker:UpdateModelPackage",
    "sagemaker:DescribeModelPackage",
    "sagemaker:ListModelPackages",
    "sagemaker:CreateMonitoringJobDefinition",
    "sagemaker:DeleteMonitoringJobDefinition",
    "sagemaker:UpdateMonitoringJobDefinition",
    "sagemaker:DescribeMonitoringJobDefinition",
    "sagemaker:ListMonitoringJobDefinitions",
    "sagemaker:CreateMonitoringJobRun",
    "sagemaker:DeleteMonitoringJobRun",
    "sagemaker:UpdateMonitoringJobRun",
    "sagemaker:DescribeMonitoringJobRun",
    "sagemaker:ListMonitoringJobRuns",
    "sagemaker:CreateTrialCard",
    "sagemaker:DeleteTrialCard",
    "sagemaker:UpdateTrialCard",
    "sagemaker:DescribeTrialCard",
    "sagemaker:ListTrialCards",
    "sagemaker:CreateFeatureGroup",
    "sagemaker:DeleteFeatureGroup",
    "sagemaker:UpdateFeatureGroup",
    "sagemaker:DescribeFeatureGroup",
    "sagemaker:ListFeatureGroups",
    "sagemaker:CreateEndpointConfig",
    "sagemaker:DeleteEndpointConfig",
    "sagemaker:UpdateEndpointConfig",
    "sagemaker:DescribeEndpointConfig",
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    "sagemaker:DeleteEndpoint",
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    "sagemaker:CreateModel",
    "sagemaker:DeleteModel",
    "sagemaker:UpdateModel",
    "sagemaker:DescribeModel",
    "sagemaker:ListModels",
    "sagemaker:CreateModelPackageGroup",
    "sagemaker:DeleteModelPackageGroup",
    "sagemaker:UpdateModelPackageGroup",
    "sagemaker:DescribeModelPackageGroup",
    "sagemaker:ListModelPackageGroups",
    "sagemaker:CreateTrialCardGroup",
    "sagemaker:DeleteTrialCardGroup",
    "sagemaker:UpdateTrialCardGroup",
    "sagemaker:DescribeTrialCardGroup",
    "sagemaker:ListTrialCardGroups",
    "sagemaker:CreateFeatureGroupDatasetAssociation",
    "sagemaker:DeleteFeatureGroupDatasetAssociation",
    "sagemaker:UpdateFeatureGroupDatasetAssociation",
    "sagemaker:DescribeFeatureGroupDatasetAssociation",
    "sagemaker:ListFeatureGroupDatasetAssociations",
    "sagemaker:CreateDatasetGroup",
    "sagemaker:DeleteDatasetGroup",
    "sagemaker:UpdateDatasetGroup",
    "sagemaker:DescribeDatasetGroup",
    "sagemaker:ListDatasetGroups",
    "sagemaker:CreateDatasetGroupDatasetAssociation",
    "sagemaker:DeleteDatasetGroupDatasetAssociation",
    "sagemaker:UpdateDatasetGroupDatasetAssociation",
    "sagemaker:DescribeDatasetGroupDatasetAssociation",
    "sagemaker:ListDatasetGroupDatasetAssociations",
    "sagemaker:CreateHyperparameterTuningJobDefinition",
    "sagemaker:DeleteHyperparameterTuningJobDefinition",
    "sagemaker:UpdateHyperparameterTuningJobDefinition",
    "sagemaker:DescribeHyperparameterTuningJobDefinition",
    "sagemaker:ListHyperparameterTuningJobDefinitions",
    "sagemaker:CreateHyperparameterTuningJobRun",
    "sagemaker:DeleteHyperparameterTuningJobRun",
    "sagemaker:UpdateHyperparameterTuningJobRun",
    "sagemaker:DescribeHyperparameterTuningJobRun",
    "sagemaker:ListHyperparameterTuningJobRuns",
    "sagemaker:CreatePredictor",
    "sagemaker:DeletePredictor",
    "sagemaker:UpdatePredictor",
    "sagemaker:DescribePredictor",
    "sagemaker:ListPredictors",
    "sagemaker:CreateModelPackageGroupTrialCardGroupAssociation",
    "sagemaker:DeleteModelPackageGroupTrialCardGroupAssociation",
    "sagemaker:UpdateModelPackageGroupTrialCardGroupAssociation",
    "sagemaker:DescribeModelPackageGroupTrialCardGroupAssociation",
    "sagemaker:ListModelPackageGroupTrialCardGroupAssociations",
    "sagemaker:CreateOptimizerTrialCardGroupAssociation",
    "sagemaker:DeleteOptimizerTrialCardGroupAssociation",
    "sagemaker:UpdateOptimizerTrialCardGroupAssociation",
    "sagemaker:DescribeOptimizerTrialCardGroupAssociation",
    "sagemaker:ListOptimizerTrialCardGroupAssociations",
    "sagemaker:CreateOptimizerTrialCardGroupDatasetGroupAssociation",
    "sagemaker:DeleteOptimizerTrialCardGroupDatasetGroupAssociation",
    "sagemaker:UpdateOptimizerTrialCardGroupDatasetGroupAssociation",
    "sagemaker:DescribeOptimizerTrialCardGroupDatasetGroupAssociation",
    "sagemaker:ListOptimizerTrialCardGroupDatasetGroupAssociations",
    "sagemaker:CreateOptimizerTrialCardGroupDatasetAssociation",
    "sagemaker:DeleteOptimizerTrialCardGroupDatasetAssociation",
    "sagemaker:UpdateOptimizerTrialCardGroupDatasetAssociation",
    "sagemaker:DescribeOptimizerTrialCardGroupDatasetAssociation",
    "sagemaker:ListOptimizerTrialCardGroupDatasetAssociations",
    "sagemaker:CreateOptimizerTrialCardGroupDataset","sagemaker:DeleteOptimizerTrialCardGroupDataset",
    "sagemaker:UpdateOptimizerTrialCardGroupDataset",
    "sagemaker:DescribeOptimizerTrialCardGroupDataset",
    "sagemaker:ListOptimizerTrialCardGroupDatasets",
    "sagemaker:CreateOptimizerTrialCardGroupDataset",
    "sagemaker:DeleteOptimizerTrialCardGroupDataset",
    "sagemaker:UpdateOptimizerTrialCardGroupDataset",
    "sagemaker:DescribeOptimizerTrialCardGroupDataset",
    "sagemaker:ListOptimizerTrialCardGroupDatasets",
    "sagemaker:CreateOptimizerTrialCardGroupDataset",
    "sagemaker:DeleteOptimizerTrialCardGroupDataset",
    "sagemaker:UpdateOptimizerTrialCardGroupDataset",
    "sagemaker:DescribeOptimizerTrialCardGroupDataset",
    "sagemaker:ListOptimizerTrialCardGroupDatasets",
    "sagemaker:CreateOptimizerTrialCardGroupDataset",
    "sagemaker:DeleteOptimizerTrialCardGroupDataset",
    "sagemaker:UpdateOptimizerTrialCardGroupDataset",
    "sagemaker:DescribeOptimizerTrialCardGroupDataset",
    "sagemaker:ListOptimizerTrialCardGroupDatasets",
    "sagemaker:CreateOptimizerTrialCardGroupDataset",
    "sagemaker:DeleteOptimizerTrialCardGroupDataset",
    "sagemaker:UpdateOptimizerTrialCardGroupDataset",
    "sagemaker:DescribeOptimizerTrialCardGroupDataset",
    "sagemaker:ListOptim"
Change the default storage size

You can change the default storage settings of your users. You can also change the default storage settings based on your organizational requirements and the needs of your users.

To change the storage size of your users, do the following:

1. Update the Amazon EBS storage settings in the domain.
2. Create a user profile and specify the storage settings within it.

Use the following AWS Command Line Interface (AWS CLI) command to update the domain.

```bash
aws --region $REGION sagemaker update-domain \
--domain-id $DOMAIN_ID \
--default-user-settings '{
   "SpaceStorageSettings": {
      "DefaultEbsStorageSettings":{
         "DefaultEbsVolumeSizeInGb":5,
         "MaximumEbsVolumeSizeInGb":100
      }
   }
}
```
Use the following AWS CLI command to create the user profile and specify the default storage settings.

```bash
aws --region $REGION sagemaker create-user-profile \
--domain-id $DOMAIN_ID \
--user-profile-name $USER_PROFILE_NAME \
--user-settings '{
    "SpaceStorageSettings": {
        "DefaultEbsStorageSettings":{
            "DefaultEbsVolumeSizeInGb":5,
            "MaximumEbsVolumeSizeInGb":100
        }
    }
}
''
```

Use the following AWS CLI commands to update the default storage settings in the user profile.

```bash
aws --region $REGION sagemaker update-user-profile \
--domain-id $DOMAIN_ID \
--user-profile-name $USER_PROFILE_NAME \
--user-settings '{
    "SpaceStorageSettings": {
        "DefaultEbsStorageSettings":{
            "DefaultEbsVolumeSizeInGb":25,
            "MaximumEbsVolumeSizeInGb":200
        }
    }
}
''
```

**Code Editor lifecycle configurations**

You can use Code Editor lifecycle configurations to automate customization for your Studio environment. This customization includes installing custom packages, configuring extensions, preloading datasets, and setting up source code repositories.

The following instructions use the AWS Command Line Interface (AWS CLI) to create, attach, debug, and detach lifecycle configurations for the CodeEditor application type:

- [Create and attach lifecycle configurations in Studio](#)
Create and attach lifecycle configurations in Studio

The following section provides AWS CLI commands to create a lifecycle configuration, attach a lifecycle configuration when creating a new user profile, and attach a lifecycle configuration when updating a user profile. For prerequisites and general steps on creating and attaching lifecycle configurations in Studio, see Create and associate a lifecycle configuration.

When creating your Studio lifecycle configuration with the create-studio-lifecycle-config command, be sure to specify that the studio-lifecycle-config-app-type is CodeEditor. The following example shows how to create a new Studio lifecycle configuration for your Code Editor application.

```
aws sagemaker create-studio-lifecycle-config \
--studio-lifecycle-config-name my-code-editor-lcc \
--studio-lifecycle-config-content $LCC_CONTENT \
--studio-lifecycle-config-app-type CodeEditor
```

Note the ARN of the newly created lifecycle configuration that is returned. When attaching a lifecycle configuration, provide this ARN within the LifecycleConfigArns list of CodeEditorAppSettings.

You can attach a lifecycle configuration when creating a user profile or domain. The following example shows how to create a new user profile with the lifecycle configuration attached. You can also create a new domain with a lifecycle configuration attached by using the create-domain command.

```
# Create a new UserProfile
aws sagemaker create-user-profile \
--domain-id domain-id \
--user-profile-name user-profile-name \
--user-settings '{
"CodeEditorAppSettings": {
   "LifecycleConfigArns": [
   lifecycle-configuration-arn-list
   ]
}
}
```
You can alternatively attach a lifecycle configuration when updating a user profile or domain. The following example shows how to update a user profile with the lifecycle configuration attached. You can also update a new domain with a lifecycle configuration attached by using the `update-domain` command.

```bash
# Update a UserProfile
aws sagemaker update-user-profile \
  --domain-id <domain-id> \
  --user-profile-name <user-profile-name> \
  --user-settings '{
    "CodeEditorAppSettings": {
      "LifecycleConfigArns": [
        <lifecycle-configuration-arn-list>
      ]
    }
  }'
```

**Debug lifecycle configurations in Studio**

For instructions on debugging lifecycle configurations in Studio, see [Debug lifecycle configurations](#).

To find the logs for a specific application, search the log streams using the following format:

```
domain-id/space-name/CodeEditor/default/LifecycleConfigOnStart
```

**Detach lifecycle configurations in Studio**

For steps on detaching lifecycle configurations in Studio, see [Detach lifecycle configurations](#).

To detach a lifecycle configuration using the AWS CLI, remove the desired lifecycle configuration from the list of lifecycle configurations attached to the resource. Then pass the list as part of the respective command:

- `update-user-profile`
- `update-domain`

For example, the following command removes all lifecycle configurations for the Code Editor application attached to the domain.

```bash
aws sagemaker update-domain --domain-id <domain-id> \
```

Create a lifecycle configuration to clone repositories into a Code Editor application

This section shows how to clone a repository and create a Code Editor application with the lifecycle configuration attached.

1. From your local machine, create a file named `my-script.sh` with the following content:

   ```bash
   #!/bin/bash
   set -eux
   ```

2. Clone the repository of your choice in your lifecycle configuration script.

   ```bash
   export REPOSITORY_URL="https://github.com/aws-samples/sagemaker-studio-lifecycle-config-examples.git"
   git -C /home/sagemaker-user clone $REPOSITORY_URL
   ```

3. After finalizing your script, create and attach your lifecycle configuration. For more information, see Create and attach lifecycle configurations in Studio.

4. Create your Code Editor application with the lifecycle configuration attached.

   ```bash
   aws sagemaker create-app
   --domain-id domain-id
   --space-name space-name
   --app-type CodeEditor
   --app-name default
   ```

   For more information about available Code Editor image ARNs, see Code Editor application instances and images.
Create a lifecycle configuration to install Code Editor extensions

This section shows how to create a lifecycle configuration to install extensions from the Open VSX Registry in your Code Editor environment.

1. From your local machine, create a file named `my-script.sh` with the following content:

   ```bash
   #!/bin/bash
   set -eux
   
   2. Within the script, install the Open VSX Registry extension of your choice:

   ```bash
   sagemaker-code-editor --install-extension AmazonEMR.emr-tools --extensions-dir /opt/amazon/sagemaker/sagemaker-code-editor-server-data/extensions
   
   You can retrieve the extension name from the URL of the extension in the Open VSX Registry. The extension name to use in the `sagemaker-code-editor` command should contain all text that follows `https://open-vsx.org/extension/` in the URL. Replace all instances of a slash (`/`) with a period (`.`). For example, `AmazonEMR/emr-tools` should be `AmazonEMR.emr-tools`.

3. After finalizing your script, create and attach your lifecycle configuration. For more information, see Create and attach lifecycle configurations in Studio.

4. Create your Code Editor application with the lifecycle configuration attached:
aws sagemaker create-app \
--domain-id domain-id \
--space-name space-name \
--app-type CodeEditor \
--app-name default \

For more information about available Code Editor image ARNs, see Code Editor application instances and images. For more information about connections and extensions, see Code Editor Connections and Extensions.

SageMaker HyperPod

SageMaker HyperPod helps you provision resilient clusters for running machine learning (ML) workloads and developing state-of-the-art models such as large language models (LLMs), diffusion models, and foundation models (FMs). It accelerates development of FMs by removing undifferentiated heavy-lifting involved in building and maintaining large-scale compute clusters powered by thousands of accelerators such as AWS Trainium and NVIDIA A100 and H100 Graphical Processing Units (GPUs). When accelerators fail, self-healing clusters automatically detect and replace the faulty hardware on the fly so that you can focus on running ML workloads for weeks and months without disruption. Additionally, with SageMaker HyperPod, you can customize your computing environment to best suit your needs and configure it with the Amazon SageMaker distributed training libraries to achieve optimal performance on AWS.

Operating clusters

You can create, configure, and maintain SageMaker HyperPod clusters graphically through the console user interface (UI) and programmatically through the AWS command line interface (CLI) or AWS SDK for Python (Boto3). With Amazon VPC, you can secure the cluster network and also take advantage of configuring your cluster with resources in your VPC, such as Amazon FSx for Lustre, which offers the fastest throughput. You can also give different IAM roles to cluster instance groups, and limit actions that your cluster resources and users can operate. To learn more, see the section called “Operate SageMaker HyperPod”.

Configuring your ML environment
SageMaker HyperPod runs the section called “SageMaker HyperPod DLAMI”, which sets up an ML environment on the HyperPod clusters. You can configure additional customizations to the DLAMI by providing lifecycle scripts to support your use case. To learn more about how to set up lifecycle scripts, see the section called “Getting started with SageMaker HyperPod” and the section called “SageMaker HyperPod lifecycle configuration best practices”.

**Scheduling jobs**

After you successfully create a HyperPod cluster, cluster users can log into the cluster nodes (such as head or controller node, log-in node, and worker node) and schedule jobs for running machine learning workloads. To learn more, see the section called “Run jobs on HyperPod clusters”.

**Resiliency against hardware failures**

SageMaker HyperPod runs health checks on cluster nodes and provides a workload auto-resume functionality. With the cluster resiliency features of HyperPod, you can resume your workload from the last checkpoint you saved, after faulty nodes are replaced with healthy ones in clusters with more than 16 nodes. To learn more, see the section called “Cluster resiliency”.

**Logging and managing clusters**

You can find SageMaker HyperPod resource utilization metrics and lifecycle logs in Amazon CloudWatch, and manage SageMaker HyperPod resources by tagging them. Each CreateCluster API run creates a distinct log stream, named in `<cluster-name>-<timestamp>` format. In the log stream, you can check the host names, the name of failed lifecycle scripts, and outputs from the failed scripts such as stdout and stderr. For more information, see the section called “Cluster management”.

**Compatible with SageMaker tools**

Using SageMaker HyperPod, you can configure clusters with AWS optimized collective communications libraries offered by SageMaker, such as the SageMaker distributed data parallelism (SMDDP) library. The SMDDP library implements the AllGather operation optimized to the AWS compute and network infrastructure for the most performant SageMaker machine learning instances powered by NVIDIA A100 GPUs. To learn more, see the section called “Schedule jobs for distributed training workloads on SageMaker HyperPod”.

**Topics**

- SageMaker HyperPod prerequisites
SageMaker HyperPod prerequisites

The following sections walk you through prerequisites you need to prepare before you get started with SageMaker HyperPod.

Topics

- SageMaker HyperPod quotas
- Set up IAM users and roles for SageMaker HyperPod users and resources
- Set up AWS Systems Manager and Run As for cluster user access control
- (Optional) Set up SageMaker HyperPod with your Amazon VPC
- (Optional) Set up SageMaker HyperPod with Amazon FSx for Lustre

SageMaker HyperPod quotas

You can create SageMaker HyperPod clusters given the quotas for `cluster usage` in your AWS account.

⚠️ Important

To learn more about SageMaker HyperPod pricing, see the section called “SageMaker HyperPod pricing” and Amazon SageMaker Pricing.
View Amazon SageMaker HyperPod quotas using the AWS Management Console

Look up the default and applied values of a *quota*, also referred to as a *limit*, for *cluster usage*, which is used for SageMaker HyperPod.

1. Open the Service Quotas console.
2. In the left navigation pane, choose AWS services.
3. From the AWS services list, search for and select Amazon SageMaker.
4. In the Service quotas list, you can see the service quota name, applied value (if it's available), AWS default quota, and whether the quota value is adjustable.
5. In the search bar, type cluster usage. This shows quotas for cluster usage, applied quotas, and the default quotas.

To increase Amazon SageMaker HyperPod quotas using the AWS Management Console

Increase your quotas at the account or resource level.

1. To increase the quota of instances for cluster usage, select the quota that you want to increase.
2. If the quota is adjustable, you can request a quota increase at either the account level or resource level based on the value listed in the Adjustability column.
3. For Increase quota value, enter the new value. The new value must be greater than the current value.
4. Choose Request.
5. To view any pending or recently resolved requests in the console, navigate to the Request history tab from the service's details page, or choose Dashboard from the navigation pane. For pending requests, choose the status of the request to open the request receipt. The initial status of a request is Pending. After the status changes to Quota requested, you see the case number with AWS Support. Choose the case number to open the ticket for your request.

To learn more about requesting a quota increase in general, see Requesting a Quota Increase in the AWS Service Quotas User Guide.

Set up IAM users and roles for SageMaker HyperPod users and resources

There are three main layers of SageMaker HyperPod users: AWS account admin, cluster administrators (such as cloud architects), and cluster users (such as machine learning scientists).
The AWS account admin should set up IAM users by attaching the right permissions or policies for cluster administrators. For cluster administrators, the AWS account admin also should create IAM roles that the cluster administrators can use for SageMaker HyperPod clusters to assume to run and communicate with necessary AWS resources, such as Amazon S3, Amazon CloudWatch, and AWS Systems Manager (SSM). Finally, cluster administrators can grant cluster users permissions to log into the SageMaker HyperPod clusters through SSM Agent.

**Topics**

- Set up IAM users for cluster administrators
- Set up IAM users for cluster users
- IAM role for SageMaker HyperPod

**Set up IAM users for cluster administrators**

Cluster administrators are cloud architects who operate and configure SageMaker HyperPod clusters, performing the tasks in the section called “Operate SageMaker HyperPod”. The following policy example includes the minimum set of permissions for cluster administrators to run the SageMaker HyperPod core APIs and manage any cluster within your AWS account.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "sagemaker:CreateCluster",
        "sagemaker:ListClusters"
      ],
      "Resource": "*"
    },
    {
      "Effect": "Allow",
      "Action": [
        "sagemaker:DeleteCluster",
        "sagemaker:DescribeCluster",
        "sagemaker:DescribeClusterNode",
        "sagemaker:ListClusterNodes",
        "sagemaker:UpdateCluster",
        "sagemaker:UpdateClusterSoftware"
      ]
    }
  ]
}
```
To grant permissions to access the SageMaker console, use the sample policy provided at [Permissions Required to Use the Amazon SageMaker Console](#).

To grant permissions to access the SSM console, use the sample policy provided at [Using the AWS Systems Manager console](#) in the *AWS Systems Manager User Guide*.

You might also consider attaching the [AmazonSageMakerFullAccess](#) policy to the IAM users; however, note that the AmazonSageMakerFullAccess policy grants permissions to the entire SageMaker API calls, features, and resources.

For guidance on IAM users in general, see [IAM users](#) in the *AWS Identity and Access Management User Guide*.

**Set up IAM users for cluster users**

Cluster users are machine learning engineers who log into and run ML workloads on SageMaker HyperPod cluster nodes provisioned by cluster administrators. For cluster users in your AWS account, you should grant the permission "ssm:StartSession" to run the SSM start-session command. The following is a policy example for IAM users.

**IAM permissions to all resources**

Add the following policy to give an IAM user SSM session permissions to connect to an SSM target for all resources.

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "ssm:StartSession",
        "ssm:TerminateSession"
      ],
      "Resource": "*"
    }
  ]
}
```
IAM role for SageMaker HyperPod

For SageMaker HyperPod clusters to run and communicate with necessary AWS resources, you need to attach the managed `AmazonSageMakerClusterInstanceRolePolicy` to the cluster instance groups. Given this AWS managed policy, SageMaker HyperPod cluster instance groups assume the role to communicate with Amazon CloudWatch, Amazon S3, and AWS Systems Manager Agent (SSM Agent). This managed policy is the minimum requirement for SageMaker HyperPod resources to run properly, so you must provide an IAM role with this policy to all instance groups. The `AmazonSageMakerClusterInstanceRolePolicy` has the following permissions:

- **logs** - Needed to allow SageMaker HyperPod to publish log streams.
- **cloudwatch** – Needed to allow SageMaker HyperPod to post CloudWatch metrics.
- **s3** - Needed to allow SageMaker HyperPod to list and retrieve files from an Amazon S3 bucket in your account with the prefix `sagemaker-`.
- **ssmmessages** - Needed to allow the SSM Agent to communicate with the SSM backend services. Principals can use SSM Agent for creating and opening control and data channels. SageMaker starts and manages the SSM Agent when it initiates a cluster instance.

**Tip**

Depending on your preference on designing the level of permissions for multiple instance groups, you can also set up multiple IAM roles and attach them to different instance groups. When you set up your cluster user access to specific SageMaker HyperPod cluster nodes, the nodes assume the role with the selective permissions you manually attached. When you, as a AWS account admin or cluster administrator, set up the cluster user access to specific cluster nodes through [AWS Systems Manager](https://aws.amazon.com/cloudformation/) (see also the section called “[Set up AWS Systems Manager and Run As for cluster user access control](https://aws.amazon.com/cloudformation/)”), the cluster nodes assume the role with the selective permissions you manually attach.

After you are done with creating IAM roles, make notes of their names and ARNs. You use the roles when creating a SageMaker HyperPod cluster, granting the correct permissions required for each instance group to communicate with necessary AWS resources.
If you want to use your own Amazon Virtual Private Cloud (VPC) instead of the default SageMaker VPC, you should add the following additional permissions to the IAM role for SageMaker HyperPod.

```
{
    "Effect": "Allow",
    "Action": [
        "ec2:CreateNetworkInterface",
        "ec2:CreateNetworkInterfacePermission",
        "ec2:DeleteNetworkInterface",
        "ec2:DeleteNetworkInterfacePermission",
        "ec2:DescribeNetworkInterfaces",
        "ec2:DescribeVpcs",
        "ec2:DescribeDhcpOptions",
        "ec2:DescribeSubnets",
        "ec2:DescribeSecurityGroups",
        "ec2:DetachNetworkInterface"
    ],
    "Resource": "*
}

{
    "Effect": "Allow",
    "Action": "ec2:CreateTags",
    "Resource": ["arn:aws:ec2:***:network-interface/**"
    ]
}
```

The following list breaks down which permissions are needed to enable SageMaker HyperPod cluster functionalities when you configure the cluster with your own Amazon VPC.

- The following ec2 permissions are required to enable configuring a SageMaker HyperPod cluster with your VPC.

```
{
    "Effect": "Allow",
    "Action": [
        "ec2:CreateNetworkInterface",
        "ec2:CreateNetworkInterfacePermission",
        "ec2:DeleteNetworkInterface",
    ]
}
```
• The following `ec2` permission is required to enable the SageMaker HyperPod auto-resume functionality.

```json
{
    "Effect": "Allow",
    "Action": [
        "ec2:DeleteNetworkInterface",
        "ec2:DescribeNetworkInterfaces",
        "ec2:DescribeVpcs",
        "ec2:DescribeDhcpOptions",
        "ec2:DescribeSubnets",
        "ec2:DescribeSecurityGroups"
    ],
    "Resource": "*"
}
```

• The following `ec2` permission allows SageMaker HyperPod to create tags on the network interfaces within your account.

```json
{
    "Effect": "Allow",
    "Action": "ec2:CreateTags",
    "Resource": [
        "arn:aws:ec2::*::*:network-interface/*"
    ],
    "Resource": "*"
}
```

### Set up AWS Systems Manager and Run As for cluster user access control

The section called “SageMaker HyperPod DLAMI” comes with AWS Systems Manager (SSM) out of the box to help you manage access to your SageMaker HyperPod cluster instance groups. This section describes how to create operating system (OS) users in your SageMaker HyperPod clusters and associate them with IAM users and roles. This is useful to authenticate SSM sessions using the credentials of the OS user account.
Enable Run As in your AWS account

As an AWS account admin or a cloud administrator, you can manage access to SageMaker HyperPod clusters at an IAM role or user level by using the Run As feature in SSM. With this feature, you can start each SSM session using the OS user associated to the IAM role or user.

To enable Run As in your AWS account, follow the steps in Turn on Run As support for Linux and macOS managed nodes. If you already created OS users in your cluster, make sure that you associate them with IAM roles or users by tagging them as guided in Option 2 of step 5 under To turn on Run As support for Linux and macOS managed nodes.

Prepare a script for setting up Linux users

To complete setting up cluster users to access a HyperPod cluster through SSM, you need to configure a script for adding users while preparing lifecycle configuration scripts for creating a HyperPod cluster. In the template repository provided in the section called “Start with base lifecycle scripts provided by HyperPod”, there is a script named add_users.sh that consumes the user data from shared_users.txt. There's also a template named shared_users_sample.txt, so refer to the sample text file to fill and create a shared_users.txt file. Note that you'll need to upload the two files as part of preparing and uploading lifecycle scripts to an S3 bucket, which you'll learn in the section called “Getting started with SageMaker HyperPod”.

(Optional) Set up SageMaker HyperPod with your Amazon VPC

If you don't provide a VPC, SageMaker HyperPod uses the default SageMaker VPC. To set up a SageMaker HyperPod cluster with your Amazon VPC, check the following items.

- If you want to use your own VPC to connect SageMaker HyperPod with AWS resources in your VPC, you need to provide the VPC name, ID, AWS Region, subnet ID, and security group ID when you create SageMaker HyperPod. If you want to create a new VPC, see Create a default VPC or Create a VPC in the Amazon Virtual Private Cloud User Guide.

- It is important that you should create all your resources in the same AWS Region and Availability Zone, and configure security group rules to allow connection between the resources in your VPC. For example, assume that you create a VPC in us-west-2. You should create a subnet in this VPC in Availability Zone us-west-2a, and create a security group that allows all incoming (inbound) traffic from inside the security group and all outbound traffic.

- You also need to ensure that your VPC has connection to Amazon Simple Storage Service (S3). If you configure a VPC, SageMaker HyperPod instance groups don't have access to the internet,
and therefore can't connect to Amazon S3 for accessing or storing files such as lifecycle scripts, training data, and model artifacts. To establish connection with Amazon S3 while using VPC, you should create a VPC endpoint. By creating a VPC endpoint, you can allow the SageMaker HyperPod instance groups to access the S3 buckets within the same VPC. We recommend that you also create a custom policy that only allows requests from your private VPC to access your S3 buckets. For more information, see [Endpoints for Amazon S3](https://docs.aws.amazon.com/AmazonS3/latest/userguide/privateconnections.html) in the [AWS PrivateLink Guide](https://docs.aws.amazon.com/AmazonS3/latest/userguide/privateconnections.html).

- If you want to create a HyperPod cluster with EFA-enabled instances, make sure that you set up a security group to allow all inbound and outbound traffic to and from the security group itself. To learn more, see [Step 1: Prepare an EFA-enabled security group](https://docs.aws.amazon.com/AmazonS3/latest/userguide/privateconnections.html) in the [Amazon EC2 User Guide](https://docs.aws.amazon.com/AWSEC2/latest/UserGuide/elastic-fabric-adapter.html).

**(Optional) Set up SageMaker HyperPod with Amazon FSx for Lustre**

To start using SageMaker HyperPod and mapping data paths between the cluster and your FSx for Lustre file system, select one of the AWS Regions supported by SageMaker HyperPod. After choosing the AWS Region you prefer, you also should determine which Availability Zone (AZ) to use. If you use SageMaker HyperPod compute nodes in AZs different from the AZs where your FSx for Lustre file system is set up within the same AWS Region, there might be communication and network overhead. We recommend that you to use the same physical AZ as the one for the SageMaker HyperPod service account to avoid any cross-AZ traffic between SageMaker HyperPod clusters and your FSx for Lustre file system. Also, make sure that you have configured it with your VPC. If you want to use Amazon FSx as the main file system for storage, you must configure SageMaker HyperPod clusters with VPC.

**Getting started with SageMaker HyperPod**

Get started with creating your first SageMaker HyperPod cluster and learn the cluster operation functionalities of SageMaker HyperPod.

You can create a SageMaker HyperPod cluster through the SageMaker console UI or the AWS CLI commands. This tutorial shows how to create a new SageMaker HyperPod cluster with Slurm, which is a popular workload scheduler software. After you go through this tutorial, you will know how to log into the cluster nodes using the AWS Systems Manager commands (`aws ssm`). After you complete this tutorial, see also [the section called “Operate SageMaker HyperPod”](https://docs.aws.amazon.com/AmazonS3/latest/userguide/privateconnections.html) to learn more about the SageMaker HyperPod basic operations, and [the section called “Run jobs on HyperPod clusters”](https://docs.aws.amazon.com/AmazonS3/latest/userguide/privateconnections.html) to learn how to schedule jobs on the provisioned cluster.

**Topics**
Using the SageMaker HyperPod console UI

Create your first SageMaker HyperPod cluster using the SageMaker HyperPod console UI.

Create your first SageMaker HyperPod cluster with Slurm

The following tutorial demonstrates how to create a new SageMaker HyperPod cluster and set it up with Slurm through the SageMaker console UI. Following the tutorial, you'll create a HyperPod cluster with three Slurm nodes, my-controller-group, my-login-group, and worker-group-1.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose HyperPod Clusters in the left navigation pane.
3. On the SageMaker HyperPod Clusters page, choose Create cluster.
4. In Step 1: Cluster settings, specify a name for the new cluster. Skip the Tags section.
5. In Step 2: Instance groups, add instance groups. Each instance group can be configured differently, and you can create a heterogeneous cluster that consists of multiple instance groups with various instance types. For lifecycle configuration scripts to run on the instance group during cluster creation, you can start with using the sample lifecycle scripts provided in the Awsome Distributed Training GitHub repository.

   a. For Instance group name, specify a name for the instance group. For this tutorial, create three instance groups named my-controller-group, my-login-group, and worker-group-1.

   b. For Select instance type, choose the instance for the instance group. For this tutorial, select ml.c5.xlarge for my-controller-group, ml.m5.4xlarge for my-login-group, and ml.trn1.32xlarge for worker-group-1. If you receive an error due to resource limit, make sure that you change the instance type to what you already have in your account or request quotas by following the instructions at the section called “SageMaker HyperPod quotas”.

   c. For Quantity, specify an integer not exceeding the instance quota for cluster usage. For this tutorial, enter 1 for all three groups.
d. For **S3 path to lifecycle script files**, enter the Amazon S3 path in which your lifecycle scripts are stored. If you don't have lifecycle scripts, go through the following substeps to use the base lifecycle scripts provided by the SageMaker HyperPod service team.

i. Clone the [Awsome Distributed Training GitHub repository](https://github.com/aws-samples/awsome-distributed-training/)

```bash
git clone https://github.com/aws-samples/awsome-distributed-training/
```

ii. Under `1.architectures/5.sagemaker_hyperpods/LifecycleScripts/base-config`, you can find a set of base lifecycle scripts. To learn more about the lifecycle scripts, see also the section called “Prepare lifecycle scripts for setting up Slurm on SageMaker HyperPod”.

iii. Write a Slurm configuration file and save it as `provisioning_params.json`. In the file, specify basic Slurm configuration parameters to properly assign Slurm nodes to the SageMaker HyperPod cluster instance groups. For example, the `provisioning_params.json` should be similar to the following based on the HyperPod cluster instance group configured through the previous steps 5a, 5b, and 5c.

```json
{
    "version": "1.0.0",
    "workload_manager": "slurm",
    "controller_group": "my-controller-group",
    "login_group": "my-login-group",
    "worker_groups": [
        {
            "instance_group_name": "worker-group-1",
            "partition_name": "partition-1"
        }
    ]
}
```

iv. Upload the scripts to your Amazon S3 bucket. Create an S3 bucket with a path in the following format: `s3://sagemaker-<unique-s3-bucket-name>/<lifecycle-script-directory>/src`. You can create this bucket using the Amazon S3 console.
Note

You must prefix sagemaker- to the S3 bucket path, because the `***` with AmazonSageMakerClusterInstanceRolePolicy only allows principals to access S3 buckets with this specific prefix.

e. For **Directory path to your on-create lifecycle script**, enter the file name of the lifecycle script under **S3 path to lifecycle script files**.

f. For **IAM role**, choose the IAM role you created using the AmazonSageMakerClusterInstanceRolePolicy from the section **the section called “IAM role for SageMaker HyperPod”**.

g. For **Threads per core** under **Advanced configuration**, specify 1 for disabling multi-threading and 2 for enabling multi-threading. To find which instance type supports multi-threading, see the reference table of **CPU cores and threads per CPU core per instance type** in the Amazon Elastic Compute Cloud User Guide.

6. In **Step 3: Advanced configuration**, set up network settings within, and in and out of, the cluster. Select your own VPC if you already have one that gives SageMaker access to your VPC. If you don’t have one but want to create a new VPC, follow the instructions at **Create a VPC** in the Amazon Virtual Private Cloud User Guide. You can leave it as **no VPC** to use the default SageMaker VPC.

7. In **Step 4: Review and create**, review the configuration you’ve set from step 1 to 3 and finish submitting the cluster creation request.

8. The new cluster should appear under **Clusters** in the main pane of the SageMaker HyperPod console. You can check the status of it displayed under the **Status** column.

9. After the status of the cluster turns to **InService**, cluster users can start logging into the cluster nodes. For more information about accessing the cluster nodes and running ML workloads, see **the section called “Run jobs on HyperPod clusters”**.

Delete the cluster and clean resources

After you have successfully tested creating a SageMaker HyperPod cluster, it continues running in the **InService** state until you delete the cluster. We recommend that you delete any clusters created using on-demand SageMaker instances when not in use to avoid incurring continued service charges based on on-demand pricing. In this tutorial, you have created a cluster that
consists of two instance groups. One of them uses a C5 instance, so make sure you delete the cluster by following the instructions at the section called “Delete a SageMaker HyperPod cluster”.

However, if you have created a cluster with reserved compute capacity, the status of the clusters does not affect service billing.

To clean up the lifecycle scripts from the S3 bucket used for this tutorial, go to the S3 bucket you used during cluster creation and remove the files entirely.

If you have tested running any workloads on the cluster, make sure if you have uploaded any data or if your job saved any artifacts to different S3 buckets or file system services such as Amazon FSx for Lustre and Amazon Elastic File System. To prevent any incurring charges, delete all artifacts and data from the storage or file system.

**Using the AWS CLI commands for the SageMaker HyperPod APIs**

Create your first SageMaker HyperPod cluster using the AWS CLI commands for HyperPod.

**Create your first SageMaker HyperPod cluster with Slurm**

The following tutorial demonstrates how to create a new SageMaker HyperPod cluster and set it up with Slurm through the [AWS CLI commands for SageMaker HyperPod](https://docs.aws.amazon.com/sagemaker/latest/dg/hyperpod-cli.html). Following the tutorial, you'll create a HyperPod cluster with three Slurm nodes, `my-controller-group`, `my-login-group`, and `worker-group-1`.

1. First, prepare and upload lifecycle scripts to an S3 bucket. During cluster creation, HyperPod runs them in each instance group. Upload lifecycle scripts to S3 using the following command.

   ```bash
   aws s3 sync ~/local-dir-to-lifecycle-scripts/* s3://sagemaker-<unique-s3-bucket-name>/<lifecycle-script-directory>/src
   ```

   **Note**

   The S3 bucket path should start with a prefix `sagemaker-`, because the [AmazonSageMakerClusterInstanceRolePolicy](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-cluster-instance-role.html) only allows access to S3 buckets that starts with the specific prefix.
If you are starting from scratch, use sample lifecycle scripts provided in the Awsome Distributed Training GitHub repository. The following sub-steps show how to download, what to modify, and how to upload the sample lifecycle scripts to an S3 bucket.

a. Download a copy of the lifecycle script samples to a directory on your local computer.

```
git clone https://github.com/aws-samples/awesome-distributed-training/
```

b. Go into the directory `1.architectures/5.sagemaker_hyperpods/LifecycleScripts/base-config`, where you can find a set of lifecycle scripts.

```
cd awsome-distributed-training/1.architectures/5.sagemaker_hyperpods/LifecycleScripts/base-config
```

To learn more about the lifecycle script samples, see the section called “Prepare lifecycle scripts for setting up Slurm on SageMaker HyperPod”.

c. Write a Slurm configuration file and save it as `provisioning_params.json`. In the file, specify basic Slurm configuration parameters to properly assign Slurm nodes to the SageMaker HyperPod cluster instance groups. In this tutorial, set up three Slurm nodes named `my-controller-group`, `my-login-group`, and `worker-group-1`, as shown in the following example configuration `provisioning_params.json`.

```
{
    "version": "1.0.0",
    "workload_manager": "slurm",
    "controller_group": "my-controller-group",
    "login_group": "my-login-group",
    "worker_groups": [
        {
            "instance_group_name": "worker-group-1",
            "partition_name": "partition-1"
        }
    ]
}
```

d. Upload the scripts to `s3://sagemaker-<unique-s3-bucket-name>/<lifecycle-script-directory>/src`. You can do so by using the S3 console, or by running the following AWS CLI S3 command.
2. Prepare a CreateCluster request file in JSON format and save as create_cluster.json. The following request example is based on the Slurm nodes defined in the provisioning_params.json in step 1c. For ExecutionRole, provide the ARN of the IAM role you created with the managed AmazonSageMakerClusterInstanceRolePolicy in the section called “Prerequisites”. If you receive an error due to resource limit, make sure that you change the instance type to what you already have in your account or request quotas by following the instructions at the section called “SageMaker HyperPod quotas”.

```json
{
    "ClusterName": "my-hyperpod-cluster",
    "InstanceGroups": [
        {
            "InstanceGroupName": "my-controller-group",
            "InstanceType": "ml.c5.xlarge",
            "InstanceCount": 1,
            "LifeCycleConfig": {
                "SourceS3Uri": "s3://sagemaker-<unique-s3-bucket-name>/lifecycle-script-directory/src",
                "OnCreate": "on_create.sh"
            },
            "ExecutionRole": "${ROLE}",
            "ThreadsPerCore": 1
        },
        {
            "InstanceGroupName": "my-login-group",
            "InstanceType": "ml.m5.4xlarge",
            "InstanceCount": 1,
            "LifeCycleConfig": {
                "SourceS3Uri": "s3://sagemaker-<unique-s3-bucket-name>/lifecycle-script-directory/src",
                "OnCreate": "on_create.sh"
            },
            "ExecutionRole": "${ROLE}",
            "ThreadsPerCore": 1
        },
        {
            "InstanceGroupName": "worker-group-1",
```
3. Run the following command to create the cluster.

```bash
aws sagemaker create-cluster --cli-input-json file://complete/path/to/create_cluster.json
```

This should return the ARN of the created cluster.

4. Run `describe-cluster` to check the status of the cluster.

```bash
aws sagemaker describe-cluster --cluster-name my-hyperpod-cluster
```

After the status of the cluster turns to **InService**, proceed to the next step.

5. Run `list-cluster-nodes` to check the details of the cluster nodes.

```bash
aws sagemaker list-cluster-nodes --cluster-name my-hyperpod-cluster
```

This returns a response, and the InstanceId is what your cluster users need for logging (aws ssm) into them. For more information about logging into the cluster nodes and running ML workloads, see the section called “Run jobs on HyperPod clusters”.

**Delete the cluster and clean resources**

After you have successfully tested creating a SageMaker HyperPod cluster, it continues running in the **InService** state until you delete the cluster. We recommend that you delete any clusters created using on-demand SageMaker capacity when not in use to avoid incurring continued service charges based on on-demand pricing. In this tutorial, you have created a cluster that consists
of two instance groups. One of them uses a C5 instance, so make sure you delete the cluster by running the following command.

```
aws sagemaker delete-cluster --cluster-name my-hyperpod-cluster
```

To clean up the lifecycle scripts from the S3 bucket used for this tutorial, go to the S3 bucket you used during cluster creation and remove the files entirely.

If you have tested running any model training workloads on the cluster, also check if you have uploaded any data or if your job has saved any artifacts to different S3 buckets or file system services such as Amazon FSx for Lustre and Amazon Elastic File System. To prevent incurring charges, delete all artifacts and data from the storage or file system.

**Operate SageMaker HyperPod**

This section walks you through how to operate SageMaker HyperPod clusters through the console UI or AWS CLI.

**Topics**

- **Using the SageMaker HyperPod console UI**
- **Using the AWS CLI**

**Using the SageMaker HyperPod console UI**

The following topics provide guidance on how to operate SageMaker HyperPod through the console UI.

**Topics**

- **Create a SageMaker HyperPod cluster**
- **Browse your SageMaker HyperPod clusters**
- **View details of each SageMaker HyperPod cluster**
- **Edit a SageMaker HyperPod cluster**
- **Delete a SageMaker HyperPod cluster**
Create a SageMaker HyperPod cluster

See the following instructions on creating a new SageMaker HyperPod cluster through the SageMaker HyperPod console UI.

2. Choose **HyperPod Clusters** in the left navigation pane.
3. In the SageMaker HyperPod landing page, choose **Create cluster**.
4. In **Step 1: Cluster settings**, set up basic information for the cluster.
   a. For **Cluster name**, specify a name for the new cluster.
   b. For **Tags**, add key and value pairs to the new cluster and manage the cluster as an AWS resource. To learn more, see [Tagging your AWS resources](https).
5. In **Step 2: Instance groups**, choose **Create instance group**. Each instance group can be configured differently, and you can create a heterogeneous cluster that consists of multiple instance groups with various instance types. In the **Create an instance group** configuration pop-up window, fill the instance group configuration information.
   a. For **Instance group name**, specify a name for the instance group.
   b. For **Select instance type**, choose the instance for the instance group.
   c. For **Quantity**, specify an integer not exceeding the instance quota for cluster usage.
   d. For **Amazon S3 path to lifecycle script files**, enter the S3 path in which your lifecycle scripts are stored.
   e. For **Directory path to your on-create lifecycle script**, enter the file name of the lifecycle script under **S3 path to lifecycle script files**.
   f. For **IAM role**, choose the IAM role you have created for SageMaker HyperPod resources, following the section **Set up IAM users and roles for SageMaker HyperPod users and resources**.
   g. For **Advanced configuration**, you can set Threads per core, specify 1 for disabling multi-threading and 2 for enabling multi-threading. To find which instance type supports multi-threading, see the reference table of **CPU cores and threads per CPU core per instance type** in the Amazon EC2 User Guide.
6. In **Step 3: Advanced configuration**, configure optional network settings within cluster and in-and-out of the cluster. Select your own VPC if you already have one that gives SageMaker access to your resources under the VPC. If you want to create a new VPC, see [Create a default VPC](https).
VPC or Create a VPC in the Amazon Virtual Private Cloud User Guide. If you don't make any selections, it picks up the default VPC of your account.

Note
If you want to use your own VPC, you should add additional permissions to the IAM role for SageMaker HyperPod clusters. To learn more, see the section called “(Optional) Set up SageMaker HyperPod with your Amazon VPC”.

7. In Step 4: Review and create, review the configuration you have set from Step 1 to Step 3 and finish submitting the cluster creation request.

Browse your SageMaker HyperPod clusters

Under Clusters on the SageMaker HyperPod console main page, all created clusters should appear listed under the Clusters section, which provides a summary view of clusters, their ARNs, status, and creation time.

View details of each SageMaker HyperPod cluster

Under Clusters on the console main page, the cluster Names are activated as links. Choose the cluster name link to see details of each cluster.

Edit a SageMaker HyperPod cluster

1. Under Clusters, choose the cluster you want to update.
2. Choose Actions button, and choose Edit cluster.
3. In the Edit <your-cluster> page, you can edit the configurations of existing instance groups, add more instance groups, and change tags for the cluster. After making changes, choose Submit. Note that currently you cannot reduce or delete existing instance groups.
   a. In the Configure instance groups section, you can add more instance groups by choosing Create cluster group.
   b. In the Configure instance groups section, you can choose one of the instance groups, and choose Edit to change its configuration.
   c. In the Tags section, you can update tags for the cluster.
Delete a SageMaker HyperPod cluster

1. Under **Clusters**, choose the cluster you want to delete.

2. Choose **Actions**, and choose **Delete cluster**.

3. In the pop-up window for cluster deletion, review the cluster information carefully to confirm that you chose the right cluster to delete.

4. After you reviewed the cluster information, choose **Yes, delete cluster**.

5. In the text field to confirm this deletion, type **delete**.

6. Choose **Delete** on the lower right corner of the pop-up window to finish sending the cluster deletion request.

Using the AWS CLI

The following topics provide guidance on writing SageMaker HyperPod API request files in JSON format and run them using the AWS CLI commands.

**Topics**

- [Create a new cluster](#)
- [Describe a cluster](#)
- [List details of cluster nodes](#)
- [Describe details of a cluster node](#)
- [List clusters](#)
- [Update cluster configuration](#)
- [Update the SageMaker HyperPod platform software of a cluster](#)
- [Delete a cluster](#)

Create a new cluster

1. Prepare lifecycle configuration scripts and upload them to an S3 bucket, such as `s3://sagemaker-<your-s3-bucket>/<lifecycle-script-directory>/src/`. The following step 2 assumes that there’s an entry point script named `on_create.sh` in the specified S3 bucket.
Important

Make sure that you set the S3 path to start with s3://sagemaker-. The section called “IAM role for SageMaker HyperPod” has the managed AmazonSageMakerClusterInstanceRolePolicy attached, which allows access to S3 buckets with the specific prefix sagemaker-.

2. Prepare a CreateCluster API request file in JSON format. The following request example is based on the nodes needed for groups defined in the provisioning_params.json in Step 1.2. The minimum requirement for Slurm is one controller and one worker group. For ExecutionRole, provide the ARN of the IAM role you created with the managed AmazonSageMakerClusterInstanceRolePolicy from the section the section called “IAM role for SageMaker HyperPod”.

```json
// create_cluster.json
{
    // Required
    "ClusterName": "your-hyperpod-cluster",
    // Required
    "InstanceGroups": [
        {
            "InstanceGroupName": "controller-group",
            "InstanceType": "ml.m5.xlarge",
            "InstanceCount": 1,
            "LifeCycleConfig": {
                "SourceS3Uri": "s3://sagemaker-<your-s3-bucket>/<lifecycle-script-directory>/src/",
                "OnCreate": "on_create.sh"
            },
            "ExecutionRole": "arn:aws:iam::111122223333:role/iam-role-for-cluster",
            "ThreadsPerCore": 1
        },
        {
            "InstanceGroupName": "worker-group-1",
            "InstanceType": "ml.p4d.xlarge",
            "InstanceCount": 1,
            "LifeCycleConfig": {
                "SourceS3Uri": "s3://sagemaker-<your-s3-bucket>/<lifecycle-script-directory>/src/",
                "OnCreate": "on_create.sh"
            }
        }
    ]
}
```
Depending on how you designed the cluster structure through your lifecycle scripts, you can add and configure multiple instance groups under the `InstanceGroups` request parameter.

For the `Tags` request parameter, you can add custom tags for managing the SageMaker HyperPod cluster as an AWS resource. You can add tags to your cluster in the same way you add them in other AWS services that support tagging. To learn more about tagging AWS resources in general, see [Tagging AWS Resources User Guide](#).

For the `VpcConfig` request parameter, specify the information of a VPC you want to use. For more information, see [the section called “(Optional) Set up SageMaker HyperPod with your Amazon VPC”](#).

3. Run the following command to submit the `CreateCluster` API request.

```bash
aws sagemaker create-cluster \
   --cli-input-json file://complete/path/to/create_cluster.json
```

This should return the ARN of the new cluster.
Describe a cluster

Run `describe-cluster` to check the status of the cluster. You can specify either the name or the ARN of the cluster.

```
aws sagemaker describe-cluster --cluster-name your-hyperpod-cluster
```

After the status of the cluster turns to **InService**, proceed to the next step. Using this API, you can also retrieve failure messages from running other HyperPod API operations.

List details of cluster nodes

Run `list-cluster-nodes` to check the key information of the cluster nodes.

```
aws sagemaker list-cluster-nodes --cluster-name your-hyperpod-cluster
```

This returns a response, and the InstanceId is what you need to use for logging (using `aws ssm`) into them.

Describe details of a cluster node

Run `describe-cluster-node` to retrieve details of a cluster node. You can get the cluster node ID from list-cluster-nodes output. You can specify either the name or the ARN of the cluster.

```
aws sagemaker describe-cluster-node \
   --cluster-name your-hyperpod-cluster \
   --node-id i-111222333444555aa
```

List clusters

Run `list-clusters` to list all clusters in your account.

```
aws sagemaker list-clusters
```

You can also add additional flags to filter the list of clusters down. To learn more about what this command runs at low level and additional flags for filtering, see the [ListClusters](#) API reference.

Update cluster configuration

Run `update-cluster` to update the configuration of a cluster.
1. Create an `UpdateCluster` request file in JSON format. Make sure that you specify the right cluster name and instance group name to update. You can change the instance type, the number of instances, the lifecycle configuration entrypoint script, and the path to the script.

   a. For `ClusterName`, specify the name of the cluster you want to update.
   
   b. For `InstanceGroupName`
      
      i. To update an existing instance group, specify the name of the instance group you want to update.
      
      ii. To add a new instance group, specify a new name not existing in your cluster.
   
   c. For `InstanceType`
      
      i. To update an existing instance group, you must match the instance type you initially specified to the group.
      
      ii. To add a new instance group, specify an instance type you want to configure the group with.
   
   d. For `InstanceCount`
      
      i. To update an existing instance group, specify an integer greater than the current number of instances. Currently, you can only increase the number of instances.
      
      ii. To add a new instance group, specify an integer greater or equal to 1.
   
   e. For `LifeCycleConfig`, you can change both `SourceS3Uri` and `OnCreat` values as you want to update the instance group.
   
   f. For `ExecutionRole`
      
      i. For updating an existing instance group, keep using the same IAM role you attached during cluster creation.
      
      ii. For adding a new instance group, specify an IAM role you want to attach.
   
   g. For `TreadsPerCore`
      
      i. For updating an existing instance group, keep using the same value you specified during cluster creation.
      
      ii. For adding a new instance group, you can choose any value from the allowed options per instance type. For more information, search the instance type and see the **Valid treads per core** column in the reference table at [CPU cores and threads per CPU core per instance type](https://docs.aws.amazon.com/AWSEC2/latest/UserGuide/cpu-cores-threads.html) in the *Amazon EC2 User Guide*.

The following code snippet is a JSON request file template you can use. For more information about the request syntax and parameters of this API, see the [UpdateCluster API reference](https://docs.aws.amazon.com/sagemaker/latest/dg/API_UpdateCluster.html).
// update_cluster.json
{
  // Required
  "ClusterName": "name-of-cluster-to-update",
  // Required
  "InstanceGroups": [
    {
      "InstanceGroupName": "name-of-instance-group-to-update",
      "InstanceType": "ml.m5.xlarge",
      "InstanceCount": 1,
      "LifeCycleConfig": {
        "SourceS3Uri": "s3://sagemaker-<your-s3-bucket>/<lifecycle-script-directory>/src/",
        "OnCreate": "on_create.sh"
      },
      "ExecutionRole": "arn:aws:iam::111122223333:role/iam-role-for-cluster",
      "ThreadsPerCore": 1
    },
    // add more blocks of instance groups as needed
    { ... }
  ]
}

2. Run the following `update-cluster` command to submit the request.

```
aws sagemaker update-cluster \
  --cli-input-json file://complete/path/to/update_cluster.json
```

**Update the SageMaker HyperPod platform software of a cluster**

Run `update-cluster-software` to update existing clusters with software and security patches provided by the SageMaker HyperPod service. For `--cluster-name`, specify either the name or the ARN of the cluster to update.

⚠️ **Important**

Note that you must back up your work before running this API. The patching process replaces the root volume with the updated AMI, which means that your previous data stored in the instance root volume will be lost. Make sure that you back up your data from
the instance root volume to Amazon S3 or Amazon FSx for Lustre. For more information, see the section called “Use the backup script provided by SageMaker HyperPod”.

```
aws sagemaker update-cluster-software --cluster-name your-hyperpod-cluster
```

This command calls the UpdateClusterSoftware API. After the API call, SageMaker HyperPod updates the cluster instances to use the latest the section called “SageMaker HyperPod DLAMI” and runs your lifecycle scripts in the S3 bucket that you specified during cluster creation or update. The SageMaker HyperPod service team regularly rolls out new the section called “SageMaker HyperPod DLAMI”’s for enhancing security and improving user experiences. We recommend that you always keep updating to the latest SageMaker HyperPod DLAMI. For future SageMaker HyperPod DLAMI updates for security patching, follow up with the section called “HyperPod release notes”.

Tip

If the security patch fails, you can retrieve failure messages by running the DescribeCluster API as instructed at the section called “Describe a cluster”.

Note

You can only run this API programatically. The patching functionality is not implemented in the SageMaker HyperPod console UI.

Use the backup script provided by SageMaker HyperPod

SageMaker HyperPod provides a script to back up and restore your data at 1.architectures/5.sagemaker-hyperpod/patching-backup.sh in the Awsome Distributed Training GitHub repository. The script provides the following two functions.

To back up data to an S3 bucket before patching

```
sudo bash patching-backup.sh --create <s3-backup-bucket-path>
```
After you run the command, the script checks squeue if there are queued jobs, stops Slurm if there's no job in the queue, backs up mariadb, and copies local items on disc defined under LOCAL_ITEMS. You can add more files and directories to LOCAL_ITEMS.

```bash
# Define files and directories to back up.
LOCAL_ITEMS=(
    "/var/spool/slurmd"
    "/var/spool/slurmctld"
    "/etc/systemd/system/slurmctld.service"
    "/home/ubuntu/backup_slurm_acct_db.sql"
    # ... Add more items as needed
)
```

Also, you can add custom code to the provided script to back up any applications for your use case.

To restore data from an S3 bucket after patching

```bash
sudo bash patching-backup.sh --restore <s3-buckup-bucket-path>
```

Delete a cluster

Run delete-cluster to delete a cluster. You can specify either the name or the ARN of the cluster.

```bash
aws sagemaker delete-cluster --cluster-name your-hyperpod-cluster
```

SageMaker HyperPod lifecycle configuration best practices

SageMaker HyperPod offers always up-and-running compute clusters, which are highly customizable as you can write lifecycle scripts to tell SageMaker HyperPod how to set up the cluster resources. The following topics are best practices for preparing lifecycle scripts to set up SageMaker HyperPod clusters with open source workload manager tools.

Prepare lifecycle scripts for setting up Slurm on SageMaker HyperPod

The following topics discuss how to prepare lifecycle scripts to set up Slurm on SageMaker HyperPod.

Topics
- High-level overview
• Start with base lifecycle scripts provided by HyperPod
• What particular configurations HyperPod manages in Slurm configuration files
• Mount Amazon FSx for Lustre to your HyperPod cluster
• Validate the JSON configuration files before running create-cluster
• Develop lifecycle scripts interactively on a cluster node
• Update a cluster with new or updated lifecycle scripts
• Considerations

High-level overview

The following procedure is the main flow of provisioning a HyperPod cluster and setting it up with Slurm. The steps are put in order of a **bottom-up** approach.

1. Plan how you want to create Slurm nodes on a HyperPod cluster. For example, if you want to configure two Slurm nodes, you'll need to set up two instance groups in a HyperPod cluster.

2. Prepare a `provisioning_params.json` file, which is a [the section called “Configuration form for provisioning Slurm nodes on HyperPod”](#). `provisioning_params.json` should contain Slurm node configuration information to be provisioned on the HyperPod cluster. This should reflect the design of Slurm nodes from Step 1.

3. Prepare a set of lifecycle scripts to set up Slurm on HyperPod to install software packages and set up an environment in the cluster for your use case. You should structure the lifecycle scripts to collectively run in order in a central Python script (`lifecycle_script.py`), and write an entrypoint shell script (`on_create.sh`) to run the Python script. The entrypoint shell script is what you need to provide to a HyperPod cluster creation request later in Step 5.

   Also, note that you should write the scripts to expect `resource_config.json` that will be generated by HyperPod during cluster creation. `resource_config.json` contains HyperPod cluster resource information such as IP addresses, instance types, and ARNs, and is what you need to use for configuring Slurm.

4. Collect all the files from the previous steps into a folder.

```bash
### lifecycle_files // your local folder
    ### provisioning_params.json
    ### on_create.sh
    ### lifecycle_script.py
    ### ... // more setup scripts to be fed into lifecycle_script.py
```
5. Upload all the files to an S3 bucket. Copy and keep the S3 bucket path. Note that you should create an S3 bucket path starting with `sagemaker-` because you need to choose an the section called “IAM role for SageMaker HyperPod” attached with `AmazonSageMakerClusterInstanceRolePolicy`, which only allows S3 bucket paths starting with the prefix `sagemaker-`. The following command is an example command to upload all the files to S3.

```bash
aws s3 cp --recursive ./lifecycle_files s3://sagemaker-hyperpod-lifecycle/src
```

6. Prepare a HyperPod cluster creation request. If you use the AWS CLI, write a cluster creation request in JSON format (`create_cluster.json`). If you use the SageMaker console UI, fill the Create a cluster request form in the HyperPod console UI. At this stage, make sure that you create instance groups in the same structure that you planned in Step 1 and 2. Also, make sure that you specify the S3 bucket from Step 5 in the request forms.

7. Submit the cluster creation request. HyperPod provisions a cluster based on the request, creates a `resource_config.json` file in the HyperPod cluster instances, and sets up Slurm on the cluster running the lifecycle scripts.

The following section walks you through and dives deep into details on how to organize configuration files and lifecycle scripts to work properly during HyperPod cluster creation.

**Start with base lifecycle scripts provided by HyperPod**

This section walks you through every component of the basic flow of setting up Slurm on HyperPod in a *top-down* approach. It starts from preparing a HyperPod cluster creation request to run the CreateCluster API, and dives deep into the hierarchical structure down to lifecycle scripts. Use the sample lifecycle scripts provided in the [Awesome Distributed Training GitHub repository](https://github.com/aws-samples/awsome-distributed-training/). Clone the repository by running the following command.

```bash
git clone https://github.com/aws-samples/awsome-distributed-training/
```

The base lifecycle scripts for setting up a Slurm cluster on SageMaker HyperPod are available at [1.architectures/5.sagemaker_hyperpods/LifecycleScripts/base-config](https://github.com/aws-samples/awsome-distributed-training/1.architectures/5.sagemaker_hyperpods/LifecycleScripts/base-config).

The following flowchart shows a detailed overview of how you should design the base lifecycle scripts. The descriptions below the diagram and the procedural guide explain how they work during the HyperPod CreateCluster API call.
Figure: A detailed flowchart of HyperPod cluster creation and the structure of lifecycle scripts. (1) The dashed arrows are directed to where the boxes are "called into" and shows the flow of configuration files and lifecycle scripts preparation. It starts from preparing `provisioning_params.json` and lifecycle scripts. These are then coded in `lifecycle_script.py` for a collective execution in order. And the execution of the `lifecycle_script.py` script is done by the `on_create.sh` shell script, which to be run in the HyperPod instance terminal. (2) The solid arrows show the main HyperPod cluster creation flow and how the boxes are "called into" or "submitted to". `on_create.sh` is required for cluster creation request, either in `create_cluster.json` or the **Create a cluster** request form in the console UI. After you submit the request, and HyperPod runs the `CreateCluster API` based on the given configuration information from the request and the lifecycle scripts. (3) The dotted arrow indicates that the HyperPod platform creates `resource_config.json` in the cluster instances during cluster...
resource provisioning. resource_config.json contains HyperPod cluster resource information such as the cluster ARN, instance types, and IP addresses. It is important to note that you should prepare the lifecycle scripts to expect the resource_config.json file during cluster creation. For more information, see the procedural guide below.

The following procedural guide explains what happens during HyperPod cluster creation and how the base lifecycle scripts are designed.

1. create_cluster.json – To submit a HyperPod cluster creation request, you prepare a CreateCluster request file in JSON format. In this best practices example, we assume that the request file is named create_cluster.json. Write create_cluster.json to provision a HyperPod cluster with instance groups. The best practice is to add the same number of instance groups as the number of Slurm nodes you plan to configure on the HyperPod cluster. Make sure that you give distinctive names to the instance groups that you'll assign to Slurm nodes you plan to set up.

   Also, you are required to specify an S3 bucket path to store your entire set of configuration files and lifecycle scripts to the field name InstanceGroups.LifeCycleConfig.SourceS3Uri in the CreateCluster request form, and specify the file name of an entrypoint shell script (assume that it's named on_create.sh) to InstanceGroups.LifeCycleConfig.OnCreate.

   **Note**
   If you are using the Create a cluster submission form in the HyperPod console UI, the console manages filling and submitting the CreateCluster request on your behalf, and runs the CreateCluster API in the backend. In this case, you don't need to create create_cluster.json; instead, make sure that you specify the correct cluster configuration information to the Create a cluster submission form.

2. on_create.sh – For each instance group, you need to provide an entrypoint shell script, on_create.sh, to run commands, run scripts to install software packages, and set up the HyperPod cluster environment with Slurm. The two things you need to prepare are a provisioning_params.json required by HyperPod for setting up Slurm and a set of lifecycle scripts for installing software packages. This script should be written to find and run the following files as shown in the sample script at on_create.sh.
Note

Make sure that you upload the entire set of lifecycle scripts to the S3 location you specify in create_cluster.json. You should also place your provisioning_params.json in the same location.

a. provisioning_params.json – This is a **section called “Configuration form for provisioning Slurm nodes on HyperPod”**. The on_create.sh script finds this JSON file and defines environment variable for identifying the path to it. Through this JSON file, you can configure Slurm nodes and storage options such as Amazon FSx for Lustre for Slurm to communicate with. In provisioning_params.json, make sure that you assign the HyperPod cluster instance groups using the names you specified in create_cluster.json to the Slurm nodes appropriately based on how you plan to set them up.

The following diagram shows an example of how the two JSON configuration files create_cluster.json and provisioning_params.json should be written to assign HyperPod instance groups to Slurm nodes. In this example, we assume a case of setting up three Slurm nodes: controller (management) node, log-in node (which is optional), and compute (worker) node.
Figure: Direct comparison between create_cluster.json for HyperPod cluster creation and provisioning_params.json for Slurm configuration. The number of instance groups in create_cluster.json should match with the number of nodes you want to configure as Slurm nodes. In case of the example in the figure, three Slurm nodes will be configured on a HyperPod cluster of three instance groups. You should assign the HyperPod cluster instance groups to Slurm nodes by specifying the instance group names accordingly.

b. resource_config.json – During cluster creation, the lifecycle_script.py script is written to expect a resource_config.json file from HyperPod. This file contains information about the cluster, such as instance types and IP addresses.

When you run the CreateCluster API, HyperPod creates a resource configuration file at /opt/ml/config/resource_config.json based on the create_cluster.json file. The file path is saved to the environment variable named SAGEMAKER RESOURCE CONFIG PATH.
Important

The resource_config.json file is auto-generated by the HyperPod platform, and you DO NOT need to create it. The following code is to show an example of resource_config.json that would be created from the cluster creation based on create_cluster.json in the previous step, and to help you understand what happens in the backend and how an auto-generated resource_config.json would look.

```json
{
    "ClusterConfig": {
        "ClusterName": "your-hyperpod-cluster"
    },
    "InstanceGroups": [
        {
            "Name": "controller-machine",
            "InstanceType": "ml.c5.xlarge",
            "Instances": [
                {
                    "InstanceName": "controller-machine-1",
                    "AgentIpAddress": "111.222.333.444",
                    "CustomerIpAddress": "111.222.333.444",
                    "InstanceId": "i-12345abcedfg67890"
                }
            ]
        },
        {
            "Name": "login-group",
            "InstanceType": "ml.m5.xlarge",
            "Instances": [
                {
                    "InstanceName": "login-group-1",
                    "AgentIpAddress": "111.222.333.444",
                    "CustomerIpAddress": "111.222.333.444",
                    "InstanceId": "i-12345abcedfg67890"
                }
            ]
        }
    ]
}
```
c. lifecycle_script.py – This is the main Python script that collectively runs lifecycle scripts setting up Slurm on the HyperPod cluster while being provisioned. This script reads in provisioning_params.json and resource_config.json from the paths that are specified or identified in on_create.sh, passes the relevant information to each lifecycle script, and then runs the lifecycle scripts in order.

Lifecycle scripts are a set of scripts that you have a complete flexibility to customize to install software packages and set up necessary or custom configurations during cluster creation, such as setting up Slurm, creating users, installing Conda or Docker.
The sample `lifecycle_script.py` script is prepared to run other base lifecycle scripts in the repository, such as launching Slurm daemons (`start_slurm.sh`), mounting Amazon FSx for Lustre (`mount_fsx.sh`), and setting up MariaDB accounting (`setup_mariadb_accounting.sh`) and RDS accounting (`setup_rds_accounting.sh`). You can also add more scripts, package them under the same directory, and add code lines to `lifecycle_script.py` to let HyperPod run the scripts. For more information about the base lifecycle scripts, see 3.1 Lifecycle scripts in the README.md file in the Awsome Distributed Training GitHub repository.

In addition to the default setups, more scripts for installing the following software are available under the `utils` folder. The `lifecycle_script.py` file is already prepared to include code lines for running the installation scripts, so see the following items to search those lines and uncomment to activate them.

i. To install Docker and Enroot, search and uncomment the following code lines.

```
ExecuteBashScript("./utils/install.docker.sh").run()
ExecuteBashScript("./utils/install_enroot.pyxis.sh").run(node_type)
```

ii. To install DCGM Exporter and EFA node exporter, search and uncomment the following code lines. This also requires having Docker installed.

```
ExecuteBashScript("./utils/install_dcmg_exporter.sh").run()
ExecuteBashScript("./utils/install_efa_node_exporter.sh").run()
```

iii. To install Slurm exporter and Prometheus on the controller node, search and uncomment the following code lines.

```
ExecuteBashScript("./utils/install_slurm_exporter.sh").run()
ExecuteBashScript("./utils/install_prometheus.sh").run()
```

To learn more about the base lifecycle scripts and dive deep, see 3.1 Lifecycle scripts in README.md file under the HyperPod folder in the Awsome Distributed Training GitHub repository.

3. Make sure that you upload all configuration files and setup scripts from Step 2 to the S3 bucket you provide in the CreateCluster request in Step 1. For example, assume that your `create_cluster.json` has the following.

```
"LifeCycleConfig": {
```
"SourceS3URI": "s3://sagemaker-hyperpod-lifecycle/src",
"OnCreate": "on_create.sh"
}

Then, your "s3://sagemaker-hyperpod-lifecycle/src" should contain on_create.sh, lifecycle_script.py, provisioning_params.json, and all other setup scripts. Assume that you have prepared the files in a local folder as follows.

```bash
### lifecycle_files // your local folder
### provisioning_params.json
### on_create.sh
### lifecycle_script.py
### ... // more setup scrips to be fed into lifecycle_script.py
```

To upload the files, use the S3 command as follows.

```bash
aws s3 cp --recursive ./lifecycle_scripts s3://sagemaker-hyperpod-lifecycle/src
```

### What particular configurations HyperPod manages in Slurm configuration files

When you create a Slurm cluster on HyperPod, the HyperPod agent sets up the `slurm.conf` and `gres.conf` files at `/opt/slurm/etc/` to manage the Slurm cluster based on your HyperPod cluster creation request and lifecycle scripts. The following list shows which specific parameters the HyperPod agent handles and overwrites.

⚠️ **Important**

We strongly recommend that you **do not** change these parameters managed by HyperPod.

- In `slurm.conf`, HyperPod sets up the following basic parameters: `ClusterName`, `SlurmctldHost`, `PartitionName`, and `NodeName`.

  Also, to enable the **the section called “Auto-resume”** functionality, HyperPod requires the `TaskPlugin` and `SchedulerParameters` parameters set as follows. The HyperPod agent sets up these two parameters with the required values by default.

```bash
TaskPlugin=task/none
```
SchedulerParameters=permit_job_expansion

- In `gres.conf`, HyperPod manages NodeName for GPU nodes.

**Mount Amazon FSx for Lustre to your HyperPod cluster**

To mount an Amazon FSx for Lustre shared file system to your HyperPod cluster, set up the following.

1. Use your Amazon VPC.
   a. For HyperPod cluster instances to communicate within your VPC, make sure that you attach the [section called "(Optional) Additional permissions for using SageMaker HyperPod with Amazon Virtual Private Cloud"](https://docs.aws.amazon.com/sagemaker/latest/dg/hyperpod-vpc.html) to the IAM role for SageMaker HyperPod.
   b. In `create_cluster.json`, include the following VPC information.

   ```json
   "VpcConfig": {
     "SecurityGroupIds": [ "string" ],
     "Subnets": [ "string" ]
   }
   ```

   For more tips about setting up Amazon VPC, see [the section called "(Optional) Set up SageMaker HyperPod with your Amazon VPC".](https://docs.aws.amazon.com/sagemaker/latest/dg/hyperpod-vpc.html)

2. To finish configuring Slurm with Amazon FSx for Lustre, specify the Amazon FSx DNS name and Amazon FSx mount name in `provisioning_params.json` as shown in the figure in the [section called "Start with base lifecycle scripts provided by HyperPod" section](https://docs.aws.amazon.com/sagemaker/latest/dg/hyperpod-lifecycle-scripts.html). You can find the Amazon FSx information either from the Amazon FSx for Lustre console in your account or by running the following AWS CLI command, `aws fsx describe-file-systems`.

   ```
   "fsx_dns_name": "fs-12345678a90b01cde.fsx.us-west-2.amazonaws.com",
   "fsx_mountname": "1abcdefg"
   ```

**Validate the JSON configuration files before running `create-cluster`**

Use the configuration validation script `validate-config.py`. This script parses and compares your HyperPod cluster configuration JSON file and Slurm configuration JSON file, and identifies if there's any resource misconfiguration between the two files and also across Amazon EC2, Amazon VPC, and Amazon FSx. For example, to validate the `create_cluster.json` and
provisioning_params.json files from the the section called “Start with base lifecycle scripts provided by HyperPod” section, run the validation script as follows.

```bash
python3 validate-config.py --cluster-config create_cluster.json --provisioning-parameters provisioning_params.json
```

The following is an example output of a successful validation.

```bash
##  Validated instance group name worker-group-1 is correct ...
##  Validated subnet subnet-012345abcdef67890 ...
##  Validated security group sg-012345abcdef67890 ingress rules ...
##  Validated security group sg-012345abcdef67890 egress rules ...
##  Validated FSx Lustre DNS name fs-012345abcdef67890.fsx.us-east-1.amazonaws.com
##  Validated FSx Lustre mount name dzfijbev
# Cluster Validation succeeded
```

### Develop lifecycle scripts interactively on a cluster node

This section explains how you can interactively develop lifecycle scripts without repeatedly creating and deleting a HyperPod cluster.

1. Create a HyperPod cluster with the base lifecycle scripts.
2. Log in to a cluster node.
3. Develop a script (`configure_xyz.sh`) by editing and running it repeatedly on the node.
   a. HyperPod runs the lifecycle scripts as the root user, so we recommend that you run the `configure_xyz.sh` as the root user while developing to make sure that the script is tested under the same condition while run by HyperPod.
4. Integrate the script into `lifecycle_script.py` by adding a code line similar to the following.

   ```python
   ExecuteBashScript("./utils/define_xyz.sh").run()
   ```

5. Upload the updated lifecycle scripts to the S3 bucket that you initially used for uploading the base lifecycle scripts.
6. Test the integrated version of `lifecycle_script.py` by creating a new HyperPod cluster.

### Update a cluster with new or updated lifecycle scripts

There are three ways to update the HyperPod software.
• The `UpdateClusterSoftware` API for patching the HyperPod software re-runs the lifecycle scripts on the entire instance group.
• The `UpdateCluster` API only runs the lifecycle scripts for new instance groups.
• You can also run lifecycle scripts directly in the HyperPod instances.

**Considerations**

Consider the following when using SageMaker HyperPod.

• HyperPod runs the section called “SageMaker HyperPod DLAMI” on each instance of a cluster, and the AMI has pre-installed software packages complying compatibilities between them and HyperPod functionalities. Note that if you reinstall any of the pre-installed packages, you are responsible for installing compatible packages and note that some HyperPod functionalities might not work as expected.

**Run jobs on SageMaker HyperPod clusters**

This section is for cluster users such as machine learning (ML) scientists. It provides examples of running ML workloads on provisioned SageMaker HyperPod clusters. Depending on how you have set up the environment on your HyperPod cluster, there are many ways to run ML workloads on HyperPod clusters. Examples of running ML workloads on HyperPod clusters are provided in the [Awesome Distributed Training GitHub repository](https://github.com/aws-deep-learning-ops/awesome-distributed-training). The following topics walk you through how to log in to the provisioned HyperPod clusters and get you started with running sample ML workloads.

**Topics**

• [Access the SageMaker HyperPod cluster nodes](#)
• [Schedule a Slurm job on a SageMaker HyperPod cluster](#)
• [Schedule jobs for distributed training workloads on SageMaker HyperPod](#)

**Access the SageMaker HyperPod cluster nodes**

You can access your InService cluster through AWS Systems Manager (SSM) by running the AWS CLI command `aws ssm start-session` with the SageMaker HyperPod cluster host name in format of `sagemaker-cluster:[cluster-id]_[instance-group-name]_[instance-id]`. You can retrieve the cluster ID and instance ID from the [SageMaker HyperPod console](https://console.aws.amazon.com/sagemaker/home#hyperpods) or by running the [AWS CLI commands for SageMaker HyperPod](https://docs.aws.amazon.com/cli/latest/reference/sagemaker.html). For example, if your cluster
ID is aa1bbbb222, the cluster node name is controller-group, and the cluster node ID is i-1112233344455aa, the SSM start-session command should be the following.

```
aws ssm start-session \
  --target sagemaker-cluster:aa11bbbbb222_controller-group-i-1112233344455aa \
  --region us-west-2
```

### Note

If you haven't set up AWS Systems Manager, follow the instructions provided at the section called “Set up AWS Systems Manager and Run As for cluster user access control”.

Schedule a Slurm job on a SageMaker HyperPod cluster

You can launch training jobs using the standard Slurm `sbatch` or `srun` commands. For example, to launch an 8-node training job, you can run `srun -N 8 --exclusive train.sh` SageMaker HyperPod supports training in a range of environments, including conda, venv, docker, and enviroot. You can configure an ML environment by running lifecycle scripts on your SageMaker HyperPod clusters. You also have an option to attach a shared file system such as FSx, which can also be used as a virtual environment.

The following example shows how to run a job for training Llama-2 with the Fully Sharded Data Parallelism (FSDP) technique on a SageMaker HyperPod cluster with a shared Amazon FSx file system. You can also find more examples from the [Awsome Distributed Training GitHub repository](https://github.com/aws-samples/awsome-distributed-training).

### Tip

All SageMaker HyperPod examples are available in the `3.test_cases` folder of the [Awsome Distributed Training GitHub repository](https://github.com/aws-samples/awsome-distributed-training).

1. Clone the [Awsome Distributed Training GitHub repository](https://github.com/aws-samples/awsome-distributed-training), and copy the training job examples to your shared Amazon FSx file system.

```
TRAINING_DIR=/fsx/users/my-user/fsdp
git clone https://github.com/aws-samples/awsome-distributed-training/
cp -R awsome-distributed-training/3.test_cases/10.FSDP $TRAINING_DIR
```
2. Create a conda environment on your shared Amazon FSx file system. Make sure that the file system is accessible to all nodes in the cluster.

```bash
#!/usr/bin/env bash
set -ex

wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh
chmod +x Miniconda3-latest-Linux-x86_64.sh
./Miniconda3-latest-Linux-x86_64.sh -b -f -p ./miniconda3
source ./miniconda3/bin/activate

conda create -y -n pt_fsdp python=3.10

source activate pt_fsdp

# Install PyTorch
pip install torch torchvision torchaudio
pip install packaging transformers accelerate ninja tensorboard h5py datasets

# create output dir
mkdir tensorboard
mkdir checkpoints
```

Save this script to a shared volume. This tutorial assumes that it is saved as `/fsx/users/my_user/create_env.sh`.

3. Build the virtual environment by launching a single node slurm job.

```bash
srun -N 1 /fsx/users/my_user/create_env.sh
```

4. After the environment is built, you can launch a training job by pointing to the environment path on the shared volume. You can launch both single-node and multi-node training jobs with the same setup. To launch a job, create a job launcher script (also called an entry point script) as follows.

```bash
#!/usr/bin/env bash
set -ex

ENV_PATH=/fsx/users/my_user/pytorch_env
TORCHRUN=$ENV_PATH/bin/torchrun
TRAINING_SCRIPT=/fsx/users/my_user/pt_train.py
```
WORLD_SIZE_JOB=${SLURM_NTASKS}
RANK_NODE=${SLURM_NODEID}
PROC_PER_NODE=8
MASTER_ADDR=`scontrol show hostnames \${SLURM_JOB_NODELIST} | head -n 1`
MASTER_PORT=$(expr 10000 + $(echo -n \${SLURM_JOBID} | tail -c 4))

DIST_ARGS="--nproc_per_node=${PROC_PER_NODE} \
    --nnodes=${WORLD_SIZE_JOB}  \
    --node_rank=${RANK_NODE}  \
    --master_addr=${MASTER_ADDR}  \
    --master_port=${MASTER_PORT}"

$TORCHRUN $DIST_ARGS $TRAINING_SCRIPT

Tip
If you want to make your training job more resilient against hardware failures by using the auto-resume capability of SageMaker HyperPod, you need to properly set up the environment variable MASTER_ADDR in the entrypoint script. To learn more, see the section called “Auto-resume”.

This tutorial assumes that this script is saved as /fsx/users/my_user/train.sh.

5. With this script in the shared volume at /fsx/users/my_user/train.sh, run the following `srun` command to schedule the Slurm job.

```
cd /fsx/users/my_user/
srun -N 8 train.sh
```

Schedule jobs for distributed training workloads on SageMaker HyperPod

The SageMaker distributed data parallelism (SMDDP) library is a collective communication library that improves compute performance of distributed data parallel training. The SMDDP library addresses communications overhead of the key collective communication operations by offering the following for SageMaker HyperPod.
1. The library offers AllGather optimized for AWS. AllGather is a key operation used in sharded data parallel training, which is a memory-efficient data parallelism technique offered by popular libraries such as the SageMaker model parallelism (SMP) library, DeepSpeed Zero Redundancy Optimizer (ZeRO), and PyTorch Fully Sharded Data Parallelism (FSDP).

2. The library performs optimized node-to-node communication by fully utilizing the AWS network infrastructure and the SageMaker ML instance topology.

Using SMDDP on a SageMaker HyperPod

The following are the training environment requirements for using the SMDDP library on SageMaker HyperPod.

- libstdc++ runtime version greater than 3.
- PyTorch 2.0.1 with cuda 11.8
- Python 3.10.x
- ml.p4d.24xlarge and ml.p4de.24xlarge, which are supported instance types by the SMDDP library
- imdsv2 enabled on training host

Depending on where you run the distributed training job, you can use the SMDDP library as follows.

To install the SMDDP library on SageMaker HyperPod DLAMI

- pip install --no-cache-dir https://smdataparallel.s3.amazonaws.com/binary/pytorch/2.0.1/cu118/2023-11-17/smdistributed_dataparallel-2.0.2-cp310-cp310-linux_x86_64.whl

Note

If working in a conda environment, ensure that PyTorch is installed via conda install rather than pip.

conda install pytorch==2.0.1 torchvision==0.15.2 torchaudio==2.0.2 pytorch-cuda=11.8 -c pytorch -c nvidia
To use the SMDDP library on a Docker container

- The SMDDP library is pre-installed on the SageMaker Deep Learning Containers (DLCs). To find the list of SageMaker framework DLCs for PyTorch with the SMDDP library, see Supported Frameworks in the SageMaker data parallelism library documentation. You can also bring your own Docker container with required dependencies installed to use the SMDDP library. To learn more about setting up a custom Docker container to use the SMDDP library, see also the section called “Create your own docker container with the library”.

⚠️ Important

To use the SMDDP library in a Docker container, you must bind mount the `/var/log` directory from the host machine onto `/var/log` in the container. This can be done by adding the following option when running your container.

```
docker run <OTHER_OPTIONS> -v /var/log:/var/log ...
```

Adapt your PyTorch training script to utilize the SMDDP library

Starting from the SageMaker distributed data parallelism (SMDDP) library v1.4.0, you can use the library as a backend option for the PyTorch distributed package. To use the SMDDP AllReduce and AllGather collective operations, you only need to import the SMDDP library at the beginning of your training script and set SMDDP as the the backend of PyTorch distributed modules during process group initialization. With the single line of backend specification, you can keep all the native PyTorch distributed modules and the entire training script unchanged. The following code snippets show how to use the SMDDP library as the backend of PyTorch-based distributed training packages: PyTorch distributed data parallel (DDP), PyTorch fully sharded data parallelism (FSDP), DeepSpeed, and Megatron-DeepSpeed.

For PyTorch DDP or FSDP

Initialize the process group as follows.

```python
import torch.distributed as dist
import smdistributed.dataparallel.torch.torch_smddp

dist.init_process_group(backend="smddp")
```
Note
(For PyTorch DDP jobs only) The `smddp` backend currently does not support creating subprocess groups with the `torch.distributed.new_group()` API. You also cannot use the `smddp` backend concurrently with other process group backends such as NCCL and Gloo.

For DeepSpeed or Megatron-DeepSpeed

Initialize the process group as follows.

```python
import deepspeed
import smdistributed.dataparallel.torch.torch_smddp
deepspeed.init_distributed(dist_backend="smddp")
```

Note
To use SMDDP AllGather with the `mpirun`-based launchers (smdistributed and pytorchddp) in the section called “Step 2: Launch a distributed training job”, you also need to set the following environment variable in your training script.

```bash
export SMDATAPARALLEL_OPTIMIZE_SDP=true
```

For general guidance on writing a PyTorch FSDP training script, see Advanced Model Training with Fully Sharded Data Parallel (FSDP) in the PyTorch documentation.

For general guidance on writing a PyTorch DDP training script, see Getting started with distributed data parallel in the PyTorch documentation.

SageMaker HyperPod cluster resiliency

SageMaker HyperPod provides the following cluster resiliency features.

Topics
- Cluster health check
- Auto-resume
Cluster health check

This section describes the set of health checks that SageMaker HyperPod uses to regularly monitor cluster instance health for issues with devices such as accelerators (GPU and Trainium cores) and networking (EFA).

<table>
<thead>
<tr>
<th>Category</th>
<th>Utility name</th>
<th>Instance type compatibility</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accelerator</td>
<td>DCGM policies</td>
<td>GPU</td>
<td>Each instance in the cluster continuously monitors all GPU-related policies including XID errors with NVIDIA DCGM.</td>
</tr>
<tr>
<td></td>
<td>NVIDIA SMI</td>
<td>GPU</td>
<td>nvidia-smi utility is a well-known CLI to manage and monitor GPUs. The built-in health checker parses the output from nvidia-smi to determine the health of the instance.</td>
</tr>
<tr>
<td></td>
<td>Neuron sysfs</td>
<td>Trainium</td>
<td>For Trainium-powered instances, the health of the Neuron devices is determined by reading counters from Neuron sysfs propagated directly by the Neuron driver.</td>
</tr>
<tr>
<td>Network</td>
<td>EFA</td>
<td>GPU and Trainium</td>
<td>To aid in the diagnostic of Elastic Fabric Adaptor (EFA) devices, the EFA health checker runs a series of connectivity tests using all available EFA cards within the instance.</td>
</tr>
<tr>
<td>---------</td>
<td>-----</td>
<td>------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Stress</td>
<td>DCGM Diagnostic</td>
<td>GPU</td>
<td><strong>DCGM diagnostics</strong> level 2 is used to exercise the GPUs in the system and put them under pressure to get a thorough insight of the health.</td>
</tr>
<tr>
<td>Stress</td>
<td>CPU stress</td>
<td>GPU and Trainium</td>
<td>CPU health is determined using the <strong>Linux stress</strong> tool, which runs multiple threads to achieve 100% CPU utilization and perform I/O operations.</td>
</tr>
</tbody>
</table>

**Auto-resume**

This section describes how to run a training job with the SageMaker HyperPod auto-resume functionality, which provides a zero-touch resiliency infrastructure to automatically recover a training job from the last saved checkpoint in the event of a hardware failure for clusters with more than 16 nodes.

With the auto-resume functionality, if a job fails due to a hardware failure or any transient issues in-between training, SageMaker HyperPod auto-resume starts the node replacement workflow and restarts the job after the faulty nodes are replaced.
Using the SageMaker HyperPod auto-resume functionality with Slurm

When you use SageMaker HyperPod auto-resume with Slurm, you should run the job inside an exclusive allocation acquired either by using `salloc` or `sbatch`. In any case, you need to modify the entrypoint script to make sure that all setup steps run in a single `srun` command when resuming the job. Through the entrypoint script, it is important to set up the environment on the replaced node to be consistent with the environment that the job step was running before it was stopped. The following procedure shows how to prepare an entrypoint script to keep the environment consistent and run it as a single `srun` command.

Tip

If you use `sbatch`, you can keep the batch script simple by creating a separate script for setting up the environment and using a single `srun` command.

1. Create a script using the following code example and save it as `train_auto_resume.sh`. This script deploys training environment setups assuming that there is no manual configuration previously made to the replaced node. This ensures that the environment is node-agnostic, so that when a node is replaced, the same environment is provisioned on the node before resuming the job.

Note

The following code example shows how to discover the Slurm node list associated with the job. Do not use the `$SLURM_JOB_NODELIST` environment variable provided by Slurm, because its value might be outdated after SageMaker HyperPod auto-resumes the job. The following code example shows how to define a new `NODE_LIST` variable to replace `SLURM_JOB_NODELIST`, and then set up the `MASTER_NODE` and `MASTER_ADDR` variables off of the `NODE_LIST` variable.

```
#!/bin/bash

# Filename: train_auto_resume.sh
# Sample containerized script to launch a training job with a single srun which can be auto-resumed.
```

```
# Place your training environment setup here.
# Example: Install conda, docker, activate virtual env, etc.

# Get the list of nodes for a given job
NODE_LIST=$(scontrol show jobid=$SLURM_JOBID | \
    # Show details of the SLURM job
    awk -F= '/NodeList=/{print $2}' | \
    # Extract NodeList field
    grep -v Exc)                         
# Exclude nodes marked as excluded

# Determine the master node from the node list
MASTER_NODE=$(scontrol show hostname $NODE_LIST | \
    # Convert node list to hostnames
    head -n 1)                            
# Select the first hostname as master node

# Get the master node address
MASTER_ADDR=$(scontrol show node=$MASTER_NODE | \
    # Show node information
    awk -F= '/NodeAddr=/{print $2}' | \ 
    # Extract NodeAddr
    awk '{print $1}')                   
# Print the first part of NodeAddr

# Torchrun command to launch the training job
torchrun_cmd="torchrun --nnodes=$SLURM_NNODES \ 
    --nproc_per_node=1 \ 
    --node_rank=$SLURM_NODE \ 
    --master-addr=$MASTER_ADDR \ 
    --master_port=1234 \ 
    <your_training_script.py>"

# Execute the torchrun command in the 'pytorch' Conda environment,
# streaming output live
/opt/conda/bin/conda run --live-stream -n pytorch $torchrun_cmd

💡 Tip
You can use the preceding script to add more commands for installing any additional dependencies for your job. However, we recommend that you keep the dependency installation scripts to the set of lifecycle scripts that are used during cluster creation. If you use a virtual environment hosted on a shared directory, you can also utilize this script to activate the virtual environment.
2. Launch the job with SageMaker HyperPod auto-resume enabled by adding the flag `--auto-resume=1` to indicate that the `srun` command should be automatically retried in case of hardware failure.

Note

If you have set up a resource allocation using `sbatch` or `salloc`, you can run multiple `srun` commands within the allocation. In the event of a failure, the SageMaker HyperPod auto-resume functionality only operates in the current job step of the `srun` command with the flag `--auto-resume=1`. In other words, activating auto-resume in an `srun` command doesn't apply to other `srun` commands launched within a resource allocation session.

The following are `srun` command examples with `auto-resume` enabled.

Using `sbatch`

Because most of the logic for setting up the environment is already in `train_auto_resume.sh`, the batch script should be simple and similar to the following code example. Assume that the following batch script is saved as `batch.sh`.

```bash
#!/bin/bash
#SBATCH --nodes 2
#SBATCH --exclusive
srun --auto-resume=1 train_auto_resume.sh
```

Run the preceding batch script using the following command.

```
sbatch batch.sh
```

Using `salloc`

Start by acquiring an exclusive allocation, and run the `srun` command with the `--auto-resume` flag and the entrypoint script.

```
salloc -N 2 --exclusive
srun --auto-resume=1 train_auto_resume.sh
```
How to replace a faulty node not being auto-resumed by HyperPod

The HyperPod auto-resume functionality monitors if the state of your Slurm nodes turns to fail or down. You can check the state of Slurm nodes by running `sinfo`.

If you have a node stuck with an issue but not being fixed by the HyperPod auto-resume functionality, we recommend you to run the following command to change the state of the node to fail.

```
scontrol update node=<ip-ipv4> state=fail reason="Action:Replace"
```

In the preceding command example, replace `<ip-ipv4>` with the Slurm node name (host name) of the faulty instance you want to replace.

After running this command, the node should go into the fail state, waits for the currently running jobs to finish, is replaced with a healthy instance, and is recovered with the same host name. This process takes time depending on the available instances in your Availability Zone and the time it takes to run your lifecycle scripts. During the update and replacement processes, avoid changing the state of the node manually again or restarting the Slurm controller; doing so can lead to a replacement failure. If the node does not get recovered nor turn to the idle state after a long time, contact AWS Support.

If the faulty node is continuously stuck in the fail state, the last resort you might try is to manually force change the node state to down. This requires administrator privileges (sudo permissions).

⚠️ **Warning**

Proceed carefully before you run the following command as it forces kill all jobs, and you might lose all unsaved work.

```
scontrol update node=<ip-ipv4> state=down reason="Action:Replace"
```

SageMaker HyperPod cluster management

The following topics discuss logging and managing SageMaker HyperPod clusters.
Logging SageMaker HyperPod events

All events and logs from SageMaker HyperPod are saved to Amazon CloudWatch under the log group name /aws/sagemaker/Clusters/[ClusterName]/[ClusterID]. Every call to the CreateCluster API creates a new log group. The following list contains all of the available log streams collected in each log group.

<table>
<thead>
<tr>
<th>Log Group Name</th>
<th>Log Stream Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>/aws/sagemaker/Clusters/[ClusterName]/[ClusterID]</td>
<td>LifecycleConfig/[instance-group-name]/[instance-id]</td>
</tr>
</tbody>
</table>

Logging SageMaker HyperPod at instance level

You can access the LifecycleScript logs published to CloudWatch during cluster instance configuration. Every instance within the created cluster generates a separate log stream, distinguishable by the LifecycleConfig/[instance-group-name]/[instance-id] format.

All logs that are written to /var/log/provision/provisioning.log are uploaded to the preceding CloudWatch stream. Sample LifecycleScripts at 1.architectures/5.sagemaker_hyperpods/LifecycleScripts/base-config redirect their stdout and stderr to this location. If you are using your custom scripts, write your logs to the /var/log/provision/provisioning.log location for them to be available in CloudWatch.

Tagging resources

AWS Tagging system helps manage, identify, organize, search for, and filter resources. SageMaker HyperPod supports tagging, so you can manage the clusters as an AWS resource. During cluster creation or editing an existing cluster, you can add or edit tags for the cluster. To learn more about tagging in general, see Tagging your AWS resources.

Using the SageMaker HyperPod console UI

When you are creating a new cluster and editing a cluster, you can add, remove, or edit tags.

Using the SageMaker HyperPod APIs

When you write a CreateCluster or UpdateCluster API request file in JSON format, edit the Tags section.
Using the AWS CLI tagging commands for SageMaker

To tag a cluster

Use `aws sagemaker add-tags` as follows.

```
aws sagemaker add-tags --resource-arn cluster_ARN --tags Key=string,Value=string
```

To untag a cluster

Use `aws sagemaker delete-tags` as follows.

```
aws sagemaker delete-tags --resource-arn cluster_ARN --tag-keys "tag_key"
```

To list tags for a resource

Use `aws sagemaker list-tags` as follows.

```
aws sagemaker list-tags --resource-arn cluster_ARN
```

SageMaker HyperPod references

Find more information and references about using SageMaker HyperPod in the following topics.

Topics

- [SageMaker HyperPod pricing](#)
- [SageMaker HyperPod APIs](#)
- [SageMaker HyperPod forms](#)
- [SageMaker HyperPod DLAMI](#)
- [SageMaker HyperPod API permissions reference](#)
- [SageMaker HyperPod commands in AWS CLI](#)
- [SageMaker HyperPod Python modules in AWS SDK for Python (Boto3)](#)

SageMaker HyperPod pricing

The following topics provide information about SageMaker HyperPod pricing. To find more details on price per hour for using SageMaker HyperPod instances, see also [Amazon SageMaker Pricing](#).

Capacity requests
You can allocate on-demand or reserved compute capacity with SageMaker for use on SageMaker HyperPod. On-demand cluster creation allocates available capacity from the SageMaker on-demand capacity pool. Alternatively, you can request reserved capacity to ensure access by submitting a ticket for a quota increase. Inbound capacity requests are prioritized by SageMaker and you receive an estimated time for capacity allocation.

**Service billing**

When you provision a compute capacity on SageMaker HyperPod, you are billed for the duration of the capacity allocation. SageMaker HyperPod billing appears in your anniversary bills with a line item for the type of capacity allocation (on-demand, reserved), the instance type, and the time spent on using the instance.

To submit a ticket for a quota increase, see the section called “SageMaker HyperPod quotas”.

**SageMaker HyperPod APIs**

The following list is a full set of SageMaker HyperPod APIs for submitting action requests in JSON format to SageMaker through AWS CLI or AWS SDK for Python (Boto3).

- [CreateCluster](#)
- [DeleteCluster](#)
- [DescribeCluster](#)
- [DescribeClusterNode](#)
- [ListClusterNodes](#)
- [ListClusters](#)
- [UpdateCluster](#)
- [UpdateClusterSoftware](#)

**SageMaker HyperPod forms**

To configure the Slurm workload manager tool on HyperPod, you should create a Slurm configuration file required by HyperPod using the provided form.

**Configuration form for provisioning Slurm nodes on HyperPod**

The following code is the Slurm configuration form you should prepare to properly set up Slurm nodes on your HyperPod cluster. You should complete this form and upload it as part of a set of
lifecycle scripts during cluster creation. To learn how this form should be prepared throughout HyperPod cluster creation processes, see the section called “SageMaker HyperPod lifecycle configuration best practices”.

// Save as provisioning_params.json.
{
    "version": "1.0.0",
    "workload_manager": "slurm",
    "controller_group": "string",
    "login_group": "string",
    "worker_groups": [
        
        "instance_group_name": "string",
        "partition_name": "string"
    ],
    "fsx_dns_name": "string",
    "fsx_mountname": "string"
}

- **version** – Required. This is the version of the HyperPod provisioning parameter form. Keep it to 1.0.0.
- **workload_manager** – Required. This is for specifying which workload manager to be configured on the HyperPod cluster. Keep it to slurm.
- **controller_group** – Required. This is for specifying the name of the HyperPod cluster instance group you want to assign to Slurm controller (head) node.
- **login_group** – Optional. This is for specifying the name of the HyperPod cluster instance group you want to assign to Slurm login node.
- **worker_groups** – Required. This is for setting up Slurm worker (compute) nodes on the HyperPod cluster.
  - **instance_group_name** – Required. This is for specifying the name of the HyperPod instance group you want to assign to Slurm worker (compute) node.
  - **partition_name** – Required. This is for specifying the partition name to the node.
- **fsx_dns_name** – Optional. If you want to set up your Slurm nodes on the HyperPod cluster to communicate with Amazon FSx, specify the FSx DNS name.
- **fsx_mountname** – Optional. If you want to set up your Slurm nodes on the HyperPod cluster to communicate with Amazon FSx, specify the FSx mount name.
SageMaker HyperPod DLAMI

The SageMaker HyperPod agent runs a SageMaker HyperPod DLAMI, which is built on top of AWS Deep Learning Base GPU AMI (Ubuntu 20.04).

The SageMaker HyperPod DLAMI is bundled with additional packages to support open source tools such as Slurm and dependencies, and SageMaker HyperPod cluster software packages to support features such as cluster health check and auto-resume. To follow up with HyperPod software updates that the HyperPod service team distributes through the DLAMI, see the section called “HyperPod release notes”.

SageMaker HyperPod API permissions reference

When you are setting up access control for allowing to run SageMaker HyperPod API operations and writing a permissions policy that you can attach to IAM users for cloud administrators, use the following table as a reference.

<table>
<thead>
<tr>
<th>Amazon SageMaker API Operations</th>
<th>Required Permissions (API Actions)</th>
<th>Resources</th>
</tr>
</thead>
</table>
## ListClusterNodes

<table>
<thead>
<tr>
<th>Action</th>
<th>Service</th>
<th>ARN</th>
</tr>
</thead>
<tbody>
<tr>
<td>ListClusterNodes</td>
<td>sagemaker::ListClusterNodes</td>
<td>arn:aws:sagemaker:region:account-id :cluster/ cluster-id</td>
</tr>
</tbody>
</table>

## ListClusters

<table>
<thead>
<tr>
<th>Action</th>
<th>Service</th>
<th>ARN</th>
</tr>
</thead>
<tbody>
<tr>
<td>ListClusters</td>
<td>sagemaker::ListClusters</td>
<td>arn:aws:sagemaker:region:account-id :cluster/ cluster-id</td>
</tr>
</tbody>
</table>

## UpdateCluster

<table>
<thead>
<tr>
<th>Action</th>
<th>Service</th>
<th>ARN</th>
</tr>
</thead>
<tbody>
<tr>
<td>UpdateCluster</td>
<td>sagemaker::UpdateCluster</td>
<td>arn:aws:sagemaker:region:account-id :cluster/ cluster-id</td>
</tr>
</tbody>
</table>

## UpdateClusterSoftware

<table>
<thead>
<tr>
<th>Action</th>
<th>Service</th>
<th>ARN</th>
</tr>
</thead>
<tbody>
<tr>
<td>UpdateClusterSoftware</td>
<td>sagemaker::UpdateClusterSoftware</td>
<td>arn:aws:sagemaker:region:account-id :cluster/ cluster-id</td>
</tr>
</tbody>
</table>

For a complete list of permissions and resource types for SageMaker APIs, see [Actions, resources, and condition keys for Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/aws-service-authorization-reference.html) in the AWS Service Authorization Reference.

### SageMaker HyperPod commands in AWS CLI

The following are the AWS CLI commands for SageMaker HyperPod to run the core HyperPod API operations.

- `create-cluster`
- `delete-cluster`
- `describe-cluster`
- `describe-cluster-node`
- `list-cluster-nodes`
- `list-clusters`
- `list-cluster-node`
- `update-cluster`
- `update-cluster-software`
SageMaker HyperPod Python modules in AWS SDK for Python (Boto3)

The following are the methods of the AWS SDK for Python (Boto3) client for SageMaker to run the core HyperPod API operations.

- create_cluster
- delete_cluster
- describe_cluster
- describe_cluster_node
- list_cluster_nodes
- list_clusters
- update_cluster
- update_cluster_software

SageMaker HyperPod FAQ

Use the following frequently asked questions to troubleshoot problems with using SageMaker HyperPod.

Q. Why can I not find log groups of my SageMaker HyperPod cluster in Amazon CloudWatch?

By default, agent logs and instance start-up logs are sent to the HyperPod platform account's CloudWatch. In case of user lifecycle scripts, lifecycle configuration logs are sent to your account’s CloudWatch.

If you use the sample lifecycle scripts provided by the HyperPod service team, you can expect to find the lifecycle configuration logs written to /var/log/provision/provisioning.log, and you wouldn’t encounter this problem.

However, if you use custom paths for collecting logs from lifecycle provisioning and can’t find the log groups appearing in your account's CloudWatch, it might be due to mismatches in the log file paths specified in your lifecycle scripts and what the CloudWatch agent running on the HyperPod cluster instances looks for. In this case, it means that you need to properly set up your lifecycle scripts to send logs to the CloudWatch agent, and also set up the CloudWatch agent configuration accordingly. To resolve the problem, choose one of the following options.

- **Option 1**: Update your lifecycle scripts to write logs to /var/log/provision/provisioning.log.
• **Option 2:** Update the CloudWatch agent to look for your custom paths for logging lifecycle provisioning.

1. Each HyperPod cluster instance contains a CloudWatch agent configuration file in JSON format at /opt/aws/amazon-cloudwatch-agent/sagemaker_cwagent_config.json. In the configuration file, find the field name logs.logs_collected.files.collect_list.file_path. With the default setup by HyperPod, the key-value pair should be "file_path": "/var/log/provision/provisioning.log" as documented at the section called “Logging SageMaker HyperPod at instance level”. The following code snippet shows how the JSON file looks with the HyperPod default configuration.

   ```json
   "logs": {
     "logs_collected": {
       "files": {
         "collect_list": [
           {
             "file_path": "/var/log/provision/provisioning.log",
             "log_group_name": "/aws/sagemaker/Clusters/[ClusterName]/[ClusterID]",
             "log_stream_name": "LifecycleConfig/[InstanceGroupName]/{instance_id}"
           },
           "retention_in_days": -1
         ]
       },
       "force_flush_interval": 3
     }
   }
   ```

2. Replace the value for the "file_path" field name with the custom path you use in your lifecycle scripts. For example, if you have set up your lifecycle scripts to write to /var/log/custom-provision/custom-provisioning.log, update the value to match with it as follows.

   ```json
   "file_path": "/var/log/custom-provision/custom-provisioning.log"
   ```

3. Restart the CloudWatch agent with the configuration file to finish applying the custom path. For example, the following CloudWatch command shows how to restart the CloudWatch agent with the CloudWatch agent configuration file from step 1. For more information, see also Troubleshooting the CloudWatch agent.
Q. What particular configurations does HyperPod manage in Slurm configuration files such as `slurm.conf` and `gres.conf`?

When you create a Slurm cluster on HyperPod, the HyperPod agent sets up the `slurm.conf` and `gres.conf` files at `/opt/slurm/etc/` to manage the Slurm cluster based on your HyperPod cluster creation request and lifecycle scripts. The following list shows what specific parameters the HyperPod agent handles and overwrites.

⚠️ **Important**

We strongly recommend that you DON'T change these parameters managed by HyperPod.

- In `slurm.conf`, HyperPod sets up the following basic parameters: `ClusterName`, `SlurmctldHost`, `PartitionName`, and `NodeName`.

  Also, to enable the [section called “Auto-resume”](https://www.example.com) functionality, HyperPod requires the `TaskPlugin` and `SchedulerParameters` parameters set as follows. The HyperPod agent sets up these two parameters with the required values by default.

```bash
TaskPlugin=task/none
SchedulerParameters=permit_job_expansion
```

- In `gres.conf`, HyperPod manages `NodeName` for GPU nodes.

Amazon SageMaker HyperPod release notes

See the following release notes to track the latest updates for Amazon SageMaker HyperPod.

SageMaker HyperPod release notes: March 14, 2024

HyperPod software patch
The HyperPod service team distributes software patches through the section called “SageMaker HyperPod DLAMI”. See the following details about the latest HyperPod DLAMI.

- Upgraded Slurm to v23.11.1
- Added OpenPMIx v4.2.6 for enabling Slurm with PMIx.
- Built upon the AWS Deep Learning Base GPU AMI (Ubuntu 20.04) released on 2023-10-26
- A complete list of pre-installed packages in this HyperPod DLAMI in addition to the base AMI
  - Slurm: v23.11.1
  - OpenPMIx: v4.2.6
  - Munge: v0.5.15
  - aws-neuronx-dkms: v2.*
  - aws-neuronx-collectives: v2.*
  - aws-neuronx-runtime-lib: v2.*
  - aws-neuronx-tools: v2.*
  - SageMaker HyperPod software packages to support features such as cluster health check and auto-resume

Upgrade steps

- Run the following command to call the UpdateClusterSoftware API to update your existing HyperPod clusters with the latest HyperPod DLAMI. To find more instructions, see the section called “Update the SageMaker HyperPod platform software of a cluster”.

⚠ Important

Back up your work before running this API. The patching process replaces the root volume with the updated AMI, which means that your previous data stored in the instance root volume will be lost. Make sure that you back up your data from the instance root volume to Amazon S3 or Amazon FSx for Lustre. For more information, see the section called “Use the backup script provided by SageMaker HyperPod”.

aws sagemaker update-cluster-software --cluster-name your-cluster-name
Note

Note that you should run the AWS CLI command to update your HyperPod cluster. Updating the HyperPod software through SageMaker HyperPod console UI is currently not available.

Improvements

- HyperPod now properly supports passing partition names provided through `provisioning_params.json` and creates partitions appropriately based on provided inputs. For more information about `provisioning_params.json`, see the section called “SageMaker HyperPod forms” and the section called “SageMaker HyperPod lifecycle configuration best practices”.

SageMaker HyperPod release notes: February 15, 2024

New features

- Added a new `update-cluster-software` API for SageMaker HyperPod security patching. When security patches become available, we recommend you to update existing SageMaker HyperPod clusters in your account by running `aws sagemaker update-cluster-software --cluster-name your-cluster-name`. To follow up with future security patches, keep tracking this Amazon SageMaker HyperPod release notes page. To learn how the `update-cluster-software` API works, see the section called “Update the SageMaker HyperPod platform software of a cluster”.

SageMaker HyperPod release notes: November 29, 2023

New features


HyperPod software patch

The HyperPod service team distributes software patches through the section called “SageMaker HyperPod DLAMI”. See the following details about the latest HyperPod DLAMI.
Use generative AI in SageMaker notebook environments

Jupyter AI is an open-source extension of JupyterLab integrating generative AI capabilities into Jupyter notebooks. Through the Jupyter AI chat interface and magic commands, users experiment with code generated from natural language instructions, explain existing code, ask questions about their local files, generate entire notebooks, and more. The extension connects Jupyter notebooks with large language models (LLMs) that users can use to generate text, code, or images, and to ask questions about their own data. Jupyter AI supports generative model providers such as AI21, Anthropic, AWS (SageMaker JumpStart and Amazon Bedrock), Cohere, and OpenAI.

The extension's package is included in Amazon SageMaker Distribution version 1.2 and onwards. Amazon SageMaker Distribution is a Docker environment for data science and scientific computing used as the default image of JupyterLab notebook instances. Users of different IPython environments can install Jupyter AI manually.

In this section, we provide an overview of Jupyter AI capabilities and demonstrate how to configure models provided by SageMaker JumpStart or Amazon Bedrock from JupyterLab or Studio Classic notebooks. For more in-depth information on the Jupyter AI project, refer to its documentation. Alternatively, you can refer to the blog post Generative AI in Jupyter for an overview and examples of key Jupyter AI capabilities.

Before using Jupyter AI and interacting with your LLMs, make sure that you satisfy the following prerequisites:
After completing the prerequisite steps, you can proceed to **Use Jupyter AI in JupyterLab or Studio Classic**.

**Topics**

- **Install Jupyter AI**
- **Jupyter AI Features**
- **Configure your model provider**
- **Use Jupyter AI in JupyterLab or Studio Classic**

**Install Jupyter AI**

For **Amazon SageMaker Distribution** users, we recommend selecting the SageMaker Distribution image version 1.2 or later. No further installation is necessary. Users of JupyterLab in Studio can choose the version of their Amazon SageMaker Distribution when creating a space.

For users of other IPython environments, the version of the recommended Jupyter AI package depends on the version of JupyterLab they are using.

The Jupyter AI distribution consists of two packages.

- **jupyter_ai**: This package provides a JupyterLab extension and a native chat user interface (UI). It acts as a conversational assistant using the large language model of your choice.
• jupyter_ai_magics: This package provides the IPython %%ai and %ai magic commands with which you can invoke a large language model (LLM) from your notebook cells.

**Note**

Installing jupyter_ai also installs jupyter_ai_magics. However, you can install jupyter_ai_magics independently without JupyterLab or jupyter_ai. The magic commands %%ai and %ai work in any IPython kernel environment. If you only install jupyter_ai_magics, you can't use the chat UI.

For users of JupyterLab 3, in particular Studio Classic users, we recommend installing jupyter_ai version 1.5.x or any later 1.x version. However, we highly recommend using Jupyter AI with JupyterLab 4. The jupyter_ai version compatible with JupyterLab 3 may not allow users to set additional model parameters such as temperature, top-k and top-p sampling, max tokens or max length, or user acceptance license agreements.

For users of JupyterLab 4 environments that do not use SageMaker Distribution, we recommend installing jupyter_ai version 2.5.x or any later 2.x version.

See the installation instructions in the Installation section of Jupyter AI documentation.

**Jupyter AI Features**

You can access Jupyter AI capabilities through two distinct methods: using the chat UI or using magic commands within notebooks.

**From the chat user interface AI assistant**

The chat interface connects you with Jupyternaut, a conversational agent that uses the language model of your choice.

After launching a JupyterLab application installed with Jupyter AI, you can access the chat interface by choosing the chat icon in the left navigation panel. First-time users are prompted to configure their model. See Configure your model provider in the chat UI for configuration instructions.
Using the chat UI, you can:

- **Answer questions**: For instance, you can ask Jupyternaut to create a Python function that adds CSV files to an Amazon S3 bucket. Subsequently, you can refine your answer with a follow-up question, such as adding a parameter to the function to choose the path where the files are written.

- **Interact with files in JupyterLab**: You can include a portion of your notebook in your prompt by selecting it. Then, you can either replace it with the model's suggested answer or manually copy the answer to your clipboard.

- **Generate entire notebooks** from prompts: By starting your prompt with `/generate`, you trigger a notebook generation process in the background without interrupting your use of Jupyternaut. A message containing the link to the new file is displayed upon completion of the process.

- **Learn from and ask questions about local files**: Using the `/learn` command, you can teach an embedding model of your choice about local files and then ask questions about those files using the `/ask` command. Jupyter AI stores the embedded content in a local **FAISS vector database**, then uses retrieval-augmented generation (RAG) to provide answers based on what it has learned. To erase all previously learned information from your embedding model, use `/learn -d`.

For a complete list of features and detailed instructions on their usage, see the [Jupyter AI chat interface](#) documentation. To learn about how to configure access to a model in Jupyternaut, see [Configure your model provider in the chat UI](#).

### From notebook cells

Using `%ai` and `%ai` magic commands, you can interact with the language model of your choice from your notebook cells or any IPython command line interface. The `%ai` command applies your instructions to the entire cell, whereas `%ai` apply them to the specific line.

The following example illustrates an `%ai` magic command invoking an Anthropic Claude model to output an HTML file containing the image of a white square with black borders.

```markdown
%%ai anthropic:claude-v1.2 -f html
Create a square using SVG with a black border and white fill.
```

To learn about the syntax of each command, use `%ai help`. To list the providers and models supported by the extension, run `%ai list`. 
For a complete list of features and detailed instructions on their usage, see the Jupyter AI magic commands documentation. In particular, you can customize the output format of your model using the `-f` or `--format` parameter, allow variable interpolation in prompts, including special `In` and `Out` variables, and more.

To learn about how to configure the access to a model, see Configure your model provider in a notebook.

**Configure your model provider**

---

**Note**

In this section, we assume that the language and embedding models that you plan to use are already deployed. For models provided by AWS, you should already have the ARN of your SageMaker endpoint or access to Amazon Bedrock. For other model providers, you should have the API key used to authenticate and authorize requests to your model.

Jupyter AI supports a wide range of model providers and language models, refer to the list of its supported models to stay updated on the latest available models. For information on how to deploy a model provided by SageMaker JumpStart, see Deploy a Model in the SageMaker JumpStart documentation. You need to request access to Amazon Bedrock to use it as your model provider.

---

The configuration of Jupyter AI varies depending on whether you are using the chat UI or magic commands.

**Configure your model provider in the chat UI**

---

**Note**

You can configure several LLMs and embedding models following the same instructions. However, you must configure at least one Language model.
To configure your chat UI

1. In JupyterLab, access the chat interface by choosing the chat icon
   ![Chat Icon](image)
in the left navigation panel.

2. Choose the configuration icon
   ![Configuration Icon](image)
in the top right corner of the left pane. This opens the Jupyter AI configuration panel.

3. Fill out the fields related to your service provider.

   - **For models provided by SageMaker JumpStart or Amazon Bedrock**
     - In the language model dropdown list, select `sagemaker-endpoint` for models deployed with SageMaker JumpStart or `bedrock` for models managed by Amazon Bedrock.
     - The parameters differ based on whether your model is deployed on SageMaker or Amazon Bedrock.
     - For models deployed with SageMaker JumpStart:
       - Enter the name of your endpoint in **Endpoint name**, and then the AWS Region in which your model is deployed in **Region name**. To retrieve the ARN of the SageMaker endpoints, navigate to [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/) and then choose Inference and Endpoints in the left menu.
       - Paste the JSON of the **Request schema** tailored to your model, and the corresponding **Response path** for parsing the model's output.

     - (Optional) Select an **embedding model** to which you have access. Embedding models are used to capture additional information from local documents, enabling the text generation model to respond to questions within the context of those documents.
• Choose **Save Changes** and navigate to the left arrow icon

  in the top left corner of the left pane. This opens the Jupyter AI chat UI. You can start
  interacting with your model.

• **For models hosted by third-party providers**
  
  • In the **language model** dropdown list, select your provider ID. You can find the details of
    each provider, including their ID, in Jupyter AI list of model providers.
  
  • (Optional) Select an **embedding model** to which you have access. Embedding models
    are used to capture additional information from local documents, enabling the text
    generation model to respond to questions within the context of those documents.
  
  • Insert your models' API keys.

• Choose **Save Changes** and navigate to the left arrow icon

  in the top left corner of the left pane. This opens the Jupyter AI chat UI. You can start
  interacting with your model.

The following snapshot is an illustration of the chat UI configuration panel set to invoke a Flan-t5-
small model provided by SageMaker JumpStart and deployed in SageMaker.
Language model

Specify an endpoint name as the model ID. In addition, you must specify a region name, request schema, and response path. For more information, see the documentation about SageMaker endpoints deployment and about using magic commands with SageMaker endpoints.

- Region name (required)
  - us-west-2

- Request schema (required)
  - `{"inputs": "<prompt>"}`

- Response path (required)
  - `[0]."generated_text"`

Embedding model

- Embedding model
  - None

API Keys

Input

When writing a message, press Enter to:

- Send the message
- Start a new line (use Shift+Enter to send)

Save Changes
Pass extra model parameters and custom parameters to your request

Your model may need extra parameters, like a customized attribute for user agreement approval or adjustments to other model parameters such as temperature or response length. We recommend configuring these settings as a start up option of your JupyterLab application using a Lifecycle Configuration. For information on how to create a Lifecycle Configuration and attach it to your domain, or to a user profile from the SageMaker console, see Create and associate a lifecycle configuration. You can choose your LCC script when creating a space for your JupyterLab application.

Use the following JSON schema to configure your extra parameters:

```json
{
   "AiExtension": {
      "model_parameters": {
         "<provider_id>:<model_id>": { Dictionary of model parameters which is unpacked and passed as-is to the provider.}
      }
   }
}
```

The following script is an example of a JSON configuration file that you can use when creating a JupyterLab application LCC to set the maximum length of an Al21 Labs Jurassic-2 model deployed on Amazon Bedrock. Increasing the length of the model's generated response can prevent the systematic truncation of your model's response.

```bash
#!/bin/bash
set -eux

mkdir -p /home/sagemaker-user/.jupyter

json='{"AiExtension": {"model_parameters": {"bedrock:ai21.j2-mid-v1": {"model_kwargs": {"maxTokens": 200}}}}}'

# equivalent to %ai bedrock:ai21.j2-mid-v1 -m {"model_kwargs":{"maxTokens":200}}

# File path
file_path="/home/sagemaker-user/.jupyter/jupyter_jupyter_ai_config.json"

#jupyter --paths
```
The following script is an example of a JSON configuration file for creating a JupyterLab application LCC used to set additional model parameters for an Anthropic Claude model deployed on Amazon Bedrock.

```bash
#!/bin/bash
set -eux

mkdir -p /home/sagemaker-user/.jupyter

json='{"AiExtension": {"model_parameters": {"bedrock:anthropic.claude-v2":
{"model_kwargs":{"temperature":0.1,"top_p":0.5,"top_k":250,"max_tokens_to_sample":2000}}}}}

# equivalent to %%ai bedrock:anthropic.claude-v2 -m {"model_kwargs":
{"temperature":0.1,"top_p":0.5,"top_k":250,"max_tokens_to_sample":2000}}

# File path
file_path="/home/sagemaker-user/.jupyter/jupyter_jupyter_ai_config.json"

#jupyter --paths

# Write JSON to file
echo "json" > "$file_path"

# Confirmation message
echo "JSON written to $file_path"

restart-jupyter-server

# Waiting for 30 seconds to make sure the Jupyter Server is up and running
sleep 30
```
Once you have attached your LCC to your domain, or user profile, add your LCC to your space when launching your JupyterLab application. To ensure that your configuration file is updated by the LCC, run `more ~/.jupyter/jupyter_jupyter_ai_config.json` in a terminal. The content of the file should correspond to the content of the JSON file passed to the LCC.

**Configure your model provider in a notebook**

**To invoke a model via Jupyter AI within JupyterLab or Studio Classic notebooks using the %%ai and %ai magic commands**

1. Install the client libraries specific to your model provider in your notebook environment. For example, when using OpenAI models, you need to install the `openai` client library. You can find the list of the client libraries required per provider in the *Python package(s)* column of the Jupyter AI **Model providers list**.

2. • **For models hosted by AWS**

   Ensure that your execution role has the permission to invoke your SageMaker endpoint for models provided by SageMaker JumpStart or that you have access to Amazon Bedrock.

   • **For models hosted by third-party providers**

   Export your provider’s API key in your notebook environment using environment variables. You can use the following magic command. Replace the `provider_API_key` in the command by the environment variable found in the *Environment variable* column of the Jupyter AI **Model providers list** for your provider.

   ```bash
   %env provider_API_key=your_API_key
   ```
Use Jupyter AI in JupyterLab or Studio Classic

Use language models from the chat UI

Compose your message in the chat UI text box to start interacting with your model. To clear the message history, use the `/clear` command.

⚠️ Note
Clearing the message history does not erase the chat context with the model provider.

Use language models from notebook cells

Before using the `%%ai` and `%ai` commands to invoke a language model, load the IPython extension by running the following command in a JupyterLab or Studio Classic notebook cell.

```
%load_ext jupyter_ai_magics
```

• For models hosted by AWS:
  • To invoke a model deployed in SageMaker, pass the string `sagemaker-endpoint:endpoint-name` to the `%%ai` magic command with the required parameters below, then add your prompt in the following lines.

The following table lists the required and optional parameters when invoking models hosted by SageMaker or Amazon Bedrock.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter</th>
<th>Short Version</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Request schema</td>
<td>--request-schema</td>
<td>-q</td>
<td><strong>Required</strong>: The JSON object the endpoint expects, with the prompt being substituted into any value that matches the string literal <code>&lt;prompt&gt;</code>.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter</td>
<td>Short Version</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-------------------</td>
<td>---------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Region name</td>
<td>--region-name</td>
<td>-n</td>
<td><strong>Required:</strong> The AWS Region where the model is deployed.</td>
</tr>
<tr>
<td>Response path</td>
<td>--response-path</td>
<td>-p</td>
<td><strong>Required:</strong> A JSONPath string used to extract the language model's output from the JSON response of the endpoint.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter</td>
<td>Short Version</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------</td>
<td>----------------------------</td>
<td>---------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Extra model parameters</td>
<td>--model-parameters</td>
<td>-m</td>
<td>Optional: A JSON value specifying additional parameters to be passed to the model. The accepted value is parsed into a dictionary, unpacked, and directly passed to the provider class. This is useful when the endpoint or the model requires custom parameters. For example, in Llama 2 models when accepting the End User License Agreement (EULA) is necessary, you can pass the EULA acceptance to the endpoint using <code>-m {}&quot;endpoint_kwargs&quot; :{&quot;Custom Attributes&quot;:&quot;accept_eula=true&quot;} </code>. Alternatively, you can use the <code>-m</code> parameter to pass extra model parameters.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter</td>
<td>Short Version</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
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<td>---------------</td>
<td>-------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>parameters, such as setting the maximum number of tokens for a model's generated response. For example, when working with an AI21 Labs Jurassic model: <code>-m {&quot;model_kwargs&quot;:{&quot;maxTokens&quot;:256}}</code>.</td>
</tr>
</tbody>
</table>

| Output format | --format | -f | Optional: The IPython display used to render the output. It can be any of the following values [code|html|image|json|markdown|math|markdown|text], provided that the invoked model supports the specified format. |

The following command invokes a **Llama2-7b** model hosted by SageMaker.

```bash
%%sagemaker-endpoint:jumpstart-dft-meta-textgeneration-llama-2-7b -q
{"inputs":"<prompt>"","parameters":
{"max_new_tokens":64,"top_p":0.9,"temperature":0.6,"return_full_text":false}}
```
The following example invokes a Flan-t5-small model hosted by SageMaker.

```
%%sagemaker-endpoint:hf-text2text-flan-t5-small --request-schema={"inputs":"<prompt>"} --parameters={"num_return_sequences":4} --region-name=us-west-2 --response-path=[0]["generated_text"] -f text
What is the atomic number of Hydrogen?
```

- To invoke a model deployed in Amazon Bedrock, pass the string `bedrock:model-name` to the `%%%ai` magic command with any optional parameter defined in the list of parameters for invoking models hosted by SageMaker JumpStart or Amazon Bedrock, then add your prompt in the following lines.

The following example invokes an AI21 Labs Jurassic-2 model hosted by Amazon Bedrock.

```
%%bedrock:ai21.j2-mid-v1 -m{"model_kwargs":{"maxTokens":256}} -f code
Write a function in python implementing a bubble sort.
```

- For models hosted by third-party providers

To invoke a model hosted by third-party providers, pass the string `provider-id:model-name` to the `%%%ai` magic command with an optional Output format, then add your prompt in the following lines. You can find the details of each provider, including their ID, in the Jupyter AI list of model providers.

The following command asks an Anthropic Claude model to output an HTML file containing the image of a white square with black borders.

```
%%anthropic:claude-v1.2 -f html
Create a square using SVG with a black border and white fill.
```
Label data with a human-in-the-loop

To train a machine learning model, you need a large, high-quality, labeled dataset. You can label your data using Amazon SageMaker Ground Truth. Choose from one of the Ground Truth built-in task types or create your own custom labeling workflow. To improve the accuracy of your data labels and reduce the total cost of labeling your data, use Ground Truth enhanced data labeling features like automated data labeling and annotation consolidation.

Topics

- Use Amazon SageMaker Ground Truth to Label Data
- Use Amazon SageMaker Ground Truth Plus to Label Data
- Create and Manage Workforces
- Crowd HTML Elements Reference
- Using Amazon Augmented AI for Human Review

Use Amazon SageMaker Ground Truth to Label Data

To train a machine learning model, you need a large, high-quality, labeled dataset. Ground Truth helps you build high-quality training datasets for your machine learning models. With Ground Truth, you can use workers from either Amazon Mechanical Turk, a vendor company that you choose, or an internal, private workforce along with machine learning to enable you to create a labeled dataset. You can use the labeled dataset output from Ground Truth to train your own models. You can also use the output as a training dataset for an Amazon SageMaker model.

Depending on your ML application, you can choose from one of the Ground Truth built-in task types to have workers generate specific types of labels for your data. You can also build a custom labeling workflow to provide your own UI and tools to workers labeling your data. To learn more about the Ground Truth built in task types, see Built-in Task Types. To learn how to create a custom labeling workflow, see Creating Custom Labeling Workflows.

In order to automate labeling your training dataset, you can optionally use automated data labeling, a Ground Truth process that uses machine learning to decide which data needs to be labeled by humans. Automated data labeling may reduce the labeling time and manual effort required. For more information, see Automate Data Labeling. To create a custom labeling workflow, see Creating Custom Labeling Workflows.
Use either pre-built or custom tools to assign the labeling tasks for your training dataset. A *labeling UI template* is a webpage that Ground Truth uses to present tasks and instructions to your workers. The SageMaker console provides built-in templates for labeling data. You can use these templates to get started, or you can build your own tasks and instructions by using our HTML 2.0 components. For more information, see [Creating Custom Labeling Workflows](#).

Use the workforce of your choice to label your dataset. You can choose your workforce from:

- The Amazon Mechanical Turk workforce of over 500,000 independent contractors worldwide.
- A private workforce that you create from your employees or contractors for handling data within your organization.
- A vendor company that you can find in the AWS Marketplace that specializes in data labeling services.

For more information, see [Create and Manage Workforces](#).

You store your datasets in Amazon S3 buckets. The buckets contain three things: The data to be labeled, an input manifest file that Ground Truth uses to read the data files, and an output manifest file. The output file contains the results of the labeling job. For more information, see [Use Input and Output Data](#).

Events from your labeling jobs appear in Amazon CloudWatch under the `/aws/sagemaker/LabelingJobs` group. CloudWatch uses the labeling job name as the name for the log stream.

**Are You a First-time User of Ground Truth?**

If you are a first-time user of Ground Truth, we recommend that you do the following:

1. **Read Getting started**—This section walks you through setting up your first Ground Truth labeling job.
2. **Explore other topics**—Depending on your needs, do the following:
   - **Explore built-in task types**— Use built-in task types to streamline the process of creating a labeling job. See [Built-in Task Types](#) to learn more about Ground Truth built-in task types.
   - **Manage your labeling workforce**—Create new work teams and manage your existing workforce. For more information, see [Create and Manage Workforces](#).
   - **Learn about streaming labeling jobs**— Create a streaming labeling job and send new dataset objects to workers in real time using a perpetually running labeling job. Workers continuously...
receive new data objects to label as long as the labeling job is active and new objects are being sent to it. To learn more, see Ground Truth Streaming Labeling Jobs.

3. See the Reference—This section describes operations to automate Ground Truth operations.

Getting started

This video shows you how to setup and use Amazon SageMaker Ground Truth. (Length: 9:37)

To get started using Amazon SageMaker Ground Truth, follow the instructions in the following sections. The sections here explain how to use the console to create a labeling job, assign a public or private workforce, and send the labeling job to your workforce. You can also learn how to monitor the progress of a labeling job.

If you want to create a custom labeling workflow, see Creating Custom Labeling Workflows for instructions.

Before you create a labeling job, you must upload your dataset to an Amazon S3 bucket. For more information, see Use Input and Output Data.

Topics

- Step 1: Before You Begin
- Step 2: Create a Labeling Job
- Step 3: Select Workers
- Step 4: Configure the Bounding Box Tool
- Step 5: Monitoring Your Labeling Job

Step 1: Before You Begin

Before you begin using the SageMaker console to create a labeling job, you must set up the dataset for use. Do this:

1. Save two images at publicly available HTTP URLs. The images are used when creating instructions for completing a labeling task. The images should have an aspect ratio of around 2:1. For this exercise, the content of the images is not important.

2. Create an Amazon S3 bucket to hold the input and output files. The bucket must be in the same Region where you are running Ground Truth. Make a note of the bucket name because you use it during step 2.
Ground Truth requires all S3 buckets that contain labeling job input image data have a CORS policy attached. To learn more about this change, see CORS Permission Requirement.

3. You can create an IAM role or let SageMaker create a role with the AmazonSageMakerFullAccess IAM policy. Refer to Creating IAM roles and assign the following permissions policy to the user that is creating the labeling job:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "sagemakergroundtruth",
            "Effect": "Allow",
            "Action": [
                "cognito-idp:CreateGroup",
                "cognito-idp:CreateUserPool",
                "cognito-idp:CreateUserPoolDomain",
                "cognito-idp:AdminCreateUser",
                "cognito-idp:CreateUserPoolClient",
                "cognito-idp:AdminAddUserToGroup",
                "cognito-idp:DescribeUserPoolClient",
                "cognito-idp:DescribeUserPool",
                "cognito-idp:UpdateUserPool"
            ],
            "Resource": "*"
        }
    ]
}
```

Next

**Step 2: Create a Labeling Job**

In this step you use the console to create a labeling job. You tell Amazon SageMaker Ground Truth the Amazon S3 bucket where the manifest file is stored and configure the parameters for the job. For more information about storing data in an Amazon S3 bucket, see Use Input and Output Data.
To create a labeling job

2. From the left navigation, choose Labeling jobs.
3. Choose Create labeling job to start the job creation process.
4. In the Job overview section, provide the following information:
   - **Job name** – Give the labeling job a name that describes the job. This name is shown in your job list. The name must be unique in your account in an AWS Region.
   - **Label attribute name** – Leave this unchecked as the default value is the best option for this introductory job.
   - **Input data setup** – Select Automated data setup. This option allows you to automatically connect to your input data in S3.
   - **S3 location for input datasets** – Enter the S3 location where you added the images in step 1.
   - **S3 location for output datasets** – The location where your output data is written in S3.
   - **Data type** – Use the drop down menu to select Image. Ground Truth will use all images found in the S3 location for input datasets as input for your labeling job.
   - **IAM role** – Create or choose an IAM role with the AmazonSageMakerFullAccess IAM policy attached.
5. In the Task type section, for the Task category field, choose Image.
6. In the Task selection choose Bounding box.
7. Choose Next to move on to configuring your labeling job.

Next

Step 3: Select Workers

In this step you choose a workforce for labeling your dataset. It is recommended that you create a private workforce to test Amazon SageMaker Ground Truth. Use email addresses to invite the members of your workforce. If you create a private workforce in this step you won’t be able to import your Amazon Cognito user pool later. If you want to create a private workforce using an Amazon Cognito user pool, see Manage a Private Workforce (Amazon Cognito) and use the Mechanical Turk workforce instead in this tutorial.
To create a private workforce:

1. In the Workers section, choose Private.

2. If this is your first time using a private workforce, in the Email addresses field, enter up to 100 email addresses. The addresses must be separated by a comma. You should include your own email address so that you are part of the workforce and can see data object labeling tasks.

3. In the Organization name field, enter the name of your organization. This information is used to customize the email sent to invite a person to your private workforce. You can change the organization name after the user pool is created through the console.

4. In the Contact email field enter an email address that members of the workforce use to report problems with the task.

If you add yourself to the private workforce, you will receive an email that looks similar to the following. Amazon, Inc. is replaced by the organization you enter in step 3 of the preceding procedure. Select the link in the email to log in using the temporary password provided. If prompted, change your password. When you successfully log in, you see the worker portal where your labeling tasks appear.
You're invited to work on a labeling project.

You will need this user name and temporary password to log in the first time.

User name: user1234
Temporary password: password

Open the link below to log in:
https://yourlabelingurl.com

After you log in with your temporary password, you are required to create a new one. If you have any questions, please contact help@amazon.com.

Tip

You can find the link to your private workforce's worker portal in the Labeling workforces section of the Ground Truth area of the SageMaker console. To see the link, select the Private tab. The link is under the Labeling portal sign-in URL header in Private workforce summary.

If you choose to use the Amazon Mechanical Turk workforce to label the dataset, you are charged for labeling tasks completed on the dataset.
To use the Amazon Mechanical Turk workforce:

1. In the **Workers** section, choose **Public**.
2. Set a **Price per task**.
3. If applicable, choose **The dataset does not contain adult content** to acknowledge that the sample dataset has no adult content. This information enables Amazon SageMaker Ground Truth to warn external workers on Mechanical Turk that they might encounter potentially offensive content in your dataset.
4. Choose the check box next to the following statement to acknowledge that the sample dataset does not contain any personally identifiable information (PII). This is a requirement to use Mechanical Turk with Ground Truth. If your input data does contain PII, use the private workforce for this tutorial.

You understand and agree that the Amazon Mechanical Turk workforce consists of independent contractors located worldwide and that you should not share confidential information, personal information or protected health information with this workforce.

Next

**Step 4: Configure the Bounding Box Tool**

Finally you configure the bounding box tool to give instructions to your workers. You can configure a task title that describes the task and provides high-level instructions for the workers. You can provide both quick instructions and full instructions. Quick instructions are displayed next to the image to be labeled. Full instructions contain detailed instructions for completing the task. In this example, you only provide quick instructions. You can see an example of full instructions by choosing **Full instructions** at the bottom of the section.

**To configure the bounding box tool**

1. In the **Task description** field type in brief instructions for the task. For example:

   **Draw a box around any objects in the image.**

   Replace *objects* with the name of an object that appears in your images.
2. In the **Labels** field, type a category name for the objects that the worker should draw a bounding box around. For example, if you are asking the worker to draw boxes around football players, you could use "Football Player" in this field.

3. The **Short instructions** section enables you to create instructions that are displayed on the page with the image that your workers are labeling. We suggest that you include an example of a correctly drawn bounding box and an example of an incorrectly drawn box. To create your own instructions, use these steps:
   a. Select the text between **GOOD EXAMPLE** and the image placeholder. Replace it with the following text:

   **Draw the box around the object with a small border.**

   b. Select the first image placeholder and delete it.

   c. Choose the image button and then enter the HTTPS URL of one of the images that you created in step 1. It is also possible to embed images directly in the short instructions section, however this section has a quota of 100 kilobytes (including text). If your images and text exceed 100 kilobytes, you receive an error.

   d. Select the text between **BAD EXAMPLE** and the image placeholder. Replace it with the following text:

   **Don't make the bounding box too large or cut into the object.**

   e. Select the second image placeholder and delete it.

   f. Choose the image button and then enter the HTTPS URL of the other image that you created in step 1.

4. Select **Preview** to preview the worker UI. The preview opens in a new tab, and so if your browser blocks pop ups you may need to manually enable the tab to open. When you add one or more annotations to the preview and then select **Submit** you can see a preview of the output data your annotation would create.

5. After you have configured and verified your instructions, select **Create** to create the labeling job.

If you used a private workforce, you can navigate to the worker portal that you logged into in **Step 3: Select Workers** of this tutorial to see your labeling tasks. The tasks may take a few minutes to appear.
Step 5: Monitoring Your Labeling Job

After you create your labeling job, you see a list of all the jobs that you have created. You can use this list to monitor the status of your labeling jobs. The list has the following fields:

- **Name** – The name that you assigned the job when you created it.
- **Status** – The completion status of the job. The status can be one of Complete, Failed, In progress, or Stopped.
- **Labeled objects/total** – Shows the total number of objects in the labeling job and how many of them have been labeled.
- **Creation time** – The date and time that you created the job.

You can also clone, chain, or stop a job. Select a job and then select one of the following from the **Actions** menu:

- **Clone** – Creates a new labeling job with the configuration copied from the selected job. You can clone a job when you want to change to the job and run it again. For example, you can clone a job that was sent to a private workforce so that you can send it to the Amazon Mechanical Turk workforce. Or you can clone a job to rerun it against a new dataset stored in the same location as the original job.
- **Chain** – Creates a new labeling job that can build upon the data and models (if any) of a stopped, failed, or completed job. For more information about the use cases and how to use it, see [Chaining Labeling Jobs](#).
- **Stop** – Stops a running job. You cannot restart a stopped job. You can clone a job to start over or chain the job to continue from where it left off. Labels for any already labeled objects are written to the output file location. For more information, see [Output Data](#).

Label Images

Use Ground Truth to label images. Select one of the following built-in task types to learn more about that task type. Each page includes instructions to help you create a labeling job using that task type.
Tip

To learn more about supported file types and input data quotas, see Input Data.

Topics

• Bounding Box
• Image Semantic Segmentation
• Auto-Segmentation Tool
• Image Classification (Single Label)
• Image Classification (Multi-label)
• Image Label Verification

Bounding Box

The images used to train a machine learning model often contain more than one object. To classify and localize one or more objects within images, use the Amazon SageMaker Ground Truth bounding box labeling job task type. In this context, localization means the pixel-location of the bounding box.

You create a bounding box labeling job using the Ground Truth section of the Amazon SageMaker console or the CreateLabelingJob operation.

Important

For this task type, if you create your own manifest file, use "source-ref" to identify the location of each image file in Amazon S3 that you want labeled. For more information, see Input Data.

Creating a Bounding Box Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) to learn how to create a bounding box labeling job in the SageMaker console. In Step 10, choose Image from the Task category drop down menu, and choose Bounding box as the task type.
Ground Truth provides a worker UI similar to the following for labeling tasks. When you create the labeling job with the console, you specify instructions to help workers complete the job and up to 50 labels that workers can choose from.

Create a Bounding Box Labeling Job (API)

To create a bounding box labeling job, use the SageMaker API operation `CreateLabelingJob`. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of `CreateLabelingJob`.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with `PRE-BoundingBox`. To find the pre-annotation Lambda ARN for your Region, see `PreHumanTaskLambdaArn`.
- Annotation-consolidation Lambda functions for this task type end with `ACS-BoundingBox`. To find the annotation-consolidation Lambda ARN for your Region, see `AnnotationConsolidationLambdaArn`.

Label Images 1376
The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.

```python
response = client.create_labeling_job(
    LabelingJobName='example-bounding-box-labeling-job,
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*,
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::*:workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/worker-task-template.html'
        },
        'PreHumanTaskLambdaArn': 'arn:aws:lambda:us-east-1:432418664414:function:PRE-BoundingBox',
        'TaskKeywords': [
            'Bounding Box',
        ],
        'TaskTitle': 'Bounding Box task',
        'TaskDescription': 'Draw bounding boxes around objects in an image',
        'NumberOfHumanWorkersPerDataObject': 123,
        'TaskTimeLimitInSeconds': 123
    }
)
```
Provide a Template for Bounding Box Labeling Jobs

If you create a labeling job using the API, you must supply a worker task template in `UiTemplateS3Uri`. Copy and modify the following template. Only modify the `short-instructions`, `full-instructions`, and `header`. Upload this template to S3, and provide the S3 URI for this file in `UiTemplateS3Uri`.

```xml
<full-instructions header="Bounding box instructions">  
  <ol>
    <li><strong>Inspect</strong> the image</li>
    <li><strong>Determine</strong> if the specified label is/are visible in the picture</li>
    <li><strong>Outline</strong> each instance of the specified label in the image using the provided “Box” tool</li>
  </ol>
  <ul>
    <li>Boxes should fit tight around each object</li>
    <li>Do not include parts of the object are overlapping or that cannot be seen, even though you think you can interpolate the whole shape</li>
    <li>Avoid including shadows</li>
    <li>If the target is off screen, draw the box up to the edge of the image</li>
  </ul>
</full-instructions>
```
Bounding Box Output Data

Once you have created a bounding box labeling job, your output data will be located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the **Output dataset location** field of the **Job overview** section of the console.

For example, the output manifest file of a successfully completed single-class bounding box task will contain the following:

```json
[
  {
    "boundingBox": {
      "boundingBoxes": [
        {
          "height": 2832,
          "label": "bird",
          "left": 681,
          "top": 599,
          "width": 1364
        }
      ],
      "inputImageProperties": {
        "height": 3726,
        "width": 2662
      }
    }
  }
]
```

The **boundingBoxes** parameter identifies the location of the bounding box drawn around an object identified as a "bird" relative to the top-left corner of the image which is taken to
be the (0,0) pixel-coordinate. In the previous example, **left** and **top** identify the location of the pixel in the top-left corner of the bounding box relative to the top-left corner of the image. The dimensions of the bounding box are identified with **height** and **width**. The `inputImageProperties` parameter gives the pixel-dimensions of the original input image.

When you use the bounding box task type, you can create single- and multi-class bounding box labeling jobs. The output manifest file of a successfully completed multi-class bounding box will contain the following:

```
[
  {
    "boundingBox": {
      "boundingBoxes": [
        {
          "height": 938,
          "label": "squirrel",
          "left": 316,
          "top": 218,
          "width": 785
        },
        {
          "height": 825,
          "label": "rabbit",
          "left": 1930,
          "top": 2265,
          "width": 540
        },
        {
          "height": 1174,
          "label": "bird",
          "left": 748,
          "top": 2113,
          "width": 927
        },
        {
          "height": 893,
          "label": "bird",
          "left": 1333,
          "top": 847,
          "width": 736
        }
      ],
      "inputImageProperties": {
```
To learn more about the output manifest file that results from a bounding box labeling job, see **Bounding Box Job Output**.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see **Output Data**.

**Image Semantic Segmentation**

To identify the contents of an image at the pixel level, use an Amazon SageMaker Ground Truth semantic segmentation labeling task. When assigned a semantic segmentation labeling job, workers classify pixels in the image into a set of predefined labels or classes. Ground Truth supports single and multi-class semantic segmentation labeling jobs.

Images that contain large numbers of objects that need to be segmented require more time. To help workers (from a private or vendor workforce) label these objects in less time and with greater accuracy, Ground Truth provides an AI-assisted auto-segmentation tool. For information, see **Auto-Segmentation Tool**.

You create a semantic segmentation labeling job using the Ground Truth section of the Amazon SageMaker console or the **CreateLabelingJob** operation.

⚠️ **Important**

For this task type, if you create your own manifest file, use "source-ref" to identify the location of each image file in Amazon S3 that you want labeled. For more information, see **Input Data**.

**Creating a Semantic Segmentation Labeling Job (Console)**

You can follow the instructions **Create a Labeling Job (Console)** to learn how to create a semantic segmentation labeling job in the SageMaker console. In Step 10, choose **Image** from the **Task category** drop down menu, and choose **Semantic segmentation** as the task type.
Ground Truth provides a worker UI similar to the following for labeling tasks. When you create the labeling job with the console, you specify instructions to help workers complete the job and labels that workers can choose from.

Create a Semantic Segmentation Labeling Job (API)

To create a semantic segmentation labeling job, use the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with PRE-SemanticSegmentation. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn.
- Annotation-consolidation Lambda functions for this task type end with ACS-SemanticSegmentation. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.
The following is an example of an **AWS Python SDK (Boto3) request** to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.

```python
response = client.create_labeling_job(
    LabelingJobName='example-semantic-segmentation-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation',
                'FreeOfAdultContent'
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*',
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::*:workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/worker-task-template.html'
        },
        'TaskKeywords': [
            'Semantic Segmentation',
        ],
        'TaskTitle': 'Semantic segmentation task',
        'TaskDescription': 'For each category provided, segment out each relevant object using the color associated with that category',
        'NumberOfHumanWorkersPerDataObject': 123,
    }
)
```
Provide a Template for Semantic Segmentation Labeling Jobs

If you create a labeling job using the API, you must supply a worker task template in UiTemplateS3Uri. Copy and modify the following template. Only modify the short-instructions, full-instructions, and header.

Upload this template to S3, and provide the S3 URI for this file in UiTemplateS3Uri.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
    <crowd-semantic-segmentation
        name="crowd-semantic-segmentation"
        src="{{ task.input.taskObject | grant_read_access }}"
        header="Please segment out all pedestrians."
        labels="{{ task.input.labels | to_json | escape }}"
    >
        <full-instructions header="Segmentation instructions">
            <ol>
                <li><strong>Read</strong> the task carefully and inspect the image.</li>
                <li><strong>Read</strong> the options and review the examples provided to understand more about the labels.</li>
                <li><strong>Choose</strong> the appropriate label that best suits an object and paint that object using the tools provided.</li>
            </ol>
        </full-instructions>
        <short-instructions>
            <h2>Good example</h2>
            <p>Enter description to explain a correctly done segmentation</p>
            <h2>Bad example</h2>
            <p>Enter description of an incorrectly done segmentation</p>
        </short-instructions>
    </crowd-semantic-segmentation>
</crowd-form>
```
Semantic Segmentation Output Data

Once you have created a semantic segmentation labeling job, your output data will be located in the Amazon S3 bucket specified in the `S3OutputPath` parameter when using the API or in the `Output dataset location` field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see [Output Data](#).

To see an example of an output manifest file for a semantic segmentation labeling job, see [3D Point Cloud Semantic Segmentation Output](#).

Auto-Segmentation Tool

Image segmentation is the process of dividing an image into multiple segments, or sets of labeled pixels. In Amazon SageMaker Ground Truth, the process of identifying all pixels that fall under a given label involves applying a colored filler, or "mask", over those pixels. Some labeling job tasks contain images with a large numbers of objects that need to be segmented. To help workers label these objects in less time and with greater accuracy, Ground Truth provides an auto-segmentation tool for segmentation tasks assigned to private and vendor workforces. This tool uses a machine learning model to automatically segment individual objects in the image with minimal worker input. Workers can refine the mask generated by the auto-segmentation tool using other tools found in the worker console. This helps workers complete image segmentation tasks faster and more accurately, resulting in lower cost and higher label quality.

Note

The auto-segmentation tool is available for segmentation tasks that are sent to a private workforce or vendor workforce. It isn't available for tasks sent to the public workforce (Amazon Mechanical Turk).
Tool Preview

When workers are assigned a labeling job that provides the auto-segmentation tool, they are provided with detailed instructions on how to use the tool. For example, a worker might see the following in the worker console:

Workers can use **View full instructions** to learn how to use the tool. Workers will need to place a point on four extreme-points (top-most, bottom-most, left-most, and right-most points) of the object of interest, and the tool will automatically generate a mask for the object. Workers can further-refine the mask using the other tools provided, or by using the auto-segment tool on smaller portions of the object that were missed.

**Tool Availability**

The auto-segmentation tool automatically appears in your workers' consoles if you create a semantic segmentation labeling job using the Amazon SageMaker console. While creating a semantic segmentation job in the SageMaker console, you will be able to preview the tool while
creating worker instructions. To learn how to create a semantic segmentation labeling job in the SageMaker console, see Getting started.

If you are creating a custom instance segmentation labeling job in the SageMaker console or creating an instance- or semantic-segmentation labeling job using the Ground Truth API, you need to create a custom task template to design your worker console and instructions. To include the auto-segmentation tool in your worker console, ensure that the following conditions are met in your custom task template:

- For semantic segmentation labeling jobs created using the API, the <crowd-semantic-segmentation> is present in the task template. For custom instance segmentation labeling jobs, the <crowd-instance-segmentation> tag is present in the task template.
- The task is assigned to a private workforce or vendor workforce.
- The images to be labeled are Amazon Simple Storage Service Amazon S3) objects that have been pre-signed for the Worker so that they can access it. This is true if the task template includes the grant_read_access filter. For information about the grant_read_access filter, see Adding automation with Liquid.

The following is an example of a custom task template for a custom instance segmentation labeling job, which includes the <crowd-instance-segmentation/> tag and the grant_read_access Liquid filter.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
  <crowd-instance-segmentation
    name="crowd-instance-segmentation"
    src="{{ task.input.taskObject | grant_read_access }}"
    labels="['Car','Road']"
  >
    <full-instructions header="Segmentation instructions">
      Segment each instance of each class of objects in the image.
    </full-instructions>
  </crowd-instance-segmentation>

  <short-instructions>
    <p>Segment each instance of each class of objects in the image.</p>

    <h3 style="color: green">GOOD EXAMPLES</h3>
    <img src="path/to/image.jpg" style="width: 100%">
    <p>Good because A, B, C.</p>

    <h3 style="color: red">BAD EXAMPLES</h3>
</crowd-form>
```
Image Classification (Single Label)

Use an Amazon SageMaker Ground Truth image classification labeling task when you need workers to classify images using predefined labels that you specify. Workers are shown images and are asked to choose one label for each image.

You can create an image classification labeling job using the Ground Truth section of the Amazon SageMaker console or the `CreateLabelingJob` operation.

**Important**

For this task type, if you create your own manifest file, use "source-ref" to identify the location of each image file in Amazon S3 that you want labeled. For more information, see Input Data.

Create an Image Classification Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) to learn how to create a image classification labeling job in the SageMaker console. In Step 10, choose Image from the Task category drop down menu, and choose Image Classification (Single Label) as the task type.

Ground Truth provides a worker UI similar to the following for labeling tasks. When you create the labeling job with the console, you specify instructions to help workers complete the job and labels that workers can choose from.
Create an Image Classification Labeling Job (API)

To create an image classification labeling job, use the SageMaker API operation `CreateLabelingJob`. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of `CreateLabelingJob`.

Follow the instructions on [Create a Labeling Job (API)](https://docs.aws.amazon.com/sagemaker/latest/dg/create-labeling-job.html) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with `PRE-ImageMultiClass`. To find the pre-annotation Lambda ARN for your Region, see [PreHumanTaskLambdaArn](https://docs.aws.amazon.com/sagemaker/latest/dg/prehuman-task-lambda-arn.html).
Annotation-consolidation Lambda functions for this task type end with ACS-ImageMultiClass. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.

The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.

```python
response = client.create_labeling_job(
    LabelingJobName='example-image-classification-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*',
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/worker-task-template.html'
        },
        'PreHumanTaskLambdaArn': 'arn:aws:lambda:us-east-1:432418664414:function:PRE-ImageMultiClass,
        'TaskKeywords': ['Image classification',
```

Label Images
Provide a Template for Image Classification Labeling Jobs

If you create a labeling job using the API, you must supply a worker task template in UiTemplateS3Uri. Copy and modify the following template. Only modify the short-instructions, full-instructions, and header.

Upload this template to S3, and provide the S3 URI for this file in UiTemplateS3Uri.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
  <crowd-image-classifier
    name="crowd-image-classifier"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="please classify"
    categories="{{ task.input.labels | to_json | escape }}"
  >
    <full-instructions header="Image classification instructions">
      <ol>
        <li><strong>Read</strong> the task carefully and inspect the image.</li>
        <li><strong>Read</strong> the options and review the examples provided to understand more about the labels.</li>
        <li><strong>Choose</strong> the appropriate label that best suits the image.</li>
      </ol>
  </full-instructions>
</crowd-form>
```
Image Classification Output Data

Once you have created an image classification labeling job, your output data will be located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the Output dataset location field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see Output Data.

To see an example of an output manifest file from an image classification labeling job, see Classification Job Output.

Image Classification (Multi-label)

Use an Amazon SageMaker Ground Truth multi-label image classification labeling task when you need workers to classify multiple objects in an image. For example, the following image features a dog and a cat. You can use multi-label image classification to associate the labels "dog" and "cat" with this image.
When working on a multi-label image classification task, workers should choose all applicable labels, but must choose at least one. When creating a job using this task type, you can provide up to 50 label-categories.
When creating a labeling job in the console, Ground Truth doesn't provide a "none" category for when none of the labels applies to an image. To provide this option to workers, include a label similar to "none" or "other" when you create a multi-label image classification job.

To restrict workers to choosing a single label for each image, use the Image Classification (Single Label) task type.

⚠️ Important

For this task type, if you create your own manifest file, use "source-ref" to identify the location of each image file in Amazon S3 that you want labeled. For more information, see Input Data.

Create a Multi-Label Image Classification Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) to learn how to create a multi-label image classification labeling job in the SageMaker console. In Step 10, choose Image from the Task category drop down menu, and choose Image Classification (Multi-label) as the task type.

Ground Truth provides a worker UI similar to the following for labeling tasks. When you create a labeling job in the console, you specify instructions to help workers complete the job and labels that workers can choose from.
Create a Multi-Label Image Classification Labeling Job (API)

To create a multi-label image classification labeling job, use the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with PRE-ImageMultiClassMultiLabel. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn.
- Annotation-consolidation Lambda functions for this task type end with ACS-ImageMultiClassMultiLabel. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.
The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.

```python
response = client.create_labeling_job(
    LabelingJobName='example-multi-label-image-classification-labeling-job,
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*,
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::*:workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/worker-task-template.html'
        },
        'TaskKeywords': [
            'Image Classification',
        ],
        'TaskTitle': 'Multi-label image classification task',
        'TaskDescription': 'Select all labels that apply to the images shown',
        'NumberOfHumanWorkersPerDataObject': 123,
        'TaskTimeLimitInSeconds': 123,
    }
)
```
Provide a Template for Multi-label Image Classification

If you create a labeling job using the API, you must supply a worker task template in `UiTemplateS3Uri`. Copy and modify the following template. Only modify the `short-instructions`, `full-instructions`, and header.

Upload this template to S3, and provide the S3 URI for this file in `UiTemplateS3Uri`.

```
<crowd-form>
  <crowd-image-classifier-multi-select
    name="crowd-image-classifier-multi-select"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="Please identify all classes in image"
    categories="{{ task.input.labels | to_json | escape }}"
  >
  <full-instructions header="Multi Label Image classification instructions">
    <ol>
      <li><strong>Read</strong> the task carefully and inspect the image.</li>
      <li><strong>Read</strong> the options and review the examples provided to understand more about the labels.</li>
      <li><strong>Choose</strong> the appropriate labels that best suit the image.</li>
    </ol>
  </full-instructions>
  <short-instructions>
    <h3><span style="color: rgb(0, 138, 0);">Good example</span></h3>
    <p>Enter description to explain the correct label to the workers</p>
    <h3><span style="color: rgb(230, 0, 0);">Bad example</span></h3>
    <p>Enter description of an incorrect label</p>
  </short-instructions>
</crowd-form>
```
Multi-label Image Classification Output Data

Once you have created a multi-label image classification labeling job, your output data will be located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the Output dataset location field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see Output Data.

To see an example of output manifest files for multi-label image classification labeling job, see Multi-label Classification Job Output.

Image Label Verification

Building a highly accurate training dataset for your machine learning (ML) algorithm is an iterative process. Typically, you review and continuously adjust your labels until you are satisfied that they accurately represent the ground truth, or what is directly observable in the real world.

You can use an Amazon SageMaker Ground Truth image label verification task to direct workers to review a dataset's labels and improve label accuracy. Workers can indicate if the existing labels are correct or rate label quality. They can also add comments to explain their reasoning. Amazon SageMaker Ground Truth supports label verification for Bounding Box and Image Semantic Segmentation labels.

You create an image label verification labeling job using the Ground Truth section of the Amazon SageMaker console or the CreateLabelingJob operation.

Ground Truth provides a worker console similar to the following for labeling tasks. When you create the labeling job with the console, you can modify the images and content that are shown. To learn how to create a labeling job using the Ground Truth console, see Create a Labeling Job (Console).
You can create a label verification labeling job using the SageMaker console or API. To learn how to create a labeling job using the Ground Truth API operation CreateLabelingJob, see Create a Labeling Job (API).

Use Ground Truth to Label Text

Use Ground Truth to label text. Select one of the following built in task types to learn more about that task type. Each page includes instructions to help you create a labeling job using that task type.

Tip

To learn more about supported file types and input data quotas, see Input Data.
Topics

- Named Entity Recognition
- Text Classification (Single Label)
- Text Classification (Multi-label)

Named Entity Recognition

To extract information from unstructured text and classify it into predefined categories, use an Amazon SageMaker Ground Truth named entity recognition (NER) labeling task. Traditionally, NER involves sifting through text data to locate noun phrases, called named entities, and categorizing each with a label, such as "person," "organization," or "brand." You can broaden this task to label longer spans of text and categorize those sequences with predefined labels that you specify.

When tasked with a named entity recognition labeling job, workers apply your labels to specific words or phrases within a larger text block. They choose a label, then apply it by using the cursor to highlight the part of the text to which the label applies. The Ground Truth named entity recognition tool supports overlapping annotations, in-context label selection, and multi-label selection for a single highlight. Also, workers can use their keyboards to quickly select labels.

You can create a named entity recognition labeling job using the Ground Truth section of the Amazon SageMaker console or the CreateLabelingJob operation.

⚠️ Important

If you manually create an input manifest file, use "source" to identify the text that you want labeled. For more information, see Input Data.

Create a Named Entity Recognition Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) to learn how to create a named entity recognition labeling job in the SageMaker console. In Step 10, choose Text from the Task category drop down menu, and choose Named entity recognition as the task type.

Ground Truth provides a worker UI similar to the following for labeling tasks. When you create the labeling job with the console, you specify instructions to help workers complete the job and labels that workers can choose from.
Create a Named Entity Recognition Labeling Job (API)

To create a named entity recognition labeling job, using the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with PRE-NamedEntityRecognition. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn.

- Annotation-consolidation Lambda functions for this task type end with ACS-NamedEntityRecognition. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.
You must provide the following ARN for `HumanTaskUiArn`:

```
```

Replace `aws-region` with the AWS Region you use to create the labeling job. For example, use `us-west-1` if you create a labeling job in US West (N. California).

Provide worker instructions in the label category configuration file using the `instructions` parameter. You can use a string, or HTML markup language in the `shortInstruction` and `fullInstruction` fields. For more details, see [Provide Worker Instructions in a Label Category Configuration File](#).

```
"instructions": {"shortInstruction":"<h1>Add header</h1><p>Add Instructions</p>"},
"fullInstruction": "<p>Add additional instructions.</p>"
```

The following is an example of an [AWS Python SDK (Boto3) request](#) to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.

```python
response = client.create_labeling_job(
    LabelingJobName='example-ner-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': ['FreeOfPersonallyIdentifiableInformation','FreeOfAdultContent'],
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*',
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
```
Provide Worker Instructions in a Label Category Configuration File

You must provide worker instructions in the label category configuration file you identify with the LabelCategoryConfigS3Uri parameter in CreateLabelingJob. You can use these instructions to provide details about the task you want workers to perform and help them use the tool efficiently.

You provide short and long instructions using shortInstruction and fullInstruction in the instructions parameter, respectively. To learn more about these instruction types, see Creating Instruction Pages.
The following is an example of a label category configuration file with instructions that can be used for a named entity recognition labeling job.

```json
{
    "document-version": "2018-11-28",
    "labels": [
        {
            "label": "label1",
            "shortDisplayName": "L1"
        },
        {
            "label": "label2",
            "shortDisplayName": "L2"
        },
        {
            "label": "label3",
            "shortDisplayName": "L3"
        },
        {
            "label": "label4",
            "shortDisplayName": "L4"
        },
        {
            "label": "label5",
            "shortDisplayName": "L5"
        }
    ],
    "instructions": {
        "shortInstruction": "<p>Enter description of the labels that workers have to choose from</p><br><p>Add examples to help workers understand the label</p>",
        "fullInstruction": "<ol>
        <li><strong>Read</strong> the text carefully.</li>
        <li><strong>Highlight</strong> words, phrases, or sections of the text.</li>
        <li><strong>Choose</strong> the label that best matches what you have highlighted.</li>
        <li>To <strong>change</strong> a label, choose highlighted text and select a new label.</li>
        <li>To <strong>remove</strong> a label from highlighted text, choose the X next to the abbreviated label name on the highlighted text.</li>
    </ol>
```
<li>You can select all of a previously highlighted text, but not a portion of it.</li>
</ol>

Named Entity Recognition Output Data

Once you have created a named entity recognition labeling job, your output data will be located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the Output dataset location field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see Output Data.

Text Classification (Single Label)

To categorize articles and text into predefined categories, use text classification. For example, you can use text classification to identify the sentiment conveyed in a review or the emotion underlying a section of text. Use Amazon SageMaker Ground Truth text classification to have workers sort text into categories that you define.

You create a text classification labeling job using the Ground Truth section of the Amazon SageMaker console or the CreateLabelingJob operation.

⚠️ Important

If you manually create an input manifest file, use "source" to identify the text that you want labeled. For more information, see Input Data.

Create a Text Classification Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) to learn how to create a text classification labeling job in the SageMaker console. In Step 10, choose Text from the Task category drop down menu, and choose Text Classification (Single Label) as the task type.

Ground Truth provides a worker UI similar to the following for labeling tasks. When you create the labeling job with the console, you specify instructions to help workers complete the job and labels that workers can choose from.
Create a Text Classification Labeling Job (API)

To create a text classification labeling job, use the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with PRE-TextMultiClass. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn.
- Annotation-consolidation Lambda functions for this task type end with ACS-TextMultiClass. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.

The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.
response = client.create_labeling_job(
    LabelingJobName='example-text-classification-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iams::*:role/*',
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::*:workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/worker-task-template.html'
        },
        'TaskKeywords': ['Text classification'],
        'TaskTitle': 'Text classification task',
        'TaskDescription': 'Carefully read and classify this text using the categories provided.',
        'NumberOfHumanWorkersPerDataObject': 123,
        'TaskTimeLimitInSeconds': 123,
        'TaskAvailabilityLifetimeInSeconds': 123,
        'MaxConcurrentTaskCount': 123,
        'AnnotationConsolidationConfig': {
        },
Provide a Template for Text Classification Labeling Jobs

If you create a labeling job using the API, you must supply a worker task template in `UiTemplateS3Uri`. Copy and modify the following template. Only modify the `short-instructions`, `full-instructions`, and header.

Upload this template to S3, and provide the S3 URI for this file in `UiTemplateS3Uri`.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
  <crowd-classifier
    name="crowd-classifier"
    categories="{{ task.input.labels | to_json | escape }}"
    header="classify text"
  >
    <classification-target style="white-space: pre-wrap">
      {{ task.input.taskObject }}
    </classification-target>
    <full-instructions header="Classifier instructions">
      <ol>
        <li><strong>Read</strong> the text carefully.</li>
        <li><strong>Read</strong> the examples to understand more about the options.</li>
        <li><strong>Choose</strong> the appropriate labels that best suit the text.</li>
      </ol>
    </full-instructions>
    <short-instructions>
      <p>Enter description of the labels that workers have to choose from</p>
      <p>Add examples to help workers understand the label</p>
    </short-instructions>
  </crowd-classifier>
</crowd-form>
```
Text Classification Output Data

Once you have created a text classification labeling job, your output data will be located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the Output dataset location field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see Output Data.

To see an example of an output manifest files from a text classification labeling job, see Classification Job Output.

Text Classification (Multi-label)

To categorize articles and text into multiple predefined categories, use the multi-label text classification task type. For example, you can use this task type to identify more than one emotion conveyed in text.

When working on a multi-label text classification task, workers should choose all applicable labels, but must choose at least one. When creating a job using this task type, you can provide up to 50 label categories.

Amazon SageMaker Ground Truth doesn't provide a "none" category for when none of the labels applies. To provide this option to workers, include a label similar to "none" or "other" when you create a multi-label text classification job.

To restrict workers to choosing a single label for each document or text selection, use the Text Classification (Single Label) task type.

⚠️ Important

If you manually create an input manifest file, use "source" to identify the text that you want labeled. For more information, see Input Data.

Create a Multi-Label Text Classification Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) to learn how to create a multi-label text classification labeling job in the Amazon SageMaker console. In Step 10, choose Text from the Task category drop down menu, and choose Text Classification (Multi-label) as the task type.
Ground Truth provides a worker UI similar to the following for labeling tasks. When you create the labeling job with the console, you specify instructions to help workers complete the job and labels that workers can choose from.

Create a Multi-Label Text Classification Labeling Job (API)

To create a multi-label text classification labeling job, use the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Pre-annotation Lambda functions for this task type end with PRE-TextMultiClassMultiLabel. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn.
- Annotation-consolidation Lambda functions for this task type end with ACS-TextMultiClassMultiLabel. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.
The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region. All parameters in red should be replaced with your specifications and resources.

```python
response = client.create_labeling_job(
    LabelingJobName='example-multi-label-text-classification-labeling-job,
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation','FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*,
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::*:workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/custom-worker-task-template.html'
        },
        'PreHumanTaskLambdaArn': 'arn:aws:lambda::function:PRE-TextMultiClassMultiLabel,
        'TaskKeywords': ['Text Classification',
                        ],
        'TaskTitle': 'Multi-label text classification task',
        'TaskDescription': 'Select all labels that apply to the text shown',
        'NumberOfHumanWorkersPerDataObject': 123,
        'TaskTimeLimitInSeconds': 123,
    }
)
```
'TaskAvailabilityLifetimeInSeconds': 123,
'MaxConcurrentTaskCount': 123,
'AnnotationConsolidationConfig': {
    'AnnotationConsolidationLambdaArn': 'arn:aws:lambda:us-east-1:432418664414:function:ACS-TextMultiClassMultiLabel'
},
'Tags':[{
    'Key': 'string',
    'Value': 'string'
}]

Create a Template for Multi-label Text Classification

If you create a labeling job using the API, you must supply a worker task template in `UiTemplateS3Uri`. Copy and modify the following template. Only modify the `short-instructions`, `full-instructions`, and `header`.

Upload this template to S3, and provide the S3 URI for this file in `UiTemplateS3Uri`.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
  <crowd-classifier-multi-select
    name="crowd-classifier-multi-select"
    categories="{"task.input.labels | to_json | escape }"
    header="Please identify all classes in the below text"
  >
    <classification-target style="white-space: pre-wrap">
      {{ task.input.taskObject }}
    </classification-target>
  </full-instructions header="Classifier instructions">  
    <ol>
      <li><strong>Read</strong> the text carefully.</li>
      <li><strong>Read</strong> the examples to understand more about the options.</li>
      <li><strong>Choose</strong> the appropriate labels that best suit the text.</li>
    </ol>
  </full-instructions>
  <short-instructions>
    <p>Enter description of the labels that workers have to choose from</p>
    <p>Add examples to help workers understand the label</p>
  </short-instructions>
</crowd-form>
```
Multi-label Text Classification Output Data

Once you have created a multi-label text classification labeling job, your output data will be located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the Output dataset location field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see Output Data.

To see an example of output manifest files for multi-label text classification labeling job, see Multi-label Classification Job Output.

Label Videos and Video Frames

You can use Ground Truth to classify videos and annotate video frames (still images extracted from videos) using one of the three built-in video task types. These task types streamline the process of creating video and video frame labeling jobs using the Amazon SageMaker console, API, and language-specific SDKs.

- Video clip classification – Enable workers to classify videos into categories you specify. For example, you can use this task type to have workers categorize videos into topics like sports, comedy, music, and education. To learn more, see Video Classification.

- Video frame labeling jobs – Enable workers to annotate video frames extracted from a video using bounding boxes, polylines, polygons or keypoint annotation tools. Ground Truth offers two built-in task types to label video frames:
  - Video frame object detection: Enable workers to identify and locate objects in video frames.
  - Video frame object tracking: Enable workers to track the movement of objects across video frames.
  - Video frame adjustment jobs: Have workers adjust labels, label category attributes, and frame attributes from a previous video frame object detection or object tracking labeling job.
  - Video frame verification jobs: Have workers verify labels, label category attributes, and frame attributes from a previous video frame object detection or object tracking labeling job.
If you have video files, you can use the Ground Truth automatic frame extraction tool to extract video frames from your videos. To learn more, see Video Frame Input Data.

Tip

To learn more about supported file types and input data quotas, see Input Data.

Topics

- Video Classification
- Label Video Frames
- Worker Instructions

Video Classification

Use an Amazon SageMaker Ground Truth video classification labeling task when you need workers to classify videos using predefined labels that you specify. Workers are shown videos and are asked to choose one label for each video.

You create a video classification labeling job using the Ground Truth section of the Amazon SageMaker console or the CreateLabelingJob operation.

Your video files must be encoded in a format that is supported by the browser used by the work team that labels your data. It is recommended that you verify that all video file formats in your input manifest file display correctly using the worker UI preview. You can communicate supported browsers to your workers using worker instructions. To see supported file formats, see Supported Data Formats.

Important

For this task type, if you create your own manifest file, use "source-ref" to identify the location of each video file in Amazon S3 that you want labeled. For more information, see Input Data.
Create a Video Classification Labeling Job (Console)

You can follow the instructions in Create a Labeling Job (Console) to learn how to create a video classification labeling job in the SageMaker console. In step 10, choose Video from the Task category dropdown list, and choose Video Classification as the task type.

Ground Truth provides a worker UI similar to the following for labeling tasks. When you create a labeling job in the console, you specify instructions to help workers complete the job and labels from which workers can choose.

Create a Video Classification Labeling Job (API)

This section covers details you need to know when you create a labeling job using the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Follow the instructions on Create a Labeling Job (API) and do the following while you configure your request:

- Use a pre-annotation Lambda function that ends with PRE-VideoClassification. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn.
• Use an annotation-consolidation Lambda function that ends with ACS-VideoClassification. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.

The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region.

```python
response = client.create_labeling_job(
    LabelingJobName='example-video-classification-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                'FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*',
    LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:region::workteam/private-crowd/*',
        'UiConfig': {
            'UiTemplateS3Uri': 's3://bucket/path/worker-task-template.html'
        },
        'PreHumanTaskLambdaArn': 'arn:aws:lambda:us-east-1:432418664414:function:PRE-VideoClassification',
        'TaskKeywords': [
            'Video Classification',
        ],
    },
)
'TaskTitle': 'Video classification task',
'TaskDescription': 'Select a label to classify this video',
'NumberOfHumanWorkersPerDataObject': 123,
'TaskTimeLimitInSeconds': 123,
'TaskAvailabilityLifetimeInSeconds': 123,
'MaxConcurrentTaskCount': 123,
'AnnotationConsolidationConfig': {
    'AnnotationConsolidationLambdaArn': 'arn:aws:lambda:us-east-1:432418664414:function:ACS-VideoClassification',
},
'Tags':[
    {
        'Key': 'string',
        'Value': 'string'
    },
]

Provide a Template for Video Classification

If you create a labeling job using the API, you must supply a worker task template in UiTemplateS3Uri. Copy and modify the following template by modifying the short-instructions, full-instructions, and header. Upload this template to Amazon S3, and provide the Amazon S3 URI to this file in UiTemplateS3Uri.

<crowd-form>
    <crowd-classifier
        name="crowd-classifier"
        categories="{{ task.input.labels | to_json | escape }}"
        header="Please classify video"
    >
    <classification-target>
        <video width="100%" controls/>
        <source src="{{ task.input.taskObject | grant_read_access }}" type="video/mp4"/>
        <source src="{{ task.input.taskObject | grant_read_access }}" type="video/webm"/>
        <source src="{{ task.input.taskObject | grant_read_access }}" type="video/ogg"/>
    </classification-target>
</crowd-form>
<full-instructions header="Video classification instructions">
  <ol>
    <li><strong>Read</strong> the task carefully and inspect the video.</li>
    <li><strong>Read</strong> the options and review the examples provided to understand more about the labels.</li>
    <li><strong>Choose</strong> the appropriate label that best suits the video.</li>
  </ol>
</full-instructions>

<short-instructions>
  <h3><span style="color: rgb(0, 138, 0);">Good example</span></h3>
  <p>Enter description to explain the correct label to the workers</p>
  <p><img src="https://d7evko5405gb7.cloudfront.net/fe4fed9b-660c-4477-9294-2c66a15d6bbe/src/images/quick-instructions-example-placeholder.png" style="max-width:100%"></p>
  <h3><span style="color: rgb(230, 0, 0);">Bad example</span></h3>
  <p>Enter description of an incorrect label</p>
  <p><img src="https://d7evko5405gb7.cloudfront.net/fe4fed9b-660c-4477-9294-2c66a15d6bbe/src/images/quick-instructions-example-placeholder.png" style="max-width:100%"></p>
</short-instructions>

Video Classification Output Data

Once you have created a video classification labeling job, your output data is located in the Amazon S3 bucket specified in the S3OutputPath parameter when using the API or in the Output dataset location field of the Job overview section of the console.

To learn more about the output manifest file generated by Ground Truth and the file structure the Ground Truth uses to store your output data, see Output Data.

To see an example of output manifest files for video classification labeling jobs, see Classification Job Output.
Label Video Frames

You can use Ground Truth built-in video frame task types to have workers annotate video frames using bounding boxes, polylines, polygons or keypoints. A video frame is a sequence of images that have been extracted from a video.

If you do not have video frames, you can provide video files (MP4 files) and use the Ground Truth automated frame extraction tool to extract video frames. To learn more, see Provide Video Files.

You can use the following built-in video task types to create video frame labeling jobs using the Amazon SageMaker console, API, and language-specific SDKs.

- **Video frame object detection** – Use this task type when you want workers to identify and locate objects in sequences of video frames. You provide a list of categories, and workers can select one category at a time and annotate objects which the category applies to in all frames. For example, you can use this task to ask workers to identify and localize various objects in a scene, such as cars, bikes, and pedestrians.

- **Video frame object tracking** – Use this task type when you want workers to track the movement of instances of objects across sequences of video frames. When a worker adds an annotation to a single frame, that annotation is associated with a unique instance ID. The worker adds annotations associated with the same ID in all other frames to identify the same object or person. For example, a worker can track the movement of a vehicle across a sequences of video frames by drawing bounding boxes associated with the same ID around the vehicle in each frame that it appears.

Use the following topics to learn more about these built-in task types and to how to create a labeling job using each task type. See Task Types to learn more about the annotations tools (bounding boxes, polylines, polygons and keypoints) available for these task types.

Before you create a labeling job, we recommend that you review Video Frame Labeling Job Overview.

Topics

- Video Frame Object Detection
- Video Frame Object Tracking
- Video Frame Labeling Job Overview
Video Frame Object Detection

You can use the video frame object detection task type to have workers identify and locate objects in a sequence of video frames (images extracted from a video) using bounding boxes, polylines, polygons or keypoint annotation tools. The tool you choose defines the video frame task type you create. For example, you can use a bounding box video frame object detection task type workers to identify and localize various objects in a series of video frames, such as cars, bikes, and pedestrians.

You can create a video frame object detection labeling job using the Amazon SageMaker Ground Truth console, the SageMaker API, and language-specific AWS SDKs. To learn more, see Create a Video Frame Object Detection Labeling Job and select your preferred method. See Task Types to learn more about the annotations tools you can choose from when you create a labeling job.

Ground Truth provides a worker UI and tools to complete your labeling job tasks: Preview the Worker UI.

You can create a job to adjust annotations created in a video object detection labeling job using the video object detection adjustment task type. To learn more, see Create Video Frame Object Detection Adjustment or Verification Labeling Job.

Preview the Worker UI

Ground Truth provides workers with a web user interface (UI) to complete your video frame object detection annotation tasks. You can preview and interact with the worker UI when you create a labeling job in the console. If you are a new user, we recommend that you create a labeling job through the console using a small input dataset to preview the worker UI and ensure your video frames, labels, and label attributes appear as expected.

The UI provides workers with the following assistive labeling tools to complete your object detection tasks:

- For all tasks, workers can use the Copy to next and Copy to all features to copy an annotation to the next frame or to all subsequent frames respectively.
- For tasks that include the bounding box tools, workers can use a Predict next feature to draw a bounding box in a single frame, and then have Ground Truth predict the location of boxes with the same label in all other frames. Workers can then make adjustments to correct predicted box locations.
Create a Video Frame Object Detection Labeling Job

You can create a video frame object detection labeling job using the SageMaker console or the `CreateLabelingJob` API operation.

This section assumes that you have reviewed the Video Frame Labeling Job Overview and have chosen the type of input data and the input dataset connection you are using.

Create a Labeling Job (Console)

You can follow the instructions in Create a Labeling Job (Console) to learn how to create a video frame object tracking job in the SageMaker console. In step 10, choose Video - Object detection from the Task category dropdown list. Select the task type you want by selecting one of the cards in Task selection.
Create a Labeling Job (API)

You create an object detection labeling job using the SageMaker API operation `CreateLabelingJob`. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of `CreateLabelingJob`. 
Create a Labeling Job (API) provides an overview of the CreateLabelingJob operation. Follow these instructions and do the following while you configure your request:

- You must enter an ARN for HumanTaskUiArn. Use `arn:aws:sagemaker:<region>:394669845002:human-task-ui/VideoObjectDetection`. Replace `<region>` with the AWS Region in which you are creating the labeling job.

  Do not include an entry for the UiTemplateS3Uri parameter.

- Your LabelAttributeName must end in -ref. For example, `video-od-labels-ref`.

- Your input manifest file must be a video frame sequence manifest file. You can create this manifest file using the SageMaker console, or create it manually and upload it to Amazon S3. For more information, see Input Data Setup.

- You can only use private or vendor work teams to create video frame object detection labeling jobs.

- You specify your labels, label category and frame attributes, the task type, and worker instructions in a label category configuration file. Specify the task type (bounding boxes, polylines, polygons or keypoint) using annotationType in your label category configuration file. For more information, see Create a Labeling Category Configuration File with Label Category and Frame Attributes to learn how to create this file.

- You need to provide pre-defined ARNs for the pre-annotation and post-annotation (ACS) Lambda functions. These ARNs are specific to the AWS Region you use to create your labeling job.

  - To find the pre-annotation Lambda ARN, refer to PreHumanTaskLambdaArn. Use the Region in which you are creating your labeling job to find the correct ARN that ends with PRE-VideoObjectDetection.

  - To find the post-annotation Lambda ARN, refer to AnnotationConsolidationLambdaArn. Use the Region in which you are creating your labeling job to find the correct ARN that ends with ACS-VideoObjectDetection.

- The number of workers specified in NumberOfHumanWorkersPerDataObject must be 1.

- Automated data labeling is not supported for video frame labeling jobs. Do not specify values for parameters in LabelingJobAlgorithmsConfig.

- Video frame object tracking labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs in TaskTimeLimitInSeconds (up to 7 days, or 604,800 seconds).
The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region.

```python
response = client.create_labeling_job(
    LabelingJobName='example-video-od-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://DOC-EXAMPLE-BUCKET/path/video-frame-sequence-input-manifest.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': ['FreeOfPersonallyIdentifiableInformation', 'FreeOfAdultContent'],
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://DOC-EXAMPLE-BUCKET/prefix/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*',
    LabelCategoryConfigS3Uri='s3://bucket/prefix/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:us-east-1:*:workteam/private-crowd/*',
        'UiConfig': {
        },
        'TaskKeywords': ['Video Frame Object Detection'],
        'TaskTitle': 'Video frame object detection task',
        'TaskDescription': 'Classify and identify the location of objects and people in video frames',
```
Create Video Frame Object Detection Adjustment or Verification Labeling Job

You can create an adjustment and verification labeling job using the Ground Truth console or CreateLabelingJob API. To learn more about adjustment and verification labeling jobs, and to learn how to create one, see Verify and Adjust Labels.

Output Data Format

When you create a video frame object detection labeling job, tasks are sent to workers. When these workers complete their tasks, labels are written to the Amazon S3 output location you specified when you created the labeling job. To learn about the video frame object detection output data format, see Video Frame Object Detection Output. If you are a new user of Ground Truth, see Output Data to learn more about the Ground Truth output data format.

Video Frame Object Tracking

You can use the video frame object tracking task type to have workers track the movement of objects in a sequence of video frames (images extracted from a video) using bounding boxes, polylines, polygons or keypoint annotation tools. The tool you choose defines the video frame task type you create. For example, you can use a bounding box video frame object tracking task type to ask workers to track the movement of objects, such as cars, bikes, and pedestrians by drawing boxes around them.

You provide a list of categories, and each annotation that a worker adds to a video frame is identified as an instance of that category using an instance ID. For example, if you provide the label category car, the first car that a worker annotates will have the instance ID car:1. The second car
the worker annotates will have the instance ID car:2. To track an object's movement, the worker adds annotations associated with the same instance ID around to object in all frames.

You can create a video frame object tracking labeling job using the Amazon SageMaker Ground Truth console, the SageMaker API, and language-specific AWS SDKs. To learn more, see Create a Video Frame Object Detection Labeling Job and select your preferred method. See Task Types to learn more about the annotations tools you can choose from when you create a labeling job.

Ground Truth provides a worker UI and tools to complete your labeling job tasks: Preview the Worker UI.

You can create a job to adjust annotations created in a video object detection labeling job using the video object detection adjustment task type. To learn more, see Create Video Frame Object Detection Adjustment or Verification Labeling Job.

Preview the Worker UI

Ground Truth provides workers with a web user interface (UI) to complete your video frame object tracking annotation tasks. You can preview and interact with the worker UI when you create a labeling job in the console. If you are a new user, we recommend that you create a labeling job through the console using a small input dataset to preview the worker UI and ensure your video frames, labels, and label attributes appear as expected.

The UI provides workers with the following assistive labeling tools to complete your object tracking tasks:

- For all tasks, workers can use the Copy to next and Copy to all features to copy an annotation with the same unique ID to the next frame or to all subsequent frames respectively.
- For tasks that include the bounding box tools, workers can use a Predict next feature to draw a bounding box in a single frame, and then have Ground Truth predict the location of boxes with the same unique ID in all other frames. Workers can then make adjustments to correct predicted box locations.

Create a Video Frame Object Tracking Labeling Job

You can create a video frame object tracking labeling job using the SageMaker console or the CreateLabelingJob API operation.

This section assumes that you have reviewed the Video Frame Labeling Job Overview and have chosen the type of input data and the input dataset connection you are using.
Create a Labeling Job (Console)

You can follow the instructions in Create a Labeling Job (Console) to learn how to create a video frame object tracking job in the SageMaker console. In step 10, choose Video - Object tracking from the Task category dropdown list. Select the task type you want by selecting one of the cards in Task selection.
Create a Labeling Job (API)

You create an object tracking labeling job using the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Create a Labeling Job (API) provides an overview of the CreateLabelingJob operation. Follow these instructions and do the following while you configure your request:

- You must enter an ARN for HumanTaskUiArn. Use arn:aws:sagemaker:<region>:394669845002:human-task-ui/VideoObjectTracking. Replace <region> with the AWS Region in which you are creating the labeling job.

  Do not include an entry for the UiTemplateS3Uri parameter.

- Your LabelAttributeName must end in -ref. For example, ot-labels-ref.

- Your input manifest file must be a video frame sequence manifest file. You can create this manifest file using the SageMaker console, or create it manually and upload it to Amazon S3. For more information, see Input Data Setup. If you create a streaming labeling job, the input manifest file is optional.

- You can only use private or vendor work teams to create video frame object detection labeling jobs.

- You specify your labels, label category and frame attributes, the task type, and worker instructions in a label category configuration file. Specify the task type (bounding boxes, polylines, polygons or keypoint) using annotationType in your label category configuration file. For more information, see Create a Labeling Category Configuration File with Label Category and Frame Attributes to learn how to create this file.

- You need to provide pre-defined ARNs for the pre-annotation and post-annotation (ACS) Lambda functions. These ARNs are specific to the AWS Region you use to create your labeling job.

  To find the pre-annotation Lambda ARN, refer to PreHumanTaskLambdaArn. Use the Region in which you are creating your labeling job to find the correct ARN that ends with PRE-VideoObjectTracking.

  To find the post-annotation Lambda ARN, refer to AnnotationConsolidationLambdaArn. Use the Region in which you are creating your labeling job to find the correct ARN that ends with ACS-VideoObjectTracking.
• The number of workers specified in NumberOfHumanWorkersPerDataObject must be 1.

• Automated data labeling is not supported for video frame labeling jobs. Do not specify values for parameters in LabelingJobAlgorithmsConfig.

• Video frame object tracking labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs in TaskTimeLimitInSeconds (up to 7 days, or 604,800 seconds).

The following is an example of an AWS Python SDK (Boto3) request to create a labeling job in the US East (N. Virginia) Region.

```python
response = client.create_labeling_job(
    LabelingJobName='example-video-ot-labeling-job,
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://DOC-EXAMPLE-BUCKET/path/video-frame-sequence-input-manifest.json'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': ['FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': 's3://DOC-EXAMPLE-BUCKET/prefix/file-to-store-output-data',
        'KmsKeyId': 'string'
    },
    RoleArn='arn:aws:iam::*:role/*,
    LabelCategoryConfigS3Uri='s3://bucket/prefix/label-categories.json',
    StoppingConditions={
        'MaxHumanLabeledObjectCount': 123,
        'MaxPercentageOfInputDatasetLabeled': 123
    },
    HumanTaskConfig={
        'WorkteamArn': 'arn:aws:sagemaker:us-east-1:*:workteam/private-crowd/*',
        'UiConfig': {
            'HumanTaskUiArn': 'arn:aws:sagemaker:us-east-1:394669845002:human-task-ui/VideoObjectTracking'
        }
    }
)
Create a Video Frame Object Tracking Adjustment or Verification Labeling Job

You can create an adjustment and verification labeling job using the Ground Truth console or CreateLabelingJob API. To learn more about adjustment and verification labeling jobs, and to learn how create one, see Verify and Adjust Labels.

Output Data Format

When you create a video frame object tracking labeling job, tasks are sent to workers. When these workers complete their tasks, labels are written to the Amazon S3 output location you specified when you created the labeling job. To learn about the video frame object tracking output data format, see Video Frame Object Tracking Output. If you are a new user of Ground Truth, see Output Data to learn more about the Ground Truth output data format.

Video Frame Labeling Job Overview

Use this page to learn about the object detection and object tracking video frame labeling jobs. The information on this page applies to both of these built-in task types.
The video frame labeling job is unique because of the following:

- You can either provide data objects that are ready to be annotated (video frames), or you can provide video files and have Ground Truth automatically extract video frames.
- Workers have the ability to save work as they go.
- You cannot use the Amazon Mechanical Turk workforce to complete your labeling tasks.
- Ground Truth provides a worker UI, as well as assistive and basic labeling tools, to help workers complete your tasks. You do not need to provide a worker task template.

Use the following topics to learn more.

Topics

- Input Data
- Job Completion Times
- Task Types
- Workforces
- Worker User Interface (UI)
- Video Frame Job Permission Requirements

Input Data

The video frame labeling job uses sequences of video frames. A single sequence is a series of images that have been extracted from a single video. You can either provide your own sequences of video frames, or have Ground Truth automatically extract video frame sequences from your video files. To learn more, see Provide Video Files.

Ground Truth uses sequence files to identify all images in a single sequence. All of the sequences that you want to include in a single labeling job are identified in an input manifest file. Each sequence is used to create a single worker task. You can automatically create sequence files and an input manifest file using Ground Truth automatic data setup. To learn more, see Automated Video Frame Input Data Setup.

To learn how to manually create sequence files and an input manifest file, see Create a Video Frame Input Manifest File.
Job Completion Times

Video and video frame labeling jobs can take workers hours to complete. You can set the total amount of time that workers can work on each task when you create a labeling job. The maximum time you can set for workers to work on tasks is 7 days. The default value is 3 days.

We strongly recommend that you create tasks that workers can complete within 12 hours. Workers must keep the worker UI open while working on a task. They can save work as they go and Ground Truth saves their work every 15 minutes.

When using the SageMaker CreateLabelingJob API operation, set the total time a task is available to workers in the TaskTimeLimitInSeconds parameter of HumanTaskConfig.

When you create a labeling job in the console, you can specify this time limit when you select your workforce type and your work team.

Task Types

When you create a video object tracking or video object detection labeling job, you specify the type of annotation that you want workers to create while working on your labeling task. The annotation type determines the type of output data Ground Truth returns and defines the task type for your labeling job.

If you are creating a labeling job using the API operation CreateLabelingJob, you specify the task type using the label category configuration file parameter annotationType. To learn more, see Create a Labeling Category Configuration File with Label Category and Frame Attributes.

The following task types are available for both video object tracking or video object detection labeling jobs:

- **Bounding box** – Workers are provided with tools to create bounding box annotations. A bounding box is a box that a worker draws around an object to identify the pixel-location and label of that object in the frame.

- **Polyline** – Workers are provided with tools to create polyline annotations. A polyline is defined by the series of ordered x, y coordinates. Each point added to the polyline is connected to the previous point by a line. The polyline does not have to be closed (the start point and end point do not have to be the same) and there are no restrictions on the angles formed between lines.

- **Polygon** – Workers are provided with tools to create polygon annotations. A polygon is a closed shape defined by a series of ordered x, y coordinates. Each point added to the polygon
is connected to the previous point by a line and there are no restrictions on the angles formed between lines. Two lines (sides) of the polygon cannot cross. The start and end point of a polygon must be the same.

- **Keypoint** – Workers are provided with tools to create keypoint annotations. A keypoint is a single point associated with an x, y coordinate in the video frame.

**Workforces**

When you create a video frame labeling job, you need to specify a work team to complete your annotation tasks. You can choose a work team from a private workforce of your own workers, or from a vendor workforce that you select in the AWS Marketplace. You cannot use the Amazon Mechanical Turk workforce for video frame labeling jobs.

To learn more about vendor workforces, see [Managing Vendor Workforces](#).

To learn how to create and manage a private workforce, see [Use a Private Workforce](#).

**Worker User Interface (UI)**

Ground Truth provides a worker user interface (UI), tools, and assistive labeling features to help workers complete your video labeling tasks. You can preview the worker UI when you create a labeling job in the console.

When you create a labeling job using the API operation `CreateLabelingJob`, you must provide an ARN provided by Ground Truth in the parameter `HumanTaskUiArn` to specify the worker UI for your task type. You can use `HumanTaskUiArn` with the SageMaker `RenderUiTemplate` API operation to preview the worker UI.

You provide worker instructions, labels, and optionally, attributes that workers can use to provide more information about labels and video frames. These attributes are referred to as label category attributes and frame attributes respectively. They are all displayed in the worker UI.

**Label Category and Frame Attributes**

When you create a video object tracking or video object detection labeling job, you can add one or more **label category attributes** and **frame attributes**:

- **Label category attribute** – A list of options (strings), a free form text box, or a numeric field associated with one or more labels. It is used by workers to provide metadata about a label.
• **Frame attribute** – A list of options (strings), a free form text box, or a numeric field that appears on each video frame a worker is sent to annotate. It is used by workers to provide metadata about video frames.

Additionally, you can use label and frame attributes to have workers verify labels in a video frame label verification job.

Use the following sections to learn more about these attributes. To learn how to add label category and frame attributes to a labeling job, use the **Create Labeling Job** sections on the task type page of your choice.

**Label Category Attributes**

Add label category attributes to labels to give workers the ability to provide more information about the annotations they create. A label category attribute is added to an individual label, or to all labels. When a label category attribute is applied to all labels it is referred to as a *global label category attribute*.

For example, if you add the label category *car*, you might also want to capture additional data about your labeled cars, such as if they are occluded or the size of the car. You can capture this metadata using label category attributes. In this example, if you added the attribute *occluded* to the car label category, you can assign *partial*, *completely*, *no* to the *occluded* attribute and enable workers to select one of these options.

When you create a label verification job, you add labels category attributes to each label you want workers to verify.

**Frame level Attributes**

Add frame attributes to give workers the ability to provide more information about individual video frames. Each frame attribute you add appears on all frames.

For example, you can add a number-frame attribute to have workers identify the number of objects they see in a particular frame.

In another example, you may want to provide a free-form text box to give workers the ability to provide an answer to a question.

When you create a label verification job, you can add one or more frame attributes to ask workers to provide feedback on all labels in a video frame.
Worker Instructions

You can provide worker instructions to help your workers complete your video frame labeling tasks. You might want to cover the following topics when writing your instructions:

- Best practices and things to avoid when annotating objects.
- The label category attributes provided (for object detection and object tracking tasks) and how to use them.
- How to save time while labeling by using keyboard shortcuts.

You can add your worker instructions using the SageMaker console while creating a labeling job. If you create a labeling job using the API operation CreateLabelingJob, you specify worker instructions in your label category configuration file.

In addition to your instructions, Ground Truth provides a link to help workers navigate and use the worker portal. View these instructions by selecting the task type on Worker Instructions.

Declining Tasks

Workers are able to decline tasks.

Workers decline a task if the instructions are not clear, input data is not displaying correctly, or if they encounter some other issue with the task. If the number of workers per dataset object (NumberOfHumanWorkersPerDataObject) decline the task, the data object is marked as expired and will not be sent to additional workers.

Video Frame Job Permission Requirements

When you create a video frame labeling job, in addition to the permission requirements found in Assign IAM Permissions to Use Ground Truth, you must add a CORS policy to your S3 bucket that contains your input manifest file.

Add a CORS Permission Policy to S3 Bucket

When you create a video frame labeling job, you specify buckets in S3 where your input data and manifest file are located and where your output data will be stored. These buckets may be the same. You must attach the following Cross-origin resource sharing (CORS) policy to your input and output buckets. If you use the Amazon S3 console to add the policy to your bucket, you must use the JSON format.
To learn how to add a CORS policy to an S3 bucket, see [How do I add cross-domain resource sharing with CORS?](https://docs.aws.amazon.com/AmazonS3/latest/userguide/CORS_index.html) in the Amazon Simple Storage Service User Guide.
Worker Instructions

This topic provides an overview of the Ground Truth worker portal and the tools available to complete your video frame labeling task. First, select the type of task you are working on from Topics.

⚠️ Important
It is recommended that you complete your task using a Google Chrome or Firefox web browser.

For adjustment jobs, select the original labeling job task type that produced the labels you are adjusting. Review and adjust the labels in your task as needed.

Topics
- Work on Video Frame Object Tracking Tasks
- Work on Video Frame Object Detection Tasks

Work on Video Frame Object Tracking Tasks

Video frame object tracking tasks require you to track the movement of objects across video frames. A video frame is a still image from a video scene.

You can use the worker UI to navigate between video frames and use the tools provided to identify unique objects and track their movement from one from to the next. Use this page to learn how to navigate your worker UI, use the tools provided, and complete your task.

It is recommended that you complete your task using a Google Chrome or Firefox web browser.

⚠️ Important
If you see annotations have already been added to one or more video frames when you open your task, adjust those annotations and add additional annotations as needed.

Topics
- Your Task
When you work on a video frame object tracking task, you need to select a category from the **Label category** menu on the right side of your worker portal to start annotating. After you've chosen a category, use the tools provided to annotate the objects that the category applies to. This annotation will be associated with a unique label ID that should only be used for that object. Use this same label ID to create additional annotations for the same object in all of the video frames that it appears in. Refer to **Tool Guide** to learn more about the tools provided.

After you've added a label, you may see a downward pointing arrow next to the label in the **Labels** menu. Select this arrow and then select one option for each label attribute you see to provide more information about that label.

You may see frame attributes under the **Labels** menu. These attributes will appear on each frame in your task. Use these attribute prompts to enter additional information about each frame.
After you've added a label, you can quickly add and edit a label category attribute value by using the downward pointing arrow next to the label in the **Labels** menu. If you select the pencil icon next to the label in the **Labels** menu, the **Edit instance** menu will appear. You can edit the label ID, label category, and label category attributes using this menu.

To edit an annotation, select the label of the annotation that you want to edit in the **Labels** menu or select the annotation in the frame. When you edit or delete an annotation, the action will only modify the annotation in a single frame.

If you are working on a task that includes a bounding box tool, use the predict next icon to predict the location of all bounding boxes that you have drawn in a frame in the next frame. If you select a single box and then select the predict next icon, only that box will be predicted in the next frame. If you have not added any boxes to the current frame, you will receive an error. You must add at least one box to the frame before using this feature.

After you've used the predict next icon, review the location of each box in the next frame and make adjustments to the box location and size if necessary.
For all other tools, you can use the **Copy to next** and **Copy to all** tools to copy your annotations to the next or all frames respectively.

**Navigate the UI**

You can navigate between video frames using the navigation bar in the bottom-left corner of your UI.

Use the play button to automatically move through the entire sequence of frames.

Use the next frame and previous frame buttons to move forward or back one frame at a time. You can also input a frame number to navigate to that frame.

You can zoom in to and out of all video frames. Once you have zoomed into a video frame, you can move around in that frame using the move icon. When you set a new view in a single video frame by zooming and moving within that frame, all video frames are set to the same view. You can reset all video frames to their original view using the fit screen icon. For additional view options, see Icons Guide.

When you are in the worker UI, you see the following menus:

- **Instructions** – Review these instructions before starting your task. Additionally, select **More instructions** and review these instructions.
- **Shortcuts** – Use this menu to view keyboard shortcuts that you can use to navigate video frames and use the tools provided.
- **Help** – Use this option to refer back to this documentation.

**Bulk Edit Label and Frame Attributes**

You can bulk edit label attributes and frame attributes (attributes).

When you bulk edit an attribute, you specify one or more ranges of frames that you want to apply the edit to. The attribute you select is edited in all frames in that range, including the start and end frames you specify. When you bulk edit label attributes, the range you specify **must** contain the label that the label attribute is attached to. If you specify frames that do not contain this label, you will receive an error.

To bulk edit an attribute you **must** specify the desired value for the attribute first. For example, if you want to change an attribute from Yes to No, you must select No, and then perform the bulk edit.
You can also specify a new value for an attribute that has not been filled in and then use the bulk edit feature to fill in that value in multiple frames. To do this, select the desired value for the attribute and complete the following procedure.

**To bulk edit a label or attribute:**

1. Use your mouse to right click the attribute you want to bulk edit.
2. Specify the range of frames you want to apply the bulk edit to using a dash (-) in the text box. For example, if you want to apply the edit to frames one through ten, enter 1-10. If you want to apply the edit to frames two to five, eight to ten and twenty enter 2-5, 8-10, 20.
3. Select **Confirm**.

If you get an error message, verify that you entered a valid range and that the label associated with the label attribute you are editing (if applicable) exists in all frames specified.

You can quickly add a label to all previous or subsequent frames using the **Duplicate to previous frames** and **Duplicate to next frames** options in the **Label** menu at the top of your screen.

**Tool Guide**

Your task will include one or more tools. The tool provided dictates the type of annotations you will create to identify and track objects. Use the following table to learn more about each tool provided.

<table>
<thead>
<tr>
<th>Tool</th>
<th>Icon</th>
<th>Action</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bounding box</td>
<td><img src="BoundingBoxIcon.png" alt="Icon" /></td>
<td>Add a bounding box annotation.</td>
<td>Choose this icon to add a bounding box. Each bounding box you add is associated with the category you choose from the Label category drop down menu. Select the bounding box or its associated label to adjust it.</td>
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<tr>
<td>Tool</td>
<td>Icon</td>
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</tr>
<tr>
<td>Bounding box</td>
<td><img src="image" alt="Icon" /></td>
<td>Predict bounding boxes in the next frame.</td>
<td>Select a bounding box, and then choose this icon to predict the location of that box in the next frame. You can select the icon multiple times in a row to automatically detect the location of box in multiple frames. For example, select this icon 5 times to predict the location of a bounding box in the next 5 frames.</td>
</tr>
<tr>
<td>Keypoint</td>
<td><img src="image" alt="Icon" /></td>
<td>Add a keypoint annotation.</td>
<td>Choose this icon to add a keypoint. Click on an object the image to place the keypoint at that location. Each keypoint you add is associated with the category you choose from the Label category drop down menu. Select a keypoint or its associated label to adjust it.</td>
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<td>Tool</td>
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<tr>
<td>Polyline</td>
<td><img src="image" alt="Polyline Icon" /></td>
<td>Add a polyline annotation.</td>
<td>Choose this icon to add a polyline. To add a polyline, continuously click around the object of interest to add new points. To stop drawing a polyline, select the last point that you placed a second time (this point will be green), or press <strong>Enter</strong> on your keyboard. Each point added to the polyline is connected to the previous point by a line. The polyline does not have to be closed (the start point and end point do not have to be the same) and there are no restrictions on the angles formed between lines. Each polyline you add is associated with the category you choose from the Label category drop down menu. Select</td>
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</table>
| Polygon  | ![Polygon Icon](image) | Add a polygon annotation.  | Choose this icon to add a polygon. To add a polygon, continuously click around the object of interest to add new points. To stop drawing the polygon, select the start point (this point will be green).  
A polygon is a closed shape defined by a series of points that you place. Each point added to the polygon is connected to the previous point by a line and there are no restrictions on the angles formed between lines. The start and end point must be the same.  
Each polygon you add is associated with the category you choose from the Label category drop down menu. Select the polygon or its associated label to adjust it. |
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</tr>
</thead>
<tbody>
<tr>
<td>Copy to Next</td>
<td><img src="image" alt="Copy to Next Icon" /></td>
<td>Copy annotations to the next frame.</td>
<td>If one or more annotations are selected in the current frame, those annotations are copied to the next frame. If no annotations are selected, all annotations in the current frame will be copied to the next frame.</td>
</tr>
<tr>
<td>Copy to All</td>
<td><img src="image" alt="Copy to All Icon" /></td>
<td>Copy annotations to all subsequent frames.</td>
<td>If one or more annotations are selected in the current frame, those annotations are copied to all subsequent frames. If no annotations are selected, all annotations in the current frame will be copied to all subsequent frames.</td>
</tr>
</tbody>
</table>

**Icons Guide**

Use this table to learn about the icons you see in your UI. You can automatically select some of these icons using the keyboard shortcuts found in the **Shortcuts** menu.
<table>
<thead>
<tr>
<th>Icon</th>
<th>Action</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td><img src="#" alt="Sun Icon" /></td>
<td>brightness</td>
<td>Choose this icon to adjust the brightness of all video frames.</td>
</tr>
<tr>
<td><img src="#" alt="Moon Icon" /></td>
<td>contrast</td>
<td>Choose this icon to adjust the contrast of all video frames.</td>
</tr>
<tr>
<td><img src="#" alt="Plus Icon" /></td>
<td>zoom in</td>
<td>Choose this icon to zoom into all of the video frames.</td>
</tr>
<tr>
<td><img src="#" alt="Minus Icon" /></td>
<td>zoom out</td>
<td>Choose this icon to zoom out of all of the video frames.</td>
</tr>
<tr>
<td><img src="#" alt="Move Icon" /></td>
<td>move screen</td>
<td>After you've zoomed into a video frame, choose this icon to move around in that video frame. You can move around the video frame using your mouse by clicking and dragging the frame in the direction you want it to move. This will change the view in all view frames.</td>
</tr>
<tr>
<td><img src="#" alt="Fit Screen Icon" /></td>
<td>fit screen</td>
<td>Reset all video frames to their original position.</td>
</tr>
<tr>
<td><img src="#" alt="Undo Icon" /></td>
<td>undo</td>
<td>Undo an action. You can use this icon to remove a bounding box that you just added, or to undo an adjustment you made to a bounding box.</td>
</tr>
<tr>
<td><img src="#" alt="Redo Icon" /></td>
<td>redo</td>
<td>Redo an action that was undone using the undo icon.</td>
</tr>
<tr>
<td><img src="#" alt="Delete Icon" /></td>
<td>delete label</td>
<td>Delete a label. This will delete the bounding box associated with the label in a single frame.</td>
</tr>
<tr>
<td><img src="#" alt="Show or Hide Label Icon" /></td>
<td>show or hide label</td>
<td>Select this icon to show a label that has been hidden. If this icon has a slash through it, select it to hide the label.</td>
</tr>
</tbody>
</table>
### Icon | Action | Description
--- | --- | ---
[Icon] | edit label | Select this icon to open the **Edit instance** menu. Use this menu to edit a label category, ID, and to add or edit label attributes.

### Shortcuts

The keyboard shortcuts listed in the **Shortcuts** menu can help you quickly select icons, undo and redo annotations, and use tools to add and edit annotations. For example, once you add a bounding box, you can use `P` to quickly predict the location of that box in subsequent frames.

Before you start your task, it is recommended that you review the **Shortcuts** menu and become acquainted with these commands.

### Release, Stop and Resume, and Decline Tasks

When you open the labeling task, three buttons on the top right allow you to decline the task (**Decline task**), release it (**Release task**), and stop and resume it at a later time (**Stop and resume later**). The following list describes what happens when you select one of these options:

- **Decline task**: You should only decline a task if something is wrong with the task, such as unclear video frame images or an issue with the UI. If you decline a task, you will not be able to return to the task.

- **Release Task**: Use this option to release a task and allow others to work on it. When you release a task, you lose all work done on that task and other workers on your team can pick it up. If enough workers pick up the task, you may not be able to return to it. When you select this button and then select **Confirm**, you are returned to the worker portal. If the task is still available, its status will be **Available**. If other workers pick it up, it will disappear from your portal.

- **Stop and resume later**: You can use the **Stop and resume later** button to stop working and return to the task at a later time. You should use the **Save** button to save your work before you select **Stop and resume later**. When you select this button and then select **Confirm**, you are returned to the worker portal, and the task status is **Stopped**. You can select the same task to resume work on it.

Be aware that the person that creates your labeling tasks specifies a time limit in which all tasks must be completed by. If you do not return to and complete this task within that time...
limit, it will expire and your work will not be submitted. Contact your administrator for more information.

**Saving Your Work and Submitting**

You should periodically save your work using the **Save** button. Ground Truth will automatically save your work every 15 minutes.

When you open a task, you must complete your work on it before pressing **Submit**.

**Work on Video Frame Object Detection Tasks**

Video frame object detection tasks required you to classify and identify the location of objects in video frames using annotations. A video frame is a still image from a video scene.

You can use the worker UI to navigate between video frames and create annotations to identify objects of interest. Use the sections on this page to learn how to navigate your worker UI, use the tools provided, and complete your task.

It is recommended that you complete your task using a Google Chrome web browser.

⚠️ **Important**

If you see annotations have already been added to one or more video frames when you open your task, adjust those annotations and add additional annotations as needed.

**Topics**

- **Your Task**
- **Navigate the UI**
- **Bulk Edit Label and Frame Attributes**
- **Tool Guide**
- **UI Icon Guide**
- **Shortcuts**
- **Release, Stop and Resume, and Decline Tasks**
- **Saving Your Work and Submitting**
Your Task

When you work on a video frame object detection task, you need to select a category from the **Label category** menu on the right side of your worker portal to start annotating. After you've chosen a category, draw annotations around objects that this category applies to. To learn more about the tools you see in your worker UI, refer to the **Tool Guide**.

After you've added a label, you may see a downward pointing arrow next to the label in the **Labels** menu. Select this arrow and then select one option for each label attribute you see to provide more information about that label.

You may see frame attributes under the **Labels** menu. These attributes will appear on each frame in your task. Use these attribute prompts to enter additional information about each frame.

To edit an annotation, select the label of the annotation that you want to edit in the **Labels** menu or select the annotation in the frame. When you edit or delete an annotation, the action will only modify the annotation in a single frame.
If you are working on a task that includes a bounding box tool, use the predict next icon to predict the location of all bounding boxes that you have drawn in a frame in the next frame. If you select a single box and then select the predict next icon, only that box will be predicted in the next frame. If you have not added any boxes to the current frame, you will receive an error. You must add at least one box to the frame before using this feature.

Note

The predict next feature will not overwrite manually created annotations. It will only add annotations. If you use predict next and as a result have more than one bounding box around a single object, delete all but one box. Each object should only be identified with a single box.

After you've used the predict next icon, review the location of each box in the next frame and make adjustments to the box location and size if necessary.

For all other tools, you can use the Copy to next and Copy to all tools to copy your annotations to the next or all frames respectively.

Navigate the UI

You can navigate between video frames using the navigation bar in the bottom-left corner of your UI.

Use the play button to automatically play through multiple frames.

Use the next frame and previous frame buttons to move forward or back one frame at a time. You can also input a frame number to navigate to that frame.

You can zoom in to and out of all video frames. Once you have zoomed into a video frame, you can move around in that frame using the move icon. When you navigate to a new view in a single video frame by zooming and moving within that frame, all video frames are set to the same view. You can reset all video frames to their original view using the fit screen icon. To learn more, see UI Icon Guide.

When you are in the worker UI, you see the following menus:

- Instructions – Review these instructions before starting your task. Additionally, select More instructions and review these instructions.
• **Shortcuts** – Use this menu to view keyboard shortcuts that you can use to navigate video frames and use the annotation tools provided.

• **Help** – Use this option to refer back to this documentation.

If you

**Bulk Edit Label and Frame Attributes**

You can bulk edit label attributes and frame attributes (attributes).

When you bulk edit an attribute, you specify one or more ranges of frames that you want to apply the edit to. The attribute you select is edited in all frames in that range, including the start and end frames you specify. When you bulk edit label attributes, the range you specify must contain the label that the label attribute is attached to. If you specify frames that do not contain this label, you will receive an error.

To bulk edit an attribute you must specify the desired value for the attribute first. For example, if you want to change an attribute from *Yes* to *No*, you must select *No*, and then perform the bulk edit.

You can also specify a new value for an attribute that has not been filled in and then use the bulk edit feature to fill in that value in multiple frames. To do this, select the desired value for the attribute and complete the following procedure.

**To bulk edit a label or attribute:**

1. Use your mouse to right click the attribute you want to bulk edit.
2. Specify the range of frames you want to apply the bulk edit to using a dash (–) in the text box. For example, if you want to apply the edit to frames one through ten, enter 1-10. If you want to apply the edit to frames two to five, eight to ten and twenty enter 2-5, 8-10, 20.
3. Select **Confirm**.

If you get an error message, verify that you entered a valid range and that the label associated with the label attribute you are editing (if applicable) exists in all frames specified.

You can quickly add a label to all previous or subsequent frames using the **Duplicate to previous frames** and **Duplicate to next frames** options in the **Label** menu at the top of your screen.
Tool Guide

Your task will include one or more tools. The tool provided dictates the type of annotations you will create to identify and label objects. Use the following table to learn more about the tool or tools you may see in your worker UI.

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<thead>
<tr>
<th>Tool</th>
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<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bounding box</td>
<td><img src="image" alt="Icon" /></td>
<td>Add a bounding box annotation.</td>
<td>Choose this icon to add a bounding box. Each bounding box you add is associated with the category you choose from the Label category drop down menu. Select the bounding box or its associated label to adjust it.</td>
</tr>
<tr>
<td>Predict next</td>
<td><img src="image" alt="Icon" /></td>
<td>Predict bounding boxes in the next frame.</td>
<td>Select a bounding box, and then choose this icon to predict the location of that box in the next frame. You can select the icon multiple times in a row to automatically detect the location of box in multiple frames. For example, select this icon 5 times to predict the location of a bounding box in the next 5 frames.</td>
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<tr>
<td>Tool</td>
<td>Icon</td>
<td>Action</td>
<td>Description</td>
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<tr>
<td>Keypoint</td>
<td><img src="image" alt="Keypoint Icon" /></td>
<td>Add a keypoint annotation.</td>
<td>Choose this icon to add a keypoint. Click on an object the image to place the keypoint at that location. Each keypoint you add is associated with the category you choose from the Label category drop down menu. Select a keypoint or its associated label to adjust it.</td>
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<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Polyline</td>
<td><img src="image" alt="Polyline Icon" /></td>
<td>Add a polyline annotation.</td>
<td>Choose this icon to add a polyline. To add a polyline, continuously click around the object of interest to add new points. To stop drawing a polyline, select the last point that you placed a second time (this point will be green), or press <strong>Enter</strong> on your keyboard. Each point added to the polyline is connected to the previous point by a line. The polyline does not have to be closed (the start point and end point do not have to be the same) and there are no restrictions on the angles formed between lines. Each polyline you add is associated with the category you choose from the Label category drop down menu. Select</td>
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<td>the polyline or its associated label to adjust it.</td>
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</tbody>
</table>
| Polygon | ![Polygon Icon](image) | Add a polygon annotation. | Choose this icon to add a polygon. To add a polygon, continuously click around the object of interest to add new points. To stop drawing the polygon, select the start point (this point will be green).

A polygon is a closed shape defined by a series of points that you place. Each point added to the polygon is connected to the previous point by a line and there are no restrictions on the angles formed between lines. Two lines (sides) of the polygon cannot cross. A line will become red if it violates this condition. The start and end point must be the same.

Each polygon you add is associated with the category you choose from the
<table>
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<tbody>
<tr>
<td>Copy to Next</td>
<td><img src="image" alt="Copy to Next Icon" /></td>
<td>Copy annotations to the next frame.</td>
<td>If one or more annotations are selected in the current frame, those annotations are copied to the next frame. If no annotations are selected, all annotations in the current frame will be copied to the next frame.</td>
</tr>
<tr>
<td>Copy to All</td>
<td><img src="image" alt="Copy to All Icon" /></td>
<td>Copy annotations to all subsequent frames.</td>
<td>If one or more annotations are selected in the current frame, those annotations are copied to all subsequent frames. If no annotations are selected, all annotations in the current frame will be copied to all subsequent frames.</td>
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</table>
# UI Icon Guide

Use this table to learn about the icons you see in your worker task portal. You can automatically select these icons using the keyboard shortcuts found in the **Shortcuts** menu.

<table>
<thead>
<tr>
<th>Icon</th>
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</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="brightness" /></td>
<td>Choose this icon to adjust the brightness of all video frames.</td>
</tr>
<tr>
<td><img src="image" alt="contrast" /></td>
<td>Choose this icon to adjust the contrast of all video frames.</td>
</tr>
<tr>
<td><img src="image" alt="zoom in" /></td>
<td>Choose this icon to zoom into all of the video frames.</td>
</tr>
<tr>
<td><img src="image" alt="zoom out" /></td>
<td>Choose this icon to zoom out of all of the video frames.</td>
</tr>
<tr>
<td><img src="image" alt="move screen" /></td>
<td>After you've zoomed into a video frame, choose this icon to move around in that video frame. You can move around in the video frame using your mouse by clicking and dragging the frame in the direction you want it to move. This will change the view in all view frames.</td>
</tr>
<tr>
<td><img src="image" alt="fit screen" /></td>
<td>Reset all video frames to their original position.</td>
</tr>
<tr>
<td><img src="image" alt="undo" /></td>
<td>Undo an action. You can use this icon to remove a bounding box that you just added, or to undo an adjustment you made to a bounding box.</td>
</tr>
<tr>
<td><img src="image" alt="redo" /></td>
<td>Redo an action that was undone using the undo icon.</td>
</tr>
<tr>
<td>Icon</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
</tr>
<tr>
<td><img src="image" alt="delete label icon" /></td>
<td>Delete a label. This will delete the bounding box associated with the label in a single frame.</td>
</tr>
<tr>
<td><img src="image" alt="show or hide label icon" /></td>
<td>Select this icon to show a label that has been hidden. If this icon has a slash through it, select it to hide the label.</td>
</tr>
</tbody>
</table>

**Shortcuts**

The keyboard shortcuts listed in the **Shortcuts** menu can help you quickly select icons, undo and redo annotations, and use tools to add and edit annotations. For example, once you add a bounding box, you can use P to quickly predict the location of that box in subsequent frames.

Before you start your task, it is recommended that you review the **Shortcuts** menu and become acquainted with these commands.

**Release, Stop and Resume, and Decline Tasks**

When you open the labeling task, three buttons on the top right allow you to decline the task (**Decline task**), release it (**Release task**), and stop and resume it at a later time (**Stop and resume later**). The following list describes what happens when you select one of these options:

- **Decline task**: You should only decline a task if something is wrong with the task, such as unclear video frame images or an issue with the UI. If you decline a task, you will not be able to return to the task.

- **Release Task**: Use this option to release a task and allow others to work on it. When you release a task, you lose all work done on that task and other workers on your team can pick it up. If enough workers pick up the task, you may not be able to return to it. When you select this button and then select **Confirm**, you are returned to the worker portal. If the task is still available, its status will be **Available**. If other workers pick it up, it will disappear from your portal.

- **Stop and resume later**: You can use the **Stop and resume later** button to stop working and return to the task at a later time. You should use the **Save** button to save your work before you select **Stop and resume later**. When you select this button and then select **Confirm**, you are returned to the worker portal, and the task status is **Stopped**. You can select the same task to resume work on it.
Be aware that the person that creates your labeling tasks specifies a time limit in which all tasks must be completed by. If you do not return to and complete this task within that time limit, it will expire and your work will not be submitted. Contact your administrator for more information.

Saving Your Work and Submitting

You should periodically save your work. Ground Truth automatically saves your work every 15 minutes.

When you open a task, you must complete your work before pressing Submit.

Use Ground Truth to Label 3D Point Clouds

Create a 3D point cloud labeling job to have workers label objects in 3D point clouds generated from 3D sensors like Light Detection and Ranging (LiDAR) sensors and depth cameras, or generated from 3D reconstruction by stitching images captured by an agent like a drone.

3D Point Clouds

Point clouds are made up of three-dimensional (3D) visual data that consists of points. Each point is described using three coordinates, typically x, y, and z. To add color or variations in point intensity to the point cloud, points may be described with additional attributes, such as i for intensity or values for the red (r), green (g), and blue (b) 8-bit color channels. When you create a Ground Truth 3D point cloud labeling job, you can provide point cloud and, optionally, sensor fusion data.

The following image shows a single, 3D point cloud scene rendered by Ground Truth and displayed in the semantic segmentation worker UI.
LiDAR

A Light Detection and Ranging (LiDAR) sensor is a common type of sensor used to collect measurements that are used to generate point cloud data. LiDAR is a remote sensing method that uses light in the form of a pulsed laser to measure the distances of objects from the sensor. You can provide 3D point cloud data generated from a LiDAR sensor for a Ground Truth 3D point cloud labeling job using the raw data formats described in Accepted Raw 3D Data Formats.

Sensor Fusion

Ground Truth 3D point cloud labeling jobs include a sensor fusion feature that supports video camera sensor fusion for all task types. Some sensors come with multiple LiDAR devices and video cameras that capture images and associate them with a LiDAR frame. To help annotators visually complete your tasks with high confidence, you can use the Ground Truth sensor fusion feature to project annotations (labels) from a 3D point cloud to 2D camera images and vice versa using 3D scanner (such as LiDAR) extrinsic matrix and camera extrinsic and intrinsic matrices. To learn more, see Sensor Fusion.

Label 3D Point Clouds

Ground Truth provides a user interface (UI) and tools that workers use to label or annotate 3D point clouds. When you use the object detection or semantic segmentation task types, workers can
annotate a single point cloud frame. When you use object tracking, workers annotate a sequence of frames. You can use object tracking to track object movement across all frames in a sequence.

The following demonstrates how a worker would use the Ground Truth worker portal and tools to annotate a 3D point cloud for an object detection task. For similar visual examples of other task types, see 3D Point Cloud Task types.

Assistive Labeling Tools for Point Cloud Annotation

Ground Truth offers assistive labeling tools to help workers complete your point cloud annotation tasks faster and with more accuracy. For details about assistive labeling tools that are included in the worker UI for each task type, select a task type and refer to the View the Worker Task Interface section of that page.

Next Steps

You can create six types of tasks when you use Ground Truth 3D point cloud labeling jobs. Use the topics in 3D Point Cloud Task types to learn more about these task types and to learn how to create a labeling job using the task type of your choice.

The 3D point cloud labeling job is different from other Ground Truth labeling modalities. Before creating a labeling job, we recommend that you read 3D Point Cloud Labeling Jobs Overview. Additionally, review input data quotas in 3D Point Cloud and Video Frame Labeling Job Quotas.
For an end-to-end demo using the SageMaker API and AWS Python SDK (boto 3) to create a 3D point cloud labeling job, see `create-3D-pointcloud-labeling-job.ipynb` in the SageMaker Examples notebook tab.

⚠️ Important
If you use a notebook instance created before June 5th, 2020 to run this notebook, you must stop and restart that notebook instance for the notebook to work.

Topics

- **3D Point Cloud Task types**
- **3D Point Cloud Labeling Jobs Overview**
- **Worker Instructions**

### 3D Point Cloud Task types

You can use Ground Truth 3D point cloud labeling modality for a variety of use cases. The following list briefly describes each 3D point cloud task type. For additional details and instructions on how to create a labeling job using a specific task type, select the task type name to see its task type page.

- **3D point cloud object detection** – Use this task type when you want workers to locate and classify objects in a 3D point cloud by adding and fitting 3D cuboids around objects.

- **3D point cloud object tracking** – Use this task type when you want workers to add and fit 3D cuboids around objects to track their movement across a sequence of 3D point cloud frames. For example, you can use this task type to ask workers to track the movement of vehicles across multiple point cloud frames.

- **3D point cloud semantic segmentation** – Use this task type when you want workers to create a point-level semantic segmentation mask by painting objects in a 3D point cloud using different colors where each color is assigned to one of the classes you specify.

- 3D point cloud adjustment task types – Each of the task types above has an associated adjustment task type that you can use to audit and adjust annotations generated from a 3D point cloud labeling job. Refer to the task type page of the associated type to learn how to create an adjustment labeling job for that task.
3D Point Cloud Object Detection

Use this task type when you want workers to classify objects in a 3D point cloud by drawing 3D cuboids around objects. For example, you can use this task type to ask workers to identify different types of objects in a point cloud, such as cars, bikes, and pedestrians.

For this task type, the data object that workers label is a single point cloud frame. Ground Truth renders a 3D point cloud using point cloud data you provide. You can also provide camera data to give workers more visual information about scenes in the frame, and to help workers draw 3D cuboids around objects.

Ground Truth provides workers with tools to annotate objects with 9 degrees of freedom (x,y,z,rx,ry,rz,l,w,h) in three dimensions in both 3D scene and projected side views (top, side, and back). If you provide sensor fusion information (like camera data), when a worker adds a cuboid to identify an object in the 3D point cloud, the cuboid shows up and can be modified in the 2D images. After a cuboid has been added, all edits made to that cuboid in the 2D or 3D scene are projected into the other view.

You can create a job to adjust annotations created in a 3D point cloud object detection labeling job using the 3D point cloud object detection adjustment task type.

If you are a new user of the Ground Truth 3D point cloud labeling modality, we recommend you review 3D Point Cloud Labeling Jobs Overview. This labeling modality is different from other Ground Truth task types, and this page provides an overview of important details you should be aware of when creating a 3D point cloud labeling job.

Topics

- View the Worker Task Interface
- Create a 3D Point Cloud Object Detection Labeling Job
- Create a 3D Point Cloud Object Detection Adjustment or Verification Labeling Job
- Output Data Format

View the Worker Task Interface

Ground Truth provides workers with a web portal and tools to complete your 3D point cloud object detection annotation tasks. When you create the labeling job, you provide the Amazon Resource Name (ARN) for a pre-built Ground Truth worker UI in the HumanTaskUiArn parameter. When you
create a labeling job using this task type in the console, this worker UI is automatically used. You can preview and interact with the worker UI when you create a labeling job in the console. If you are a new user, it is recommended that you create a labeling job using the console to ensure your label attributes, point cloud frames, and if applicable, images, appear as expected.

The following is a GIF of the 3D point cloud object detection worker task interface. If you provide camera data for sensor fusion in the world coordinate system, images are matched up with scenes in the point cloud frame. These images appear in the worker portal as shown in the following GIF.

Worker can navigate in the 3D scene using their keyboard and mouse. They can:

- Double click on specific objects in the point cloud to zoom into them.
- Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
- Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

Once a worker places a cuboid in the 3D scene, a side-view will appear with the three projected side views: top, side, and back. These side-views show points in and around the placed cuboid and help workers refine cuboid boundaries in that area. Workers can zoom in and out of each of those side-views using their mouse.

The following video demonstrates movements around the 3D point cloud and in the side-view.
Additional view options and features are available in the **View** menu in the worker UI. See the [worker instruction page](#) for a comprehensive overview of the Worker UI.

### Assistive Labeling Tools

Ground Truth helps workers annotate 3D point clouds faster and more accurately using machine learning and computer vision powered assistive labeling tools for 3D point cloud object tracking tasks. The following assistive labeling tools are available for this task type:

- **Snapping** – Workers can add a cuboid around an object and use a keyboard shortcut or menu option to have Ground Truth's autofit tool snap the cuboid tightly around the object.

- **Set to ground** – After a worker adds a cuboid to the 3D scene, the worker can automatically snap the cuboid to the ground. For example, the worker can use this feature to snap a cuboid to the road or sidewalk in the scene.

- **Multi-view labeling** – After a worker adds a 3D cuboid to the 3D scene, a side panel displays front, side, and top perspectives to help the worker adjust the cuboid tightly around the object. In all of these views, the cuboid includes an arrow that indicates the orientation, or heading of the object. When the worker adjusts the cuboid, the adjustment will appear in real time on all of the views (that is, 3D, top, side, and front).
• **Sensor fusion** – If you provide data for sensor fusion, workers can adjust annotations in the 3D scenes and in 2D images, and the annotations will be projected into the other view in real time. Additionally, workers will have the option to view the direction the camera is facing and the camera frustum.

• **View options** – Enables workers to easily hide or view cuboids, label text, a ground mesh, and additional point attributes like color or intensity. Workers can also choose between perspective and orthogonal projections.

### Create a 3D Point Cloud Object Detection Labeling Job

You can create a 3D point cloud labeling job using the SageMaker console or API operation, `CreateLabelingJob`. To create a labeling job for this task type you need the following:

- A single-frame input manifest file. To learn how to create this type of manifest file, see [Create a Point Cloud Frame Input Manifest File](#). If you are a new user of Ground Truth 3D point cloud labeling modalities, you may also want to review [Accepted Raw 3D Data Formats](#).

- A work team from a private or vendor workforce. You cannot use Amazon Mechanical Turk for video frame labeling jobs. To learn how to create workforces and work teams, see [Create and Manage Workforces](#).

Additionally, make sure that you have reviewed and satisfied the [Assign IAM Permissions to Use Ground Truth](#).

Use one of the following sections to learn how to create a labeling job using the console or an API.

### Create a Labeling Job (Console)

You can follow the instructions [Create a Labeling Job (Console)](#) in order to learn how to create a 3D point cloud object detection labeling job in the SageMaker console. While you are creating your labeling job, be aware of the following:

- Your input manifest file must be a single-frame manifest file. For more information, see [Create a Point Cloud Frame Input Manifest File](#).

- Optionally, you can provide label category and frame attributes. Workers can assign one or more of these attributes to annotations to provide more information about that object. For example, you might want to use the attribute *occluded* to have workers identify when an object is partially obstructed.
• Automated data labeling and annotation consolidation are not supported for 3D point cloud labeling tasks.

• 3D point cloud object detection labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs when you select your work team (up to 7 days, or 604800 seconds).

Create a Labeling Job (API)

This section covers details you need to know when you create a labeling job using the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Create a Labeling Job (API), provides an overview of the CreateLabelingJob operation. Follow these instructions and do the following while you configure your request:

• You must enter an ARN for HumanTaskUiArn. Use arn:aws:sagemaker:<region>:394669845002:human-task-ui/PointCloudObjectDetection. Replace <region> with the AWS Region you are creating the labeling job in.

There should not be an entry for the UiTemplateS3Uri parameter.

• Your input manifest file must be a single-frame manifest file. For more information, see Create a Point Cloud Frame Input Manifest File.

• You specify your labels, label category and frame attributes, and worker instructions in a label category configuration file. To learn how to create this file, see Create a Labeling Category Configuration File with Label Category and Frame Attributes.

• You need to provide pre-defined ARNs for the pre-annotation and post-annotation (ACS) Lambda functions. These ARNs are specific to the AWS Region you use to create your labeling job.

• To find the pre-annotation Lambda ARN, refer to PreHumanTaskLambdaArn. Use the Region you are creating your labeling job in to find the correct ARN. For example, if you are creating your labeling job in us-east-1, the ARN will be arn:aws:lambda:us-east-1:432418664414:function:PRE-3DPointCloudObjectDetection.

• To find the post-annotation Lambda ARN, refer to AnnotationConsolidationLambdaArn. Use the Region you are creating your labeling job in to find the correct ARN. For example,
if you are creating your labeling job in us-east-1, the ARN will be arn:aws:lambda:us-east-1:432418664414:function:ACS-3DPointCloudObjectDetection.

- The number of workers specified in NumberOfHumanWorkersPerDataObject must be 1.
- Automated data labeling is not supported for 3D point cloud labeling jobs. You should not specify values for parameters in LabelingJobAlgorithmsConfig.
- 3D point cloud object detection labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs in TaskTimeLimitInSeconds (up to 7 days, or 604,800 seconds).

Create a 3D Point Cloud Object Detection Adjustment or Verification Labeling Job

You can create an adjustment or verification labeling job using the Ground Truth console or CreateLabelingJob API. To learn more about adjustment and verification labeling jobs, and to learn how create one, see Verify and Adjust Labels.

When you create an adjustment labeling job, your input data to the labeling job can include labels, and yaw, pitch, and roll measurements from a previous labeling job or external source. In the adjustment job, pitch, and roll will be visualized in the worker UI, but cannot be modified. Yaw is adjustable.

Ground Truth uses Tait-Bryan angles with the following intrinsic rotations to visualize yaw, pitch and roll in the worker UI. First, rotation is applied to the vehicle according to the z-axis (yaw). Next, the rotated vehicle is rotated according to the intrinsic y'-axis (pitch). Finally, the vehicle is rotated according to the intrinsic x''-axis (roll).

Output Data Format

When you create a 3D point cloud object detection labeling job, tasks are sent to workers. When these workers complete their tasks, labels are written to the Amazon S3 bucket you specified when you created the labeling job. The output data format determines what you see in your Amazon S3 bucket when your labeling job status (LabelingJobStatus) is Completed.

If you are a new user of Ground Truth, see Output Data to learn more about the Ground Truth output data format. To learn about the 3D point cloud object detection output data format, see 3D Point Cloud Object Detection Output.

3D Point Cloud Object Tracking
Use this task type when you want workers to add and fit 3D cuboids around objects to track their movement across 3D point cloud frames. For example, you can use this task type to ask workers to track the movement of vehicles across multiple point cloud frames.

For this task type, the data object that workers label is a sequence of point cloud frames. A sequence is defined as a temporal series of point cloud frames. Ground Truth renders a series of 3D point cloud visualizations using a sequence you provide and workers can switch between these 3D point cloud frames in the worker task interface.

Ground Truth provides workers with tools to annotate objects with 9 degrees of freedom: \((x,y,z,rx,ry,rz,l,w,h)\) in three dimensions in both 3D scene and projected side views (top, side, and back). When a worker draws a cuboid around an object, that cuboid is given a unique ID, for example \(\text{Car:1}\) for one car in the sequence and \(\text{Car:2}\) for another. Workers use that ID to label the same object in multiple frames.

You can also provide camera data to give workers more visual information about scenes in the frame, and to help workers draw 3D cuboids around objects. When a worker adds a 3D cuboid to identify an object in either the 2D image or the 3D point cloud, and the cuboid shows up in the other view.

You can adjust annotations created in a 3D point cloud object detection labeling job using the 3D point cloud object tracking adjustment task type.

If you are a new user of the Ground Truth 3D point cloud labeling modality, we recommend you review [3D Point Cloud Labeling Jobs Overview](#). This labeling modality is different from other Ground Truth task types, and this page provides an overview of important details you should be aware of when creating a 3D point cloud labeling job.

### Topics

- [View the Worker Task Interface](#)
- [Create a 3D Point Cloud Object Tracking Labeling Job](#)
- [Create a 3D Point Cloud Object Tracking Adjustment or Verification Labeling Job](#)
- [Output Data Format](#)

### View the Worker Task Interface

Ground Truth provides workers with a web portal and tools to complete your 3D point cloud object tracking annotation tasks. When you create the labeling job, you provide the Amazon Resource
Name (ARN) for a pre-built Ground Truth UI in the HumanTaskUiArn parameter. When you create a labeling job using this task type in the console, this UI is automatically used. You can preview and interact with the worker UI when you create a labeling job in the console. If you are a new user, it is recommended that you create a labeling job using the console to ensure your label attributes, point cloud frames, and if applicable, images, appear as expected.

The following is a GIF of the 3D point cloud object tracking worker task interface and demonstrates how the worker can navigate the point cloud frames in the sequence. The annotating tools are a part of the worker task interface. They are not available for the preview interface.

![Worker Interface GIF]

Once workers add a single cuboid, that cuboid is replicated in all frames of the sequence with the same ID. Once workers adjust the cuboid in another frame, Ground Truth will interpolate the movement of that object and adjust all cuboids between the manually adjusted frames. The following GIF demonstrates this interpolation feature. In the navigation bar on the bottom-left, red-areas indicate manually adjusted frames.
If you provide camera data for sensor fusion, images are matched up with scenes in point cloud frames. These images appear in the worker portal as shown in the following GIF.

Worker can navigate in the 3D scene using their keyboard and mouse. They can:

- Double click on specific objects in the point cloud to zoom into them.
- Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
- Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

Once a worker places a cuboids in the 3D scene, a side-view will appear with the three projected side views: top, side, and back. These side-views show points in and around the placed cuboid and help workers refine cuboid boundaries in that area. Workers can zoom in and out of each of those side-views using their mouse.

The following video demonstrates movements around the 3D point cloud and in the side-view.
Additional view options and features are available. See the worker instruction page for a comprehensive overview of the Worker UI.

**Worker Tools**

Workers can navigate through the 3D point cloud by zooming in and out, and moving in all directions around the cloud using the mouse and keyboard shortcuts. If workers click on a point in the point cloud, the UI will automatically zoom into that area. Workers can use various tools to draw 3D cuboid around objects. For more information, see Assistive Labeling Tools.

After workers have placed a 3D cuboid in the point cloud, they can adjust these cuboids to fit tightly around cars using a variety of views: directly in the 3D cuboid, in a side-view featuring three zoomed-in perspectives of the point cloud around the box, and if you include images for sensor fusion, directly in the 2D image.

View options that enable workers to easily hide or view label text, a ground mesh, and additional point attributes. Workers can also choose between perspective and orthogonal projections.

**Assistive Labeling Tools**
Ground Truth helps workers annotate 3D point clouds faster and more accurately using UX, machine learning and computer vision powered assistive labeling tools for 3D point cloud object tracking tasks. The following assistive labeling tools are available for this task type:

- **Label autofill** – When a worker adds a cuboid to a frame, a cuboid with the same dimensions and orientation is automatically added to all frames in the sequence.

- **Label interpolation** – After a worker has labeled a single object in two frames, Ground Truth uses those annotations to interpolate the movement of that object between those two frames. Label interpolation can be turned on and off.

- **Bulk label and attribute management** – Workers can add, delete, and rename annotations, label category attributes, and frame attributes in bulk.
  
  - Workers can manually delete annotations for a given object before or after a frame. For example, a worker can delete all labels for an object after frame 10 if that object is no longer located in the scene after that frame.
  
  - If a worker accidentally bulk deletes all annotations for an object, they can add them back. For example, if a worker deletes all annotations for an object before frame 100, they can bulk add them to those frames.
  
  - Workers can rename a label in one frame and all 3D cuboids assigned that label are updated with the new name across all frames.
  
  - Workers can use bulk editing to add or edit label category attributes and frame attributes in multiple frames.

- **Snapping** – Workers can add a cuboid around an object and use a keyboard shortcut or menu option to have Ground Truth's autofit tool snap the cuboid tightly around the object's boundaries.

- **Fit to ground** – After a worker adds a cuboid to the 3D scene, the worker can automatically snap the cuboid to the ground. For example, the worker can use this feature to snap a cuboid to the road or sidewalk in the scene.

- **Multi-view labeling** – After a worker adds a 3D cuboid to the 3D scene, a side-panel displays front and two side perspectives to help the worker adjust the cuboid tightly around the object. Workers can annotation the 3D point cloud, the side panel and the adjustments appear in the other views in real time.

- **Sensor fusion** – If you provide data for sensor fusion, workers can adjust annotations in the 3D scenes and in 2D images, and the annotations will be projected into the other view in real time.

- **Auto-merge cuboids** – Workers can automatically merge two cuboids across all frames if they determine that cuboids with different labels actually represent a single object.
View options – Enables workers to easily hide or view label text, a ground mesh, and additional point attributes like color or intensity. Workers can also choose between perspective and orthogonal projections.

Create a 3D Point Cloud Object Tracking Labeling Job

You can create a 3D point cloud labeling job using the SageMaker console or API operation, CreateLabelingJob. To create a labeling job for this task type you need the following:

- A sequence input manifest file. To learn how to create this type of manifest file, see Create a Point Cloud Sequence Input Manifest. If you are a new user of Ground Truth 3D point cloud labeling modalities, we recommend that you review Accepted Raw 3D Data Formats.
- A work team from a private or vendor workforce. You cannot use Amazon Mechanical Turk for 3D point cloud labeling jobs. To learn how to create workforces and work teams, see Create and Manage Workforces.

Additionally, make sure that you have reviewed and satisfied the Assign IAM Permissions to Use Ground Truth.

To learn how to create a labeling job using the console or an API, see the following sections.

Create a Labeling Job (API)

This section covers details you need to know when you create a labeling job using the SageMaker API operation CreateLabelingJob. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the See Also section of CreateLabelingJob.

Create a Labeling Job (API) provides an overview of the CreateLabelingJob operation. Follow these instructions and do the following while you configure your request:

- You must enter an ARN for HumanTaskUiArn. Use arn:aws:sagemaker:<region>:394669845002:human-task-ui/PointCloudObjectTracking. Replace <region> with the AWS Region you are creating the labeling job in.

  There should not be an entry for the UiTemplateS3Uri parameter.

- Your LabelAttributeName must end in -ref. For example, ot-labels-ref.
• Your input manifest file must be a point cloud frame sequence manifest file. For more information, see Create a Point Cloud Sequence Input Manifest.

• You specify your labels, label category and frame attributes, and worker instructions in a label category configuration file. For more information, see Create a Labeling Category Configuration File with Label Category and Frame Attributes to learn how to create this file.

• You need to provide pre-defined ARNs for the pre-annotation and post-annotation (ACS) Lambda functions. These ARNs are specific to the AWS Region you use to create your labeling job.
  
  To find the pre-annotation Lambda ARN, refer to PreHumanTaskLambdaArn. Use the Region you are creating your labeling job in to find the correct ARN that ends with PRE-3DPointCloudObjectTracking.

  To find the post-annotation Lambda ARN, refer to AnnotationConsolidationLambdaArn. Use the Region you are creating your labeling job in to find the correct ARN that ends with ACS-3DPointCloudObjectTracking.

• The number of workers specified in NumberOfHumanWorkersPerDataObject should be 1.

• Automated data labeling is not supported for 3D point cloud labeling jobs. You should not specify values for parameters in LabelingJobAlgorithmsConfig.

• 3D point cloud object tracking labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs in TaskTimeLimitInSeconds (up to 7 days, or 604,800 seconds).

Create a Labeling Job (Console)

You can follow the instructions Create a Labeling Job (Console) in order to learn how to create a 3D point cloud object tracking labeling job in the SageMaker console. While you are creating your labeling job, be aware of the following:

• Your input manifest file must be a sequence manifest file. For more information, see Create a Point Cloud Sequence Input Manifest.

• Optionally, you can provide label category attributes. Workers can assign one or more of these attributes to annotations to provide more information about that object. For example, you might want to use the attribute occluded to have workers identify when an object is partially obstructed.

• Automated data labeling and annotation consolidation are not supported for 3D point cloud labeling tasks.
• 3D point cloud object tracking labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs when you select your work team (up to 7 days, or 604800 seconds).

Create a 3D Point Cloud Object Tracking Adjustment or Verification Labeling Job

You can create an adjustment and verification labeling job using the Ground Truth console or CreateLabelingJob API. To learn more about adjustment and verification labeling jobs, and to learn how create one, see Verify and Adjust Labels.

When you create an adjustment labeling job, your input data to the labeling job can include labels, and yaw, pitch, and roll measurements from a previous labeling job or external source. In the adjustment job, pitch, and roll will be visualized in the worker UI, but cannot be modified. Yaw is adjustable.

Ground Truth uses Tait-Bryan angles with the following intrinsic rotations to visualize yaw, pitch and roll in the worker UI. First, rotation is applied to the vehicle according to the z-axis (yaw). Next, the rotated vehicle is rotated according to the intrinsic y'-axis (pitch). Finally, the vehicle is rotated according to the intrinsic x''-axis (roll).

Output Data Format

When you create a 3D point cloud object tracking labeling job, tasks are sent to workers. When these workers complete their tasks, their annotations are written to the Amazon S3 bucket you specified when you created the labeling job. The output data format determines what you see in your Amazon S3 bucket when your labeling job status (LabelingJobStatus) is Completed.

If you are a new user of Ground Truth, see Output Data to learn more about the Ground Truth output data format. To learn about the 3D point cloud object tracking output data format, see 3D Point Cloud Object Tracking Output.

3D Point Cloud Semantic Segmentation

Semantic segmentation involves classifying individual points of a 3D point cloud into pre-specified categories. Use this task type when you want workers to create a point-level semantic segmentation mask for 3D point clouds. For example, if you specify the classes car, pedestrian, and bike, workers select one class at a time, and color all of the points that this class applies to the same color in the point cloud.
For this task type, the data object that workers label is a single point cloud frame. Ground Truth generates a 3D point cloud visualization using point cloud data you provide. You can also provide camera data to give workers more visual information about scenes in the frame, and to help workers paint objects. When a worker paints an object in either the 2D image or the 3D point cloud, the paint shows up in the other view.

You can adjust annotations created in a 3D point cloud object detection labeling job using the 3D point cloud semantic segmentation adjustment task type.

If you are a new user of the Ground Truth 3D point cloud labeling modality, we recommend you review [3D Point Cloud Labeling Jobs Overview](#). This labeling modality is different from other Ground Truth task types, and this topic provides an overview of important details you should be aware of when creating a 3D point cloud labeling job.

**Topics**

- [View the Worker Task Interface](#)
- [Create a 3D Point Cloud Semantic Segmentation Labeling Job](#)
- [Create a 3D Point Cloud Semantic Segmentation Adjustment or Verification Labeling Job](#)
- [Output Data Format](#)

**View the Worker Task Interface**

Ground Truth provides workers with a web portal and tools to complete your 3D point cloud semantic segmentation annotation tasks. When you create the labeling job, you provide the Amazon Resource Name (ARN) for a pre-built Ground Truth UI in the `HumanTaskUiArn` parameter. When you create a labeling job using this task type in the console, this UI is automatically used. You can preview and interact with the worker UI when you create a labeling job in the console. If you are a new use, it is recommended that you create a labeling job using the console to ensure your label attributes, point cloud frames, and if applicable, images, appear as expected.

The following is a GIF of the 3D point cloud semantic segmentation worker task interface. If you provide camera data for sensor fusion, images are matched with scenes in the point cloud frame. Workers can paint objects in either the 3D point cloud or the 2D image, and the paint appears in the corresponding location in the other medium. These images appear in the worker portal as shown in the following GIF.
Worker can navigate in the 3D scene using their keyboard and mouse. They can:

- Double click on specific objects in the point cloud to zoom into them.
- Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
- Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

The following video demonstrates movements around the 3D point cloud. Workers can hide and re-expand all side views and menus. In this GIF, the side-views and menus have been collapsed.
The following GIF demonstrates how a worker can label multiple objects quickly, refine painted objects using the Unpaint option and then view only points that have been painted.
Additional view options and features are available. See the worker instruction page for a comprehensive overview of the Worker UI.

**Worker Tools**

Workers can navigate through the 3D point cloud by zooming in and out, and moving in all directions around the cloud using the mouse and keyboard shortcuts. When you create a semantic segmentation job, workers have the following tools available to them:

- A paint brush to paint and unpaint objects. Workers paint objects by selecting a label category and then painting in the 3D point cloud. Workers unpaint objects by selecting the Unpaint option from the label category menu and using the paint brush to erase paint.
- A polygon tool that workers can use to select and paint an area in the point cloud.
- A background paint tool, which enables workers to paint behind objects they have already annotated without altering the original annotations. For example, workers might use this tool to paint the road after painting all of the cars on the road.
- View options that enable workers to easily hide or view label text, a ground mesh, and additional point attributes like color or intensity. Workers can also choose between perspective and orthogonal projections.

**Create a 3D Point Cloud Semantic Segmentation Labeling Job**

You can create a 3D point cloud labeling job using the SageMaker console or API operation, CreateLabelingJob. To create a labeling job for this task type you need the following:

- A single-frame input manifest file. To learn how to create this type of manifest file, see Create a Point Cloud Frame Input Manifest File. If you are a new user of Ground Truth 3D point cloud labeling modalities, we recommend that you review Accepted Raw 3D Data Formats.
- A work team from a private or vendor workforce. You cannot use Amazon Mechanical Turk workers for 3D point cloud labeling jobs. To learn how to create workforces and work teams, see Create and Manage Workforces.
- A label category configuration file. For more information, see Create a Labeling Category Configuration File with Label Category and Frame Attributes.

Additionally, make sure that you have reviewed and satisfied the Assign IAM Permissions to Use Ground Truth.
Use one of the following sections to learn how to create a labeling job using the console or an API.

**Create a Labeling Job (Console)**

You can follow the instructions **Create a Labeling Job (Console)** in order to learn how to create a 3D point cloud semantic segmentation labeling job in the SageMaker console. While you are creating your labeling job, be aware of the following:

- Your input manifest file must be a single-frame manifest file. For more information, see [Create a Point Cloud Frame Input Manifest File](#).
- Automated data labeling and annotation consolidation are not supported for 3D point cloud labeling tasks.
- 3D point cloud semantic segmentation labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs when you select your work team (up to 7 days, or 604800 seconds).

**Create a Labeling Job (API)**

This section covers details you need to know when you create a labeling job using the SageMaker API operation `CreateLabelingJob`. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the **See Also** section of `CreateLabelingJob`.

The page, **Create a Labeling Job (API)**, provides an overview of the `CreateLabelingJob` operation. Follow these instructions and do the following while you configure your request:

- You must enter an ARN for `HumanTaskUiArn`. Use `arn:aws:sagemaker:<region>:394669845002:human-task-ui/PointCloudSemanticSegmentation`. Replace `<region>` with the AWS Region you are creating the labeling job in.
  
  There should not be an entry for the `UiTemplateS3Uri` parameter.

- Your `LabelAttributeName` must end in `-ref`. For example, `ss-labels-ref`.
- Your input manifest file must be a single-frame manifest file. For more information, see [Create a Point Cloud Frame Input Manifest File](#).
- You specify your labels and worker instructions in a label category configuration file. See [Create a Labeling Category Configuration File with Label Category and Frame Attributes](#) to learn how to create this file.
• You need to provide a pre-defined ARNs for the pre-annotation and post-annotation (ACS) Lambda functions. These ARNs are specific to the AWS Region you use to create your labeling job.

• To find the pre-annotation Lambda ARN, refer to `PreHumanTaskLambdaArn`. Use the Region you are creating your labeling job in to find the correct ARN. For example, if you are creating your labeling job in us-east-1, the ARN will be `arn:aws:lambda:us-east-1:432418664414:function:PRE-3DPointCloudSemanticSegmentation`.

• To find the post-annotation Lambda ARN, refer to `AnnotationConsolidationLambdaArn`. Use the Region you are creating your labeling job in to find the correct ARN. For example, if you are creating your labeling job in us-east-1, the ARN will be `arn:aws:lambda:us-east-1:432418664414:function:ACS-3DPointCloudSemanticSegmentation`.

• The number of workers specified in `NumberOfHumanWorkersPerDataObject` should be 1.

• Automated data labeling is not supported for 3D point cloud labeling jobs. You should not specify values for parameters in `LabelingJobAlgorithmsConfig`.

• 3D point cloud semantic segmentation labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs in `TaskTimeLimitInSeconds` (up to 7 days, or 604800 seconds).

**Create a 3D Point Cloud Semantic Segmentation Adjustment or Verification Labeling Job**

You can create an adjustment and verification labeling job using the Ground Truth console or `CreateLabelingJob` API. To learn more about adjustment and verification labeling jobs, and to learn how create one, see [Verify and Adjust Labels](#).

**Output Data Format**

When you create a 3D point cloud semantic segmentation labeling job, tasks are sent to workers. When these workers complete their tasks, their annotations are written to the Amazon S3 bucket you specified when you created the labeling job. The output data format determines what you see in your Amazon S3 bucket when your labeling job status (`LabelingJobStatus`) is Completed.

If you are a new user of Ground Truth, see [Output Data](#) to learn more about the Ground Truth output data format. To learn about the 3D point cloud object detection output data format, see [3D Point Cloud Semantic Segmentation Output](#).
3D-2D Point Cloud Object Tracking

Use this task type when you want workers to link 3D point cloud annotations with 2D images annotations and also link 2D image annotations among various cameras. Currently, Ground Truth supports cuboids for annotation in a 3D point cloud and bounding boxes for annotation in 2D videos. For example, you can use this task type to ask workers to link the movement of a vehicle in 3D point cloud with its 2D video. Using 3D-2D linking, you can easily correlate point cloud data (like the distance of a cuboid) to video data (bounding box) for up to 8 cameras.

Ground Truth provides workers with tools to annotate cuboids in a 3D point cloud and bounding boxes in up to 8 cameras using the same annotation UI. Workers can also link various bounding boxes for the same object across different cameras. For example, a bounding box in camera1 can be linked to a bounding box in camera2. This lets you to correlate an object across multiple cameras using a unique ID.

Note

Currently, SageMaker does not support creating a 3D-2D linking job using the console. To create a 3D-2D linking job using the SageMaker API, see Create a Labeling Job (API).

Topics

- View the Worker Task Interface
- Input Data Format
- Create a 3D-2D Point Cloud Object Tracking Labeling Job
- Output Data

View the Worker Task Interface

Ground Truth provides workers with a web portal and tools to complete your 3D-2D object tracking annotation tasks. When you create the labeling job, you provide the Amazon Resource Name (ARN) for a pre-built Ground Truth UI in the HumanTaskUiArn parameter. To use the UI when you create a labeling job for this task type using the API, you need to provide the HumanTaskUiArn. You can preview and interact with the worker UI when you create a labeling job through the API. The annotating tools are a part of the worker task interface. They are not available for the preview interface. The following image demonstrates the worker task interface used for the 3D-2D point cloud object tracking annotation task.
When interpolation is enabled by default. After a worker adds a single cuboid, that cuboid is replicated in all frames of the sequence with the same ID. If the worker adjusts the cuboid in another frame, Ground Truth interpolates the movement of that object and adjust all cuboids between the manually adjusted frames. Additionally, using the camera view section, a cuboid can be shown with a projection (using to B button for "toggle labels" in the camera view) that provides the worker with a reference from the camera images. The accuracy of the cuboid to image projection is based on accuracy of calibrations captured in the extrinsic and intrinsic data.

If you provide camera data for sensor fusion, images are matched up with scenes in point cloud frames. Note that the camera data should be time synchronized with the point cloud data to ensure an accurate depiction of point cloud to imagery over each frame in the sequence as shown in the following image.
The manifest file holds the extrinsic and intrinsic data and the pose to allow the cuboid projection on the camera image to be shown by using the P button.

Worker can navigate in the 3D scene using their keyboard and mouse. They can:

- Double click on specific objects in the point cloud to zoom into them.
- Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
- Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

Once a worker places a cuboids in the 3D scene, a side-view appears with the three projected side views: top, side, and front. These side-views show points in and around the placed cuboid and help workers refine cuboid boundaries in that area. Workers can zoom in and out of each of those side-views using their mouse.

The worker should first select the cuboid to draw a corresponding bounding box on any of the camera views. This links the cuboid and the bounding box with a common name and unique ID.

The worker can also first draw a bounding box, select it and draw the corresponding cuboid to link them.

Additional view options and features are available. See the worker instruction page for a comprehensive overview of the Worker UI.
Worker Tools

Workers can navigate through the 3D point cloud by zooming in and out, and moving in all directions around the cloud using the mouse and keyboard shortcuts. If workers click on a point in the point cloud, the UI automatically zooms into that area. Workers can use various tools to draw 3D cuboid around objects. For more information, see Assistive Labeling Tools in the following discussion.

After workers have placed a 3D cuboid in the point cloud, they can adjust these cuboids to fit tightly around cars using a variety of views: directly in the 3D point cloud, in a side-view featuring three zoomed-in perspectives of the point cloud around the box, and if you include images for sensor fusion, directly in the 2D image.

Additional view options enable workers to easily hide or view label text, a ground mesh, and additional point attributes. Workers can also choose between perspective and orthogonal projections.

Assistive Labeling Tools

Ground Truth helps workers annotate 3D point clouds faster and more accurately using UX, machine learning and computer vision powered assistive labeling tools for 3D point cloud object tracking tasks. The following assistive labeling tools are available for this task type:

- **Label autofill** – When a worker adds a cuboid to a frame, a cuboid with the same dimensions, orientation and xyz position is automatically added to all frames in the sequence.

- **Label interpolation** – After a worker has labeled a single object in two frames, Ground Truth uses those annotations to interpolate the movement of that object between all the frames. Label interpolation can be turned on and off. It is on by default. For example, if a worker working with 5 frames adds a cuboid in frame 2, it is copied to all the 5 frames. If the worker then makes adjustments in frame 4, frame 2 and 4 now act as two points, through which a line is fit. The cuboid is then interpolated in frames 1,3 and 5.

- **Bulk label and attribute management** – Workers can add, delete, and rename annotations, label category attributes, and frame attributes in bulk.

- Workers can manually delete annotations for a given object before and after a frame, or in all frames. For example, a worker can delete all labels for an object after frame 10 if that object is no longer located in the scene after that frame.
• If a worker accidentally bulk deletes all annotations for an object, they can add them back. For example, if a worker deletes all annotations for an object before frame 100, they can bulk add them to those frames.

• Workers can rename a label in one frame and all 3D cuboids assigned that label are updated with the new name across all frames.

• Workers can use bulk editing to add or edit label category attributes and frame attributes in multiple frames.

• **Snapping** – Workers can add a cuboid around an object and use a keyboard shortcut or menu option to have Ground Truth's autofit tool snap the cuboid tightly around the object's boundaries.

• **Fit to ground** – After a worker adds a cuboid to the 3D scene, the worker can automatically snap the cuboid to the ground. For example, the worker can use this feature to snap a cuboid to the road or sidewalk in the scene.

• **Multi-view labeling** – After a worker adds a 3D cuboid to the 3D scene, a side-panel displays front and two side perspectives to help the worker adjust the cuboid tightly around the object. Workers can annotation the 3D point cloud, the side panel and the adjustments appear in the other views in real time.

• **Sensor fusion** – If you provide data for sensor fusion, workers can adjust annotations in the 3D scenes and in 2D images, and the annotations are projected into the other view in real time. To learn more about the data for sensor fusion, see [Understand Coordinate Systems and Sensor Fusion](#).

• **Auto-merge cuboids** – Workers can automatically merge two cuboids across all frames if they determine that cuboids with different labels actually represent a single object.

• **View options** – Enables workers to easily hide or view label text, a ground mesh, and additional point attributes like color or intensity. Workers can also choose between perspective and orthogonal projections.

### Input Data Format

You can create a 3D-2D object tracking job using the SageMaker API operation, `CreateLabelingJob`. To create a labeling job for this task type you need the following:

• A sequence input manifest file. To learn how to create this type of manifest file, see [Create a Point Cloud Sequence Input Manifest](#). If you are a new user of Ground Truth 3D point cloud labeling modalities, we recommend that you review [Accepted Raw 3D Data Formats](#).
You specify your labels, label category and frame attributes, and worker instructions in a label category configuration file. For more information, see Create a Labeling Category Configuration File with Label Category and Frame Attributes to learn how to create this file. The following is an example showing a label category configuration file for creating a 3D-2D object tracking job.

```json
{
   "document-version": "2020-03-01",
   "categoryGlobalAttributes": [
      {
         "name": "Occlusion",
         "description": "global attribute that applies to all label categories",
         "type": "string",
         "enum": [
            "Partial",
            "Full"
         ]
      }
   ],
   "labels": [
      {
         "label": "Car",
         "attributes": [
            {
               "name": "Type",
               "type": "string",
               "enum": [
                  "SUV",
                  "Sedan"
               ]
            }
         ]
      },
      {
         "label": "Bus",
         "attributes": [
            {
               "name": "Size",
               "type": "string",
               "enum": [
                  "Large",
                  "Medium",
                  "Small"
               ]
            }
         ]
      }
   ]
}
```
Create a 3D-2D Point Cloud Object Tracking Labeling Job

You can create a 3D-2D point cloud labeling job using the SageMaker API operation, `CreateLabelingJob`. To create a labeling job for this task type you need the following:

- A work team from a private or vendor workforce. You cannot use Amazon Mechanical Turk for 3D point cloud labeling jobs. To learn how to create workforces and work teams, see Create and Manage Workforces.

- Add a CORS policy to an S3 bucket that contains input data in the Amazon S3 console. To set the required CORS headers on the S3 bucket that contains your input images in the S3 console, follow the directions detailed in CORS Permission Requirement.

- Additionally, make sure that you have reviewed and satisfied the Assign IAM Permissions to Use Ground Truth.
To learn how to create a labeling job using the API, see the following sections.

**Create a Labeling Job (API)**

This section covers details you need to know when you create a 3D-2D object tracking labeling job using the SageMaker API operation `CreateLabelingJob`. This API defines this operation for all AWS SDKs. To see a list of language-specific SDKs supported for this operation, review the **See Also** section of `CreateLabelingJob`.

**Create a Labeling Job (API)** provides an overview of the `CreateLabelingJob` operation. Follow these instructions and do the following while you configure your request:

- You must enter an ARN for `HumanTaskUiArn`. Use `arn:aws:sagemaker:<region>:394669845002:human-task-ui/PointCloudObjectTracking`. Replace `<region>` with the AWS Region you are creating the labeling job in. There should not be an entry for the `UiTemplateS3Uri` parameter.
- Your `LabelAttributeName` must end in `-ref`. For example, `ot-labels-ref`.
- Your input manifest file must be a point cloud frame sequence manifest file. For more information, see [Create a Point Cloud Sequence Input Manifest](#). You also need to provide a label category configuration file as mentioned above.
- You need to provide pre-defined ARNs for the pre-annotation and post-annotation (ACS) Lambda functions. These ARNs are specific to the AWS Region you use to create your labeling job.
  - To find the pre-annotation Lambda ARN, refer to `PreHumanTaskLambdaArn`. Use the Region you are creating your labeling job in to find the correct ARN that ends with `PRE-3DPointCloudObjectTracking`.
  - To find the post-annotation Lambda ARN, refer to `AnnotationConsolidationLambdaArn`. Use the Region you are creating your labeling job in to find the correct ARN that ends with `ACS-3DPointCloudObjectTracking`.
- The number of workers specified in `NumberOfHumanWorkersPerDataObject` should be 1.
- Automated data labeling is not supported for 3D point cloud labeling jobs. You should not specify values for parameters in `LabelingJobAlgorithmsConfig`.
- 3D-2D object tracking labeling jobs can take multiple hours to complete. You can specify a longer time limit for these labeling jobs in `TaskTimeLimitInSeconds` (up to 7 days, or 604,800 seconds).
Note

After you have successfully created a 3D-2D object tracking job, it shows up on the console under labeling jobs. The task type for the job is displayed as **Point Cloud Object Tracking**.

Output Data

When you create a 3D-2D object tracking labeling job, tasks are sent to workers. When these workers complete their tasks, their annotations are written to the Amazon S3 bucket you specified when you created the labeling job. The output data format determines what you see in your Amazon S3 bucket when your labeling job status (**LabelingJobStatus**) is **Completed**.

If you are a new user of Ground Truth, see **Output Data** to learn more about the Ground Truth output data format. To learn about the 3D-2D point cloud object tracking output data format, see **3D-2D Object Tracking Point Cloud Object Tracking Output**.

3D Point Cloud Labeling Jobs Overview

This topic provides an overview of the unique features of a Ground Truth 3D point cloud labeling job. You can use the 3D point cloud labeling jobs to have workers label objects in a 3D point cloud generated from a 3D sensors like LiDAR and depth cameras or generated from 3D reconstruction by stitching images captured by an agent like a drone.

Job Pre-processing Time

When you create a 3D point cloud labeling job, you need to provide an **input manifest file**. The input manifest file can be:

- A **frame input manifest file** that has a single point cloud frame on each line.
- A **sequence input manifest file** that has a single sequence on each line. A sequence is defined as a temporal series of point cloud frames.

For both types of manifest files, **job pre-processing time** (that is, the time before Ground Truth starts sending tasks to your workers) depends on the total number and size of point cloud frames you provide in your input manifest file. For frame input manifest files, this is the number of lines in your manifest file. For sequence manifest files, this is the number of frames in each sequence multiplied by the total number of sequences, or lines, in your manifest file.
Additionally, the number of points per point cloud and the number of fused sensor data objects (like images) factor into job pre-processing times. On average, Ground Truth can pre-process 200 point cloud frames in approximately 5 minutes. If you create a 3D point cloud labeling job with a large number of point cloud frames, you might experience longer job pre-processing times. For example, if you create a sequence input manifest file with 4 point cloud sequences, and each sequence contains 200 point clouds, Ground Truth pre-processes 800 point clouds and so your job pre-processing time might be around 20 minutes. During this time, your labeling job status is InProgress.

While your 3D point cloud labeling job is pre-processing, you receive CloudWatch messages notifying you of the status of your job. To identify these messages, search for 3D_POINT_CLOUD_PROCESSING_STATUS in your labeling job logs.

For **frame input manifest files**, your CloudWatch logs will have a message similar to the following:

```
{
    "labeling-job-name": "example-point-cloud-labeling-job",
    "event-name": "3D_POINT_CLOUD_PROCESSING_STATUS",
    "event-log-message": "datasetObjectId from: 0 to 10, status: IN_PROGRESS"
}
```

The event log message, datasetObjectId from: 0 to 10, status: IN_PROGRESS identifies the number of frames from your input manifest that have been processed. You receive a new message every time a frame has been processed. For example, after a single frame has processed, you receive another message that says datasetObjectId from: 1 to 10, status: IN_PROGRESS.

For **sequence input manifest files**, your CloudWatch logs will have a message similar to the following:

```
{
    "labeling-job-name": "example-point-cloud-labeling-job",
    "event-name": "3D_POINT_CLOUD_PROCESSING_STATUS",
    "event-log-message": "datasetObjectId: 0, status: IN_PROGRESS"
}
```

The event log message, datasetObjectId from: 0, status: IN_PROGRESS identifies the number of sequences from your input manifest that have been processed. You receive a new message every time a sequence has been processed. For example, after a single sequence
has processed, you receive a message that says `datasetObjectId from: 1, status: IN_PROGRESS` as the next sequence begins processing.

**Job Completion Times**

3D point cloud labeling jobs can take workers hours to complete. You can set the total amount of time that workers can work on each task when you create a labeling job. The maximum time you can set for workers to work on tasks is 7 days. The default value is 3 days.

It is strongly recommended that you create tasks that workers can complete within 12 hours. Workers must keep the worker UI open while working on a task. They can save work as they go and Ground Truth will save their work every 15 minutes.

When using the SageMaker `CreateLabelingJob` API operation, set the total time a task is available to workers in the `TaskTimeLimitInSeconds` parameter of `HumanTaskConfig`.

When you create a labeling job in the console, you can specify this time limit when you select your workforce type and your work team.

**Workforces**

When you create a 3D point cloud labeling job, you need to specify a work team that will complete your point cloud annotation tasks. You can choose a work team from a private workforce of your own workers, or from a vendor workforce that you select in the AWS Marketplace. You cannot use the Amazon Mechanical Turk workforce for 3D point cloud labeling jobs.

To learn more about vendor workforce, see [Managing Vendor Workforces](#).

To learn how to create and manage a private workforce, see [Use a Private Workforce](#).

**Worker User Interface (UI)**

Ground Truth provides a worker user interface (UI), tools, and assistive labeling features to help workers complete your 3D point cloud labeling tasks.

You can preview the worker UI when you create a labeling job in the console.

When you create a labeling job using the API operation `CreateLabelingJob`, you must provide an ARN provided by Ground Truth in the parameter `HumanTaskUiArn` to specify the worker UI for your task type. You can use `HumanTaskUiArn` with the SageMaker `RenderUiTemplate` API operation to preview the worker UI.
You provide worker instructions, labels, and optionally, label category attributes that are displayed in the worker UI.

**Label Category Attributes**

When you create a 3D point cloud object tracking or object detection labeling job, you can add one or more label category attributes. You can add frame attributes to all 3D point cloud task types:

- **Label category attribute** – A list of options (strings), a free form text box, or a numeric field associated with one or more labels. It is used by workers to provide metadata about a label.

- **Frame attribute** – A list of options (strings), a free form text box, or a numeric field that appears on each point cloud frame a worker is sent to annotate. It is used by workers to provide metadata about frames.

Additionally, you can use label and frame attributes to have workers verify labels in a 3D point cloud label verification job.

Use the following sections to learn more about these attributes. To learn how to add label category and frame attributes to a labeling job, use the Create Labeling Job section on the task type page of your choice.

**Label Category Attributes**

Add label category attributes to labels to give workers the ability to provide more information about the annotations they create. A label category attribute is added to an individual label, or to all labels. When a label category attribute is applied to all labels it is referred to as a global label category attribute.

For example, if you add the label category *car*, you might also want to capture additional data about your labeled cars, such as if they are occluded or the size of the car. You can capture this metadata using label category attributes. In this example, if you added the attribute *occluded* to the car label category, you can assign *partial, completely, no* to the *occluded* attribute and enable workers to select one of these options.

When you create a label verification job, you add labels category attributes to each label you want workers to verify.
Frame Attributes

Add frame attributes to give workers the ability to provide more information about individual point cloud frames. You can specify up to 10 frame attributes, and these attributes will appear on all frames.

For example, you can add a frame attribute that allows workers to enter a number. You may want to use this attribute to have workers identify the number of objects they see in a particular frame.

In another example, you may want to provide a free-form text box to give workers the ability to provide a free form answer to a question.

When you create a label verification job, you can add one or more frame attributes to ask workers to provide feedback on all labels in a point cloud frame.

Worker Instructions

You can provide worker instructions to help your workers complete your point cloud labeling tasks. You might want to use these instructions to do the following:

- Best practices and things to avoid when annotating objects.
- Explanation of the label category attributes provided (for object detection and object tracking tasks), and how to use them.
- Advice on how to save time while labeling by using keyboard shortcuts.

You can add your worker instructions using the SageMaker console while creating a labeling job. If you create a labeling job using the API operation CreateLabelingJob, you specify worker instructions in your label category configuration file.

In addition to your instructions, Ground Truth provides a link to help workers navigate and use the worker portal. View these instructions by selecting the task type on Worker Instructions.

Declining Tasks

Workers are able to decline tasks.

Workers decline a task if the instructions are not clear, input data is not displaying correctly, or if they encounter some other issue with the task. If the number of workers per dataset object (NumberOfHumanWorkersPerDataObject) decline the task, the data object is marked as expired and will not be sent to additional workers.
3D Point Cloud Labeling Job Permission Requirements

When you create a 3D point cloud labeling job, in addition to the permission requirements found in Assign IAM Permissions to Use Ground Truth, you must add a CORS policy to your S3 bucket that contains your input manifest file.

Add a CORS Permission Policy to S3 Bucket

When you create a 3D point cloud labeling job, you specify buckets in S3 where your input data and manifest file are located and where your output data will be stored. These buckets may be the same. You must attach the following Cross-origin resource sharing (CORS) policy to your input and output buckets. If you use the Amazon S3 console to add the policy to your bucket, you must use the JSON format.

**JSON**

```json
[
  {
    "AllowedHeaders": ["*"]
  },
  "AllowedMethods": [
    "GET",
    "HEAD",
    "PUT"
  ],
  "AllowedOrigins": ["*"]
}
```

**XML**

```xml
<?xml version="1.0" encoding="UTF-8"?>
<CORSConfiguration xmlns="http://s3.amazonaws.com/doc/2006-03-01/">
  <CORSRule>
    <AllowedOrigin>*</AllowedOrigin>
  </CORSRule>
</CORSConfiguration>
```
<AllowedMethod>GET</AllowedMethod>
<AllowedMethod>HEAD</AllowedMethod>
<AllowedMethod>PUT</AllowedMethod>
<MaxAgeSeconds>3000</MaxAgeSeconds>
<ExposeHeader>Access-Control-Allow-Origin</ExposeHeader>
<AllowedHeader>*</AllowedHeader>
</CORSRule>
</CORSConfiguration>

To learn how to add a CORS policy to an S3 bucket, see [How do I add cross-domain resource sharing with CORS?](https://docs.aws.amazon.com/AmazonS3/latest/user-guide/cors.html) in the Amazon Simple Storage Service User Guide.

### Worker Instructions

This topic provides an overview of the Ground Truth worker portal and the tools available to complete your 3D Point Cloud labeling task. First, select the type of task you are working on from **Topics**.

For adjustment jobs, select the original labeling job task type that produced the labels you are adjusting. Review and adjust the labels in your task as needed.

⚠️ **Important**

It is recommended that you complete your task using a Google Chrome or Firefox web browser.

#### Topics

- [3D Point Cloud Semantic Segmentation](#)
- [3D Point Cloud Object Detection](#)
- [3D Point Cloud Object Tracking](#)

#### 3D Point Cloud Semantic Segmentation

Use this page to become familiarize with the user interface and tools available to complete your 3D point cloud semantic segmentation task.

#### Topics
Your Task

When you work on a 3D point cloud semantic segmentation task, you need to select a category from the Annotations menu on the right side of your worker portal using the drop down menu Label Categories. After you've selected a category, use the paint brush and polygon tools to paint each object in the 3D point cloud that this category applies to. For example, if you select the category Car, you would use these tools to paint all of the cars in the point cloud. The following video demonstrates how to use the paint brush tool to paint an object.

If you see one or more images in your worker portal, you can paint in the images or paint in the 3D point cloud and the paint will show up in the other medium.

You may see frame attributes under the Labels menu. Use these attribute prompts to enter additional information about the point cloud.
Important

If you see that objects have already been painted when you open the task, adjust those annotations.

The following video includes an image that can be annotated. You may not see an image in your task.
After you’ve painted one or more objects using a label category, you can select that category from the Label Category menu on the right to only view points painted for that category.
Navigate the UI

You can navigate in the 3D scene using their keyboard and mouse. You can:

- Double click on specific objects in the point cloud to zoom into them.
- Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
- Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

The following video demonstrates movements around the 3D point cloud and in the side-view. You can hide and re-expand all side views using the full screen icon. In this GIF, the side-views and menus have been collapsed.

When you are in the worker UI, you see the following menus:

- **Instructions** – Review these instructions before starting your task.
- **Shortcuts** – Use this menu to view keyboard shortcuts that you can use to navigate the point cloud and use the annotation tools provided.
- **View** – Use this menu to toggle different view options on and off. For example, you can use this menu to add a ground mesh to the point cloud, and to choose the projection of the point cloud.
• **3D Point Cloud** – Use this menu to add additional attributes to the points in the point cloud, such as color, and pixel intensity. Note that some or all of these options may not be available.

• **Paint** – Use this menu to modify the functionality of the paint brush.

When you open a task, the move scene icon is on, and you can move around the point cloud using your mouse and the navigation buttons in the point cloud area of the screen. To return to the original view you see when you first opened the task, choose the reset scene icon.

After you select the paint icon, you can add paint to the point cloud and images (if included). You must select the move scene icon again to move to another area in the 3D point cloud or image.

To collapse all panels on the right and make the 3D point cloud full screen, select the full screen icon.

For the camera images and side-panels, you have the following view options:

• **C** – View the camera angle on point cloud view.

• **F** – View the frustum, or field of view, of the camera used to capture that image on point cloud view.

• **P** – View the point cloud overlaid on the image.

**Icon Guide**

Use this table to learn about the icons available in your worker task portal.

<table>
<thead>
<tr>
<th>Icon</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="brush" /></td>
<td>brush</td>
<td>Choose this icon to turn on the brush tool. To use with this tool, choose and move over the objects that you want to paint with your mouse. After you choose it, everything you paint be associated with the category you chose.</td>
</tr>
<tr>
<td><img src="image" alt="polygon" /></td>
<td>polygon</td>
<td>Choose this icon to use the polygon paint tool. Use this tool to draw polygons around objects that you want to paint. After you choose it, everything you draw a polygon around will be associated with the category you have chosen.</td>
</tr>
<tr>
<td>Icon</td>
<td>Name</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-----------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><img src="image" alt="reset scene icon" /></td>
<td>reset scene</td>
<td>Choose this icon to reset the view of the point cloud, side panels, and if applicable, all images to their original position when the task was first opened.</td>
</tr>
<tr>
<td><img src="image" alt="move scene icon" /></td>
<td>move scene</td>
<td>Choose this icon to move the scene. By default, this icon will be selected when you first start a task.</td>
</tr>
<tr>
<td><img src="image" alt="full screen icon" /></td>
<td>full screen</td>
<td>Choose this icon to make the 3D point cloud visualization full screen, and to collapse all side panels.</td>
</tr>
<tr>
<td>Icon</td>
<td>Name</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>------</td>
<td>-------------</td>
</tr>
<tr>
<td><img src="image" alt="ruler icon" /></td>
<td>ruler</td>
<td>Use this icon to measure distances, in meters, in the point cloud. You may want to use this tool if your instructions ask you to annotate all objects in a given distance from the center of the cuboid or the object used to capture data. When you select this icon, you can place the starting point (first marker) anywhere in the point cloud by selecting it with your mouse. The tool will automatically use interpolation to place a marker on the closest point within threshold distance to the location you select, otherwise the marker will be placed on ground. If you place a starting point by mistake, you can use the Escape key to revert marker placement. After you place the first marker, you see a dotted line and a dynamic label that indicates the distance you have moved away from the first marker. Click somewhere else on the point cloud to place a second marker. When you place the second marker, the dotted line becomes solid, and the distance is set. After you set a distance, you can edit it by selecting either marker. You can delete a ruler by selecting anywhere on the ruler and using the Delete key on your keyboard.</td>
</tr>
</tbody>
</table>

**Shortcuts**

The shortcuts listed in the **Shortcuts** menu can help you navigate the 3D point cloud and use the paint tool.

Before you start your task, it is recommended that you review the **Shortcuts** menu and become acquainted with these commands.
Release, Stop and Resume, and Decline Tasks

When you open the labeling task, three buttons on the top right allow you to decline the task (Decline task), release it (Release task), and stop and resume it at a later time (Stop and resume later). The following list describes what happens when you select one of these options:

- **Decline task**: You should only decline a task if something is wrong with the task, such as an issue with the 3D point cloud, images or the UI. If you decline a task, you will not be able to return to the task.

- **Release Task**: If you release a task, you lose all work done on that task. When the task is released, other workers on your team can pick it up. If enough workers pick up the task, you may not be able to return to it. When you select this button and then select Confirm, you are returned to the worker portal. If the task is still available, its status will be Available. If other workers pick it up, it will disappear from your portal.

- **Stop and resume later**: You can use the Stop and resume later button to stop working and return to the task at a later time. You should use the Save button to save your work before you select Stop and resume later. When you select this button and then select Confirm, you are returned to the worker portal, and the task status is Stopped. You can select the same task to resume work on it.

  Be aware that the person that creates your labeling tasks specifies a time limit in which all tasks must be completed by. If you do not return to and complete this task within that time limit, it will expire and your work will not be submitted. Contact your administrator for more information.

Saving Your Work and Submitting

You should periodically save your work. Ground Truth will automatically save your work every 15 minutes.

When you open a task, you must complete your work on it before pressing Submit.

3D Point Cloud Object Detection

Use this page to become familiarized with the user interface and tools available to complete your 3D point cloud object detection task.

Topics

- **Your Task**
Your Task

When you work on a 3D point cloud object detection task, you need to select a category from the **Annotations** menu on the right side of your worker portal using the **Label Categories** menu. After you've chosen a category, use the add cuboid and fit cuboid tools to fit a cuboid around objects in the 3D point cloud that this category applies to. After you place a cuboid, you can modify its dimensions, location, and orientation directly in the point cloud, and the three panels shown on the right.

If you see one or more images in your worker portal, you can also modify cuboids in the images or in the 3D point cloud and the edits will show up in the other medium.

If you see cuboids have already been added to the 3D point cloud when you open your task, adjust those cuboids and add additional cuboids as needed.

To edit a cuboid, including moving, re-orienting, and changing cuboid dimensions, you must use shortcut keys. You can see a full list of shortcut keys in the **Shortcuts** menu in your UI. The following are important key-combinations that you should become familiar with before starting your labeling task.

<table>
<thead>
<tr>
<th>Mac Command</th>
<th>Windows Command</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cmd + Drag</td>
<td>Ctrl + Drag</td>
<td>Modify the dimensions of the cuboid.</td>
</tr>
<tr>
<td>Option + Drag</td>
<td>Alt + Drag</td>
<td>Move the cuboid.</td>
</tr>
<tr>
<td>Shift + Drag</td>
<td>Shift + Drag</td>
<td>Rotate the cuboid.</td>
</tr>
<tr>
<td>Option + O</td>
<td>Alt + O</td>
<td>Fit the cuboid tightly around the points it has been drawn around. Before using the</td>
</tr>
<tr>
<td>Mac Command</td>
<td>Windows Command</td>
<td>Action</td>
</tr>
<tr>
<td>-------------</td>
<td>-----------------</td>
<td>--------</td>
</tr>
<tr>
<td>Option + G</td>
<td>Alt + G</td>
<td>Set the cuboid to the ground.</td>
</tr>
</tbody>
</table>

Individual labels may have one or more label attributes. If a label has a label attribute associated with it, it will appear when you select the downward pointing arrow next to the label from the **Label Id** menu. Fill in required values for all label attributes.

You may see frame attributes under the **Labels** menu. Use these attribute prompts to enter additional information about each frame.

**Navigate the UI**

You can navigate in the 3D scene using your keyboard and mouse. You can:
• Double click on specific objects in the point cloud to zoom into them.
• You can use the [ and ] keys on your keyboard to zoom into and move from one label to the next. If no label is selected, when you select [ or ], the UI will zoom into the first label in the **Label Id** list.
• Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
• Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

Once you place a cuboids in the 3D scene, a side-view will appear with three projected views: top, side, and back. These side-views show points in and around the placed cuboid and help workers refine cuboid boundaries in that area. Workers can zoom in and out of each of those side-views using their mouse.

The following video demonstrates movements around the 3D point cloud and in the side-view.

When you are in the worker UI, you see the following menus:

• **Instructions** – Review these instructions before starting your task.
• **Shortcuts** – Use this menu to view keyboard shortcuts that you can use to navigate the point cloud and use the annotation tools provided.
• **Label** – Use this menu to modify a cuboid. First, select a cuboid, and then choose an option from this menu. This menu includes assistive labeling tools like setting a cuboid to the ground and automatically fitting the cuboid to the object's boundaries.

• **View** – Use this menu to toggle different view options on and off. For example, you can use this menu to add a ground mesh to the point cloud, and to choose the projection of the point cloud.

• **3D Point Cloud** – Use this menu to add additional attributes to the points in the point cloud, such as color, and pixel intensity. Note that these options may not be available.

When you open a task, the move scene icon is on, and you can move around the point cloud using your mouse and the navigation buttons in the point cloud area of the screen. To return to the original view you see when you first opened the task, choose the reset scene icon. Resetting the view will not modify your annotations.

After you select the add cuboid icon, you can add cuboids to the 3D point cloud visualization. Once you've added a cuboid, you can adjust it in the three views (top, side, and front) and in the images (if included).

You must choose the move scene icon again to move to another area in the 3D point cloud or image.

To collapse all panels on the right and make the 3D point cloud full-screen, choose the full screen icon.
If camera images are included, you may have the following view options:

- **C** – View the camera angle on point cloud view.
- **F** – View the frustum, or field of view, of the camera used to capture that image on point cloud view.
- **P** – View the point cloud overlaid on the image.
- **B** – View cuboids in the image.

The following video demonstrates how to use these view options. The **F** option is used to view the field of view of the camera (the gray area), the **C** options shows the direction the camera is facing and angle of the camera (blue lines), and the **B** option is used to view the cuboid.

![Video Demonstration](image)

**Icon Guide**

Use this table to learn about the icons you see in your worker task portal.

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Add Cuboid Icon" /></td>
<td>add cuboid</td>
</tr>
<tr>
<td><img src="image" alt="Edit Cuboid Icon" /></td>
<td>edit cuboid</td>
</tr>
</tbody>
</table>

Label 3D Point Clouds
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>orientation. After a cuboid is added, it automatically switches to edit cuboid mode.</td>
</tr>
<tr>
<td>ruler</td>
<td>Use this icon to measure distances, in meters, in the point cloud. You may want to use this tool if your instructions ask you to annotate all objects in a given distance from the center of the cuboid or the object used to capture data.</td>
</tr>
<tr>
<td></td>
<td>When you select this icon, you can place the starting point (first marker) anywhere in the point cloud by selecting it with your mouse. The tool will automatically use interpolation to place a marker on the closest point within threshold distance to the location you select, otherwise the marker will be placed on ground. If you place a starting point by mistake, you can use the Escape key to revert marker placement.</td>
</tr>
<tr>
<td></td>
<td>After you place the first marker, you see a dotted line and a dynamic label that indicates the distance you have moved away from the first marker. Click somewhere else on the point cloud to place a second marker. When you place the second marker, the dotted line becomes solid, and the distance is set.</td>
</tr>
<tr>
<td></td>
<td>After you set a distance, you can edit it by selecting either marker. You can delete a ruler by selecting anywhere on the ruler and using the Delete key on your keyboard.</td>
</tr>
<tr>
<td>reset scene</td>
<td>Choose this icon to reset the view of the point cloud, side panels, and if applicable, all images to their original position when the task was first opened.</td>
</tr>
<tr>
<td>move scene</td>
<td>Choose this icon to move the scene. By default, this icon is chosen when you first start a task.</td>
</tr>
<tr>
<td>Icon</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
</tr>
<tr>
<td>![full screen]</td>
<td>Choose this icon to make the 3D point cloud visualization full screen, and to collapse all side panels.</td>
</tr>
<tr>
<td>![show labels]</td>
<td>Show labels in the 3D point cloud visualization, and if applicable, in images.</td>
</tr>
<tr>
<td>![hide labels]</td>
<td>Hide labels in the 3D point cloud visualization, and if applicable, in images.</td>
</tr>
<tr>
<td>![delete labels]</td>
<td>Delete a label.</td>
</tr>
</tbody>
</table>

**Shortcuts**

The shortcuts listed in the **Shortcuts** menu can help you navigate the 3D point cloud and use tools to add and edit cuboids.

Before you start your task, it is recommended that you review the **Shortcuts** menu and become acquainted with these commands. You need to use some of the 3D cuboid controls to edit your cuboid.

**Release, Stop and Resume, and Decline Tasks**

When you open the labeling task, three buttons on the top right allow you to decline the task (**Decline task**), release it (**Release task**), and stop and resume it at a later time (**Stop and resume later**). The following list describes what happens when you select one of these options:

- **Decline task**: You should only decline a task if something is wrong with the task, such as an issue with the 3D point cloud, images or the UI. If you decline a task, you will not be able to return to the task.

- **Release Task**: If you release a task, you loose all work done on that task. When the task is released, other workers on your team can pick it up. If enough workers pick up the task, you may not be able to return to it. When you select this button and then select **Confirm**, you are returned to the worker portal. If the task is still available, its status will be **Available**. If other workers pick it up, it will disappear from your portal.
• **Stop and resume later**: You can use the **Stop and resume later** button to stop working and return to the task at a later time. You should use the **Save** button to save your work before you select **Stop and resume later**. When you select this button and then select **Confirm**, you are returned to the worker portal, and the task status is **Stopped**. You can select the same task to resume work on it.

Be aware that the person that creates your labeling tasks specifies a time limit in which all tasks must be completed by. If you do not return to and complete this task within that time limit, it will expire and your work will not be submitted. Contact your administrator for more information.

**Saving Your Work and Submitting**

You should periodically save your work. Ground Truth will automatically save your work every 15 minutes.

When you open a task, you must complete your work on it before pressing **Submit**.

**3D Point Cloud Object Tracking**

Use this page to become familiarize with the user interface and tools available to complete your 3D point cloud object detection task.

**Topics**

- **Your Task**
- **Navigate the UI**
- **Bulk Edit Label Category and Frame Attributes**
- **Icon Guide**
- **Shortcuts**
- **Release, Stop and Resume, and Decline Tasks**
- **Saving Your Work and Submitting**

**Your Task**

When you work on a 3D point cloud object tracking task, you need to select a category from the **Annotations** menu on the right side of your worker portal using the **Label Categories** menu. After you've selected a category, use the add cuboid and fit cuboid tools to fit a cuboid around objects
in the 3D point cloud that this category applies to. After you place a cuboid, you can modify its location, dimensions, and orientation directly in the point cloud, and the three panels shown on the right. If you see one or more images in your worker portal, you can also modify cuboids in the images or in the 3D point cloud and the edits will show up in the other medium.

⚠️ **Important**

If you see cuboids have already been added to the 3D point cloud frames when you open your task, adjust those cuboids and add additional cuboids as needed.

To edit a cuboid, including moving, re-orienting, and changing cuboid dimensions, you must use shortcut keys. You can see a full list of shortcut keys in the **Shortcuts** menu in your UI. The following are important key-combinations that you should become familiar with before starting your labeling task.

<table>
<thead>
<tr>
<th>Mac Command</th>
<th>Windows Command</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cmd + Drag</td>
<td>Ctrl + Drag</td>
<td>Modify the dimensions of the cuboid.</td>
</tr>
<tr>
<td>Option + Drag</td>
<td>Alt + Drag</td>
<td>Move the cuboid.</td>
</tr>
<tr>
<td>Shift + Drag</td>
<td>Shift + Drag</td>
<td>Rotate the cuboid.</td>
</tr>
<tr>
<td>Option + O</td>
<td>Alt + O</td>
<td>Fit the cuboid tightly around the points it has been drawn around. Before using the option, make sure the cuboid fully-surrounds the object of interest.</td>
</tr>
<tr>
<td>Option + G</td>
<td>Alt + G</td>
<td>Set the cuboid to the ground.</td>
</tr>
</tbody>
</table>

When you open your task, two frames will be loaded. If your task includes more than two frames, you need to use the navigation bar in the lower-left corner, or the load frames icon to load additional frames. You should annotate and adjust labels in all frames before submitting.
After you fit a cuboid tightly around the boundaries of an object, navigate to another frame using the navigation bar in the lower-left corner of the UI. If that same object has moved to a new location, add another cuboid and fit it tightly around the boundaries of the object. Each time you manually add a cuboid, you see the frame sequence bar in the lower-left corner of the screen turn red where that frame is located temporally in the sequence.

Your UI automatically infers the location of that object in all other frames after you've placed a cuboid. This is called interpolation. You can see the movement of that object, and the inferred and manually created cuboids using the arrows. Adjust inferred cuboids as needed. The following video demonstrates how to navigate between frames. The following video shows how, if you add a cuboid in one frame, and then adjust it in another, your UI will automatically infer the location of the cuboid in all of the frames in-between.

![Image of 3D Point Clouds](image.png)

### Tip

You can turn off the automatic cuboid interpolation across frames using the 3D Point Cloud menu item. Select **3D Point Cloud** from the top-menu, and then select **Interpolate Cuboids Across Frames**. This will uncheck this option and stop cuboid interpolation. You can reselect this item to turn cuboid interpolation back on. Turning cuboid interpolation off will not impact cuboids that have already been interpolated across frames.
Individual labels may have one or more label attributes. If a label has a label attribute associated with it, it will appear when you select the downward pointing arrow next to the label from the **Label Id** menu. Fill in required values for all label attributes.

You may see frame attributes under the **Label Id** menu. These attributes will appear on each frame in your task. Use these attribute prompts to enter additional information about each frame.

### Frame 1 attributes

**Is the point cloud clearly visible?**

*Visible: frame attribute that applies to all frames*

- Yes
- No

**Describe Issues**

*Issues: frame attribute that applies to all frames*

**Number of Cars Labeled**

*Cars labeled: frame attribute that applies to all frames*

---

**Navigate the UI**

You can navigate in the 3D scene using your keyboard and mouse. You can:

- Double click on specific objects in the point cloud to zoom into them.
- You can use the [ and ] keys on your keyboard to zoom into and move from one label to the next. If no label is selected, when you select [ or ], the UI will zoom into the first label in the **Label Id** list.
- Use a mouse-scroller or trackpad to zoom in and out of the point cloud.
• Use both keyboard arrow keys and Q, E, A, and D keys to move Up, Down, Left, Right. Use keyboard keys W and S to zoom in and out.

Once you place a cuboids in the 3D scene, a side-view will appear with three projected views: top, side, and back. These side-views show points in and around the placed cuboid and help workers refine cuboid boundaries in that area. Workers can zoom in and out of each of those side-views using their mouse.

The following video demonstrates movements around the 3D point cloud and in the side-view.

When you are in the worker UI, you see the following menus:

• **Instructions** – Review these instructions before starting your task.
• **Shortcuts** – Use this menu to view keyboard shortcuts that you can use to navigate the point cloud and use the annotation tools provided.
• **Label** – Use this menu to modify a cuboid. First, select a cuboid, and then choose an option from this menu. This menu includes assistive labeling tools like setting a cuboid to the ground and automatically fitting the cuboid to the object's boundaries.
• **View** – Use this menu to toggle different view options on and off. For example, you can use this menu to add a ground mesh to the point cloud, and to choose the projection of the point cloud.
• **3D Point Cloud** – Use this menu to add additional attributes to the points in the point cloud, such as color, and pixel intensity. Note that these options may not be available.
When you open a task, the move scene icon is on, and you can move around the point cloud using your mouse and the navigation buttons in the point cloud area of the screen. To return to the original view you see when you first opened the task, choose the reset scene icon.

After you select the add cuboid icon, you can add cuboids to the point cloud and images (if included). You must select the move scene icon again to move to another area in the 3D point cloud or image.

To collapse all panels on the right and make the 3D point cloud full-screen, choose the full screen icon.

If camera images are included, you may have the following view options:

- **C** – View the camera angle on point cloud view.
- **F** – View the frustum, or field of view, of the camera used to capture that image on point cloud view.
- **P** – View the point cloud overlaid on the image.
- **B** – View cuboids in the image.

The following video demonstrates how to use these view options. The **F** option is used to view the field of view of the camera (the gray area), the **C** options shows the direction the camera is facing and angle of the camera (blue lines), and the **B** option is used to view the cuboid.
Delete Cuboids

You can select a cuboid or label ID and:

- Delete an individual cuboid in the current frame you are viewing.
- Delete all cuboids with that label ID before or after the frame you are viewing.
- Delete all cuboids with that label ID in all frames.

A common use-case for cuboid deletion is if the object leaves the scene.

You can use one or more of these options to delete both manually placed and interpolated cuboids with the same label ID.

- To delete all cuboids before or after the frame you are currently on, select the cuboid, select the Label menu item at the top of the UI and then select one of Delete in previous frames or Delete in next frames. Use the Shortcuts menu to see the shortcut keys you can use for these options.
- To delete a label in all frames, select Delete in all frames from the Labels menu, or use the shortcut Shift + Delete on your keyboard.
- To delete an individual cuboid from a single frame, select the cuboid and either select the trashcan icon next to that label ID in the Label ID sidebar on the right or use the Delete key on your keyboard to delete that cuboid.

If you have manually placed more than one cuboid with the same label in different frames, when you delete one of the manually placed cuboids, all interpolated cuboids adjust. This adjustment happens because the UI uses manually placed cuboids as anchor points when calculating the location of interpolated cuboid. When you remove one of these anchor points, the UI must recalculate the position of interpolated cuboids.

If you delete a cuboid from a frame, but later decide that you want to get it back, you can use the Duplicate to previous frames or Duplicate to next frames options in the Label menu to copy the cuboid into all the previous or all of the following frames, respectively.
Bulk Edit Label Category and Frame Attributes

You can bulk edit label attributes and frame attributes.

When you bulk edit an attribute, you specify one or more ranges of frames that you want to apply the edit to. The attribute you select is edited in all frames in that range, including the start and end frames you specify. When you bulk edit label attributes, the range you specify must contain the label that the label attribute is attached to. If you specify frames that do not contain this label, you will receive an error.

To bulk edit an attribute you must specify the desired value for the attribute first. For example, if you want to change an attribute from Yes to No, you must select No, and then perform the bulk edit.

You can also specify a new value for an attribute that has not been filled in and then use the bulk edit feature to fill in that value in multiple frames. To do this, select the desired value for the attribute and complete the following procedure.

**To bulk edit a label or attribute:**

1. Use your mouse to right click the attribute you want to bulk edit.
2. Specify the range of frames you want to apply the bulk edit to using a dash (-) in the text box. For example, if you want to apply the edit to frames one through ten, enter 1-10. If you want to apply the edit to frames two to five, eight to ten and twenty enter 2-5, 8-10, 20.
3. Select **Confirm**.

If you get an error message, verify that you entered a valid range and that the label associated with the label attribute you are editing (if applicable) exists in all frames specified.

You can quickly add a label to all previous or subsequent frames using the **Duplicate to previous frames** and **Duplicate to next frames** options in the **Label** menu at the top of your screen.

**Icon Guide**

Use this table to learn about the icons you see in your worker task portal.
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>add cuboid</strong> Choose this icon to add a cuboid. Each cuboid you add is associated with the category you chose.</td>
</tr>
<tr>
<td></td>
<td><strong>edit cuboid</strong> Choose this icon to edit a cuboid. After you add a cuboid, you can edit its dimensions, location, and orientation. After a cuboid is added, it automatically switches to edit cuboid mode.</td>
</tr>
<tr>
<td></td>
<td><strong>ruler</strong> Use this icon to measure distances, in meters, in the point cloud. You may want to use this tool if your instructions ask you to annotate all objects in a given distance from the center of the cuboid or the object used to capture data.</td>
</tr>
</tbody>
</table>

When you select this icon, you can place the starting point (first marker) anywhere in the point cloud by selecting it with your mouse. The tool will automatically use interpolation to place a marker on the closest point within threshold distance to the location you select, otherwise the marker will be placed on ground. If you place a starting point by mistake, you can use the Escape key to revert marker placement.

After you place the first marker, you see a dotted line and a dynamic label that indicates the distance you have moved away from the first marker. Click somewhere else on the point cloud to place a second marker. When you place the second marker, the dotted line becomes solid, and the distance is set.

After you set a distance, you can edit it by selecting either marker. You can delete a ruler by selecting anywhere on the ruler and using the Delete key on your keyboard.
<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="reset scene" /></td>
<td>Choose this icon to reset the view of the point cloud, side panels, and if applicable, all images to their original position when the task was first opened.</td>
</tr>
<tr>
<td><img src="image" alt="move scene" /></td>
<td>Choose this icon to move the scene. By default, this icon is chosen when you first start a task.</td>
</tr>
<tr>
<td><img src="image" alt="full screen" /></td>
<td>Choose this icon to make the 3D point cloud visualization full screen and to collapse all side panels.</td>
</tr>
<tr>
<td><img src="image" alt="load frames" /></td>
<td>Choose this icon to load additional frames.</td>
</tr>
<tr>
<td><img src="image" alt="hide labels" /></td>
<td>Hide labels in the 3D point cloud visualization, and if applicable, in images.</td>
</tr>
<tr>
<td><img src="image" alt="show labels" /></td>
<td>Show labels in the 3D point cloud visualization, and if applicable, in images.</td>
</tr>
<tr>
<td><img src="image" alt="delete labels" /></td>
<td>Delete a label. This option can only be used to delete labels you have manually created or adjusted.</td>
</tr>
</tbody>
</table>

**Shortcuts**

The shortcuts listed in the **Shortcuts** menu can help you navigate the 3D point cloud and use tools to add and edit cuboids.

Before you start your task, it is recommended that you review the **Shortcuts** menu and become acquainted with these commands. You need to use some of the 3D cuboid controls to edit your cuboid.
Release, Stop and Resume, and Decline Tasks

When you open the labeling task, three buttons on the top right allow you to decline the task (Decline task), release it (Release task), and stop and resume it at a later time (Stop and resume later). The following list describes what happens when you select one of these options:

- **Decline task**: You should only decline a task if something is wrong with the task, such as an issue with the 3D point clouds, images or the UI. If you decline a task, you will not be able to return to the task.

- **Release Task**: Use this option to release a task and allow others to work on it. When you release a task, you lose all work done on that task and other workers on your team can pick it up. If enough workers pick up the task, you may not be able to return to it. When you select this button and then select Confirm, you are returned to the worker portal. If the task is still available, its status will be Available. If other workers pick it up, it will disappear from your portal.

- **Stop and resume later**: You can use the Stop and resume later button to stop working and return to the task at a later time. You should use the Save button to save your work before you select Stop and resume later. When you select this button and then select Confirm, you are returned to the worker portal, and the task status is Stopped. You can select the same task to resume work on it.

Be aware that the person that creates your labeling tasks specifies a time limit in which all tasks must be completed by. If you do not return to and complete this task within that time limit, it will expire and your work will not be submitted. Contact your administrator for more information.

Saving Your Work and Submitting

You should periodically save your work. Ground Truth will automatically save your work every 15 minutes.

When you open a task, you must complete your work on it before pressing Submit.

Verify and Adjust Labels

When the labels on a dataset need to be validated, Amazon SageMaker Ground Truth provides functionality to have workers verify that labels are correct or to adjust previous labels.

These types of jobs fall into two distinct categories:
• **Label verification** — Workers indicate if the existing labels are correct, or rate their quality, and can add comments to explain their reasoning. Workers will not be able to modify or adjust labels.

If you create a 3D point cloud or video frame label adjustment or verification job, you can choose to make label category attributes (not supported for 3D point cloud semantic segmentation) and frame attributes editable by workers.

• **Label adjustment** — Workers adjust prior annotations and, if applicable, label category and frame attributes to correct them.

The following Ground Truth [built-in task types](https://docs.aws.amazon.com/sagemaker/latest/dg/labeling-builtin-task-types.html) support adjustment and verification labeling jobs:

- Bounding box
- Semantic segmentation
- 3D point cloud object detection, 3D point cloud object tracking, and 3D point cloud semantic segmentation
- All video frame object detection and video frame object tracking task types — bounding box, polyline, polygon and keypoint

**Tip**

For 3D point cloud and video frame labeling verification jobs, it is recommended that you add new label category attributes or frame attributes to the labeling job. Workers can use these attributes to verify individual labels or the entire frame. To learn more about label category and frame attributes, see [Worker User Interface (UI)](https://docs.aws.amazon.com/sagemaker/latest/dg/worker-ui-3d-point-cloud.html) for 3D point cloud and [Worker User Interface (UI)](https://docs.aws.amazon.com/sagemaker/latest/dg/worker-ui-video-frame.html) for video frame.

You can start a label verification and adjustment jobs using the SageMaker console or the API.

**Topics**

- [Requirements to Create Verification and Adjustment Labeling Jobs](https://docs.aws.amazon.com/sagemaker/latest/dg/labeling-verify-adjust-requirements.html)
- [Create a Label Verification Job (Console)](https://docs.aws.amazon.com/sagemaker/latest/dg/labeling-verify-adjust-create-label-verification-job.html)
- [Create a Label Adjustment Job (Console)](https://docs.aws.amazon.com/sagemaker/latest/dg/labeling-verify-adjust-create-label-adjustment-job.html)
- [Start a Label Verification or Adjustment Job (API)](https://docs.aws.amazon.com/sagemaker/latest/dg/labeling-verify-adjust-start-label-verification-or-adjustment-job.html)
- [Label Verification and Adjustment Data in the Output Manifest](https://docs.aws.amazon.com/sagemaker/latest/dg/labeling-verify-adjust-data-output-manifest.html)
• **Cautions and Considerations**

### Requirements to Create Verification and Adjustment Labeling Jobs

To create a label verification or adjustment job, the following criteria must be satisfied.

- **For non streaming labeling jobs:** The input manifest file you use must contain the label attribute name (LabelAttributeName) of the labels that you want adjusted. When you chain a successfully completed labeling job, the output manifest file is used as the input manifest file for the new, chained job. To learn more about the format of the output manifest file Ground Truth produces for each task type, see [Output Data](#).

- **For streaming labeling jobs:** The Amazon SNS message you sent to the Amazon SNS input topic of the adjustment or verification labeling job must contain the label attribute name of the labels you want adjusted or verified. To see an example of how you can create an adjustment or verification labeling job with streaming labeling jobs, see this [Jupyter Notebook example](#) in GitHub.

- **The task type of the verification or adjustment labeling job must be the same as the task type of the original job unless you are using the Image Label Verification task type to verify bounding box or semantic segmentation image labels.** See the next bullet point for more details about the video frame task type requirements.

- **For video frame annotation verification and adjustment jobs,** you must use the same annotation task type used to create the annotations from the previous labeling job. For example, if you create a video frame object detection job to have workers draw bounding boxes around objects, and then you create a video object detection adjustment job, you must specify *bounding boxes* as the annotation task type. To learn more video frame annotation task types, see [Task Types](#).

- **The task type you select for the adjustment or verification labeling job must support an audit workflow.** The following Ground Truth **built-in task types** support adjustment and verification labeling jobs: bounding box, semantic segmentation, 3D point cloud object detection, 3D point cloud object tracking, and 3D point cloud semantic segmentation, and all video frame object detection and video frame object tracking task types — bounding box, polyline, polygon and keypoint.
Create a Label Verification Job (Console)

Bounding box and semantic segmentation labeling jobs are created by choosing the Label verification task type in the console. To create a verification job for 3D point cloud and video frame task types, you must choose the same task type as the original labeling job and choose to display existing labels. Use one of the following sections to create a label verification job for your task type.

Topics

- Create an Image Label Verification Job (Console)
- Create a Point Cloud or Video Frame Label Verification Job (Console)

Create an Image Label Verification Job (Console)

Use the following procedure to create a bounding box or semantic segmentation verification job using the console. This procedure assumes that you have already created a bounding box or semantic segmentation labeling job and its status is Complete. This the labeling job that produces the labels you want verified.

To create an image label verification job:

2. Start a new labeling job by chaining a prior job or start from scratch, specifying an input manifest that contains labeled data objects.
3. In the Task type pane, select Label verification.
4. Choose Next.
5. In the Workers section, choose the type of workforce you would like to use. For more details about your workforce options see Create and Manage Workforces.
6. (Optional) After you've selected your workforce, specify the Task timeout and Task expiration time.
7. In the Existing-labels display options pane, the system shows the available label attribute names in your manifest. Choose the label attribute name that identifies the labels that you want workers to verify. Ground Truth tries to detect and populate these values by analyzing the manifest, but you might need to set the correct value.
8. Use the instructions areas of the tool designer to provide context about what the previous labelers were asked to do and what the current verifiers need to check.

You can add new labels that workers choose from to verify labels. For example, you can ask workers to verify the image quality, and provide the labels *Clear* and *Blurry*. Workers will also have the option to add a comment to explain their selection.

9. Choose **See preview** to check that the tool is displaying the prior labels correctly and presents the label verification task clearly.

10. Select **Create**. This will create and start your labeling job.

### Create a Point Cloud or Video Frame Label Verification Job (Console)

Use the following procedure to create a 3D point cloud or video frame verification job using the console. This procedure assumes that you have already created a labeling job using the task type that produces the types of labels you want to be verified and its status is Complete.

**To create an image label verification job:**

2. Start a new labeling job by **chaining** a prior job or start from scratch, specifying an input manifest that contains labeled data objects.
3. In the **Task type** pane, select the same task type as the labeling job that you chained. For example, if the original labeling job was a video frame object detection keypoint labeling job, select that task type.
4. Choose **Next**.
5. In the **Workers** section, choose the type of workforce you would like to use. For more details about your workforce options see [Create and Manage Workforces](#).
6. (Optional) After you've selected your workforce, specify the **Task timeout** and **Task expiration time**.
7. Toggle on the switch next to **Display existing labels**.
8. Select **Verification**.
9. For **Label attribute name**, choose the name from your manifest that corresponds to the labels that you want to display for verification. You will only see label attribute names for labels that match the task type you selected on the previous screen. Ground Truth tries to detect and populate these values by analyzing the manifest, but you might need to set the correct value.
10. Use the instructions areas of the tool designer to provide context about what the previous labelers were asked to do and what the current verifiers need to check.

You cannot modify or add new labels. You can remove, modify and add new label category attributes or frame attributes. It is recommended that you add new label category attributes or frame attributes to the labeling job. Workers can use these attribute to verify individual labels or the entire frame.

By default, preexisting label category attributes and frame attributes will not be editable by workers. If you want to make a label category or frame attribute editable, select the Allow workers to edit this attribute check box for that attribute.

To learn more about label category and frame attributes, see Worker User Interface (UI) for 3D point cloud and Worker User Interface (UI) for video frame.

11. Choose See preview to check that the tool is displaying the prior labels correctly and presents the label verification task clearly.

12. Select Create. This will create and start your labeling job.

Create a Label Adjustment Job (Console)

Use one of the following sections to create a label verification job for your task type.

Topics
- Create an Image Label Adjustment Job (Console)
- Create a Point Cloud or Video Frame Label Adjustment Job (Console)

Create an Image Label Adjustment Job (Console)

Use the following procedure to create a bounding box or semantic segmentation adjustment labeling job using the console. This procedure assumes that you have already created a bounding box or semantic segmentation labeling job and its status is Complete. This the labeling job that produces the labels you want adjusted.

To create an image label adjustment job (console)
2. Start a new labeling job by chaining a prior job or start from scratch, specifying an input manifest that contains labeled data objects.

3. Choose the same task type as the original labeling job.

4. Choose Next.

5. In the Workers section, choose the type of workforce you would like to use. For more details about your workforce options see Create and Manage Workforces.

6. (Optional) After you've selected your workforce, specify the Task timeout and Task expiration time.

7. Expand Existing-labels display options by selecting the arrow next to the title.

8. Check the box next to I want to display existing labels from the dataset for this job.

9. For Label attribute name, choose the name from your manifest that corresponds to the labels that you want to display for adjustment. You will only see label attribute names for labels that match the task type you selected on the previous screen. Ground Truth tries to detect and populate these values by analyzing the manifest, but you might need to set the correct value.

10. Use the instructions areas of the tool designer to provide context about what the previous labelers were tasked with doing and what the current verifiers need to check and adjust.

11. Choose See preview to check that the tool shows the prior labels correctly and presents the task clearly.

12. Select Create. This will create and start your labeling job.

Create a Point Cloud or Video Frame Label Adjustment Job (Console)

Use the following procedure to create a 3D point cloud or video frame adjustment job using the console. This procedure assumes that you have already created a labeling job using the task type that produces the types of labels you want to be verified and its status is Complete.

To create a 3D point cloud or video frame label adjustment job (console)


2. Start a new labeling job by chaining a prior job or start from scratch, specifying an input manifest that contains labeled data objects.

3. Choose the same task type as the original labeling job.

4. Toggle on the switch next to Display existing labels.
5. Select **Adjustment**.

6. For **Label attribute name**, choose the name from your manifest that corresponds to the labels that you want to display for adjustment. You will only see label attribute names for labels that match the task type you selected on the previous screen. Ground Truth tries to detect and populate these values by analyzing the manifest, but you might need to set the correct value.

7. Use the instructions areas of the tool designer to provide context about what the previous labelers were asked to do and what the current adjusters need to check.

You cannot remove or modify existing labels but you can add new labels. You can remove, modify and add new label category attributes or frame attributes.

Be default, preexisting label category attributes and frame attributes will be editable by workers. If you want to make a label category or frame attribute uneditable, deselect the **Allow workers to edit this attribute** check box for that attribute.

To learn more about label category and frame attributes, see [Worker User Interface (UI) for 3D point cloud](https://github.com/aws/sagemaker-ground-truth) and [Worker User Interface (UI) for video frame](https://github.com/aws/sagemaker-ground-truth).

8. Choose **See preview** to check that the tool shows the prior labels correctly and presents the task clearly.

9. Select **Create**. This will create and start your labeling job.

### Start a Label Verification or Adjustment Job (API)

Start a label verification or adjustment job by chaining a successfully completed job or starting a new job from scratch using the `CreateLabelingJob` operation. The procedure is almost the same as setting up a new labeling job with `CreateLabelingJob`, with a few modifications. Use the following sections to learn what modifications are required to chain a labeling job to create an adjustment or verification labeling job.

When you create an adjustment or verification labeling job using the Ground Truth API, you **must** use a different `LabelAttributeName` than the original labeling job. The original labeling job is the job used to create the labels you want adjusted or verified.

⚠️ **Important**

The label category configuration file you identify for an adjustment or verification job in `LabelCategoryConfigS3Uri` of `CreateLabelingJob` must contain the same labels...
used in the original labeling job. You can add new labels. For 3D point cloud and video frame jobs, you can add new label category and frame attributes to the label category configuration file.

Bounding Box and Semantic Segmentation

To create a bounding box or semantic segmentation label verification or adjustment job, use the following guidelines to specify API attributes for the CreateLabelingJob operation.

- Use the `LabelAttributeName` parameter to specify the output label name that you want to use for verified or adjusted labels. You must use a different `LabelAttributeName` than the one used for the original labeling job.

- If you are chaining the job, the labels from the previous labeling job to be adjusted or verified will be specified in the custom UI template. To learn how to create a custom template, see [Create Custom Worker Task Templates](#).

Identify the location of the UI template in the `UiTemplateS3Uri` parameter. SageMaker provides widgets that you can use in your custom template to display old labels. Use the `initial-value` attribute in one of the following crowd elements to extract the labels that need verification or adjustment and include them in your task template:

- `crowd-semantic-segmentation`—Use this crowd element in your custom UI task template to specify semantic segmentation labels that need to be verified or adjusted.

- `crowd-bounding-box`—Use this crowd element in your custom UI task template to specify bounding box labels that need to be verified or adjusted.

- The `LabelCategoryConfigS3Uri` parameter must contain the same label categories as the previous labeling job.

- Use the bounding box or semantic segmentation adjustment or verification lambda ARNs for `PreHumanTaskLambdaArn` and `AnnotationConsolidationLambdaArn`:

  - For bounding box, the adjustment labeling job lambda function ARNs end with `AdjustmentBoundingBox` and the verification lambda function ARNs end with `VerificationBoundingBox`.

  - For semantic segmentation, the adjustment labeling job lambda function ARNs end with `AdjustmentSemanticSegmentation` and the verification lambda function ARNs end with `VerificationSemanticSegmentation`.

Verify and Adjust Labels
3D Point Cloud and Video Frame

- Use the `LabelAttributeName` parameter to specify the output label name that you want to use for verified or adjusted labels. You must use a different `LabelAttributeName` than the one used for the original labeling job.

- You must use the human task UI Amazon Resource Name (ARN) (`HumanTaskUiArn`) used for the original labeling job. To see supported ARNs, see `HumanTaskUiArn`.

- In the label category configuration file, you must specify the label attribute name (`LabelAttributeName`) of the previous labeling job that you use to create the adjustment or verification labeling job in the `auditLabelAttributeName` parameter.

- You specify whether your labeling job is a verification or adjustment labeling job using the `editsAllowed` parameter in your label category configuration file identified by the `LabelCategoryConfigS3Uri` parameter.

  - For verification labeling jobs, you must use the `editsAllowed` parameter to specify that all labels cannot be modified. `editsAllowed` must be set to "none" in each entry in `labels`. Optionally, you can specify whether or not label categories attributes and frame attributes can be adjusted by workers.

  - Optionally, for adjustment labeling jobs, you can use the `editsAllowed` parameter to specify labels, label category attributes, and frame attributes that can or cannot be modified by workers. If you do not use this parameter, all labels, label category attributes, and frame attributes will be adjustable.

To learn more about the `editsAllowed` parameter and configuring your label category configuration file, see `Label Category Configuration File Schema`.

- Use the 3D point cloud or video frame adjustment lambda ARNs for `PreHumanTaskLambdaArn` and `AnnotationConsolidationLambdaArn` for both adjustment and verification labeling jobs:

  - For 3D point clouds, the adjustment and verification labeling job lambda function ARNs end with `Adjustment3DPointCloudSemanticSegmentation`, `Adjustment3DPointCloudObjectTracking`, and `Adjustment3DPointCloudObjectDetection` for 3D point cloud semantic segmentation, object detection, and object tracking respectively.

  - For video frames, the adjustment and verification labeling job lambda function ARNs end with `AdjustmentVideoObjectDetection` and `AdjustmentVideoObjectTracking` for video frame object detection and object tracking respectively.
Ground Truth stores the output data from a label verification or adjustment job in the S3 bucket that you specified in the S3OutputPath parameter of the CreateLabelingJob operation. For more information about the output data from a label verification or adjustment labeling job, see Label Verification and Adjustment Data in the Output Manifest.

**Label Verification and Adjustment Data in the Output Manifest**

Amazon SageMaker Ground Truth writes label verification data to the output manifest within the metadata for the label. It adds two properties to the metadata:

- A type property, with a value of "groundtruth/label-verification".
- A worker-feedback property, with an array of comment values. This property is added when the worker enters comments. If there are no comments, the field doesn't appear.

The following example output manifest shows how label verification data appears:

```json
{
    "source-ref":"S3 bucket location",
    "verify-bounding-box":"1",
    "verify-bounding-box-metadata":
    {
        "class-name": "bad",
        "confidence": 0.93,
        "type": "groundtruth/label-verification",
        "job-name": "verify-bounding-boxes",
        "human-annotated": "yes",
        "creation-date": "2018-10-18T22:18:13.527256",
        "worker-feedback": [
            {"comment": "The bounding box on the bird is too wide on the right side."},
            {"comment": "The bird on the upper right is not labeled."}
        ]
    }
}
```

The worker output of adjustment tasks resembles the worker output of the original task, except that it contains the adjusted values and an adjustment-status property with the value of adjusted or unadjusted to indicate whether an adjustment was made.

For more examples of the output of different tasks, see Output Data.
Cautions and Considerations

To get expected behavior when creating a label verification or adjustment job, carefully verify your input data.

- If you are using image data, verify that your manifest file contains hexadecimal RGB color information.
- To save money on processing costs, filter your data to ensure you are not including unwanted objects in your labeling job input manifest.
- Add required Amazon S3 permissions to ensure your input data is processed correctly.

When you create an adjustment or verification labeling job using the Ground Truth API, you must use a different LabelAttributeName than the original labeling job.

Color Information Requirements for Semantic Segmentation Jobs

To properly reproduce color information in verification or adjustment tasks, the tool requires hexadecimal RGB color information in the manifest (for example, #FFFFFF for white). When you set up a Semantic Segmentation verification or adjustment job, the tool examines the manifest to determine if this information is present. If it can't find it, Amazon SageMaker Ground Truth displays an error message and ends job setup.

In prior iterations of the Semantic Segmentation tool, category color information wasn't output in hexadecimal RGB format to the output manifest. That feature was introduced to the output manifest at the same time the verification and adjustment workflows were introduced. Therefore, older output manifests aren't compatible with this new workflow.

Filter Your Data Before Starting the Job

Amazon SageMaker Ground Truth processes all objects in your input manifest. If you have a partially labeled data set, you might want to create a custom manifest using an Amazon S3 Select query on your input manifest. Unlabeled objects individually fail, but they don't cause the job to fail, and they might incur processing costs. Filtering out objects you don't want verified reduces your costs.

If you create a verification job using the console, you can use the filtering tools provided there. If you create jobs using the API, make filtering your data part of your workflow where needed.
Creating Custom Labeling Workflows

This document guides you through the process of setting up a workflow with a custom labeling template. To learn more about starting a labeling job, see Getting started. In that section, when you choose the Task type, select Custom labeling task, and then follow this section's instructions to configure it.

Topics

- Step 1: Setting up your workforce
- Step 2: Creating your custom worker task template
- Step 3: Processing with AWS Lambda
- Demo Template: Annotation of Images with crowd-bounding-box
- Demo Template: Labeling Intents with crowd-classifier
- Custom Workflows via the API

For more information about creating custom labeling workflows, see Build a custom data labeling workflow with Amazon SageMaker Ground Truth.

Step 1: Setting up your workforce

In this step you use the console to establish which worker type to use and make the necessary sub-selections for the worker type. It assumes you have already completed the steps up to this point in the Getting started section and have chosen the Custom labeling task as the Task type.

To configure your workforce.

1. First choose an option from the Worker types. There are three types currently available:

   - **Public** uses an on-demand workforce of independent contractors, powered by Amazon Mechanical Turk. They are paid on a per-task basis.
   - **Private** uses your employees or contractors for handling data that needs to stay within your organization.
   - **Vendor** uses third party vendors that specialize in providing data labeling services, available via the AWS Marketplace.

2. If you choose the Public option, you are asked to set the number of workers per dataset object. Having more than one worker perform the same task on the same object can help
increase the accuracy of your results. The default is three. You can raise or lower that depending on the accuracy you need.

You are also asked to set a **price per task** by using a drop-down menu. The menu recommends price points based on how long it will take to complete the task.

The recommended method to determine this is to first run a short test of your task with a **private** workforce. The test provides a realistic estimate of how long the task takes to complete. You can then select the range your estimate falls within on the **Price per task** menu. If your average time is more than 5 minutes, consider breaking your task into smaller units.

**Next**

**Step 2: Creating your custom worker task template**

*A worker task template* is a file used by Ground Truth to customize the worker user interface (UI), or human task UI. You can create a worker task template using HTML, CSS, JavaScript, **Liquid template language**, and **Crowd HTML Elements**. Liquid is used to automate the template, and Crowd HTML Elements can be used to include common annotation tools and provide the logic to submit to Ground Truth.

Use the following topics to learn how you can create a worker task template. You can see a repository of example Ground Truth worker task templates on **GitHub**.

**Topics**

- **Starting with a base template**
- **Developing templates locally**
- **Using External Assets**
- **Track your variables**
- **A simple sample**
- **Adding automation with Liquid**
- **End-to-end demos**
Starting with a base template

You can use a template editor in the Ground Truth console to start creating a template. This editor includes a number of pre-designed base templates and an HTML and Crowd HTML Element autofill feature.

To access the Ground Truth custom template editor:

1. Following the instructions in Create a Labeling Job (Console) and select Custom for the labeling job Task type.
2. When you select Next, you will be able to access the template editor and base templates in the Custom labeling task setup section.
3. (Optional) Select a base template from the drop-down menu under Templates. If you prefer to create a template from scratch, choose Custom from the drop down-menu for a minimal template skeleton.

Developing templates locally

While you need to be in the console to test how your template will process incoming data, you can test the look and feel of your template's HTML and custom elements in your browser by adding this code to the top of your HTML file.

Example

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
```

This loads the necessary code to render the custom HTML elements. Use this if you want to develop your template's look and feel in your preferred editor rather than in the console.

Remember, though, this will not parse your variables. You may want to replace them with sample content while developing locally.

Using External Assets

Amazon SageMaker Ground Truth custom templates allow external scripts and style sheets to be embedded. For example, the following code block demonstrates how you would add a style sheet located at https://www.example.com/my-enhancement-styles.css to your template.
Example

```html
<script src="https://www.example.com/my-enhancement-script.js"></script>
<link rel="stylesheet" type="text/css" href="https://www.example.com/my-enhancement-styles.css"/>
```

If you encounter errors, ensure that your originating server is sending the correct MIME type and encoding headers with the assets.

For example, the MIME and encoding types for remote scripts are: application/javascript;CHARSET=UTF-8.

The MIME and encoding type for remote stylesheets are: text/css;CHARSET=UTF-8.

**Track your variables**

In the process of building the sample below, there will be a step that adds variables to it to represent the pieces of data that may change from task to task, worker to worker. If you're starting with one of the sample templates, you will need to make sure you're aware of the variables it already uses. When you create your pre-annotation AWS Lambda script, its output will need to contain values for any of those variables you choose to keep.

The values you use for the variables can come from your manifest file. All the key-value pairs in your data object are provided to your pre-annotation Lambda. If it's a simple pass-through script, matching keys for values in your data object to variable names in your template is the easiest way to pass those values through to the tasks forms your workers see.

**A simple sample**

All tasks begin and end with the `<crowd-form> </crowd-form>` elements. Like standard HTML `<form>` elements, all of your form code should go between them.

For a simple tweet-analysis task, use the `<crowd-classifier>` element. It requires the following attributes:

- **name** - the variable name to use for the result in the form output.
- **categories** - a JSON formatted array of the possible answers.
- **header** - a title for the annotation tool

As children of the `<crowd-classifier>` element, you must have three regions.
Amazon SageMaker

Developer Guide

• <classiﬁcation-target> - the text the worker will classify based on the options speciﬁed in the
categories attribute above.
• <full-instructions> - instructions that are available from the "View full instructions" link in the
tool. This can be left blank, but it is recommended that you give good instructions to get better
results.
• <short-instructions> - a more brief description of the task that appears in the tool's sidebar. This
can be left blank, but it is recommended that you give good instructions to get better results.
A simple version of this tool would look like this.
Example of using crowd-classifier
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<crowd-classifier
name="tweetFeeling"
categories="['positive','negative','neutral', 'unclear']"
header="Which term best describes this tweet?"
>
<classification-target>
My favorite football team won today!
Bring on the division finals!
</classification-target>
<full-instructions header="Sentiment Analysis Instructions">
Try to determine the sentiment the author
of the tweet is trying to express.
If none seem to match, choose "cannot determine."
</full-instructions>
<short-instructions>
Pick the term best describing the sentiment
of the tweet.
</short-instructions>
</crowd-classifier>
</crowd-form>

You can copy and paste the code into the editor in the Ground Truth labeling job creation workﬂow
to preview the tool, or try out a demo of this code on CodePen.
Creating Custom Labeling Workﬂows

1541


Adding automation with Liquid

Our custom template system uses Liquid for automation. It is an open source inline markup language. In Liquid, the text between single curly braces and percent symbols is an instruction or tag that performs an operation like control flow or iteration. Text between double curly braces is a variable or object that outputs its value.

The most common use of Liquid will be to parse the data coming from your pre-annotation Lambda and pull out the relevant variables to create the task. The taskInput object returned by your Pre-annotation Lambda will be available as the task.input object in your templates.

The properties in your manifest's data objects are passed into your Pre-annotation Lambda as the event.dataObject. A simple pass-through script simply returns that object as the taskInput object. You would represent values from your manifest as variables as follows.

Example Manifest data object

```json
{
    "source": "This is a sample text for classification",
    "labels": [ "angry", "sad", "happy", "inconclusive" ],
    "header": "What emotion is the speaker feeling?"
}
```

Example Sample HTML using variables

```html
<crowd-classifier
    name='tweetFeeling'
```
Note the addition of " | to_json" to the labels property above. That's a filter to turn the array into a JSON representation of the array. Variable filters are explained in the next section.

The following list includes two types of Liquid tags that you may find useful to automate template input data processing. If you select one of the following tag-types, you will be redirected to the Liquid documentation.

- **Control flow**: Includes programming logic operators like if/else, unless, and case/when.
- **Iteration**: Enables you to run blocks of code repeatedly using statements like for loops.

For an example of an HTML template that uses Liquid elements to create a for loop, see `translation-review-and-correction.liquid.html` in GitHub.

For more information and documentation, visit the [Liquid homepage](#).

**Variable filters**

In addition to the standard [Liquid filters](#) and actions, Ground Truth offers a few additional filters. Filters are applied by placing a pipe (`|`) character after the variable name, then specifying a filter name. Filters can be chained in the form of:

**Example**

```liquid
{{ <content> | <filter> | <filter> }}
```

**Autoescape and explicit escape**

By default, inputs will be HTML escaped to prevent confusion between your variable text and HTML. You can explicitly add the escape filter to make it more obvious to someone reading the source of your template that the escaping is being done.

**escape_once**

`escape_once` ensures that if you've already escaped your code, it doesn't get re-escaped on top of that. For example, so that `&` doesn't become `&amp;`. 

Creating Custom Labeling Workflows
skip_autoescape

skip_autoescape is useful when your content is meant to be used as HTML. For example, you might have a few paragraphs of text and some images in the full instructions for a bounding box.

Use skip_autoescape sparingly

The best practice in templates is to avoid passing in functional code or markup with skip_autoescape unless you are absolutely sure you have strict control over what's being passed. If you're passing user input, you could be opening your workers up to a Cross Site Scripting attack.

to_json

to_json will encode what you feed it to JSON (JavaScript Object Notation). If you feed it an object, it will serialize it.

grant_read_access

grant_read_access takes an S3 URI and encodes it into an HTTPS URL with a short-lived access token for that resource. This makes it possible to display to workers photo, audio, or video objects stored in S3 buckets that are not otherwise publicly accessible.

Example of the filters

Input

| auto-escape: {{ "Have you read 'James & the Giant Peach'?" }} |
| explicit escape: {{ "Have you read 'James & the Giant Peach'?" | escape }} |
| explicit escape_once: {{ "Have you read 'James &amp; the Giant Peach'?" | escape_once }} |
| skip_autoescape: {{ "Have you read 'James & the Giant Peach'?" | skip_autoescape }} |
| to_json: {{ jsObject | to_json }} |
| grant_read_access: {{ "s3://mybucket/myphoto.png" | grant_read_access }} |

Example

Output

auto-escape: Have you read &quot;James &amp; the Giant Peach&quot;?
explicit escape: Have you read &quot;James &amp; the Giant Peach&quot;
Example of an automated classification template.

To automate the simple text classification sample, replace the tweet text with a variable.

The text classification template is below with automation added. The changes/additions are highlighted in bold.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
  <crowd-classifier
    name="tweetFeeling"
    categories="['positive', 'negative', 'neutral', 'cannot determine']"
    header="Which term best describes this tweet?"
  >
    <classification-target>
      {{ task.input.source }}
    </classification-target>

    <full-instructions header="Analyzing a sentiment">
      Try to determine the feeling the author of the tweet is trying to express. If none seem to match, choose "other."
    </full-instructions>

    <short-instructions>
      Pick the term best describing the sentiment of the tweet.
    </short-instructions>
  </crowd-classifier>
</crowd-form>
```

The tweet text that was in the prior sample is now replaced with an object. The entry.taskInput object uses source (or another name you specify in your pre-annotation Lambda) as the property name for the text and it is inserted directly in the HTML by virtue of being between double curly braces.
End-to-end demos

You can view the following end-to-end demos which include sample Lambda function:

- Demo Template: Annotation of Images with crowd-bounding-box
- Demo Template: Labeling Intents with crowd-classifier

Step 3: Processing with AWS Lambda

In this step, you learn how to create and specify the two types of AWS Lambda functions that are required to create a custom labeling workflow:

- **Pre-annotation Lambda**: This function initiates for and pre-processes each data object sent to your labeling job prior to sending it to workers.
- **Post-annotation Lambda**: This function processes the results once workers submit a task. If you specify multiple workers per data object, this function may include logic to consolidate annotations.

If you are a new user of Lambda and Ground Truth, we recommend that you use the pages in this section as follows:

1. First, review Pre-annotation and Post-annotation Lambda Function Requirements.
2. Then, use the page Required Permissions To Use AWS Lambda With Ground Truth to learn about security and permission requirements to use your pre-annotation and post-annotation Lambda functions in a Ground Truth custom labeling job.
3. Next, you need to visit the Lambda console or use Lambda's APIs to create your functions. Use the section Create Lambda Functions for a Custom Labeling Workflow to learn how to create Lambda functions.
4. To learn how to test your Lambda functions, see Test Pre-Annotation and Post-Annotation Lambda Functions.
5. After you create pre-processing and post-processing Lambda functions, select them from the Lambda functions section that comes after the code editor for your custom HTML in the Ground Truth console. To learn how to use these functions in a CreateLabelingJob API request, see Create a Labeling Job (API).
For a custom labeling workflow tutorial that includes example pre-annotation and post-annotation Lambda functions, in the "Demo Template: Annotation of Images with crowd-bounding-box" document.

Topics

- Pre-annotation and Post-annotation Lambda Function Requirements
- Required Permissions To Use AWS Lambda With Ground Truth
- Create Lambda Functions for a Custom Labeling Workflow
- Test Pre-Annotation and Post-Annotation Lambda Functions

Pre-annotation and Post-annotation Lambda Function Requirements

Use this section to learn about the syntax of the requests sent to pre-annotation and post-annotation Lambda functions, and the response syntax that Ground Truth requires to run a custom labeling workflow.

Topics

- Pre-annotation Lambda
- Post-annotation Lambda

Pre-annotation Lambda

Before a labeling task is sent to the worker, your pre-annotation Lambda function is invoked.

Ground Truth sends your Lambda function a JSON-formatted request to provide details about the labeling job and the data object. The following table contains the pre-annotation request schemas. Each parameter is described below.

Data object identified with "source-ref"

```json
{
   "version": "2018-10-16",
   "labelingJobArn": "<labelingJobArn>
   "dataObject" : {
       "source-ref": "<s3Uri>
   }
}
```
### Data object identified with "source"

```json
{
    "version": "2018-10-16",
    "labelingJobArn": "<labelingJobArn>
    "dataObject" : {
        "source": "<string>
    }
}
```

- **version (string):** This is a version number used internally by Ground Truth.

- **labelingJobArn (string):** This is the Amazon Resource Name, or ARN, of your labeling job. This ARN can be used to reference the labeling job when using Ground Truth API operations such as `DescribeLabelingJob`.

- **dataObject (JSON object):** The key contains a single JSON line, either from your input manifest file or sent from Amazon SNS. The JSON line objects in your manifest can be up to 100 kilobytes in size and contain a variety of data. For a very basic image annotation job, the `dataObject` JSON may just contain a `source-ref` key, identifying the image to be annotated. If the data object (for example, a line of text) is included directly in the input manifest file, the data object is identified with `source`. If you create a verification or adjustment job, this line may contain label data and metadata from the previous labeling job.

The following table includes code block examples of a pre-annotation request. Each parameter in these example requests is explained below the tabbed table.

### Data object identified with "source-ref"

```json
{
    "version": "2018-10-16",
    "labelingJobArn": "arn:aws:sagemaker:<aws_region>:<aws_account_number>:labeling-job/<labeling_job_name>
    "dataObject" : {
        "source-ref": "s3://<input-data-bucket>/<data-object-file-name>
    }
}
```

#### Creating Custom Labeling Workflows

1548
Data object identified with "source"

```json
{
    "version": "2018-10-16",
    "labelingJobArn": "arn:aws:sagemaker:<aws_region>:<aws_account_number>:labeling-job/<labeling_job_name>"
    "dataObject": {
        "source": "Sue purchased 10 shares of the stock on April 10th, 2020"
    }
}
```

In return, Ground Truth requires a response formatted like the following:

**Example of expected return data**

```json
{
    "taskInput": <json object>,
    "isHumanAnnotationRequired": <boolean> # Optional
}
```

In the previous example, the `<json object>` needs to contain all the data your custom worker task template needs. If you're doing a bounding box task where the instructions stay the same all the time, it may just be the HTTP(S) or Amazon S3 resource for your image file. If it's a sentiment analysis task and different objects may have different choices, it is the object reference as a string and the choices as an array of strings.

**Implications of `isHumanAnnotationRequired`**

This value is optional because it defaults to `true`. The primary use case for explicitly setting it is when you want to exclude this data object from being labeled by human workers.

If you have a mix of objects in your manifest, with some requiring human annotation and some not needing it, you can include a `isHumanAnnotationRequired` value in each data object. You can add logic to your pre-annotation Lambda to dynamically determine if an object requires annotation, and set this boolean value accordingly.
Examples of Pre-annotation Lambda Functions

The following, basic pre-annotation Lambda function accesses the JSON object in dataObject from the initial request, and returns it in the taskInput parameter.

```python
import json

def lambda_handler(event, context):
    return {
        "taskInput":  event['dataObject']
    }
```

Assuming the input manifest file uses "source-ref" to identify data objects, the worker task template used in the same labeling job as this pre-annotation Lambda must include a Liquid element like the following to ingest dataObject:

```liquid
{{ task.input.source-ref | grant_read_access }}
```

If the input manifest file used source to identify the data object, the work task template can ingest dataObject with the following:

```liquid
{{ task.input.source }}
```

The following pre-annotation Lambda example includes logic to identify the key used in dataObject, and to point to that data object using taskObject in the Lambda's return statement.

```python
import json

def lambda_handler(event, context):
    # Event received
    print("Received event: " + json.dumps(event, indent=2))

    # Get source if specified
    source = event['dataObject']['source'] if "source" in event['dataObject'] else None

    # Get source-ref if specified
    source_ref = event['dataObject']['source-ref'] if "source-ref" in event['dataObject'] else None
```
# if source field present, take that otherwise take source-ref
task_object = source if source is not None else source_ref

# Build response object
output = {
    "taskInput": {
        "taskObject": task_object
    },
    "humanAnnotationRequired": "true"
}

print(output)
# If neither source nor source-ref specified, mark the annotation failed
if task_object is None:
    print("Failed to pre-process {} !".format(event["labelingJobArn"]))
    output["humanAnnotationRequired"] = "false"

return output

Post-annotation Lambda

When all workers have annotated the data object or when TaskAvailabilityLifetimeInSeconds has been reached, whichever comes first, Ground Truth sends those annotations to your post-annotation Lambda. This Lambda is generally used for Consolidate Annotations.

Tip

To see an example of a post-consolidation Lambda function, see annotation_consolidation_lambda.py in the aws-sagemaker-ground-truth-recipe GitHub repository.

The following code block contains the post-annotation request schema. Each parameter is described in the following bulleted list.

```json
{
    "version": "2018-10-16",
    "labelingJobArn": <string>,
    "labelCategories": [<string>],
```
The following code block contains an example of a post-annotation request. Each parameter in this example request is explained below the code block.

**Example of an post-annotation Lambda request**

```json
{
   "version": "2018-10-16",
   "labelCategories": ["Ex Category1","Ex Category2", "Ex Category3"],
   "labelAttributeName": "labeling-job-attribute-name",
   "roleArn": "arn:aws:iam::111122223333:role/role-name",
   "payload": {
      "s3Uri": "s3://DOC-EXAMPLE-BUCKET/annotations.json"
   }
}
```
If no worker works on the data object and `TaskAvailabilityLifetimeInSeconds` has been reached, the data object is marked as failed and not included as part of post-annotation Lambda invocation.

The following code block contains the payload schema. This is the file that is indicated by the `s3Uri` parameter in the post-annotation Lambda request payload JSON object. For example, if the previous code block is the post-annotation Lambda request, the following annotation file is located at `s3://DOC-EXAMPLE-BUCKET/annotations.json`.

Each parameter is described in the following bulleted list.

**Example of an annotation file**

```json
[
  {
    "datasetObjectId": "string",
    "dataObject": {
      "s3Uri": "string",
      "content": "string"
    },
    "annotations": [
      {
        "workerId": "string",
        "annotationData": {
          "content": "string",
          "s3Uri": "string"
        }
      }
    ]
  }
]
```

- **datasetObjectId (string)**: Identifies a unique ID that Ground Truth assigns to each data object you send to the labeling job.

- **dataObject (JSON object)**: The data object that was labeled. If the data object is included in the input manifest file and identified using the `source` key (for example, a string), `dataObject` includes a `content` key, which identifies the data object. Otherwise, the location of the data object (for example, a link or S3 URI) is identified with `s3Uri`. 
• annotations (list of JSON objects): This list contains a single JSON object for each annotation submitted by workers for that dataObject. A single JSON object contains a unique workerId that can be used to identify the worker that submitted that annotation. The annotationData key contains one of the following:
  • content (string): Contains the annotation data.
  • s3Uri (string): Contains an S3 URI that identifies the location of the annotation data.

The following table contains examples of the content that you may find in payload for different types of annotation.

### Named Entity Recognition Payload

```
[
  {
    "datasetObjectId": "1",
    "dataObject": {
      "content": "Sift 3 cups of flour into the bowl."
    },
    "annotations": [
      {
        "workerId": "private.us-west-2.ef7294f850a3d9d1",
        "annotationData": {
          "content": "{"crowd-entity-annotation":{"entities":[{"endOffset":4,"label":"verb","startOffset":0},{"endOffset":6,"label":"number","startOffset":5},{"endOffset":20,"label":"object","startOffset":15},{"endOffset":34,"label":"object","startOffset":30}]}"
        }
      }
    ]
  }
]
```

### Semantic Segmentation Payload

```
[
  {
    "datasetObjectId": "2",
    "dataObject": {
      "s3Uri": "s3://DOC-EXAMPLE-BUCKET/gt-input-data/images/bird3.jpg"
    },
    "annotations": [
```

Creating Custom Labeling Workflows
Your post-annotation Lambda function may contain logic similar to the following to loop through and access all annotations contained in the request. For a full example, see annotation_consolidation_lambda.py in the aws-sagemaker-ground-truth-recipe GitHub repository. In this GitHub example, you must add your own annotation consolidation logic.

```python
for i in range(len(annotations)):
    worker_id = annotations[i]['workerId']
    annotation_content = annotations[i]['annotationData'].get('content')
```
annotation_s3_uri = annotations[i]["annotationData'].get('s3uri')
annotation = annotation_content if annotation_s3_uri is None else
s3_client.get_object_from_s3(
    annotation_s3_uri)
annotation_from_single_worker = json.loads(annotation)

print("{} Received Annotations from worker [{}]) is [{]]"
    .format(log_prefix, worker_id, annotation_from_single_worker))

Tip

When you run consolidation algorithms on the data, you can use an AWS database service to store results, or you can pass the processed results back to Ground Truth. The data you return to Ground Truth is stored in consolidated annotation manifests in the S3 bucket specified for output during the configuration of the labeling job.

In return, Ground Truth requires a response formatted like the following:

Example of expected return data

```json
[

  {
    "datasetObjectId": <string>,
    "consolidatedAnnotation": {
      "content": {
        "<labelattributename>": {
          # ... label content
        }
      }
    }
  },

  {
    "datasetObjectId": <string>,
    "consolidatedAnnotation": {
      "content": {
        "<labelattributename>": {
          # ... label content
        }
      }
    }
  }

]```
At this point, all the data you're sending to your S3 bucket, other than the datasetObjectId, is in the content object.

When you return annotations in content, this results in an entry in your job's output manifest like the following:

**Example of label format in output manifest**

```json
{
    "source-ref"/"source" : "<s3uri or content>",
    "<labelAttributeName>": {
        # ... label content from you
    },
    "<labelAttributeName>-metadata": {
        # This will be added by Ground Truth
        "job_name": <labelingJobName>,
        "type": "groundTruth/custom",
        "human-annotated": "yes",
        "creation_date": <date> # Timestamp of when received from Post-labeling Lambda
    }
}
```

Because of the potentially complex nature of a custom template and the data it collects, Ground Truth does not offer further processing of the data.

**Required Permissions To Use AWS Lambda With Ground Truth**

You may need to configure some or all the following to create and use AWS Lambda with Ground Truth.

- You need to grant an IAM role or user (collectively, an IAM entity) permission to create the pre-annotation and post-annotation Lambda functions using AWS Lambda, and to choose them when creating the labeling job.
- The IAM execution role specified when the labeling job is configured needs permission to invoke the pre-annotation and post-annotation Lambda functions.
- The post-annotation Lambda functions may need permission to access Amazon S3.
Use the following sections to learn how to create the IAM entities and grant permissions described above.

**Topics**

- Grant Permission to Create and Select an AWS Lambda Function
- Grant IAM Execution Role Permission to Invoke AWS Lambda Functions
- Grant Post-Annotation Lambda Permissions to Access Annotation

**Grant Permission to Create and Select an AWS Lambda Function**

If you do not require granular permissions to develop pre-annotation and post-annotation Lambda functions, you can attach the AWS managed policy `AWSLambda_FullAccess` to a user or role. This policy grants broad permissions to use all Lambda features, as well as permission to perform actions in other AWS services with which Lambda interacts.

To create a more granular policy for security-sensitive use cases, refer to the documentation [Identity-based IAM policies for Lambda](#) in the to AWS Lambda Developer Guide to learn how to create an IAM policy that fits your use case.

**Policies to Use the Lambda Console**

If you want to grant an IAM entity permission to use the Lambda console, see [Using the Lambda console](#) in the AWS Lambda Developer Guide.

Additionally, if you want the user to be able to access and deploy the Ground Truth starter pre-annotation and post-annotation functions using the AWS Serverless Application Repository in the Lambda console, you must specify the `<aws-region>` where you want to deploy the functions (this should be the same AWS Region used to create the labeling job), and add the following policy to the IAM role.

```json
{
    "Version": "2012-10-17",
    "Statement": [
    {
        "Sid": "VisualEditor0",
        "Effect": "Allow",
        "Action": [
            "serverlessrepo:ListApplicationVersions",
            "serverlessrepo:GetApplication",
            "serverlessrepo:CreateApplication",
            "serverlessrepo:UpdateApplication",
            "serverlessrepo:DeleteApplication",
            "serverlessrepo:UploadApplication",
            "serverlessrepo:DownloadApplication",
            "serverlessrepo:TagApplication",
            "serverlessrepo:UntagApplication",
            "serverlessrepo:DescribeApplication",
            "serverlessrepo:DescribeApplicationRevision",
            "serverlessrepo:ListApplicationRevisions",
            "serverlessrepo:ListApplications",
            "serverlessrepo:DescribeApplicationVersion",
            "serverlessrepo:ListApplicationVersions",
            "serverlessrepo:GetApplicationVersion",
            "serverlessrepo:CreateDeployment",
            "serverlessrepo:DescribeDeployment",
            "serverlessrepo:DeleteDeployment",
            "serverlessrepo:StartHealthCheck",
            "serverlessrepo:StopHealthCheck",
            "serverlessrepo:GetHealthCheck",
            "serverlessrepo:GetHealthCheckResult",
            "serverlessrepo:CreateBucket",
            "serverlessrepo:UpdateBucket",
            "serverlessrepo:DeleteBucket",
            "serverlessrepo:GetBucket",
            "serverlessrepo:ListBucket",
            "serverlessrepo:TagBucket",
            "serverlessrepo:UntagBucket",
            "serverlessrepo:ListBucketVersions",
            "serverlessrepo:ListBucketVersion`
```
Policies to See Lambda Functions in the Ground Truth Console

To grant an IAM entity permission to view Lambda functions in the Ground Truth console when the user is creating a custom labeling job, the entity must have the permissions described in Grant IAM Permission to Use the Amazon SageMaker Ground Truth Console, including the permissions described in the section Custom Labeling Workflow Permissions.

Grant IAM Execution Role Permission to Invoke AWS Lambda Functions

If you add the IAM managed policy AmazonSageMakerGroundTruthExecution to the IAM execution role used to create the labeling job, this role has permission to list and invoke Lambda functions with one of the following strings in the function name: GtRecipe, SageMaker, Sagemaker, sagemaker, or LabelingFunction.

If the pre-annotation or post-annotation Lambda function names do not include one of the terms in the preceding paragraph, or if you require more granular permission than those in the AmazonSageMakerGroundTruthExecution managed policy, you can add a policy similar to the following to give the execution role permission to invoke pre-annotation and post-annotation functions.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": "lambda:InvokeFunction",
            "Resource": [
                "serverlessrepo:CreateCloudFormationTemplate"
            ]
        },
        {
            "Sid": "VisualEditor1",
            "Effect": "Allow",
            "Action": "serverlessrepo:SearchApplications",
            "Resource": "*"
        }
    ]
}
```
Grant Post-Annotation Lambda Permissions to Access Annotation

As described in Post-annotation Lambda, the post-annotation Lambda request includes the location of the annotation data in Amazon S3. This location is identified by the s3Uri string in the payload object. To process the annotations as they come in, even for a simple pass through function, you need to assign the necessary permissions to the post-annotation Lambda execution role to read files from the Amazon S3.

There are many ways that you can configure your Lambda to access annotation data in Amazon S3. Two common ways are:

- Allow the Lambda execution role to assume the SageMaker execution role identified in roleArn in the post-annotation Lambda request. This SageMaker execution role is the one used to create the labeling job, and has access to the Amazon S3 output bucket where the annotation data is stored.
- Grant the Lambda execution role permission to access the Amazon S3 output bucket directly.

Use the following sections to learn how to configure these options.

Grant Lambda Permission to Assume SageMaker Execution Role

To allow a Lambda function to assume a SageMaker execution role, you must attach a policy to the Lambda function's execution role, and modify the trust relationship of the SageMaker execution role to allow Lambda to assume it.

1. **Attach the following IAM policy** to your Lambda function's execution role to assume the SageMaker execution role identified in Resource. Replace 222222222222 with an AWS account ID. Replace sm-execution-role with the name of the assumed role.

```json
{
    "Version": "2012-10-17",
```
2. **Modify the trust policy** of the SageMaker execution role to include the following **Statement**. Replace `222222222222` with an [AWS account ID](https://aws.amazon.com/). Replace `my-lambda-execution-role` with the name of the assumed role.

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::222222222222:role/my-lambda-execution-role"
      },
      "Action": "sts:AssumeRole"
    }
  ]
}
```

---

**Grant Lambda Execution Role Permission to Access S3**

You can add a policy similar to the following to the post-annotation Lambda function execution role to give it S3 read permissions. Replace `DOC-EXAMPLE-BUCKET` with the name of the output bucket you specify when you create a labeling job.

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": ["s3:GetObject"],
      "Resource": "arn:aws:s3:::DOC-EXAMPLE-BUCKET/*"
    }
  ]
}
```
To add S3 read permissions to a Lambda execution role in the Lambda console, use the following procedure.

**Add S3 read permissions to post-annotation Lambda:**

1. Open the [Functions page](#) in the Lambda console.
2. Choose the name of the post-annotation function.
3. Choose **Configuration** and then choose **Permissions**.
4. Select the **Role name** and the summary page for that role opens in the IAM console in a new tab.
5. Select **Attach policies**.
6. Do one of the following:
   - Search for and select **AmazonS3ReadOnlyAccess** to give the function permission to read all buckets and objects in the account.
   - If you require more granular permissions, select **Create policy** and use the policy example in the preceding section to create a policy. Note that you must navigate back to the execution role summary page after you create the policy.
7. If you used the AmazonS3Read0nlyAccess managed policy, select **Attach policy**.
   
   If you created a new policy, navigate back to the Lambda execution role summary page and attach the policy you just created.

**Create Lambda Functions for a Custom Labeling Workflow**

You can create a Lambda function using the Lambda console, the AWS CLI, or an AWS SDK in a supported programming language of your choice. Use the AWS Lambda Developer Guide to learn more about each of these options:

- To learn how to create a Lambda function using the console, see [Create a Lambda function with the console](#).
- To learn how to create a Lambda function using the AWS CLI, see [Using AWS Lambda with the AWS Command Line Interface](#).
Select the relevant section in the table of contents to learn more about working with Lambda in the language of your choice. For example, select Working with Python to learn more about using Lambda with the AWS SDK for Python (Boto3).

Ground Truth provides pre-annotation and post-annotation templates through an AWS Serverless Application Repository (SAR) recipe. Use the following procedure to select the Ground Truth recipe in the Lambda console.

**Use the Ground Truth SAR recipe to create pre-annotation and post-annotation Lambda functions:**

1. Open the Functions page on the Lambda console.
2. Select Create function.
3. Select Browse serverless app repository.
4. In the search text box, enter aws-sagemaker-ground-truth-recipe and select that app.
5. Select Deploy. The app may take a couple of minutes to deploy.

Once the app deploys, two functions appear in the Functions section of the Lambda console: serverlessrepo-aws-sagema-GtRecipePreHumanTaskFunc-<id> and serverlessrepo-aws-sagema-GtRecipeAnnotationConsol-<id>.

6. Select one of these functions and add your custom logic in the Code section.
7. When you are finished making changes, select Deploy to deploy them.

**Test Pre-Annotation and Post-Annotation Lambda Functions**

You can test your pre-annotation and post annotation Lambda functions in the Lambda console. If you are a new user of Lambda, you can learn how to test, or invoke, your Lambda functions in the console using the Create a Lambda function tutorial with the console in the AWS Lambda Developer Guide.

You can use the sections on this page to learn how to test the Ground Truth pre-annotation and post-annotation templates provided through an AWS Serverless Application Repository (SAR).

**Topics**

- Prerequisites
- Test the Pre-annotation Lambda Function
Test the Post-Annotation Lambda Function

Prerequisites

You must do the following to use the tests described on this page.

- You need access to the Lambda console, and you need permission to create and invoke Lambda functions. To learn how to set up these permissions, see [Grant Permission to Create and Select an AWS Lambda Function](#).
- If you have not deployed the Ground Truth SAR recipe, use the procedure in [Create Lambda Functions for a Custom Labeling Workflow](#) to do so.
- To test the post-annotation Lambda function, you must have a data file in Amazon S3 with sample annotation data. For a simple test, you can copy and paste the following code into a file and save it as `sample-annotations.json` and upload this file to Amazon S3. Note the S3 URI of this file—you need this information to configure the post-annotation Lambda test.

```json
[{
  "datasetObjectId": "0",
  "dataObject": {
    "content": "To train a machine learning model, you need a large, high-quality, labeled dataset. Ground Truth helps you build high-quality training datasets for your machine learning models."},
  "annotations": [
    {
      "workerId": "private.us-west-2.0123456789",
      "annotationData": {
        "content": ""Verb: to train a machine learning model, you need a large, high-quality, labeled dataset. Ground Truth helps you build high-quality training datasets for your machine learning models."",
        "crowd-entity-annotation": {
          "entities": [
            {"endOffset": 8, "label": "verb", "startOffset": 0},
            {"endOffset": 13, "label": "object", "startOffset": 11},
            {"endOffset": 17, "label": "adjective", "startOffset": 15},
            {"endOffset": 18, "label": "adjective", "startOffset": 16}
          ]
        }
      }
    }
  ]
},
{
  "datasetObjectId": "1",
  "dataObject": {
    "content": "Sift 3 cups of flour into the bowl."},
  "annotations": [
    {
      "workerId": "private.us-west-2.0123456789",
      "annotationData": {
        "content": ""Sift 3 cups of flour into the bowl."",
        "crowd-entity-annotation": {
          "entities": [
            {"endOffset": 2, "label": "verb", "startOffset": 0},
            {"endOffset": 5, "label": "number", "startOffset": 3},
            {"endOffset": 7, "label": "object", "startOffset": 5},
            {"endOffset": 11, "label": "object", "startOffset": 9},
            {"endOffset": 16, "label": "object", "startOffset": 14}
          ]
        }
      }
    }
  ]
},
{
  "datasetObjectId": "2",
  "dataObject": {
    "content": "Jen purchased 10 shares of the stock on January 1st, 2020."},
  "annotations": [
    {
      "workerId": "private.us-west-2.0123456789",
      "annotationData": {
        "content": "Jen purchased 10 shares of the stock on January 1st, 2020.",
        "crowd-entity-annotation": {
          "entities": [
            {"endOffset": 0, "label": "person", "startOffset": 0},
            {"endOffset": 1, "label": "object", "startOffset": 0}
          ]
        }
      }
    }
  ]
}]
```
Creating Custom Labeling Workflows

You must use the directions in Grant Post-Annotation Lambda Permissions to Access Annotation to give your post-annotation Lambda function's execution role permission to assume the SageMaker execution role you use to create the labeling job. The post-annotation Lambda function uses the SageMaker execution role to access the annotation data file, sample-annotations.json, in S3.

Test the Pre-annotation Lambda Function

Use the following procedure to test the pre-annotation Lambda function created when you deployed the Ground Truth AWS Serverless Application Repository (SAR) recipe.

Test the Ground Truth SAR recipe pre-annotation Lambda function

1. Open the Functions page in the Lambda console.
2. Select the pre-annotation function that was deployed from the Ground Truth SAR recipe. The name of this function is similar to serverlessrepo-aws-sagemaker-GtRecipePreHumanTaskFunc-<id>.
3. In the Code source section, select the arrow next to Test.
4. Select Configure test event.
5. Keep the Create new test event option selected.
7. Give your test an Event name.
8. Select Create.
9. Select the arrow next to Test again and you should see that the test you created is selected, which is indicated with a dot by the event name. If it is not selected, select it.
10. Select Test to run the test.
After you run the test, you can see the **Execution results**. In the **Function logs**, you should see a response similar to the following:

```
START RequestId: cd117d38-8365-4e1a-bffb-0dcd631a878f Version: $LATEST
   Received event: {
      "version": "2018-10-16",
      "dataObject": {
         "source-ref": "s3://sagemakerexample/object_to_annotate.jpg"
      }
   }
   'taskInput': {'taskObject': 's3://sagemakerexample/object_to_annotate.jpg'},
   'isHumanAnnotationRequired': 'true'}
END RequestId: cd117d38-8365-4e1a-bffb-0dcd631a878f
REPORT RequestId: cd117d38-8365-4e1a-bffb-0dcd631a878f Duration: 0.42 ms Billed Duration: 1 ms Memory Size: 128 MB Max Memory Used: 43 MB
```

In this response, we can see the Lambda function's output matches the required pre-annotation response syntax:

```
{ 'taskInput': {'taskObject': 's3://sagemakerexample/object_to_annotate.jpg'},
   'isHumanAnnotationRequired': 'true'}
```

**Test the Post-Annotation Lambda Function**

Use the following procedure to test the post-annotation Lambda function created when you deployed the Ground Truth AWS Serverless Application Repository (SAR) recipe.

**Test the Ground Truth SAR recipe post-annotation Lambda**

1. Open the **Functions page** in the Lambda console.
2. Select the post-annotation function that was deployed from the Ground Truth SAR recipe. The name of this function is similar to `serverlessrepo-aws-sagemaker-GtRecipeAnnotationConsol-<id>`.
3. In the **Code source** section, select the arrow next to **Test**.
4. Select **Configure test event**.
5. Keep the **Create new test event** option selected.
6. Under **Event template**, select **SageMaker Ground Truth AnnotationConsolidation**.
7. Give your test an **Event name**.

8. Modify the template code provided as follows:

   - Replace the Amazon Resource Name (ARN) in `roleArn` with the ARN of the SageMaker execution role you used to create the labeling job.
   - Replace the S3 URI in `s3Uri` with the URI of the `sample-annotations.json` file you added to Amazon S3.

   After you make these modifications, your test should look similar to the following:

   ```json
   {
     "version": "2018-10-16",
     "labelAttributeName": "example-attribute",
     "roleArn": "arn:aws:iam::222222222222:role/sm-execution-role",
     "payload": {
       "s3Uri": "s3://your-bucket/sample-annotations.json"
     }
   }
   ```

9. Select **Create**.

10. Select the arrow next to **Test** again and you should see that the test you created is selected, which is indicated with a dot by the event name. If it is not selected, select it.

11. Select the **Test** to run the test.

   After you run the test, you should see a -- Consolidated Output -- section in the **Function Logs**, which contains a list of all annotations included in `sample-annotations.json`.

### Demo Template: Annotation of Images with crowd-bounding-box

When you chose to use a custom template as your task type in the Amazon SageMaker Ground Truth console, you reach the **Custom labeling task panel**. There you can choose from multiple base templates. The templates represent some of the most common tasks and provide a sample to work from as you create your customized labeling task's template. If you are not using the console, or as an additional recourse, see [Amazon SageMaker Ground Truth Sample Task UIs](#) for a repository of demo templates for a variety of labeling job task types.
This demonstration works with the BoundingBox template. The demonstration also works with the AWS Lambda functions needed for processing your data before and after the task. In the Github repository above, to find templates that work with AWS Lambda functions, look for `{{ task.input.<property name> }}` in the template.

**Topics**

- [Starter Bounding Box custom template](#)
- [Your own Bounding Box custom template](#)
- [Your manifest file](#)
- [Your pre-annotation Lambda function](#)
- [Your post-annotation Lambda function](#)
- [The output of your labeling job](#)

**Starter Bounding Box custom template**

This is the starter bounding box template that is provided.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>

  <crowd-bounding-box
    name="boundingBox"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="{{ task.input.header }}"
    labels="{{ task.input.labels | to_json | escape }}"
  >

  <!-- The <full-instructions> tag is where you will define the full instructions of your task. -->
  <full-instructions header="Bounding Box Instructions" >
  <p>Use the bounding box tool to draw boxes around the requested target of interest:</p>
  <ol>
    <li>Draw a rectangle using your mouse over each instance of the target.</li>
    <li>Make sure the box does not cut into the target, leave a 2 - 3 pixel margin</li>
    <li>When targets are overlapping, draw a box around each object, include all contiguous parts of the target in the box.</li>
  </ol>

</full-instructions>

</crowd-form>
```
The custom templates use the Liquid template language, and each of the items between double curly braces is a variable. The pre-annotation AWS Lambda function should provide an object named taskInput and that object's properties can be accessed as `{{ task.input.<property name> }}` in your template.

**Your own Bounding Box custom template**

As an example, assume you have a large collection of animal photos in which you know the kind of animal in an image from a prior image-classification job. Now you want to have a bounding box drawn around it.

In the starter sample, there are three variables: taskObject, header, and labels.

Each of these would be represented in different parts of the bounding box.

- **taskObject** is an HTTP(S) URL or S3 URI for the photo to be annotated. The added `grant_read_access` is a filter that will convert an S3 URI to an HTTPS URL with short-lived access to that resource. If you're using an HTTP(S) URL, it's not needed.
- **header** is the text above the photo to be labeled, something like "Draw a box around the bird in the photo."
• labels is an array, represented as ['item1', 'item2', ...]. These are labels that can be assigned by the worker to the different boxes they draw. You can have one or many.

Each of the variable names come from the JSON object in the response from your pre-annotation Lambda. The names above are merely suggested. Use whatever variable names make sense to you and will promote code readability among your team.

⚠️ Only use variables when necessary

If a field will not change, you can remove that variable from the template and replace it with that text, otherwise you have to repeat that text as a value in each object in your manifest or code it into your pre-annotation Lambda function.

Example: Final Customized Bounding Box Template

To keep things simple, this template will have one variable, one label, and very basic instructions. Assuming your manifest has an "animal" property in each data object, that value can be re-used in two parts of the template.

```html
<full-instructions header="Bounding Box Instructions">
<p>Draw a bounding box around the {{ task.input.animal }} in the image. If there is more than one {{ task.input.animal }} per image, draw a bounding box around the largest one.</p>
<p>The box should be tight around the {{ task.input.animal }} with no more than a couple of pixels of buffer around the edges.</p>
<p>If the image does not contain a {{ task.input.animal }}, check the <strong>Nothing to label</strong> box.</p>
</full-instructions>

<short-instructions>
<p>Draw a bounding box around the {{ task.input.animal }} in each image. If there is more than one {{ task.input.animal }} per image, draw a bounding
```
Note the re-use of `{{ task.input.animal }}` throughout the template. If your manifest had all of the animal names beginning with a capital letter, you could use `{{ task.input.animal | downcase }}`, incorporating one of Liquid's built-in filters in sentences where it needed to be presented lowercase.

**Your manifest file**

Your manifest file should provide the variable values you're using in your template. You can do some transformation of your manifest data in your pre-annotation Lambda, but if you don't need to, you maintain a lower risk of errors and your Lambda will run faster. Here's a sample manifest file for the template.

```json
{"source-ref": "<S3 image URI>", "animal": "horse"}
{"source-ref": "<S3 image URI>", "animal": "bird"}
{"source-ref": "<S3 image URI>", "animal": "dog"}
{"source-ref": "<S3 image URI>", "animal": "cat"}
```

**Your pre-annotation Lambda function**

As part of the job set-up, provide the ARN of an AWS Lambda function that can be called to process your manifest entries and pass them to the template engine.

**Naming your Lambda function**

The best practice in naming your function is to use one of the following four strings as part of the function name: `SageMaker`, `Sagemaker`, `sagemaker`, or `LabelingFunction`. This applies to both your pre-annotation and post-annotation functions.

When you're using the console, if you have AWS Lambda functions that are owned by your account, a drop-down list of functions meeting the naming requirements will be provided to choose one.

In this very basic example, you're just passing through the information from the manifest without doing any additional processing on it. This sample pre-annotation function is written for Python 3.7.
The JSON object from your manifest will be provided as a child of the event object. The properties inside the taskInput object will be available as variables to your template, so simply setting the value of taskInput to event['dataObject'] will pass all the values from your manifest object to your template without having to copy them individually. If you wish to send more values to the template, you can add them to the taskInput object.

**Your post-annotation Lambda function**

As part of the job set-up, provide the ARN of an AWS Lambda function that can be called to process the form data when a worker completes a task. This can be as simple or complex as you want. If you want to do answer consolidation and scoring as it comes in, you can apply the scoring and/or consolidation algorithms of your choice. If you want to store the raw data for offline processing, that is an option.

### Provide permissions to your post-annotation Lambda

The annotation data will be in a file designated by the s3Uri string in the payload object. To process the annotations as they come in, even for a simple pass through function, you need to assign S3Read0nly access to your Lambda so it can read the annotation files.

In the Console page for creating your Lambda, scroll to the **Execution role** panel. Select **Create a new role from one or more templates**. Give the role a name. From the **Policy templates** drop-down, choose **Amazon S3 object read-only permissions**. Save the Lambda and the role will be saved and selected.

The following sample is in Python 2.7.

```python
import json
import boto3
from urlparse import urlparse

def lambda_handler(event, context):
    return {
        "taskInput": event['dataObject']
    }
```
consolidated_labels = []

parsed_url = urlparse(event['payload']['s3Uri']);
s3 = boto3.client('s3')
textFile = s3.get_object(Bucket = parsed_url.netloc, Key = parsed_url.path[1:])
filecont = textFile['Body'].read()
annotations = json.loads(filecont);

for dataset in annotations:
    for annotation in dataset['annotations']:
        new_annotation = json.loads(annotation['annotationData']['content'])
        label = {
            'datasetObjectIds': dataset['datasetObjectIds'],
            'consolidatedAnnotation': {
                'content': {
                    'event': event['labelAttributeName'], {
                        'workerId': annotation['workerId'],
                        'boxesInfo': new_annotation,
                        'imageSource': dataset['dataObject']
                }
            }
        }
        consolidated_labels.append(label)

return consolidated_labels

The post-annotation Lambda will often receive batches of task results in the event object. That batch will be the payload object the Lambda should iterate through. What you send back will be an object meeting the API contract.

**The output of your labeling job**

You'll find the output of the job in a folder named after your labeling job in the target S3 bucket you specified. It will be in a subfolder named manifests.

For a bounding box task, the output you find in the output manifest will look a bit like the demo below. The example has been cleaned up for printing. The actual output will be a single line per record.

**Example : JSON in your output manifest**

```json
{

}
Note how the additional animal attribute from your original manifest is passed to the output manifest on the same level as the source-ref and labeling data. Any properties from your input manifest, whether they were used in your template or not, will be passed to the output manifest.

**Demo Template: Labeling Intents with crowd-classifier**

If you choose a custom template, you'll reach the [Custom labeling task panel](#). There you can select from multiple starter templates that represent some of the more common tasks. The templates provide a starting point to work from in building your customized labeling task's template.

In this demonstration, you work with the Intent Detection template, which uses the [crowd-classifier](#) element, and the AWS Lambda functions needed for processing your data before and after the task.

**Topics**

- [Starter Intent Detection custom template](#)
- [Your Intent Detection custom template](#)
- [Your pre-annotation Lambda function](#)
- [Your post-annotation Lambda function](#)
- [Your labeling job output](#)
Starter Intent Detection custom template

This is the intent detection template that is provided as a starting point.

The custom templates use the Liquid template language, and each of the items between double curly braces is a variable. The pre-annotation AWS Lambda function should provide an object named taskInput and that object's properties can be accessed as {{ task.input.<property name> }} in your template.

Your Intent Detection custom template

In the starter template, there are two variables: the task.input.labels property in the crowd-classifier element opening tag and the task.input.utterance in the classification-target region's content.

Unless you need to offer different sets of labels with different utterances, avoiding a variable and just using text will save processing time and creates less possibility of error. The template used in
this demonstration will remove that variable, but variables and filters like `to_json` are explained in more detail in the [crowd-bounding-box demonstration] article.

**Styling Your Elements**

Two parts of these custom elements that sometimes get overlooked are the `<full-instructions>` and `<short-instructions>` regions. Good instructions generate good results.

In the elements that include these regions, the `<short-instructions>` appear automatically in the "Instructions" pane on the left of the worker's screen. The `<full-instructions>` are linked from the "View full instructions" link near the top of that pane. Clicking the link opens a modal pane with more detailed instructions.

You can not only use HTML, CSS, and JavaScript in these sections, you are encouraged to if you believe you can provide a strong set of instructions and examples that will help workers complete your tasks with better speed and accuracy.

**Example Try out a sample with JSFiddle**

Try out an example `<crowd-classifier>` task. The example is rendered by JSFiddle, therefore all the template variables are replaced with hard-coded values. Click the "View full instructions" link to see a set of examples with extended CSS styling. You can fork the project to experiment with your own changes to the CSS, adding sample images, or adding extended JavaScript functionality.

**Example : Final Customized Intent Detection Template**

This uses the example `<crowd-classifier>` task, but with a variable for the `<classification-target>`. If you are trying to keep a consistent CSS design among a series of
different labeling jobs, you can include an external stylesheet using a `<link rel...>` element the same way you'd do in any other HTML document.
Creating Custom Labeling Workflows
Example: Your manifest file

If you are preparing your manifest file manually for a text-classification task like this, have your data formatted in the following manner.
This differs from the manifest file used for the "Demo Template: Annotation of Images with crowd-bounding-box" demonstration in that source-ref was used as the property name instead of source. The use of source-ref designates S3 URIs for images or other files that must be converted to HTTP. Otherwise, source should be used like it is with the text strings above.

Your pre-annotation Lambda function

As part of the job set-up, provide the ARN of an AWS Lambda that can be called to process your manifest entries and pass them to the template engine.

This Lambda function is required to have one of the following four strings as part of the function name: SageMaker, Sagemaker, sagemaker, or LabelingFunction.

This applies to both your pre-annotation and post-annotation Lambdas.

When you're using the console, if you have Lambdas that are owned by your account, a drop-down list of functions meeting the naming requirements will be provided to choose one.

In this very basic sample, where you have only one variable, it's primarily a pass-through function. Here's a sample pre-labeling Lambda using Python 3.7.

```python
import json

def lambda_handler(event, context):
    return {
        "taskInput": event['dataObject']
    }
```

The dataObject property of the event contains the properties from a data object in your manifest.

In this demonstration, which is a simple pass through, you just pass that straight through as the taskInput value. If you add properties with those values to the event['dataObject'] object, they will be available to your HTML template as Liquid variables with the format `{{ task.input.<property name> }}`.  

Creating Custom Labeling Workflows

1580
Your post-annotation Lambda function

As part of the job set up, provide the ARN of a Lambda function that can be called to process the form data when a worker completes a task. This can be as simple or complex as you want. If you want to do answer-consolidation and scoring as data comes in, you can apply the scoring or consolidation algorithms of your choice. If you want to store the raw data for offline processing, that is an option.

Set permissions for your post-annotation Lambda function

The annotation data will be in a file designated by the s3Uri string in the payload object. To process the annotations as they come in, even for a simple pass through function, you need to assign S3Readonly access to your Lambda so it can read the annotation files. In the Console page for creating your Lambda, scroll to the Execution role panel. Select Create a new role from one or more templates. Give the role a name. From the Policy templates drop-down, choose Amazon S3 object read-only permissions. Save the Lambda and the role will be saved and selected.

The following sample is for Python 3.7.

```python
import json
import boto3
from urllib.parse import urlparse

def lambda_handler(event, context):
    consolidated_labels = []

    parsed_url = urlparse(event['payload']['s3Uri']);
    s3 = boto3.client('s3')
    textFile = s3.get_object(Bucket = parsed_url.netloc, Key = parsed_url.path[1:])
    filecont = textFile['Body'].read()
    annotations = json.loads(filecont);

    for dataset in annotations:
        for annotation in dataset['annotations']:
            new_annotation = json.loads(annotation['annotationData']['content'])
            label = {
                'datasetObjectId': dataset['datasetObjectId'],
                'consolidatedAnnotation' : {
                    'content': {
```
event['labelAttributeName']: {
'workerId': annotation['workerId'],
'result': new_annotation,
'labeledContent': dataset['dataObject']
}
}
}
consolidated_labels.append(label)

return consolidated_labels

Your labeling job output

The post-annotation Lambda will often receive batches of task results in the event object. That batch will be the payload object the Lambda should iterate through.

You'll find the output of the job in a folder named after your labeling job in the target S3 bucket you specified. It will be in a subfolder named manifests.

For an intent detection task, the output in the output manifest will look a bit like the demo below. The example has been cleaned up and spaced out to be easier for humans to read. The actual output will be more compressed for machine reading.

Example: JSON in your output manifest

```json
[
{
  "datasetObjectId":"<Number representing item's place in the manifest>",
  "consolidatedAnnotation":
  {
    "content":
    {
      "<name of labeling job>":
      {
        "workerId":"private.us-east-1.XXXXXXXXXXXXXXXXXXXXXXXXX",
        "result":
        {
          "intent":
          {
            "label": "<label chosen by worker>"
          }
        }
      }
    }
  }
}
```
This should help you create and use your own custom template.

**Custom Workflows via the API**

When you have created your custom UI template (Step 2) and processing Lambda functions (Step 3), you should place the template in an Amazon S3 bucket with a file name format of: `<FileName>.liquid.html`. 
Use the **CreateLabelingJob** action to configure your task. You'll use the location of a custom template (Step 2: Creating your custom worker task template) stored in a `<filename>`.liquid.html file on S3 as the value for the UiTemplateS3Uri field in the **UiConfig** object within the **HumanTaskConfig** object.

For the AWS Lambda tasks described in Step 3: Processing with AWS Lambda, the post-annotation task's ARN will be used as the value for the AnnotationConsolidationLambdaArn field, and the pre-annotation task will be used as the value for the PreHumanTaskLambdaArn.

**Create a Labeling Job**

You can create a labeling job in the Amazon SageMaker console and by using an AWS SDK in your preferred language to run CreateLabelingJob. After a labeling job has been created, you can track worker metrics (for private workforces) and your labeling job status using CloudWatch.

Before you create a labeling job it is recommended that you review the following pages, as applicable:

- You can specify your input data using an automatic data setup in the console, or an input manifest file in either the console or when using CreateLabelingJob API. For automated data setup, see Automated Data Setup. To learn how to create an input manifest file, see Use an Input Manifest File.
- Review labeling job input data quotas: Input Data Quotas.

After you have chosen your task type, use the topics on this page to learn how to create a labeling job.

If you are a new Ground Truth user, we recommend that you start by walking through the demo in Getting started.

**Important**

Ground Truth requires all S3 buckets that contain labeling job input image data to have a CORS policy attached. To learn more, see CORS Permission Requirement.

**Topics**

- Built-in Task Types
Built-in Task Types

Amazon SageMaker Ground Truth has several built-in task types. Ground Truth provides a worker task template for built-in task types. Additionally, some built in task types support Automate Data Labeling. The following topics describe each built-in task type and demo the worker task templates that are provided by Ground Truth in the console. To learn how to create a labeling job in the console using one of these task types, select the task type page.

<table>
<thead>
<tr>
<th>Label Images</th>
<th>Label Text</th>
<th>Label Videos and Video Frames</th>
<th>Label 3D Point Clouds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bounding Box</td>
<td>Named Entity Recognition</td>
<td>Video Classification</td>
<td>3D Point Cloud Object Detection</td>
</tr>
<tr>
<td>Image Classification (Single Label)</td>
<td>Text Classification (Single Label)</td>
<td>Video Frame Object Detection</td>
<td>3D Point Cloud Object Tracking</td>
</tr>
<tr>
<td>Image Classification (Multi-label)</td>
<td>Text Classification (Multi-label)</td>
<td>Video Frame Object Tracking</td>
<td>3D Point Cloud Semantic Segmentation</td>
</tr>
<tr>
<td>Image Semantic Segmentation</td>
<td>Verify and Adjust Labels</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Note**

Each of the video frame and 3D point cloud task types has an adjustment task type that you use to verify and adjust labels from a previous labeling job. Select a video frame or 3D point cloud task type page above to learn how to adjust labels created using that task type.
Creating Instruction Pages

Create custom instructions for labeling jobs to improve your worker's accuracy in completing their task. You can modify the default instructions that are provided in the console or you can create your own. The instructions are shown to the worker on the page where they complete their labeling task.

There are two kinds of instructions:

- **Short instructions**—instructions that are shown on the same webpage where the worker completes their task. These instructions should provide an easy reference to show the worker the correct way to label an object.

- **Full instructions**—instructions that are shown on a dialog box that overlays the page where the worker completes their task. We recommend that you provide detailed instructions for completing the task with multiple examples showing edge cases and other difficult situations for labeling objects.

Create instructions in the console when you are creating your labeling job. Start with the existing instructions for the task and use the editor to modify them to suit your labeling job.

⚠️ **Note**

Once you create your labeling job, it will automatically start and you will not be able to modify your worker instructions. If you need to change your worker instructions, stop the labeling job that you created, clone it, and modify your worker instructions before creating a new job.

You can clone a labeling job in the console by selecting the labeling job and then selecting **Clone** in the **Actions** menu.

To clone a labeling job using the Amazon SageMaker API or your preferred Amazon SageMaker SDK, make a new request to the `CreateLabelingJob` operation with the same specifications as your original job after modifying your worker instructions.

**Short Instructions**

Short instructions appear on the same web page that workers use to label your data object. For example, the following is the editing page for a bounding box task. The short instructions panel is on the left.
Keep in mind that a worker will only spend seconds looking at the short instructions. Workers must be able to scan and understand your information quickly. In all cases it should take less time to understand the instructions than it takes to complete the task. Keep these points in mind:

- Your instructions should be clear and simple.
- Pictures are better than words. Create a simple illustration of your task that your workers can immediately understand.
- If you must use words, use short, concise examples.
- Your short instructions are more important than your full instructions.
The Amazon SageMaker Ground Truth console provides an editor so that you can create your short instructions. Replace the placeholder text and images with instructions for your task. Preview the worker's task page by choosing Preview. The preview will open in a new window, be sure to turn off pop-up blocking so that the window will show.

**Full Instructions**

You can provide additional instructions for your workers in a dialog box that overlays the page where workers label your data objects. Use full instructions to explain more complex tasks and to show workers the proper way to label edge cases or other difficult objects.

You can create full instructions using an editor in the Ground Truth console. As with quick instructions, keep the following in mind:

- Workers will want detailed instruction the first few times that the complete your task. Any information that they *must* have should be in the quick instructions.
- Pictures are more important than words.
- Text should be concise.
- Full instructions should supplement the short instructions. Don't repeat information that appears in the short instructions.

The Ground Truth console provides an editor so that you can create your full instructions. Replace the placeholder text and images with instructions for your task. Preview the full instruction page by choosing Preview. The preview will open in a new window, be sure to turn off pop-up blocking so that the window will show.

**Add example images to your instructions**

Images provide useful examples for your workers. To add a publicly accessible image to your instructions:

- Place the cursor where the image should go in the instructions editor.
- Click the image icon in the editor toolbar.
- Enter the URL of your image.

If your instruction image in Amazon S3 is not publicly accessible:
• As the image URL, enter: `{{ 'https://s3.amazonaws.com/your-bucket-name/image-file-name' | grant_read_access }}`.

• This renders the image URL with a short-lived, one-time access code appended so the worker's browser can display it. A broken image icon is displayed in the instructions editor, but previewing the tool displays the image in the rendered preview.

Create a Labeling Job (Console)

You can use the Amazon SageMaker console to create a labeling job for all of the Ground Truth built-in task types and custom labeling workflows. For built-in task types, we recommend that you use this page alongside the page for your task type. Each task type page includes specific details on creating a labeling job using that task type.

You need to provide the following to create a labeling job in the SageMaker console:

• An input manifest file in Amazon S3. You can place your input dataset in Amazon S3 and automatically generate a manifest file using the Ground Truth console (not supported for 3D point cloud labeling jobs).

  Alternatively, you can manually create an input manifest file. To learn how, see Input Data.

• An Amazon S3 bucket to store your output data.

• An IAM role with permission to access your resources in Amazon S3 and with a SageMaker execution policy attached. For a general solution, you can attach the managed policy, AmazonSageMakerFullAccess, to an IAM role and include `sagemaker` in your bucket name.

  For more granular policies, see the section called “IAM Permissions”.

  3D point cloud task types have additional security considerations. Learn more.

• A work team. You create a work team from a workforce made up of Amazon Mechanical Turk workers, vendors, or your own private workers. To learn more, see Create and Manage Workforces.

  You cannot use the Mechanical Turk workforce for 3D point cloud or video frame labeling jobs.

• If you are using a custom labeling workflow, you must save a worker task template in Amazon S3 and provide an Amazon S3 URI for that template. For more information, see Step 2: Creating your custom worker task template.

• (Optional) An AWS KMS key ARN if you want SageMaker to encrypt the output of your labeling job using your own AWS KMS encryption key instead of the default Amazon S3 service key.
(Optional) Existing labels for the dataset you use for your labeling job. Use this option if you want workers to adjust, or approve and reject labels.

If you want to create an adjustment or verification labeling job, you must have an output manifest file in Amazon S3 that contains the labels you want adjusted or verified. This option is only supported for bounding box and semantic segmentation image labeling jobs and 3D point cloud and video frame labeling jobs. It is recommended that you use the instructions on Verify and Adjust Labels to create a verification or adjustment labeling job.

⚠️ **Important**

Your work team, input manifest file, output bucket, and other resources in Amazon S3 must be in the same AWS Region you use to create your labeling job.

When you create a labeling job using the SageMaker console, you add worker instructions and labels to the worker UI that Ground Truth provides. You can preview and interact with the worker UI while creating your labeling job in the console. You can also see a preview of the worker UI on your built-in task type page.

**To create a labeling job (console)**

2. In the left navigation pane, choose **Labeling jobs**.
3. On the **Labeling jobs** page, choose **Create labeling job**.
4. For **Job name**, enter a name for your labeling job.
5. (Optional) If you want to identify your labels with a key, select **I want to specify a label attribute name different from the labeling job name**. If you do not select this option, the labeling job name you specified in the previous step will be used to identify your labels in your output manifest file.
6. Choose a data setup to create a connection between your input dataset and Ground Truth.
   - For **Automated data setup**:
     - Follow the instructions in Automated Data Setup for image, text, and video clip labeling jobs.
     - Follow the instructions in Automated Video Frame Input Data Setup for video frame labeling jobs.
• For **Manual data setup:**
  • For **Input dataset location**, provide the location in Amazon S3 in which your input manifest file is located. For example, if your input manifest file, manifest.json, is located in `example.bucket`, enter `s3://example-bucket/manifest.json`.
  • For **Output dataset location**, provide the location in Amazon S3 where you want Ground Truth to store the output data from your labeling job.

7. For **IAM Role**, choose an existing IAM role or create an IAM role with permission to access your resources in Amazon S3, to write to the output Amazon S3 bucket specified above, and with a SageMaker execution policy attached.

8. (Optional) For **Additional configuration**, you can specify how much of your dataset you want workers to label, and if you want SageMaker to encrypt the output data for your labeling job using an AWS KMS encryption key. To encrypt your output data, you must have the required AWS KMS permissions attached to the IAM role you provided in the previous step. For more details, see the section called “IAM Permissions”.

9. In the **Task type** section, under **Task category**, use the dropdown list to select your task category.

10. In **Task selection**, choose your task type.

11. (Optional) Provide tags for your labeling job to make it easier to find in the console later.

12. Choose **Next**.

13. In the **Workers** section, choose the type of workforce you would like to use. For more details about your workforce options see Create and Manage Workforces.

14. (Optional) After you've selected your workforce, specify the **Task timeout**. This is the maximum amount of time a worker has to work on a task.

   For 3D point cloud annotation tasks, the default task timeout is 3 days. The default timeout for text and image classification and label verification labeling jobs is 5 minutes. The default timeout for all other labeling jobs is 60 minutes.

15. (Optional) For bounding box, semantic segmentation, video frame, and 3D point cloud task types, you can select **Display existing labels** if you want to display labels for your input data set for workers to verify or adjust.

   For bounding box and semantic segmentation labeling jobs, this will create an adjustment labeling job.

   For 3D point cloud and video frame labeling jobs:
- Select **Adjustment** to create an adjustment labeling job. When you select this option, you can add new labels but you cannot remove or edit existing labels from the previous job. Optionally, you can choose label category attributes and frame attributes that you want workers to edit. To make an attribute editable, select the check box **Allow workers to edit this attribute** for that attribute.

Optionally, you can add new label category and frame attributes.

- Select **Verification** to create an adjustment labeling job. When you select this option, you cannot add, modify, or remove existing labels from the previous job. Optionally, you can choose label category attributes and frame attributes that you want workers to edit. To make an attribute editable, select the check box **Allow workers to edit this attribute** for that attribute.

We recommend that you can add new label category attributes to the labels that you want workers to verify, or add one or more frame attributes to have workers provide information about the entire frame.

For more information, see [Verify and Adjust Labels](#).

16. **Configure your workers' UI:**

- If you are using a **built-in task type**, specify workers instructions and labels.
  
  - For image classification and text classification (single and multi-label) you must specify at least two label categories. For all other built-in task types, you must specify at least one label category.

  - (Optional) If you are creating a 3D point cloud or video frame labeling job, you can specify label category attributes (not supported for 3D point cloud semantic segmentation) and frame attributes. Label category attributes can be assigned to one or more labels. Frame attributes will appear on each point cloud or video frame workers label. To learn more, see [Worker User Interface (UI)](#) for 3D point cloud and [Worker User Interface (UI)](#) for video frame.

  - (Optional) Add **Additional instructions** to help your worker complete your task.

- If you are creating a custom labeling workflow you must:
  
  - Enter a **custom template** in the code box. Custom templates can be created using a combination of HTML, the Liquid templating language and our pre-built web components. Optionally, you can choose a base-template from the drop-down menu to get started.
• Specify pre-annotation and post-annotation lambda functions. To learn how to create these functions, see Step 3: Processing with AWS Lambda.

17. (Optional) You can select See preview to preview your worker instructions, labels, and interact with the worker UI. Make sure the pop-up blocker of the browser is disabled before generating the preview.

18. Choose Create.

After you've successfully created your labeling job, you are redirected to the Labeling jobs page. The status of the labeling job you just created is In progress. This status progressively updates as workers complete your tasks. When all tasks are successfully completed, the status changes to Completed.

If an issue occurs while creating the labeling job, its status changes to Failed.

To view more details about the job, choose the labeling job name.

Next Steps

After your labeling job status changes to Completed, you can view your output data in the Amazon S3 bucket that you specified while creating that labeling job. For details about the format of your output data, see Output Data.

Create a Labeling Job (API)

To create a labeling job using the Amazon SageMaker API, you use the CreateLabelingJob operation. For specific instructions on creating a labeling job for a built-in task type, see that task type page. To learn how to create a streaming labeling job, which is a labeling job that runs perpetually, see Create a Streaming Labeling Job.

To use the CreateLabelingJob operation, you need the following:

• A worker task template (UiTemplateS3Uri) or human task UI ARN (HumanTaskUiArn) in Amazon S3.

• For 3D point cloud jobs, video object detection and tracking jobs, and NER jobs, use the ARN listed in HumanTaskUiArn for your task type.

• If you are using a built-in task type other than 3D point cloud tasks, you can add your worker instructions to one of the pre-built templates and save the template (using a .html or .liquid extension) in your S3 bucket. Find the pre-build templates on your task type page.
If you are using a custom labeling workflow, you can create a custom template and save the template in your S3 bucket. To learn how to built a custom worker template, see Step 2: Creating your custom worker task template. For custom HTML elements that you can use to customize your template, see Crowd HTML Elements Reference. For a repository of demo templates for a variety of labeling tasks, see Amazon SageMaker Ground Truth Sample Task UIs.

An input manifest file that specifies your input data in Amazon S3. Specify the location of your input manifest file in ManifestS3Uri. For information about creating an input manifest, see Input Data. If you create a streaming labeling job, this is optional. To learn how to create a streaming labeling job, see Create a Streaming Labeling Job.

An Amazon S3 bucket to store your output data. You specify this bucket, and optionally, a prefix in S3OutputPath.

A label category configuration file. Each label category name must be unique. Specify the location of this file in Amazon S3 using the LabelCategoryConfigS3Uri parameter. The format and label categories for this file depend on the task type you use:

- For image classification and text classification (single and multi-label) you must specify at least two label categories. For all other task types, the minimum number of label categories required is one.
- For named entity recognition tasks, you must provide worker instructions in this file. See Provide Worker Instructions in a Label Category Configuration File for details and an example.
- For 3D point cloud and video frame task type, use the format in Create a Labeling Category Configuration File with Label Category and Frame Attributes.
- For all other built-in task types and custom tasks, your label category configuration file must be a JSON file in the following format. Identify the labels you want to use by replacing label_1, label_2, ..., label_n with your label categories.

```json
{
    "document-version": "2018-11-28",
    "labels": [
        {"label": "label_1"},
        {"label": "label_2"},
        ...
        {"label": "label_n"}
    ]
}
```
• An AWS Identity and Access Management (IAM) role with the AmazonSageMakerGroundTruthExecution managed IAM policy attached and with permissions to access your S3 buckets. Specify this role in RoleArn. To learn more about this policy, see Use IAM Managed Policies with Ground Truth. If you require more granular permissions, see the section called “IAM Permissions”.

If your input or output bucket name does not contain sagemaker, you can attach a policy similar to the following to the role that is passed to the CreateLabelingJob operation.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "s3:GetObject"
         ],
         "Resource": [
            "arn:aws:s3:::my_input_bucket/*"
         ]
      },
      {
         "Effect": "Allow",
         "Action": [
            "s3:PutObject"
         ],
         "Resource": [
            "arn:aws:s3:::my_output_bucket/*"
         ]
      }
   ]
}
```

• A pre-annotation and post-annotation (or annotation-consolidation) AWS Lambda function Amazon Resource Name (ARN) to process your input and output data.

• Lambda functions are predefined in each AWS Region for built-in task types. To find the pre-annotation Lambda ARN for your Region, see PreHumanTaskLambdaArn. To find the annotation-consolidation Lambda ARN for your Region, see AnnotationConsolidationLambdaArn.
For custom labeling workflows, you must provide a custom pre- and post-annotation Lambda ARN. To learn how to create these Lambda functions, see Step 3: Processing with AWS Lambda.

A work team ARN that you specify in WorkteamArn. You receive a work team ARN when you subscribe to a vendor workforce or create a private workteam. If you are creating a labeling job for a video frame or point cloud task type, you cannot use the Amazon Mechanical Turk workforce. For all other task types, to use the Mechanical Turk workforce, use the following ARN. Replace region with the AWS Region you are using to create the labeling job.

arn:aws:sagemaker:region:394669845002:workteam/public-crowd/default

If you use the Amazon Mechanical Turk workforce, use the ContentClassifiers parameter in DataAttributes of InputConfig to declare that your content is free of personally identifiable information and adult content.

Ground Truth requires that your input data is free of personally identifiable information (PII) if you use the Mechanical Turk workforce. If you use Mechanical Turk and do not specify that your input data is free of PII using the FreeOfPersonallyIdentifiableInformation flag, your labeling job will fail. Use the FreeOfAdultContent flag to declare that your input data is free of adult content. SageMaker may restrict the Amazon Mechanical Turk workers that can view your task if it contains adult content.

To learn more about work teams and workforces, see Create and Manage Workforces.

If you use the Mechanical Turk workforce, you must specify the price you'll pay workers for performing a single task in PublicWorkforceTaskPrice.

To configure the task, you must provide a task description and title using TaskDescription and TaskTitle respectively. Optionally, you can provide time limits that control how long the workers have to work on an individual task (TaskTimeLimitInSeconds) and how long tasks remain in the worker portal, available to workers (TaskAvailabilityLifetimeInSeconds).

(Optional) For some task types, you can have multiple workers label a single data object by inputting a number greater than one for the NumberOfHumanWorkersPerDataObject parameter. For more information about annotation consolidation, see Consolidate Annotations.

(Optional) To create an automated data labeling job, specify one of the ARNs listed in LabelingJobAlgorithmSpecificationArn in LabelingJobAlgorithmsConfig. This ARN identifies the algorithm used in the automated data labeling job. The task type associated with this ARN must match the task type of the PreHumanTaskLambdaArn and...
AnnotationConsolidationLambdaArn you specify. Automated data labeling is supported for the following task types: image classification, bounding box, semantic segmentation, and text classification. The minimum number of objects allowed for automated data labeling is 1,250, and we strongly suggest providing a minimum of 5,000 objects. To learn more about automated data labeling jobs, see Automate Data Labeling.

- (Optional) You can provide StoppingConditions that cause the labeling job to stop if one of the conditions is met. You can use stopping conditions to control the cost of the labeling job.

Examples

The following code examples demonstrate how to create a labeling job using CreateLabelingJob. For additional examples, we recommend you use one of the Ground Truth Labeling Jobs Jupyter notebooks in the SageMaker Examples section of a SageMaker notebook instance. To learn how to use a notebook example from the SageMaker Examples, see Example Notebooks. You can also see these example notebooks on GitHub in the SageMaker Examples repository.

AWS SDK for Python (Boto3)

The following is an example of an AWS Python SDK (Boto3) request to create a labeling job for a built-in task type in the US East (N. Virginia) Region using a private workforce. Replace all red-italized text with your labeling job resources and specifications.

```python
response = client.create_labeling_job(
    LabelingJobName="example-labeling-job",
    LabelAttributeName="label",
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': "s3://bucket/path/manifest-with-input-data.json"
            }
        },
        'DataAttributes': {
            'ContentClassifiers': [
                "FreeOfPersonallyIdentifiableInformation","FreeOfAdultContent",
            ]
        }
    },
    OutputConfig={
        'S3OutputPath': "s3://bucket/path/file-to-store-output-data",
    }
)
```
'KmsKeyId': "string",
},
RoleArn="arn:aws:iam::*:role/*",
LabelCategoryConfigS3Uri="s3://bucket/path/label-categories.json",
StoppingConditions={
  'MaxHumanLabeledObjectCount': 123,
  'MaxPercentageOfInputDatasetLabeled': 123
},
HumanTaskConfig={
  'WorkteamArn': "arn:aws:sagemaker:region::workteam/private-crowd/*",
  'UiConfig': {
    'UiTemplateS3Uri': "s3://bucket/path/custom-worker-task-template.html"
  },
  'TaskKeywords': [
    "Images",
    "Classification",
    "Multi-label"
  ],
  'TaskTitle': "Multi-label image classification task",
  'TaskDescription': "Select all labels that apply to the images shown",
  'NumberOfHumanWorkersPerDataObject': 1,
  'TaskTimeLimitInSeconds': 3600,
  'TaskAvailabilityLifetimeInSeconds': 21600,
  'MaxConcurrentTaskCount': 1000,
  'AnnotationConsolidationConfig': {
    'AnnotationConsolidationLambdaArn': "arn:aws:lambda:us-east-1:432418664414:function:ACS-"
  },
  Tags=[
    {
      'Key': "string",
      'Value': "string"
    }
  ]
}

AWS CLI

The following is an example of an AWS CLI request to create a labeling job for a built-in task type in the US East (N. Virginia) Region using the Amazon Mechanical Turk workforce. For
more information, see `start-human-loop` in the *AWS CLI Command Reference*. Replace all *red-italized text* with your labeling job resources and specifications.

```bash
$ aws --region us-east-1 sagemaker create-labeling-job \
  --labeling-job-name "example-labeling-job" \
  --label-attribute-name "label" \
  --role-arn "arn:aws:iam::account-id:role/role-name" \
  --input-config '{
    "DataAttributes": {
      "ContentClassifiers": [
        "FreeOfPersonallyIdentifiableInformation",
        "FreeOfAdultContent"
      ],
    },
    "DataSource": {
      "S3DataSource": {
        "ManifestS3Uri": "s3://bucket/path/manifest-with-input-data.json"
      }
    }
  }' \
  --output-config '{
    "KmsKeyId": "",
    "S3OutputPath": "s3://bucket/path/file-to-store-output-data"
  }' \
  --human-task-config '{
    "AnnotationConsolidationConfig": {
      "AnnotationConsolidationLambdaArn": "arn:aws:lambda:us-east-1:432418664414:function:ACS-"
    },
    "TaskAvailabilityLifetimeInSeconds": 21600,
    "TaskTimeLimitInSeconds": 3600,
    "NumberOfHumanWorkersPerDataObject": 1,
    "WorkteamArn": "arn:aws:sagemaker:us-east-1:394669845002:workteam/public-crowd/default",
    "PublicWorkforceTaskPrice": {
      "AmountInUsd": {
        "Dollars": 0,
        "TenthFractionsOfACent": 6,
        "Cents": 3
      }
    }
  },
```
"TaskDescription": "Select all labels that apply to the images shown",
"MaxConcurrentTaskCount": 1000,
"TaskTitle": "Multi-label image classification task",
"TaskKeywords": [
    "Images",
    "Classification",
    "Multi-label"
],
"UiConfig": {
    "UiTemplateS3Uri": "s3://bucket/path/custom-worker-task-template.html"
}

For more information about this operation, see CreateLabelingJob. For information about how to use other language-specific SDKs, see See Also in the CreateLabelingJobs topic.

Create a Streaming Labeling Job

Streaming labeling jobs enable you to send individual data objects in real time to a perpetually running, streaming labeling job. To create a streaming labeling job, you must create an Amazon SNS input topic and specify this topic in CreateLabelingJob parameters InputConfig of SnsDataSource. Optionally, you can also create an Amazon SNS output topic and specify it in OutputConfig if you want to receive label data in real time.

⚠️ Important

If you are a new user of Ground Truth streaming labeling jobs, it is recommended that you review Ground Truth Streaming Labeling Jobs before creating a streaming labeling job.

Use the following sections to create the resources that you need and can use to create a streaming labeling job:

- Learn how to create SNS topics with the permissions required for Ground Truth streaming labeling jobs by following the steps in Create Amazon SNS Input and Output Topics. Your SNS topics must be created in the same AWS Region as your labeling job.

- See Subscribe an Endpoint to Your Amazon SNS Output Topic to learn how to set up an endpoint to receive labeling task output data at a specified endpoint each time a labeling task is completed.
To learn how to configure your Amazon S3 bucket to send notifications to your Amazon SNS input topic, see Set up Amazon S3 Bucket Event Notifications.

Optionally, add data objects that you want to have labeled as soon as the labeling job starts to your input manifest. For more information, see Create a Manifest File (Optional).

There are other resources required to create a labeling job, such as an IAM role, Amazon S3 bucket, a worker task template and label categories. These are described in the Ground Truth documentation on creating a labeling job. For more information, see Create a Labeling Job.

**Important**

When you create a labeling job you must provide an IAM execution role. Attach the AWS managed policy AmazonSageMakerGroundTruthExecution to this role to ensure it has required permissions to execute your labeling job.

When you submit a request to create a streaming labeling job, the state of your labeling job is Initializing. Once the labeling job is active, the state changes to InProgress. Do not send new data objects to your labeling job or attempt to stop your labeling job while it is in the Initializing state. Once the state changes to InProgress, you can start sending new data objects using Amazon SNS and the Amazon S3 configuration.

**Topics**

- Create Amazon SNS Input and Output Topics
- Set up Amazon S3 Bucket Event Notifications
- Create a Manifest File (Optional)
- Example: Use SageMaker API To Create Streaming Labeling Job
- Stop a Streaming Labeling Job

**Create Amazon SNS Input and Output Topics**

You need to create an Amazon SNS input to create a streaming labeling job. Optionally, you may provide an Amazon SNS output topic.

When you create an Amazon SNS topic to use in your streaming labeling job, note down the topic Amazon Resource Name (ARN). The ARN will be the input values for the parameter SnsTopicArn in InputConfig and OutputConfig when you create a labeling job.
Create an Input Topic

Your input topic is used to send new data objects to Ground Truth. To create an input topic, follow the instructions in Creating an Amazon SNS topic in the Amazon Simple Notification Service Developer Guide.

Note down your input topic ARN and use it as input for the CreateLabelingJob parameter SnsTopicArn in InputConfig.

Create an Output Topic

If you provide an output topic, it is used to send notifications when a data object is labeled. When you create a topic, you have the option to add an encryption key. Use this option to add a AWS Key Management Service customer managed key to your topic to encrypt the output data of your labeling job before it is published to your output topic.

To create an output topic, follow the instructions in Creating an Amazon SNS topic in the Amazon Simple Notification Service Developer Guide.

If you add encryption, you must attach additional permission to the topic. See Add Encryption to Your Output Topic (Optional) for more information.

⚠️ Important

To add a customer managed key to your output topic while creating a topic in the console, do not use the (Default) alias/aws/sns option. Select a customer managed key that you created.

Note down your input topic ARN and use it in your CreateLabelingJob request in the parameter SnsTopicArn in OutputConfig.

Add Encryption to Your Output Topic (Optional)

To encrypt messages published to your output topic, you need to provide an AWS KMS customer managed key to your topic. Modify the following policy and add it to your customer managed key to give Ground Truth permission to encrypt output data before publishing it to your output topic.

Replace <account_id> with the ID of the account that you are using to create your topic. To learn how to find your AWS account ID, see Finding Your AWS Account ID.
Additionally, you must modify and add the following policy to the execution role that you use to create your labeling job (the input value for RoleArn).

```json
{
    "Id": "key-console-policy",
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "Enable IAM User Permissions",
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::<account_id>:root"
            },
            "Action": "kms:*",
            "Resource": "*"
        },
        {
            "Sid": "Allow access for Key Administrators",
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::<account_id>:role/Admin"
            },
            "Action": [
                "kms:Create*",
                "kms:Describe*",
                "kms:Enable*",
                "kms:List*",
                "kms:Put*",
                "kms:Update*",
                "kms:Revoke*",
                "kms:Disable*",
                "kms:Get*",
                "kms:Delete*",
                "kms:TagResource",
                "kms:UntagResource",
                "kms:ScheduleKeyDeletion",
                "kms:CancelKeyDeletion"
            ],
            "Resource": "*"
        }
    ]
}
```
Replace `<account_id>` with the ID of the account that you are using to create your topic. Replace `<region>` with the AWS Region you are using to create your labeling job. Replace `<key_id>` with your customer managed key ID.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "sid1",
      "Effect": "Allow",
      "Action": [
        "kms:Decrypt",
        "kms:GenerateDataKey"
      ],
      "Resource": "arn:aws:kms:<region>:<account_id>:key/<key_id>"
    }
  ]
}
```

For more information on creating and securing keys, see [Creating Keys](https://docs.aws.amazon.com/AWSSimpleDB/latest/SG/using-key-policies.html) and [Using Key Policies](https://docs.aws.amazon.com/AWSSimpleDB/latest/SG/using-key-policies.html) in the AWS Key Management Service Developer Guide.

### Subscribe an Endpoint to Your Amazon SNS Output Topic

When a worker completes a labeling job task from a Ground Truth streaming labeling job, Ground Truth uses your output topic to publish output data to one or more endpoints that you specify. To receive notifications when a worker finishes a labeling task, you must subscribe an endpoint to your Amazon SNS output topic.

To learn how to add endpoints to your output topic, see [Subscribing to an Amazon SNS topic](https://docs.aws.amazon.com/AWSSimpleDB/latest/SG/subscribing-endpoints.html) in the Amazon Simple Notification Service Developer Guide.

To learn more about the output data format that is published to these endpoints, see [Output Data](https://docs.aws.amazon.com/AWSSimpleDB/latest/SG/creating-labeling-job.html).

⚠️ **Important**

If you do not subscribe an endpoint to your Amazon SNS output topic, you will not receive notifications when new data objects are labeled.
Set up Amazon S3 Bucket Event Notifications

You can add an event notification to your Amazon S3 bucket using the Amazon S3 console, API, and language specific AWS SDKs, or the AWS Command Line Interface. Set up this event to send notifications to the same Amazon SNS input topic that you specify using SnsTopicArn in InputConfig when you create a labeling job. Do not set up event notifications using the same Amazon S3 location that you specified for S3OutputPath in OutputConfig – doing so may result in unwanted data objects being processed by Ground Truth for labeling.

You decide the types of events that you want to send to your Amazon SNS topic. Ground Truth creates a labeling job when you send object creation events.

The event structure sent to your Amazon SNS input topic must be a JSON message formatted using the same structure found in Event message structure.

To see examples of how you can set up an event notification for your Amazon S3 bucket using the Amazon S3 console, AWS SDK for .NET, and AWS SDK for Java, follow this walkthrough, Walkthrough: Configure a bucket for notifications (SNS topic or SQS queue) in the Amazon Simple Storage Service User Guide.

Create a Manifest File (Optional)

When you create a streaming labeling job, you have the one time option to add objects (such as images or text) to an input manifest file that you specify in ManifestS3Uri of CreateLabelingJob. When the streaming labeling job starts, these objects are sent to workers or added to the Amazon SQS queue if the total number of objects exceed MaxConcurrentTaskCount. The results are added to the Amazon S3 path that you specify when creating the labeling job periodically as workers complete labeling tasks. Output data is sent to any endpoint that you subscribe to your output topic.

If you want to provide initial objects to be labeled, create a manifest file that identifies these objects and place it in Amazon S3. Specify the S3 URI of this manifest file in ManifestS3Uri within InputConfig.

To learn how to format your manifest file, see Input Data. To use the SageMaker console to automatically generate a manifest file (not supported for 3D point cloud task types), see Automated Data Setup.
Example: Use SageMaker API To Create Streaming Labeling Job

The following is an example of an AWS Python SDK (Boto3) request that you can use to start a streaming labeling job for a built-in task type in the US East (N. Virginia) Region. For more details about each parameter below see CreateLabelingJob. To learn how you can create a labeling job using this API and associated language specific SDKs, see Create a Labeling Job (API).

In this example, note the following parameters:

- **SnsDataSource** – This parameter appears in InputConfig and OutputConfig and is used to identify your input and output Amazon SNS topics respectively. To create a streaming labeling job, you are required to provide an Amazon SNS input topic. Optionally, you can also provide an Amazon SNS output topic.

- **S3DataSource** – This parameter is optional. Use this parameter if you want to include an input manifest file of data objects that you want labeled as soon as the labeling job starts.

- **StoppingConditions** – This parameter is ignored when you create a streaming labeling job. To learn more about stopping a streaming labeling job, see Stop a Streaming Labeling Job.

- Streaming labeling jobs do not support automated data labeling. Do not include the LabelingJobAlgorithmsConfig parameter.

```python
response = client.create_labeling_job(
    LabelingJobName= 'example-labeling-job',
    LabelAttributeName='label',
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': 's3://bucket/path/manifest-with-input-data.json'
            },
            'SnsDataSource': {
                'SnsTopicArn': 'arn:aws:sns:us-east-1:123456789012:your-sns-input-topic'
            }
        },
        'DataAttributes': {
            'ContentClassifiers': ['FreeOfPersonallyIdentifiableInformation'|'FreeOfAdultContent',
        ]
    },
    OutputConfig={
```

'S3OutputPath': 's3://bucket/path/file-to-store-output-data',
'KmsKeyId': 'string',
'SnsTopicArn': 'arn:aws:sns:us-east-1:123456789012:your-sns-output-topic'
},
RoleArn='arn:aws:iam::*:role/*',
LabelCategoryConfigS3Uri='s3://bucket/path/label-categories.json',
HumanTaskConfig={
'WorkteamArn': 'arn:aws:sagemaker:us-east-1:*:workteam/private-crowd/*',
'UiConfig': {
 'UiTemplateS3Uri': 's3://bucket/path/custom-worker-task-template.html'
},
'TaskKeywords': [
 'Example key word',
],
'TaskTitle': 'Multi-label image classification task',
'TaskDescription': 'Select all labels that apply to the images shown',
'NumberOfHumanWorkersPerDataObject': 123,
'TaskTimeLimitInSeconds': 123,
'TaskAvailabilityLifetimeInSeconds': 123,
'MaxConcurrentTaskCount': 123,
'AnnotationConsolidationConfig': {
 'AnnotationConsolidationLambdaArn': 'arn:aws:lambda:us-east-1:432418664414:function:ACS-tasktype'
}
},
Tags=[
{
'Key': 'string',
'Value': 'string'
},
]
)

Stop a Streaming Labeling Job

You can manually stop your streaming labeling job using the operation StopLabelingJob.

If your labeling job remains idle for over 10 days, it is automatically stopped by Ground Truth. In this context, a labeling job is considered idle if no objects are sent to the Amazon SNS input topic and no objects remain in your Amazon SQS queue, waiting to be labeled. For example, if no data objects are fed to the Amazon SNS input topic and all the objects fed to the labeling job are
already labeled, Ground Truth starts a timer. After the timer starts, if no items are received within a 10 day period, the labeling job is stopped.

When a labeling job is stopped, its status is STOPPING while Ground Truth cleans up labeling job resources and unsubscribes your Amazon SNS topic from your Amazon SQS queue. The Amazon SQS is *not* deleted by Ground Truth because this queue may contain unprocessed data objects. You should manually delete the queue if you want to avoid incurring additional charges from Amazon SQS. To learn more, see [Amazon SQS pricing](#).

**Create a Labeling Category Configuration File with Label Category and Frame Attributes**

When you create a 3D point cloud or video frame labeling job using the Amazon SageMaker API operation `CreateLabelingJob`, you use a label category configuration file to specify your labels and worker instructions. Optionally, you can also provide the following in your label category attribute file:

- You can provide *label category attributes* for video frame and 3D point cloud object tracking and object detection task types. Workers can use one or more attributes to give more information about an object. For example, you may want to use the attribute *occluded* to have workers identify when an object is partially obstructed. You can either specify a label category attribute for a single label using the `categoryAttributes` parameter, or for all labels using the `categoryGlobalAttributes` parameter.

- You can provide *frame attributes* for video frame and 3D point cloud object tracking and object detection task types using `frameAttributes`. When you create a frame attribute, it appears on each frame or point cloud in the worker task. In video frame labeling jobs, these are attributes that workers assign to an entire video frame. For 3D point cloud labeling jobs, these attributes are applied to a single point cloud. Use frame attributes to have workers provide more information about the scene in a specific frame or point cloud.

- For video frame labeling jobs, you use the label category configuration file to specify the task type (bounding box, polyline, polygon, or keypoint) sent to workers.

For workers, specifying values for label category attributes and frame attributes will be optional.

⚠️ **Important**

You should only provide a label attribute name in `auditLabelAttributeName` if you are running an audit job to verify or adjust labels. Use this parameter to input the...
**LabelAttributeName** used in the labeling job that generated the annotations you want your worker to adjust. When you create a labeling job in the console, if you did not specify a label attribute name, the **Name** of your job is used as the LabelAttributeName.

Topics

- Label Category Configuration File Schema
- Example: Label Category Configuration Files for 3D Point Cloud Labeling Jobs
- Example: Label Category Configuration Files for Video Frame Labeling Jobs
- Creating Worker Instructions

**Label Category Configuration File Schema**

The following table lists elements you can and must include in your label category configuration file.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>frameAttributes</td>
<td>No</td>
<td>A list of JSON objects.</td>
<td>Use this parameter to create a frame attribute that is applied to all frames or 3D point clouds in your labeling job. See the third table in this section for more information.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Required Parameters in each JSON Object:</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>name, type, description</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>minimum and maximum are required if type is &quot;number&quot;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Optional Parameters in each JSON Object:</strong></td>
<td></td>
</tr>
</tbody>
</table>

Note

The parameter annotationType is only supported for video frame labeling jobs.
<table>
<thead>
<tr>
<th><strong>Parameter</strong></th>
<th><strong>Required</strong></th>
<th><strong>Accepted Values</strong></th>
<th><strong>Description</strong></th>
</tr>
</thead>
</table>
| **categoryGlobalAttributes** | No | enum, editsAllowed, isRequired | A list of JSON objects. **Required Parameters in each JSON Object:**
- name, type
- minimum and maximum are required if type is "number"
**Optional Parameters in each JSON Object:**
- description, enum, editsAllowed, isRequired
Use this parameter to create label category attributes that are applied to all labels you specify in labels. See the third table in this section for more information.

Create a Labeling Job
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>labels</td>
<td>Yes</td>
<td>A list of up to 30 JSON objects</td>
<td>Use this parameter to specify your labels, or classes. Add one label for each class.</td>
</tr>
</tbody>
</table>

**Required Parameters in each JSON Object:**
- label

**Optional Parameters in each JSON Object:**
- categoryAttributes
- editsAllowed

To add a label category attribute to a label, add categoryAttributes to that label.

Use editsAllowed to specify whether or not a label can be edited in an adjustment labeling job. Set editsAllowed to "none" for verification labeling jobs.

See the following table for more information.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>annotationType (only supported for video frame labeling jobs)</td>
<td>No</td>
<td>String</td>
<td>Use this to specify the task type for your video frame labeling jobs. For example, for a polygon video frame object detection task, choose Polygon. If you do not specify an annotationType when you create a video frame labeling job, Ground Truth will use BoundingBox by default.</td>
</tr>
</tbody>
</table>

**Accepted Parameters:**
- BoundingBox
- Polyline
- Polygon
- Keypoint

**Default:**
- BoundingBox
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>instructions</td>
<td>No</td>
<td>A JSON object Required Parameters in each JSON Object: &quot;shortInstruction&quot;, &quot;fullInstruction&quot;</td>
<td>Use this parameter to add worker instructions to help your workers complete their tasks. For more information about worker instructions, see Worker Instructions. Short instructions must be under 255 characters and long instruction must be under 2,048 characters. For more information, see Creating Worker Instructions.</td>
</tr>
</tbody>
</table>
The following table describes the parameters that you can and must use to create a list of Labels. Each parameter should be included in a JSON object.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>label</td>
<td>Yes</td>
<td>String</td>
<td>The name of the label category that is displayed to workers. Each label category name must be unique.</td>
</tr>
<tr>
<td>categoryAttributes</td>
<td>No</td>
<td>A list of JSON objects.</td>
<td>Use this parameter to add label category attributes to specific labels you specify in labels.</td>
</tr>
<tr>
<td>auditLabelAttributeName</td>
<td>Required for adjustment and verification task types</td>
<td>String</td>
<td>Enter the <code>LabelAttribute</code>Name used in the labeling job you want to adjust annotations of. Only use this parameter if you are creating an adjustment job for video frame and 3D point cloud object detection, object tracking, or 3D point cloud semantic segmentation.</td>
</tr>
</tbody>
</table>

Create a Labeling Job
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>name, type</td>
<td></td>
<td>minimum and maximum required if type is &quot;number&quot;</td>
<td>To add one or more label category attributes to a label, include the categoryAttributes JSON object in the same labels JSON object as that label. See the following table for more information.</td>
</tr>
</tbody>
</table>

Optional Parameters in each JSON Object:
- description
- enum
- editsAllowed
- isRequired
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>editsAllowed</td>
<td>No</td>
<td>String</td>
<td>Specifies whether or not a label can be edited by workers.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Supported Values:</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;none&quot;: no modifications are not allowed.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>or</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;any&quot; (Default): all modifications are allowed.</td>
<td></td>
</tr>
</tbody>
</table>

For video frame or 3D point cloud adjustment labeling jobs, add this parameter to one or more JSON objects in the labels list to specify whether or not a worker can edit a label.

For 3D point cloud and video frame verification labeling jobs, add this parameter with the value "none" to each JSON object in the labels list. This will make all labels uneditable.

The following table describes the parameters that you can and must use to create a frame attributes using frameAttributes and label category attribute using the categoryGlobalAttributes and categoryAttributes parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>name</td>
<td>Yes</td>
<td>String</td>
<td>Use this parameter to assign a name to your label category</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>or frame attribute. This is the attribute name that workers see.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Each label category attribute name in your label category configuration file must be unique.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Global label category attributes and label specific label category attributes cannot have the same name.</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>type</td>
<td>Yes</td>
<td>String</td>
<td>Use this parameter to define the label category or frame attribute type. Use &quot;string&quot; or &quot;number&quot; for type.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;string&quot; or &quot;number&quot;</td>
<td>If you specify &quot;string&quot; for type and provide an enum value for this attribute, workers will be able to choose from one of the choices you provide.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>If you specify &quot;string&quot; for type and do not provide an enum value, workers can enter free form text.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>If you specify number for type, worker can enter a number between the minimum and maximum numbers you specify.</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>enum</td>
<td>No</td>
<td>List of strings</td>
<td>Use this parameter to define options that workers can choose from for this label category or frame attribute. Workers can choose one value specified in enum. For example, if you specify [&quot;foo&quot;, &quot;buzz&quot;, &quot;bar&quot;] for enum, workers can choose one of foo, buzz, or bar. You must specify &quot;string&quot; for type to use an enum list.</td>
</tr>
<tr>
<td>description</td>
<td>frameAttributes : Yes</td>
<td>String</td>
<td>Use this parameter to add a description of the label category or frame attribute. You can use this field to give workers more information about the attribute. This field is only required for frame attributes.</td>
</tr>
<tr>
<td></td>
<td>categoryAttributes or categoryGlobalAttributes : No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------------</td>
<td>--------------</td>
<td>-----------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>minimum and maximum</td>
<td>Required if attribute type is &quot;number&quot;</td>
<td>Integers</td>
<td>Use these parameters to specify minimum and maximum (inclusive) values workers can enter for numeric label category or frame attributes. You must specify &quot;number&quot; for type to use minimum and maximum.</td>
</tr>
<tr>
<td>editsAllowed</td>
<td>No</td>
<td>String</td>
<td>Specifies whether or not a label category or frame attribute can be edited by workers. For video frame or 3D point cloud adjustment and verification labeling jobs, add this parameter to label category and frame attribute JSON objects to specify whether or not a worker can edit an attribute.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Required Values:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;none&quot;: no modifications are not allowed.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;any&quot; (Default): all modifications are allowed.</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>----------</td>
<td>-----------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>isRequired</td>
<td>No</td>
<td>Boolean</td>
<td>Specifies whether workers are required to annotate an attribute. Workers cannot submit the job until all required attributes are annotated.</td>
</tr>
</tbody>
</table>

**Label and label category attribute quotas**

You can specify up to 10 label category attributes per class. This 10-attribute quotas includes global label category attributes. For example, if you create four global label category attributes, and then assign three label category attributes to label X, that label will have 4+3=7 label category attributes in total. For all label category and label category attribute limits, refer to the following table.

<table>
<thead>
<tr>
<th>Type</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Labels (Labels)</td>
<td>1</td>
<td>30</td>
</tr>
<tr>
<td>Label name character quota</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Label category attributes per label (sum of categoryAttributes and categoryGlobalAttributes)</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>Free form text entry label category attributes per label (sum of categoryAttributes and categoryGlobalAttributes)</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Type</td>
<td>Min</td>
<td>Max</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
<td>-----</td>
<td>-----</td>
</tr>
<tr>
<td>Frame attributes</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>Free form text entry attributes in frameAttributes</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Attribute name character quota (name)</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Attribute description character quota (description)</td>
<td>0</td>
<td>128</td>
</tr>
<tr>
<td>Attribute type characters quota (type)</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Allowed values in the enum list for a string attribute</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>Character quota for a value in enum list</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Maximum characters in free form text response for free form text frameAttributes</td>
<td>0</td>
<td>1000</td>
</tr>
<tr>
<td>Maximum characters in free form text response for free form text categoryAttributes and categoryGlobalAttributes</td>
<td>0</td>
<td>80</td>
</tr>
</tbody>
</table>
Example: Label Category Configuration Files for 3D Point Cloud Labeling Jobs

Select a tab in the following tables to see examples of 3D point cloud label category configuration files for object detection, object tracking, semantic segmentation, adjustment, and verification labeling jobs.

3D Point Cloud Object Tracking and Object Detection

The following is an example of a label category configuration file that includes label category attributes for a 3D point cloud object detection or object tracking labeling job. This example includes a two frame attributes, which will be added to all point clouds submitted to the labeling job. The Car label will include four label category attributes—X, Y, Z, and the global attribute, W.

```json
{
    "documentVersion": "2020-03-01",
    "frameAttributes": [
        {
            "name": "count players",
            "description": "How many players to you see in the scene?",
            "type": "number"
        },
        {
            "name": "select one",
            "description": "describe the scene",
            "type": "string",
            "enum": ["clear", "blurry"],
            "isRequired": true
        }
    ],
    "categoryGlobalAttributes": [
        {
            "name": "W",
            "description": "label-attributes-for-all-labels",
            "type": "string",
            "enum": ["foo", "buzz", "biz"]
        }
    ],
    "labels": [
        {
            "label": "Car",
            "categoryAttributes": [
                {
                ```
3D Point Cloud Semantic Segmentation

The following is an example of a label category configuration file for a 3D point cloud semantic segmentation labeling job.

Label category attributes are not supported for 3D point cloud semantic segmentation task types. Frame attributes are supported. If you provide label category attributes for a semantic segmentation labeling job, they will be ignored.
Select a tab in the following table to see an example of a label category configuration file for 3D point cloud verification or adjustment labeling jobs.

### 3D Point Cloud Adjustment

The following is an example of a label category configuration file for a 3D point cloud object detection or object tracking adjustment labeling job. For 3D point cloud semantic segmentation adjustment labeling jobs, categoryGlobalAttributes and categoryAttributes are not supported.

You must include auditLabelAttributeName to specify the label attribute name of the previous labeling job that you use to create the adjustment labeling job. Optionally, you can use the editsAllowed parameter to specify whether or not a label or frame attribute can be edited.

```json
{
    "documentVersion": "2020-03-01",
    "frameAttributes": [
        
```
{
    "name":"count players",
    "description":"How many players do you see in the scene?",
    "type":"number"
},
{
    "name":"select one",
    "editsAllowed":"none",
    "description":"describe the scene",
    "type":"string",
    "enum": ["clear", "blurry"]
},
"categoryGlobalAttributes": [
{
    "name":"W",
    "editsAllowed":"any",
    "description":"label-attributes-for-all-labels",
    "type":"string",
    "enum": ["foo", "buzz", "biz"]
}
],
"labels": [
{
    "label": "Car",
    "editsAllowed": "any",
    "categoryAttributes": [
    {
        "name": "X",
        "description": "enter a number",
        "type": "number"
    },
    {
        "name": "Y",
        "description": "select an option",
        "type": "string",
        "enum": ["y1", "y2"],
        "editsAllowed": "any"
    },
    {
        "name": "Z",
        "description": "submit a free-form response",
        "type": "string",
        "editsAllowed": "none"
    }
}]}
3D Point Cloud Verification

The following is an example of a label category configuration file you may use for a 3D point cloud object detection or object tracking verification labeling job. For a 3D point cloud semantic segmentation verification labeling job, categoryGlobalAttributes and categoryAttributes are not supported.

You must include auditLabelAttributeName to specify the label attribute name of the previous labeling job that you use to create the verification labeling job. Additionally, you must use the editsAllowed parameter to specify that no labels can be edited.

```json
{
    "documentVersion": "2020-03-01",
    "frameAttributes": [
        {
            "name": "count players",
            "editsAllowed": "any",
            "description": "How many players do you see in the scene?",
            "type": "number"
        },
        {
            "name": "select one",
            "editsAllowed": "any",
            "description": "describe the scene",
            "type": "string",
            "enum": ["clear", "blurry"]
        }
    ],
    "categoryGlobalAttributes": [
    ]
}
```
Create a Labeling Job

```
{
    "name": "W",
    "editsAllowed": "none",
    "description": "label-attributes-for-all-labels",
    "type": "string",
    "enum": ["foo", "buzz", "biz"]
}
]
,"labels": [
{
    "label": "Car",
    "editsAllowed": "none",
    "categoryAttributes": [
        {
            "name": "X",
            "description": "enter a number",
            "type": "number",
            "editsAllowed": "none"
        },
        {
            "name": "Y",
            "description": "select an option",
            "type": "string",
            "enum": ["y1", "y2"],
            "editsAllowed": "any"
        },
        {
            "name": "Z",
            "description": "submit a free-form response",
            "type": "string",
            "editsAllowed": "none"
        }
    ]
},
{
    "label": "Pedestrian",
    "editsAllowed": "none",
    "categoryAttributes": [...]  
}
],
"instructions": {"shortInstruction": "Draw a tight Cuboid",
"fullInstruction": "<html markup>
// include auditLabelAttributeName for label verification jobs  
"auditLabelAttributeName": "myPrevJobLabelAttributeName"
Example: Label Category Configuration Files for Video Frame Labeling Jobs

The annotation tools available to your worker and task type used depends on the value you specify for annotationType. For example, if you want workers to use key points to track changes in the pose of specific objects across multiple frames, you would specify Keypoint for the annotationType. If you do not specify an annotation type, BoundingBox will be used by default.

The following is an example of a video frame keypoint label category configuration file with label category attributes. This example includes two frame attributes, which will be added to all frames submitted to the labeling job. The Car label will include four label category attributes—X, Y, Z, and the global attribute, W.

```json
{
   "documentVersion": "2020-03-01",
   "frameAttributes": [
   {
      "name": "count players",
      "description": "How many players to you see in the scene?",
      "type": "number"
   },
   {
      "name": "select one",
      "description": "describe the scene",
      "type": "string",
      "enum": ["clear", "blurry"]
   }
   ],
   "categoryGlobalAttributes": [
   {
      "name": "W",
      "description": "label-attributes-for-all-labels",
      "type": "string",
      "enum": ["foo", "buz", "buz2"]
   }
   ],
   "labels": [
   {
      "label": "Car",
      "categoryAttributes": [
```
Select a tab from the following table to see examples of label category configuration files for video frame adjustment and verification labeling jobs.

### Video Frame Adjustment

The following is an example of a label category configuration file you may use for a video frame adjustment labeling job.

You must include `auditLabelAttributeName` to specify the label attribute name of the previous labeling job that you use to create the verification labeling job. Optionally, you can use the `editsAllowed` parameter to specify whether or not labels, label category attributes, or frame attributes can be edited.
"frameAttributes": [ 
  {
    "name": "count players",
    "editsAllowed": "none",
    "description": "How many players do you see in the scene?",
    "type": "number"
  },
  {
    "name": "select one",
    "description": "Describe the scene",
    "type": "string",
    "enum": ["clear", "blurry"]
  }
],
"categoryGlobalAttributes": [ 
  {
    "name": "W",
    "editsAllowed": "any",
    "description": "label-attributes-for-all-labels",
    "type": "string",
    "enum": ["foo", "buz", "buz2"]
  }
],
"labels": [ 
  {
    "label": "Car",
    "editsAllowed": "any",
    "categoryAttributes": [ 
      {
        "name": "X",
        "description": "Enter a number",
        "type": "number",
        "editsAllowed": "any"
      },
      {
        "name": "Y",
        "description": "Select an option",
        "type": "string",
        "enum": ["y1", "y2"],
        "editsAllowed": "any"
      },
      {
        "name": "Z",
        "description": "Submit a free-form response",
        
      }
    ]
  }
]
Video Frame Verification

The following is an example of a label category configuration file for a video frame labeling job.

You must include auditLabelAttributeName to specify the label attribute name of the previous labeling job that you use to create the verification labeling job. Additionally, you must use the editsAllowed parameter to specify that no labels can be edited.

```json
{
  "documentVersion": "2020-03-01",
  "frameAttributes": [
    {
      "name": "count players",
      "editsAllowed": "none",
      "description": "How many players do you see in the scene?",
      "type": "number"
    },
    {
      "name": "select one",
      "editsAllowed": "any",
      "description": "describe the scene",
      "type": "string",
      "enum": ["clear", "blurry"]
    }
  ]
}
```
"categoryGlobalAttributes": [
    {
      "name": "W",
      "editsAllowed": "none",
      "description": "label-attributes-for-all-labels",
      "type": "string",
      "enum": ["foo", "buz", "buz2"
    }
  ],
  "labels": [
    {
      "label": "Car",
      "editsAllowed": "none",
      "categoryAttributes": [
        {
          "name": "X",
          "description": "enter a number",
          "type": "number",
          "editsAllowed": "any"
        },
        {
          "name": "Y",
          "description": "select an option",
          "type": "string",
          "enum": ["y1", "y2"],
          "editsAllowed": "any"
        },
        {
          "name": "Z",
          "description": "submit a free-form response",
          "type": "string",
          "editsAllowed": "none"
        }
      ]
    },
    {
      "label": "Pedestrian",
      "editsAllowed": "none",
      "categoryAttributes": [...]
    }
  ],
  "annotationType": "Keypoint",
  "instructions": {
    "shortInstruction": "add example short instructions here",
    "fullInstruction": "<html markup>"}
Creating Worker Instructions

Create custom instructions for labeling jobs to improve your worker's accuracy in completing their task. Your instructions are accessible when workers select the **Instructions** menu option in the worker UI. Short instructions must be under 255 characters and long instruction must be under 2,048 characters.

There are two kinds of instructions:

- **Short instructions** – These instructions are shown to workers when they select **Instructions** in the worker UI menu. They should provide an easy reference to show the worker the correct way to label an object.

- **Full instructions** – These instructions are shown when workers select **More Instructions** in instructions the pop-up window. We recommend that you provide detailed instructions for completing the task with multiple examples showing edge cases and other difficult situations for labeling objects.

For 3D point cloud and video frame labeling jobs, you can add worker instructions to your label category configuration file. You can use a single string to create instructions or you can add HTML mark up to customize the appearance of your instructions and add images. Make sure that any images you include in your instructions are publicly available, or if your instructions are in Amazon S3, that your workers have read-access so that they can view them.

Use Input and Output Data

The input data that you provide to Amazon SageMaker Ground Truth is sent to your workers for labeling. You choose the data to send to your workers by creating a single manifest file that defines all of the data that requires labeling or by sending input data objects to an ongoing, streaming labeling job to be labeled in real time.

The output data is the result of your labeling job. The output data file, or **augmented manifest file**, contains label data for each object you send to the labeling job and metadata about the label assigned to data objects.
When you use image classification (single and multi-label), text classification (single and multi-label), object detection, and semantic segmentation built in task types to create a labeling job, you can use the resulting augmented manifest file to launch a SageMaker training job. For a demonstration of how to use an augmented manifest to train an object detection machine learning model with Amazon SageMaker, see `object_detection_augmented_manifest_training.ipynb`. For more information, see `Provide Dataset Metadata to Training Jobs with an Augmented Manifest File`.

**Topics**

- **Input Data**
- **3D Point Cloud Input Data**
- **Video Frame Input Data**
- **Output Data**

**Input Data**

The input data are the data objects that you send to your workforce to be labeled. There are two ways to send data objects to Ground Truth for labeling:

- Send a list of data objects that require labeling using an input manifest file.
- Send individual data objects in real time to a perpetually running, streaming labeling job.

If you have a dataset that needs to be labeled one time, and you do not require an ongoing labeling job, create a standard labeling job using an input manifest file.

If you want to regularly send new data objects to your labeling job after it has started, create a streaming labeling job. When you create a streaming labeling job, you can optionally use an input manifest file to specify a group of data that you want labeled immediately when the job starts. You can continuously send new data objects to a streaming labeling job as long as it is active.

**Note**

Streaming labeling jobs are only supported through the SageMaker API. You cannot create a streaming labeling job using the SageMaker console.

The following task types have special input data requirements and options:
For **3D point cloud** labeling job input data requirements, see [3D Point Cloud Input Data](#).

For **video frame** labeling job input data requirements, see [Video Frame Input Data](#).

**Topics**

- [Use an Input Manifest File](#)
- [Automated Data Setup](#)
- [Supported Data Formats](#)
- [Ground Truth Streaming Labeling Jobs](#)
- [Input Data Quotas](#)
- [Filter and Select Data for Labeling](#)

**Use an Input Manifest File**

Each line in an input manifest file is an entry containing an object, or a reference to an object, to label. An entry can also contain labels from previous jobs and for some task types, additional information.

Input data and the manifest file must be stored in Amazon Simple Storage Service (Amazon S3). Each has specific storage and access requirements, as follows:

- The Amazon S3 bucket that contains the input data must be in the same AWS Region in which you are running Amazon SageMaker Ground Truth. You must give Amazon SageMaker access to the data stored in the Amazon S3 bucket so that it can read it. For more information about Amazon S3 buckets, see [Working with Amazon S3 buckets](#).

- The manifest file must be in the same AWS Region as the data files, but it doesn't need to be in the same location as the data files. It can be stored in any Amazon S3 bucket that is accessible to the AWS Identity and Access Management (IAM) role that you assigned to Ground Truth when you created the labeling job.

**Note**

3D point cloud and video frame **task types** have different input manifest requirements and attributes.

For **3D point cloud task types**, refer to Create an Input Manifest File for a 3D Point Cloud Labeling Job.
For video frame task types, refer to Create a Video Frame Input Manifest File.

The manifest is a UTF-8 encoded file in which each line is a complete and valid JSON object. Each line is delimited by a standard line break, \n or \r\n. Because each line must be a valid JSON object, you can't have unescaped line break characters. For more information about data format, see JSON Lines.

Each JSON object in the manifest file can be no larger than 100,000 characters. No single attribute within an object can be larger than 20,000 characters. Attribute names can't begin with $ (dollar sign).

Each JSON object in the manifest file must contain one of the following keys: source-ref or source. The value of the keys are interpreted as follows:

- **source-ref** – The source of the object is the Amazon S3 object specified in the value. Use this value when the object is a binary object, such as an image.
- **source** – The source of the object is the value. Use this value when the object is a text value.

The following is an example of a manifest file for files stored in an Amazon S3 bucket:

```
{"source-ref": "S3 bucket location 1"}
{"source-ref": "S3 bucket location 2"}
...
{"source-ref": "S3 bucket location n"}
```

Use the source-ref key for image files for bounding box, image classification (single and multi-label), semantic segmentation, and video clips for video classification labeling jobs. 3D point cloud and video frame labeling jobs also use the source-ref key but these labeling jobs require additional information in the input manifest file. For more information see 3D Point Cloud Input Data and Video Frame Input Data.

The following is an example of a manifest file with the input data stored in the manifest:

```
{"source": "Lorem ipsum dolor sit amet"}
{"source": "consectetur adipiscing elit"}
...
{"source": "mollit anim id est laborum"}
```
Use the source key for single and multi-label text classification and named entity recognition labeling jobs.

You can include other key-value pairs in the manifest file. These pairs are passed to the output file unchanged. This is useful when you want to pass information between your applications. For more information, see Output Data.

Automated Data Setup

You can use the automated data setup to create manifest files for your labeling jobs in the Ground Truth console using images, videos, video frames, text (.txt) files, and comma-separated value (.csv) files stored in Amazon S3. When you use automated data setup, you specify an Amazon S3 location where your input data is stored and the input data type, and Ground Truth looks for the files that match that type in the location you specify.

Note

Ground Truth does not use an AWS KMS key to access your input data or write the input manifest file in the Amazon S3 location that you specify. The user or role that creates the labeling job must have permissions to access your input data objects in Amazon S3.

Before using the following procedure, ensure that your input images or files are correctly formatted:

- Image files – Image files must comply with the size and resolution limits listed in the tables found in Input File Size Quota.
- Text files – Text data can be stored in one or more .txt files. Each item that you want labeled must be separated by a standard line break.
- CSV files – Text data can be stored in one or more .csv files. Each item that you want labeled must be in a separate row.
- Videos – Video files can be any of the following formats: .mp4, .ogg, and .webm. If you want to extract video frames from your video files for object detection or object tracking, see Provide Video Files.
- Video frames – Video frames are images extracted from a videos. All images extracted from a single video are referred to as a sequence of video frames. Each sequence of video frames must have unique prefix keys in Amazon S3. See Provide Video Frames. For this data type, see Automated Video Frame Input Data Setup.
Important

For video frame object detection and video frame object tracking labeling jobs, see Automated Video Frame Input Data Setup to learn how to use the automated data setup.

Use these instructions to automatically set up your input dataset connection with Ground Truth.

Automatically connect your data in Amazon S3 with Ground Truth

1. Navigate to the Create labeling job page in the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.

   This link puts you in the North Virginia (us-east-1) AWS Region. If your input data is in an Amazon S3 bucket in another Region, switch to that Region. To change your AWS Region, on the navigation bar, choose the name of the currently displayed Region.

2. Select Create labeling job.

3. Enter a Job name.

4. In the section Input data setup, select Automated data setup.

5. Enter an Amazon S3 URI for S3 location for input datasets.

6. Specify your S3 location for output datasets. This is where your output data is stored.

7. Choose your Data type using the dropdown list.

8. Use the drop down menu under IAM Role to select an execution role. If you select Create a new role, specify the Amazon S3 buckets that you want grant this role permission to access. This role must have permission to access the S3 buckets you specified in Steps 5 and 6.

9. Select Complete data setup.

The following GIF demonstrates how to use the automated data setup for image data. This example will create a file, dataset-YYMMDDTHHMMSS.manifest in the Amazon S3 bucket example-groundtruth-images where YYMMDDTHHMMSS indicates the year (YY), month (MM), day (DD) and time in hours (HH), minutes (mm) and seconds (ss), that the input manifest file was created.

Supported Data Formats

When you create an input manifest file for a built-in task types manually, your input data must be in one of the following support file formats for the respective input data type. To learn about automated data setup, see Automated Data Setup.
### Tip

When you use the automated data setup, additional data formats can be used to generate an input manifest file for video frame and text based task types.

<table>
<thead>
<tr>
<th>Task Types</th>
<th>Input Data Type</th>
<th>Support Formats</th>
<th>Example Input Manifest Line</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bounding Box, Semantic Segmentation, Image Classification (Single Label and Multi-label), Verify and Adjust Labels</td>
<td>Image</td>
<td>.jpg, .jpeg, .png</td>
<td><code>{&quot;source-ref&quot;: &quot;s3://DOC-EXAMPLE-BUCKET/1/example-image.png&quot;}</code></td>
</tr>
<tr>
<td>Named Entity Recognition, Text Classification (Single and Multi-Label)</td>
<td>Text</td>
<td>Raw text</td>
<td><code>{&quot;source&quot;: &quot;Lorem ipsum dolor sit amet&quot;}</code></td>
</tr>
<tr>
<td>Video Classification</td>
<td>Video clips</td>
<td>.mp4, .ogg, and .webm</td>
<td><code>{&quot;source-ref&quot;: &quot;s3:///example-video.mp4&quot;}</code></td>
</tr>
<tr>
<td>Video Frame Object Detection, Video Frame Object Tracking (bounding boxes, polylines, polygons or keypoint)</td>
<td>Video frames and video frame sequence files (for Object Tracking)</td>
<td><strong>Video frames</strong>: .jpg, .jpeg, .png <strong>Sequence files</strong>: .json</td>
<td>Refer to Create a Video Frame Input Manifest File.</td>
</tr>
<tr>
<td>3D Point Cloud Semantic Segmentation, 3D Point Cloud Object Detection, 3D</td>
<td>Point clouds and point cloud sequence files (for Object Tracking)</td>
<td><strong>Point clouds</strong>: Binary pack format and ASCII. For more information see</td>
<td>Refer to Create an Input Manifest File for a 3D Point Cloud Labeling Job.</td>
</tr>
</tbody>
</table>
Ground Truth Streaming Labeling Jobs

If you want to perpetually send new data objects to Amazon SageMaker Ground Truth to be labeled, use a streaming labeling job. Streaming labeling jobs allow you to:

- Send new dataset objects to workers in real time using a perpetually running labeling job. Workers continuously receive new data objects to label as long as the labeling job is active and new objects are being sent to it.
- Gain visibility into the number of objects that have been queued and are waiting to be labeled. Use this information to control the flow of data objects sent to your labeling job.
- Receive label data for individual data objects in real time as workers finish labeling them.

Ground Truth streaming labeling jobs remain active until they are manually stopped or have been idle for more than 10 days. You can intermittently send new data objects to workers while the labeling job is active.

If you are a new user of Ground Truth streaming labeling jobs, it is recommended that you review How It Works.

Use Create a Streaming Labeling Job to learn how to create a streaming labeling job.

Note

Ground Truth streaming labeling jobs are only supported through the SageMaker API.

Topics

- How It Works
- Send Data to a Streaming Labeling Job
How It Works

When you create a Ground Truth streaming labeling job, the job remains active until it is manually stopped, remains idle for more than 10 days, or is unable to access input data sources. You can intermittently send new data objects to workers while it is active. A worker can continue to receive new data objects in real time as long as the total number of tasks currently available to the worker is less than the value in `MaxConcurrentTaskCount`. Otherwise, the data object is sent to a queue that Ground Truth creates on your behalf in Amazon Simple Queue Service (Amazon SQS) for later processing. These tasks are sent to workers as soon as the total number of tasks currently available to a worker falls below `MaxConcurrentTaskCount`. If a data object is not sent to a worker after 14 days, it expires. You can view the number of tasks pending in the queue and adjust the number of objects you send to the labeling job. For example, you may decrease the speed at which you send objects to the labeling job if the backlog of pending objects moves above a threshold.

Send Data to a Streaming Labeling Job

You can optionally submit input data to a streaming labeling job one time when you create the labeling job using an input manifest file. Once the labeling job has started and the state is `InProgress`, you can submit new data objects to your labeling job in real time using your Amazon SNS input topic and Amazon S3 event notifications.

Submit Data Objects When you Start the Labeling Job (One Time):

- **Use an Input Manifest File** – You can optionally specify an input manifest file Amazon S3 URI in `ManifestS3Uri` when you create the streaming labeling job. Ground Truth sends each data object in the manifest file to workers for labeling as soon as the labeling job starts. To learn more, see [Create a Manifest File (Optional)](#).

After you submit a request to create the streaming labeling job, its status will be `Initializing`. Once the labeling job is active, the state changes to `InProgress` and you can start using the real-time options to submit additional data objects for labeling.

Submit Data Objects in Real Time:
• **Send data objects using Amazon SNS messages** – You can send Ground Truth new data objects to label by sending an Amazon SNS message. You will send this message to an Amazon SNS input topic that you create and specify when you create your streaming labeling job. For more information, see [Send Data Objects Using Amazon SNS](#).

• **Send data objects by placing them in an Amazon S3 bucket** – Each time you add a new data object to an Amazon S3 bucket, you can prompt Ground Truth to process that object for labeling. To do this, you add an event notification to the bucket so that it notifies your Amazon SNS input topic each time a new object is added to (or *created in*) that bucket. For more information, see [Send Data Objects using Amazon S3](#). This option is not available for text-based labeling jobs such as text classification and named entity recognition.

**Important**

If you use the Amazon S3 configuration, do not use the same Amazon S3 location for your input data configuration and your output data. You specify the S3 prefix for your output data when you create a labeling job.

---

### Send Data Objects Using Amazon SNS

You can send data objects to your streaming labeling job using Amazon Simple Notification Service (Amazon SNS). Amazon SNS is a web service that coordinates and manages the delivery of messages to and from *endpoints* (for example, an email address or AWS Lambda function). An Amazon SNS *topic* acts as a communication channel between two or more endpoints. You use Amazon SNS to send, or *publish*, new data objects to the topic specified in the `CreateLabelingJob` parameter `SnsTopicArn` in `InputConfig`. The format of these messages is the same as a single line from an *input manifest file*.

For example, you may send a piece of text to an active text classification labeling job by publishing it to your input topic. The message that you publish may look similar to the following:

```json
{"source": "Lorem ipsum dolor sit amet"}
```

To send a new image object to an image classification labeling job, your message may look similar to the following:

```json
{"source-ref": "s3://awsexamplebucket/example-image.jpg"}
```
When Ground Truth creates your streaming labeling job, it subscribes to your Amazon SNS input topic.

**Send Data Objects using Amazon S3**

You can send one or more new data objects to a streaming labeling job by placing them in an Amazon S3 bucket that is configured with an Amazon SNS event notification. You can set up an event to notify your Amazon SNS input topic anytime a new object is created in your bucket. You must specify this same Amazon SNS input topic in the `CreateLabelingJob` parameter `SnsTopicArn` in `InputConfig`.

Anytime you configure an Amazon S3 bucket to send notifications to Amazon SNS, Ground Truth will publish a test event, "s3:TestEvent", to ensure that the topic exists and that the owner of the Amazon S3 bucket specified has permission to publish to the specified topic. It is recommended that you set up your Amazon S3 connection with Amazon SNS before starting a streaming labeling job. If you do not, this test event may register as a data object and be sent to Ground Truth for labeling.

**Important**

If you use the Amazon S3 configuration, do not use the same Amazon S3 location for your input data configuration and your output data. You specify the S3 prefix for your output data when you create a labeling job. For image-based labeling jobs, Ground Truth requires all S3 buckets to have a CORS policy attached. To learn more, see [CORS Permission Requirement](#).

Once you have configured your Amazon S3 bucket and created your labeling job, you can add objects to your bucket and Ground Truth either sends that object to workers or places it on your Amazon SQS queue.

To learn more, see [Set up Amazon S3 Bucket Event Notifications](#).
Manage Labeling Requests with an Amazon SQS Queue

When Ground Truth creates your streaming labeling job, it creates an Amazon SQS queue in the AWS account used to create the labeling job. The queue name is GroundTruth-\textit{labeling\_job\_name} where \textit{labeling\_job\_name} is the name of your labeling job, in lowercase letters. When you send data objects to your labeling job, Ground Truth either sends the data objects directly to workers or places the task in your queue to be processed at a later time. If a data object is not sent to a worker after 14 days, it expires and is removed from the queue. You can setup an alarm in Amazon SQS to detect when objects expire and use this mechanism to control the volume of objects you send to your labeling job.

Important

Modifying, deleting, or sending objects directly to the Amazon SQS queue associated with your streaming labeling job may lead to job failures.

Receive Output Data from a Streaming Labeling Job

Your Amazon S3 output bucket is periodically updated with new output data from your streaming labeling job.

Optionally, you can specify an Amazon SNS output topic. Each time a worker submits a labeled object, a notification with the output data is sent to that topic. You can subscribe an endpoint to your SNS output topic to receive notifications or trigger events when you receive output data from a labeling task. Use an Amazon SNS output topic if you want to do real time chaining to another streaming job and receive an Amazon SNS notifications each time a data object is submitted by a worker.

To learn more, see Subscribe an Endpoint to Your Amazon SNS Output Topic.
**Duplicate Message Handling**

For data objects sent in real time, Ground Truth guarantees idempotency by ensuring each unique object is only sent for labeling once, even if the input message referring to that object is received multiple times (duplicate messages). To do this, each data object sent to a streaming labeling job is assigned a *deduplication ID*, which is identified with a *deduplication key*.

If you send your requests to label data objects directly through your Amazon SNS input topic using Amazon SNS messages, you can optionally choose a custom deduplication key and deduplication IDs for your objects. For more information, see [Specify A Deduplication Key and ID in an Amazon SNS Message](#).

If you do not provide your own deduplication key, or if you use the Amazon S3 configuration to send data objects to your labeling job, Ground Truth uses one of the following for the deduplication ID:

- For messages sent directly to your Amazon SNS input topic, Ground Truth uses the SNS message ID.
- For messages that come from an Amazon S3 configuration, Ground Truth creates a deduplication ID by combining the Amazon S3 URI of the object with the *sequencer token* in the message.

### Specify A Deduplication Key and ID in an Amazon SNS Message

When you send a data object to your streaming labeling job using an Amazon SNS message, you have the option to specify your deduplication key and deduplication ID in one of the following ways. In all of these scenarios, identify your deduplication key with `dataset-objectid-attribute-name`.

**Bring Your Own Deduplication Key and ID**

Create your own deduplication key and deduplication ID by configuring your Amazon SNS message as follows. Replace `byo-key` with your key and `UniqueId` with the deduplication ID for that data object.

```json
{
    "source-ref": "s3://bucket/prefix/object1",
    "dataset-objectid-attribute-name": "byo-key",
    "byo-key": "UniqueId"
}
```
Your deduplication key can be up to 140 characters. Supported patterns include: "^[a-zA-Z0-9]-\*\^[a-zA-Z0-9]".

Your deduplication ID can be up to 1,024 characters. Supported patterns include: ^\(http[s]://([^/]+)/(.*)$.

**Use an Existing Key for your Deduplication Key**

You can use an existing key in your message as the deduplication key. When you do this, the value associated with that key is used for the deduplication ID.

For example, you can specify use the `source-ref` key as your deduplication key by formatting your message as follows:

```
{
    "source-ref": "s3://bucket/prefix/object1",
    "dataset-objectid-attribute-name": "source-ref"
}
```

In this example, Ground Truth uses "s3://bucket/prefix/object1" for the deduplication id.

**Find Deduplication Key and ID in Your Output Data**

You can see the deduplication key and ID in your output data. The deduplication key is identified by `dataset-objectid-attribute-name`.

When you use your own custom deduplication key, your output contains something similar to the following:

```
"dataset-objectid-attribute-name": "byo-key",
"byo-key": "UniqueId",
```

When you do not specify a key, you can find the deduplication ID that Ground Truth assigned to your data object as follows. The `$label-attribute-name-object-id` parameter identifies your deduplication ID.

```
{
    "source-ref": "s3://bucket/prefix/object1",
    "dataset-objectid-attribute-name": "$label-attribute-name-object-id"
}```
For `<service-generated-key>`, if the data object came through an Amazon S3 configuration, Ground Truth adds a unique value used by the service and emits a new field keyed by `$sequencer` which shows the Amazon S3 sequencer used. If object was fed to SNS directly, Ground Truth use the SNS message ID.

**Note**

Do not use the `$` character in your label attribute name.

### Input Data Quotas

Input datasets used in semantic segmentation labeling jobs have a quota of 20,000 items. For all other labeling job types, the dataset size quota is 100,000 items. To request an increase to the quota for labeling jobs other than semantic segmentation jobs, review the procedures in [AWS Service Quotas](https://aws.amazon.com/service-quotas) to request a quota increase.

Input image data for active and non-active learning labeling jobs must not exceed size and resolution quotas. *Active learning* refers to labeling job that use [automated data labeling](https://aws.amazon.com/sagemaker). *Non-active learning* refers to labeling jobs that don't use automated data labeling.

Additional quotas apply for label categories for all task types, and for input data and labeling category attributes for 3D point cloud and video frame task types.

### Input File Size Quota

Input files can't exceed the following size-quotas for both active and non-active learning labeling jobs. There is no input file size quota for videos used in [video classification](https://aws.amazon.com/sagemaker) labeling jobs.

<table>
<thead>
<tr>
<th>Labeling Job Task Type</th>
<th>Input File Size Quota</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image classification</td>
<td>40 MB</td>
</tr>
<tr>
<td>Bounding box (Object detection)</td>
<td>40 MB</td>
</tr>
</tbody>
</table>
### Labeling Job Task Type

<table>
<thead>
<tr>
<th>Labeling Job Task Type</th>
<th>Input File Size Quota</th>
</tr>
</thead>
<tbody>
<tr>
<td>Semantic segmentation</td>
<td>40 MB</td>
</tr>
<tr>
<td>Bounding box (Object detection) label adjustment</td>
<td>40 MB</td>
</tr>
<tr>
<td>Semantic segmentation label adjustment</td>
<td>40 MB</td>
</tr>
<tr>
<td>Bounding box (Object detection) label verification</td>
<td>40 MB</td>
</tr>
<tr>
<td>Semantic segmentation label verification</td>
<td>40 MB</td>
</tr>
</tbody>
</table>

### Input Image Resolution Quotas

Image file resolution refers to the number of pixels in an image, and determines the amount of detail an image holds. Image resolution quotas differ depending on the labeling job type and the SageMaker built-in algorithm used. The following table lists the resolution quotas for images used in active and non-active learning labeling jobs.

<table>
<thead>
<tr>
<th>Labeling Job Task Type</th>
<th>Resolution Quota - Non Active Learning</th>
<th>Resolution Quota - Active Learning</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image classification</td>
<td>100 million pixels</td>
<td>3840 x 2160 pixels (4 K)</td>
</tr>
<tr>
<td>Bounding box (Object detection)</td>
<td>100 million pixels</td>
<td>3840 x 2160 pixels (4 K)</td>
</tr>
<tr>
<td>Semantic segmentation</td>
<td>100 million pixels</td>
<td>1920 x 1080 pixels (1080 p)</td>
</tr>
<tr>
<td>Object detection label adjustment</td>
<td>100 million pixels</td>
<td>3840 x 2160 pixels (4 K)</td>
</tr>
<tr>
<td>Semantic segmentation label adjustment</td>
<td>100 million pixels</td>
<td>1920 x 1080 pixels (1080 p)</td>
</tr>
<tr>
<td>Object detection label verification</td>
<td>100 million pixels</td>
<td>Not available</td>
</tr>
</tbody>
</table>
Labeling Job Task Type | Resolution Quota - Non Active Learning | Resolution Quota - Active Learning
---|---|---
Semantic segmentation label verification | 100 million pixels | Not available

**Label Category Quotas**

Each labeling job task type has a quota for the number of label categories you can specify. Workers select label categories to create annotations. For example, you may specify label categories *car*, *pedestrian*, and *biker* when creating a bounding box labeling job and workers will select the *car* category before drawing bounding boxes around cars.

⚠️ **Important**

Label category names cannot exceed 256 characters.
All label categories must be unique. You cannot specify duplicate label categories.

The following label category limits apply to labeling jobs. Quotas for label categories depend on whether you use the SageMaker API operation `CreateLabelingJob` or the console to create a labeling job.

<table>
<thead>
<tr>
<th>Labeling Job Task Type</th>
<th>Label Category Quota - API</th>
<th>Label Category Quota - Console</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image classification (Multi-label)</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Image classification (Single label)</td>
<td>Unlimited</td>
<td>30</td>
</tr>
<tr>
<td>Bounding box (Object detection)</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Label verification</td>
<td>Unlimited</td>
<td>30</td>
</tr>
<tr>
<td>Labeling Job Task Type</td>
<td>Label Category Quota - API</td>
<td>Label Category Quota - Console</td>
</tr>
<tr>
<td>------------------------------------------------</td>
<td>---------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>Semantic segmentation (with active learning)</td>
<td>20</td>
<td>10</td>
</tr>
<tr>
<td>Semantic segmentation (without active learning)</td>
<td>Unlimited</td>
<td>10</td>
</tr>
<tr>
<td>Named entity recognition</td>
<td>Unlimited</td>
<td>30</td>
</tr>
<tr>
<td>Text classification (Multi-label)</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Text classification (Single label)</td>
<td>Unlimited</td>
<td>30</td>
</tr>
<tr>
<td>Video classification</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Video frame object detection</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Video frame object tracking</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>3D point cloud object detection</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>3D point cloud object tracking</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>3D point cloud semantic segmentation</td>
<td>30</td>
<td>30</td>
</tr>
</tbody>
</table>

### 3D Point Cloud and Video Frame Labeling Job Quotas

The following quotas apply for 3D point cloud and video frame labeling job input data.

<table>
<thead>
<tr>
<th>Labeling Job Task Type</th>
<th>Input Data Quota</th>
</tr>
</thead>
<tbody>
<tr>
<td>Video frame object detection</td>
<td>2,000 video frames (images) per sequence</td>
</tr>
</tbody>
</table>
When you create a video frame or 3D point cloud labeling job, you can add one or more label category attributes to each label category that you specify to have workers provide more information about an annotation.

Each label category attribute has a single label category attribute name, and a list of one or more options (values) to choose from. To learn more, see Worker User Interface (UI) for 3D point cloud labeling jobs and Worker User Interface (UI) for video frame labeling jobs.

The following quotas apply to the number of label category attributes names and values you can specify for labeling jobs.

<table>
<thead>
<tr>
<th>Labeling Job Task Type</th>
<th>Label Category Attribute (name) Quota</th>
<th>Label Category Attribute Values Quota</th>
</tr>
</thead>
<tbody>
<tr>
<td>Video frame object detection</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Video frame object tracking</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>3D point cloud object detection</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>3D point cloud object tracking</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>
Filter and Select Data for Labeling

You can use the Amazon SageMaker console to select a portion of your dataset for labeling. The data must be stored in an Amazon S3 bucket. You have three options:

- Use the full dataset.
- Choose a randomly selected sample of the dataset.
- Specify a subset of the dataset using a query.

The following options are available in the Labeling jobs section of the SageMaker console after selecting Create labeling job. To learn how to create a labeling job in the console, see Getting started. To configure the dataset that you use for labeling, in the Job overview section, choose Additional configuration.

Use the Full Dataset

When you choose to use the Full dataset, you must provide a manifest file for your data objects. You can provide the path of the Amazon S3 bucket that contains the manifest file or use the SageMaker console to create the file. To learn how to create a manifest file using the console, see Automated Data Setup.

Choose a Random Sample

When you want to label a random subset of your data, select Random sample. The dataset is stored in the Amazon S3 bucket specified in the Input dataset location field.

After you have specified the percentage of data objects that you want to include in the sample, choose Create subset. SageMaker randomly picks the data objects for your labeling job. After the objects are selected, choose Use this subset.

SageMaker creates a manifest file for the selected data objects. It also modifies the value in the Input dataset location field to point to the new manifest file.
Specify a Subset

You can specify a subset of your data objects using an Amazon S3 SELECT query on the object file names.

The SELECT statement of the SQL query is defined for you. You provide the WHERE clause to specify which data objects should be returned.

For more information about the Amazon S3 SELECT statement, see Selecting Content from Objects.

Choose Create subset to start the selection, and then choose Use this subset to use the selected data.

SageMaker creates a manifest file for the selected data objects. It also updates the value in the Input dataset location field to point to the new manifest file.

3D Point Cloud Input Data

To create a 3D point cloud labeling job, you must create an input manifest file. Use this topic to learn the formatting requirements of the input manifest file for each task type. To learn about the raw input data formats Ground Truth accepts for 3D point cloud labeling jobs, see the section Accepted Raw 3D Data Formats.

Use your labeling job task type to choose a topics on Create an Input Manifest File for a 3D Point Cloud Labeling Job to learn about the formatting requirements for each line of your input manifest file.

Topics

- Accepted Raw 3D Data Formats
- Create an Input Manifest File for a 3D Point Cloud Labeling Job
- Understand Coordinate Systems and Sensor Fusion

Accepted Raw 3D Data Formats

Ground Truth uses your 3D point cloud data to render a 3D scenes that workers annotate. This section describes the raw data formats that are accepted for point cloud data and sensor fusion data for a point cloud frame. To learn how to create an input manifest file to connect your raw input data files with Ground Truth, see Create an Input Manifest File for a 3D Point Cloud Labeling Job.
For each frame, Ground Truth supports Compact Binary Pack Format (.bin) and ASCII (.txt) files. These files contain information about the location (x, y, and z coordinates) of all points that make up that frame, and, optionally, information about the pixel color of each point for colored point clouds. When you create a 3D point cloud labeling job input manifest file, you can specify the format of your raw data in the `format` parameter.

The following table lists elements that Ground Truth supports in point cloud frame files to describe individual points.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>x</td>
<td>The x coordinate of the point.</td>
</tr>
<tr>
<td>y</td>
<td>The y coordinate of the point.</td>
</tr>
<tr>
<td>z</td>
<td>The z coordinate of the point.</td>
</tr>
<tr>
<td>i</td>
<td>The intensity of the point.</td>
</tr>
<tr>
<td>r</td>
<td>The red color channel component. An 8-bit value (0-255).</td>
</tr>
<tr>
<td>g</td>
<td>The green color channel component. An 8-bit value (0-255)</td>
</tr>
<tr>
<td>b</td>
<td>The blue color channel component. An 8-bit value (0-255)</td>
</tr>
</tbody>
</table>

Ground Truth assumes the following about your input data:

- All of the positional coordinates (x, y, z) are in meters.
- All the pose headings (qx, qy, qz, qw) are measured in Spatial Quaternions.

**Compact Binary Pack Format**

The Compact Binary Pack Format represents a point cloud as an ordered set of a stream of points. Each point in the stream is an ordered binary pack of 4-byte float values in some variant of the
form xyzirgb. The x, y, and z elements are required and additional information about that pixel can be included in a variety of ways using i, r, g, and b.

To use a binary file to input point cloud frame data to a Ground Truth 3D point cloud labeling job, enter binary/ in the format parameter for your input manifest file and replace with the order of elements in each binary pack. For example, you may enter one of the following for the format parameter.

- binary/xyz – When you use this format, your point element stream would be in the following order: x1y1z1i1x2y2z2i2...
- binary/xyzrgb – When you use this format, your point element stream would be in the following order: x1y1z1r1g1b1x2y2z2r2g2b2...
- binary/xyzirgb – When you use this format, your point element stream would be in the following order: x1y1z1i1r1g1b1x2y2z2i2r2g2b2...

When you use a binary file for your point cloud frame data, if you do not enter a value for format, the default pack format binary/xyz is used.

**ASCII Format**

The ASCII format uses a text file to represent a point cloud, where each line in the ASCII point cloud file represents a single point. Each point is a line the text file and contains white space separated values, each of which is a 4-byte float ASCII values. The x, y, and z elements are required for each point and additional information about that point can be included in a variety of ways using i, r, g, and b.

To use a text file to input point cloud frame data to a Ground Truth 3D point cloud labeling job, enter text/ in the format parameter for your input manifest file and replace with the order of point elements on each line.

For example, if you enter text/xyz for format, your text file for each point cloud frame should look similar to the following:

```
x1  y1  z1  i1
x2  y2  z2  i2
...
```

If you enter text/xyzrgb, your text file should look similar to the following:
When you use a text file for your point cloud frame data, if you do not enter a value for `format`, the default format `text/xyzi` will be used.

**Point Cloud Resolution Limits**

Ground Truth does not have a resolution limit for 3D point cloud frames. However, we recommend that you limit each point cloud frame to 500K points for optimal performance. When Ground Truth renders the 3D point cloud visualization, it must be viewable on your workers' computers, which depends on workers' computer hardware. Point cloud frames that are larger than 1 million points may not render on standard machines, or may take too long to load.

**Create an Input Manifest File for a 3D Point Cloud Labeling Job**

When you create a labeling job, you provide an input manifest file where each line of the manifest describes a unit of task to be completed by annotators. The format of your input manifest file depends on your task type.

- If you are creating a 3D point cloud **object detection** or **semantic segmentation** labeling job, each line in your input manifest file contains information about a single 3D point cloud frame. This is called a **point cloud frame input manifest**. To learn more, see [Create a Point Cloud Frame Input Manifest File](#).

- If you are creating a 3D point cloud **object tracking** labeling job, each line of your input manifest file contains a sequence of 3D point cloud frames and associated data. This is called a **point cloud sequence input manifest**. To learn more, see [Create a Point Cloud Sequence Input Manifest](#).

**Create a Point Cloud Frame Input Manifest File**

The manifest is a UTF-8 encoded file in which each line is a complete and valid JSON object. Each line is delimited by a standard line break, \n or \r\n. Because each line must be a valid JSON object, you can't have unescaped line break characters. In the single-frame input manifest file, each line in the manifest contains data for a single point cloud frame. The point cloud frame data can either be stored in binary or ASCII format (see [Accepted Raw 3D Data Formats](#)). This is the manifest file formatting required for 3D point cloud object detection and semantic segmentation. Optionally, you can also provide camera sensor fusion data for each point cloud frame.
Ground Truth supports point cloud and video camera sensor fusion in the world coordinate system for all modalities. If you can obtain your 3D sensor extrinsic (like a LiDAR extrinsic), we recommend that you transform 3D point cloud frames into the world coordinate system using the extrinsic. For more information, see Sensor Fusion.

However, if you cannot obtain a point cloud in world coordinate system, you can provide coordinates in the original coordinate system that the data was captured in. If you are providing camera data for sensor fusion, it is recommended that you provide LiDAR sensor and camera pose in the world coordinate system.

To create a single-frame input manifest file, you will identify the location of each point cloud frame that you want workers to label using the source-ref key. Additionally, you must use the source-ref-metadata key to identify the format of your dataset, a timestamp for that frame, and, optionally, sensor fusion data and video camera images.

The following example demonstrates the syntax used for an input manifest file for a single-frame point cloud labeling job. The example includes two point cloud frames. For details about each parameter, see the table following this example.

```
Important

Each line in your input manifest file must be in JSON Lines format. The following code block shows an input manifest file with two JSON objects. Each JSON object is used to point to and provide details about a single point cloud frame. The JSON objects have been expanded for readability, but you must minimize each JSON object to fit on a single line when creating an input manifest file. An example is provided under this code block.

```
"qx": -0.0211129612379595,
"qy": -0.006495469416730261,
"qz": -0.008024565904865688,
"qw": 0.9997181192298087
}
],
"prefix": "s3://awsexamplebucket/lidar_singleframe_dataset/someprefix/",
"images": [
{
"image-path": "images/frame300.bin_camera0.jpg",
"unix-timestamp": 1566861644.759115,
"fx": 847.7962624528487,
"fy": 850.0340893791985,
"cx": 576.2129134707038,
"cy": 317.2423573573745,
"k1": 0,
"k2": 0,
"k3": 0,
"k4": 0,
"p1": 0,
"p2": 0,
"skew": 0,
"position": {
"x": -2.2722515189268138,
"y": 116.86003310568965,
"z": 1.454614668542299
},
"heading": {
"qx": 0.7594754093069037,
"qy": 0.02181790885672969,
"qz": -0.02461725233103356,
"qw": -0.6496916273040025
},
"camera-model": "pinhole"
}
]

{
"source-ref": "s3://awsexamplebucket/examplefolder/frame2.bin",
"source-ref-metadata":{
"format": "binary/xyz",
"unix-timestamp": 1566861632.759133,
"ego-vehicle-pose":{
"position": {
"x": 1659
}}
}
"x": -2.7161461413869947,
"y": 116.25822288149078,
"z": 1.8348751887989483
},
"heading": {
    "qx": -0.02111296123795955,
    "qy": -0.006495469416730261,
    "qz": -0.008024565904865688,
    "qw": 0.9997181192298087
},
"prefix": "s3://awsexamplebucket/lidar_singleframe_dataset/someprefix/",
"images": [
    {
        "image-path": "images/frame300.bin_camera0.jpg",
        "unix-timestamp": 1566861644.759115,
        "fx": 847.7962624528487,
        "fy": 850.0340893791985,
        "cx": 576.2129134707038,
        "cy": 317.2423573573745,
        "k1": 0,
        "k2": 0,
        "k3": 0,
        "k4": 0,
        "p1": 0,
        "p2": 0,
        "skew": 0,
        "position": {
            "x": -2.2722515189268138,
            "y": 116.86003310568965,
            "z": 1.454614668542299
        },
        "heading": {
            "qx": 0.7594754093069037,
            "qy": 0.02181790885672969,
            "qz": -0.02461725233103356,
            "qw": -0.6496916273040025
        },
        "camera-model": "pinhole"
    }
]
When you create an input manifest file, you must collapse your JSON objects to fit on a single line. For example, the code block above would appear as follows in an input manifest file:

```json
{
    "source-ref": "s3://awsexamplebucket/examplefolder/frame1.bin",
    "source-ref-metadata": {
        "format": "binary/xyz",
        "unix-timestamp": 1566861644.759115,
        "ego-vehicle-pose": {
            "position": {
                "x": -2.716146141389947,
                "y": 116.25822288149078,
                "z": 1.8348751887989483
            },
            "heading": {
                "qx": -0.02111296123795955,
                "qy": -0.006495469416730261,
                "qz": -0.008024565904865688,
                "qw": 0.9997181192298087
            }
        },
        "prefix": "s3://awsexamplebucket/lidar_singleframe_dataset/someprefix/",
        "images": [{
            "image-path": "images/frame300.bin_camera0.jpg",
            "unix-timestamp": 1566861644.759115,
            "fx": 847.7962624528487,
            "fy": 850.0340893791985,
            "cx": 576.2129134707038,
            "cy": 317.2423573573745,
            "k1": 0,
            "k2": 0,
            "k3": 0,
            "k4": 0,
            "p1": 0,
            "p2": 0,
            "skew": 0,
            "position": {
                "x": -2.2722515189268138,
                "y": 116.86003310568965,
                "z": 1.454614668542299
            },
            "heading": {
                "qx": 0.7594754093069037,
                "qy": 0.02181790885672969,
                "qz": -0.0246172533103356,
                "qw": -0.6496916273040025
            },
            "camera-model": "pinhole"
        }],
        "source-ref": "s3://awsexamplebucket/examplefolder/frame2.bin",
        "source-ref-metadata": {
            "format": "binary/xyz",
            "unix-timestamp": 1566861632.759133,
            "ego-vehicle-pose": {
                "position": {
                    "x": -2.716146141389947,
                    "y": 116.25822288149078,
                    "z": 1.8348751887989483
                },
                "heading": {
                    "qx": -0.02111296123795955,
                    "qy": -0.006495469416730261,
                    "qz": -0.008024565904865688,
                    "qw": 0.9997181192298087
                }
            },
            "prefix": "s3://awsexamplebucket/lidar_singleframe_dataset/someprefix/",
            "images": [{
                "image-path": "images/frame300.bin_camera0.jpg",
                "unix-timestamp": 1566861632.759133,
                "fx": 847.7962624528487,
                "fy": 850.0340893791985,
                "cx": 576.2129134707038,
                "cy": 317.2423573573745,
                "k1": 0,
                "k2": 0,
                "k3": 0,
                "k4": 0,
                "p1": 0,
                "p2": 0,
                "skew": 0,
                "position": {
                    "x": -2.2722515189268138,
                    "y": 116.86003310568965,
                    "z": 1.454614668542299
                },
                "heading": {
                    "qx": 0.7594754093069037,
                    "qy": 0.02181790885672969,
                    "qz": -0.0246172533103356,
                    "qw": -0.6496916273040025
                },
                "camera-model": "pinhole"
            }]
        }
    }
}
```

The following table shows the parameters you can include in your input manifest file:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>source-ref</td>
<td>Yes</td>
<td>String</td>
<td>The Amazon S3 location of a single point cloud frame.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted string value format:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>s3://&lt;bucket-name&gt; /&lt;folder-name&gt; /point-cloud-frame-file</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>----------</td>
<td>--------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>source-ref-metadata</td>
<td>Yes</td>
<td>JSON object</td>
<td>Use this parameter to include additional information about the point cloud in source-ref, and to provide camera data for sensor fusion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted parameters:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>format, unix-timestamp,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>ego-vehicle-pose,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>position, prefix,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>images</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>---------------</td>
<td>----------</td>
<td>-----------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>format</td>
<td>No</td>
<td>String</td>
<td>Use this parameter to specify the format of your point cloud data. For more information, see <a href="#">Accepted Raw 3D Data Formats</a>.</td>
</tr>
<tr>
<td>unix-timestamp</td>
<td>Yes</td>
<td>Number</td>
<td>The unix timestamp is the number of seconds since January 1st, 1970 until the UTC time that the data was collected by a sensor.</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>ego-vehicle-pose</td>
<td>No</td>
<td>JSON object</td>
<td>The pose of the device used to collect the point cloud data. For more information about this parameter, see Include Vehicle Pose Information in Your Input Manifest.</td>
</tr>
<tr>
<td>prefix</td>
<td>No</td>
<td>String</td>
<td>The location in Amazon S3 where your metadata, such as camera images, is stored for this frame. The prefix must end with a forward slash: /.</td>
</tr>
<tr>
<td>images</td>
<td>No</td>
<td>List</td>
<td>A list of parameters describing color camera images used for sensor fusion. You can include up to 8 images in this list. For more information about the parameters required for each image, see Include Camera Data in Your Input Manifest.</td>
</tr>
</tbody>
</table>
Include Vehicle Pose Information in Your Input Manifest

Use the ego-vehicle location to provide information about the location of the vehicle used to capture point cloud data. Ground Truth use this information to compute LiDAR extrinsic matrix.

Ground Truth uses extrinsic matrices to project labels to and from the 3D scene and 2D images. For more information, see Sensor Fusion.

The following table provides more information about the position and orientation (heading) parameters that are required when you provide ego-vehicle information.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
</table>
| position  | Yes      | JSON object     | The translation vector of the ego vehicle in the world coordinate system. Required Parameters: 

x, y, and z. Enter numbers for these parameters. |
| heading   | Yes      | JSON Object     | The orientation of the frame of reference of the device or sensor mounted on the vehicle sensing the surrounding, measured in quaternions, (qx, qy, qz, qw) in the a coordinate system. Required Parameters: 

qx, qy, qz, and qw. Enter numbers for these parameters. |

Include Camera Data in Your Input Manifest

If you want to include video camera data with a frame, use the following parameters to provide information about each image. The Required column below applies when the images parameter is
included in the input manifest file under `source-ref-metadata`. You are not required to include images in your input manifest file.

If you include camera images, you must include information about the camera position and heading used to capture the images in the world coordinate system.

If your images are distorted, Ground Truth can automatically undistort them using information you provide about the image in your input manifest file, including distortion coefficients (k1, k2, k3, k4, p1, p1), the camera model and the camera intrinsic matrix. The intrinsic matrix is made up of focal length (fx, fy), and the principal point (cx, cy). See Intrinsic Matrix to learn how Ground Truth uses the camera intrinsic. If distortion coefficients are not included, Ground Truth will not undistort an image.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>image-path</td>
<td>Yes</td>
<td>String</td>
<td>The relative location, in Amazon S3 of your image file. This relative path will be appended to the path you specify in prefix.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Example of format:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>&lt;folder-name&gt;/imagefile.png&gt;</code></td>
<td></td>
</tr>
<tr>
<td>unix-timestamp</td>
<td>Yes</td>
<td>Number</td>
<td>The unix timestamp is the number of seconds since January 1st, 1970 until the UTC time that the data was collected by a camera.</td>
</tr>
<tr>
<td>camera-model</td>
<td>No</td>
<td>String:</td>
<td>The model of the camera used to capture the image. This information is used to undistort camera images.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted Values:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;pinhole&quot;, &quot;fisheye&quot;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default:</td>
<td></td>
</tr>
</tbody>
</table>

Use Input and Output Data
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;pinhole&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>fx, fy</td>
<td>Yes</td>
<td>Numbers</td>
<td>The focal length of the camera, in the x (fx) and y (fy) directions.</td>
</tr>
<tr>
<td>cx, cy</td>
<td>Yes</td>
<td>Numbers</td>
<td>The x (cx) and y (cy) coordinates of the principal point.</td>
</tr>
<tr>
<td>k1, k2, k3, k4</td>
<td>No</td>
<td>Number</td>
<td>Radial distortion coefficients. Supported for both fisheye and pinhole camera models.</td>
</tr>
<tr>
<td>p1, p2</td>
<td>No</td>
<td>Number</td>
<td>Tangential distortion coefficients. Supported for pinhole camera models.</td>
</tr>
<tr>
<td>skew</td>
<td>No</td>
<td>Number</td>
<td>A parameter to measure the skew of an image.</td>
</tr>
<tr>
<td>position</td>
<td>Yes</td>
<td>JSON object</td>
<td>The location or origin of the frame of reference of the camera mounted on the vehicle capturing images.</td>
</tr>
</tbody>
</table>

**Required Parameters:**

- x, y, and z. Enter numbers for these parameters.
### Parameter

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>heading</td>
<td>Yes</td>
<td>JSON Object</td>
<td>The orientation of the frame of reference of the camera mounted on the vehicle capturing images, measured using quaternions, ( (qx, qy, qz, qw) ), in the world coordinate system.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Required Parameter(s): qx, qy, qz, and qw. Enter numbers for these parameters.</td>
<td></td>
</tr>
</tbody>
</table>

**Point Cloud Frame Limits**

You can include up to 100,000 point cloud frames in your input manifest file. 3D point cloud labeling job have longer pre-processing times than other Ground Truth task types. For more information, see [Job Pre-processing Time](#).

**Create a Point Cloud Sequence Input Manifest**

The manifest is a UTF-8 encoded file in which each line is a complete and valid JSON object. Each line is delimited by a standard line break, \n or \r\n. Because each line must be a valid JSON object, you can't have unescaped line break characters. In the point cloud sequence input manifest file, each line in the manifest contains a sequence of point cloud frames. The point cloud data for each frame in the sequence can either be stored in binary or ASCII format. For more information, see [Accepted Raw 3D Data Formats](#). This is the manifest file formatting required for 3D point cloud object tracking. Optionally, you can also provide point attribute and camera sensor fusion data for each point cloud frame. When you create a sequence input manifest file, you must provide LiDAR and video camera sensor fusion data in a [world coordinate system](#).

The following example demonstrates the syntax used for an input manifest file when each line in the manifest is a sequence file. Each line in your input manifest file must be in [JSON Lines](#) format.

```json
{"source-ref": "s3://awsexamplebucket/example-folder/seq1.json"}
{"source-ref": "s3://awsexamplebucket/example-folder/seq2.json"}
```
The data for each sequence of point cloud frames needs to be stored in a JSON data object. The following is an example of the format you use for a sequence file. Information about each frame is included as a JSON object and is listed in the frames list. This is an example of a sequence file with two point cloud frame files, frame300.bin and frame303.bin. The ... is used to indicated where you should include information for additional frames. Add a JSON object for each frame in the sequence.

The following code block includes a JSON object for a single sequence file. The JSON object has been expanded for readability.

```json
{
    "seq-no": 1,
    "prefix": "s3://awsexamplebucket/example_lidar_sequence_dataset/seq1/",
    "number-of-frames": 100,
    "frames": [  
        {  
            "frame-no": 300,
            "unix-timestamp": 1566861644.759115,
            "frame": "example_lidar_frames/frame300.bin",
            "format": "binary/xyzi",
            "ego-vehicle-pose": {  
                "position": {  
                    "x": -2.7161461413869947,
                    "y": 116.25822288149078,
                    "z": 1.8348751887989483  
                },
                "heading": {  
                    "qx": -0.02111296123795955,
                    "qy": -0.006495469416730261,
                    "qz": -0.008024565904865688,
                    "qw": 0.9997181192298087  
                }  
            },
            "images": [  
                {  
                    "image-path": "example_images/frame300.bin_camera0.jpg",
                    "unix-timestamp": 1566861644.759115,
                    "fx": 847.7962624528487,
                    "fy": 850.0340893791985,
                    "cx": 576.2129134707038,
                    "cy": 317.2423573573745,
                    "k1": 0,
                    "k2": 0,
                }  
            ]  
        }  
    ]  
}
```
"k3": 0,
"k4": 0,
"p1": 0,
"p2": 0,
"skew": 0,
"position": {
  "x": -2.2722515189268138,
  "y": 116.86003310568965,
  "z": 1.454614668542299
},
"heading": {
  "qx": 0.7594754093069037,
  "qy": 0.02181790885672969,
  "qz": -0.02461725233103356,
  "qw": -0.6496916273040025
},
"camera-model": "pinhole"
}],

The following table provides details about the top-level parameters of a sequence file. For detailed information about the parameters required for individual frames in the sequence file, see Parameters for Individual Point Cloud Frames.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>seq-no</td>
<td>Yes</td>
<td>Integer</td>
<td>The ordered number of the sequence.</td>
</tr>
<tr>
<td>prefix</td>
<td>Yes</td>
<td>String</td>
<td>The Amazon S3 location where the</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>----------</td>
<td>----------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted Values: s3://&lt;bucket-name&gt; /&lt;prefix&gt;/</td>
<td>sequence files are located. The prefix must end with a forward slash: /.</td>
</tr>
<tr>
<td>number-of-frames</td>
<td>Yes</td>
<td>Integer</td>
<td>The total number of frames included in the sequence file. This number must match the total number of frames listed in the frames parameter in the next row.</td>
</tr>
<tr>
<td>frames</td>
<td>Yes</td>
<td>List of JSON objects</td>
<td>A list of frame data. The length of the list must equal number-of-frames. In the worker UI, frames in a sequence will be the same as the order of frames in this array. For details about the format of each frame, see <a href="#">Parameters for Individual Point Cloud Frames</a>.</td>
</tr>
</tbody>
</table>

**Parameters for Individual Point Cloud Frames**

The following table shows the parameters you can include in your input manifest file.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>frame-no</td>
<td>No</td>
<td>Integer</td>
<td>A frame number. This is an optional identifier specified by the customer to identify the frame within a sequence. It is not used by Ground Truth.</td>
</tr>
</tbody>
</table>
| unix-timestamp| Yes      | Number          | The unix timestamp is the number of seconds since January 1st, 1970 until the UTC time that the data was collected by a sensor. The timestamp for each frame must be different and timestamps must be sequential because they are used for cuboid interpolation. Ideally, this should be the real timestamp when the data was collected. If this is not available, you must use an incremental sequence of timestamps, where the first frame in your sequence file corresponds to the
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>frame</td>
<td>Yes</td>
<td>String</td>
<td>The relative location, in Amazon S3 of your sequence file. This relative path will be appended to the path you specify in prefix.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Example of format</td>
<td>example of format</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>&lt;folder-name&gt; /&lt;sequence-file.json&gt;</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>----------</td>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>format</td>
<td>No</td>
<td>String</td>
<td>Use this parameter to specify the format of your point cloud data. For more information, see <a href="#">Accepted Raw 3D Data Formats</a>.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted string values: &quot;binary/xyz&quot;,&quot;binary/xyzri&quot;,&quot;binary/xyzrgb&quot;,&quot;binary/xyzirgb&quot;,&quot;text/xyz&quot;,&quot;text/xyzri&quot;,&quot;text/xyzrgb&quot;,&quot;text/xyzirgb&quot;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default Values:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>When the file identified in source-ref has a .bin extension, binary/xyzri</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>When the file identified in source-ref has a .txt extension, text/xyzri</td>
<td></td>
</tr>
<tr>
<td>ego-vehicle-pose</td>
<td>No</td>
<td>JSON object</td>
<td>The pose of the device used to collect the point cloud data. For more information about this parameter, see <a href="#">Include Vehicle Pose Information in Your Input Manifest</a>.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>prefix</td>
<td>No</td>
<td>String</td>
<td>The location in Amazon S3 where your metadata, such as camera images, is stored for this frame. The prefix must end with a forward slash: <code>/</code>.</td>
</tr>
<tr>
<td>images</td>
<td>No</td>
<td>List</td>
<td>A list parameter describing color camera images used for sensor fusion. You can include up to 8 images in this list. For more information about the parameters required for each image, see Include Camera Data in Your Input Manifest.</td>
</tr>
</tbody>
</table>

### Include Vehicle Pose Information in Your Input Manifest

Use the ego-vehicle location to provide information about the pose of the vehicle used to capture point cloud data. Ground Truth use this information to compute LiDAR extrinsic matrices.

Ground Truth uses extrinsic matrices to project labels to and from the 3D scene and 2D images. For more information, see Sensor Fusion.

The following table provides more information about the position and orientation (heading) parameters that are required when you provide ego-vehicle information.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>position</td>
<td>Yes</td>
<td>JSON object</td>
<td>The translation vector of the ego vehicle in the world coordinate system.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Required Parameter</strong>s:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>x, y, and z. Enter numbers for these parameters.</td>
<td></td>
</tr>
<tr>
<td>heading</td>
<td>Yes</td>
<td>JSON Object</td>
<td>The orientation of the frame of reference of the device or sensor mounted on the vehicle sensing the surrounding, measured in quaternions, ((qx, qy, qz, qw)) in the a coordinate system.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Required Parameter</strong>s:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>qx, qy, qz, and qw. Enter numbers for these parameters.</td>
<td></td>
</tr>
</tbody>
</table>

### Include Camera Data in Your Input Manifest

If you want to include color camera data with a frame, use the following parameters to provide information about each image. The **Required** column in the following table applies when the images parameter is included in the input manifest file. You are not required to include images in your input manifest file.

If you include camera images, you must include information about the position and orientation (heading) of the camera used the capture the images.

If your images are distorted, Ground Truth can automatically undistort them using information you provide about the image in your input manifest file, including distortion coefficients \((k1, k2, k3, k4, p1, p1)\), camera model and focal length \((fx, fy)\), and the principal point \((cx, cy)\). To learn more about these coefficients and undistorting images, see [Camera calibration With OpenCV](#). If distortion coefficients are not included, Ground Truth will not undistort an image.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>image-path</td>
<td>Yes</td>
<td>String</td>
<td>The relative location, in Amazon S3 of your image file. This relative path will be appended to the path you specify in prefix.</td>
</tr>
<tr>
<td>unix-timestamp</td>
<td>Yes</td>
<td>Number</td>
<td>The timestamp of the image.</td>
</tr>
<tr>
<td>camera-model</td>
<td>No</td>
<td>String:</td>
<td>The model of the camera used to capture the image. This information is used to undistort camera images.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted Values:</td>
<td>&quot;pinhole&quot;, &quot;fisheye&quot;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default:</td>
<td>&quot;pinhole&quot;</td>
</tr>
<tr>
<td>fx, fy</td>
<td>Yes</td>
<td>Numbers</td>
<td>The focal length of the camera, in the x (fx) and y (fy) directions.</td>
</tr>
<tr>
<td>cx, cy</td>
<td>Yes</td>
<td>Numbers</td>
<td>The x (cx) and y (cy) coordinates of the principal point.</td>
</tr>
<tr>
<td>k1, k2, k3, k4</td>
<td>No</td>
<td>Number</td>
<td>Radial distortion coefficients. Supported for both fisheye and pinhole camera models.</td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>p1, p2</td>
<td>No</td>
<td>Number</td>
<td>Tangential distortion coefficients. Supported for <strong>pinhole</strong> camera models.</td>
</tr>
<tr>
<td>skew</td>
<td>No</td>
<td>Number</td>
<td>A parameter to measure any known skew in the image.</td>
</tr>
<tr>
<td>position</td>
<td>Yes</td>
<td>JSON Object</td>
<td>The location or origin of the frame of reference of the camera mounted on the vehicle capturing images. <strong>Required Parameters:</strong> x, y, and z. Enter numbers for these parameters.</td>
</tr>
<tr>
<td>heading</td>
<td>Yes</td>
<td>JSON Object</td>
<td>The orientation of the frame of reference of the camera mounted on the vehicle capturing images, measured using quaternions, (qx, qy, qz, qw). <strong>Required Parameters:</strong> qx, qy, qz, and qw. Enter numbers for these parameters.</td>
</tr>
</tbody>
</table>

**Sequence File and Point Cloud Frame Limits**

You can include up to 100,000 point cloud frame sequences in your input manifest file. You can include up to 500 point cloud frames in each sequence file.

Keep in mind that 3D point cloud labeling job have longer pre-processing times than other Ground Truth task types. For more information, see [Job Pre-processing Time](#).
Understand Coordinate Systems and Sensor Fusion

Point cloud data is always located in a coordinate system. This coordinate system may be local to the vehicle or the device sensing the surroundings, or it may be a world coordinate system. When you use Ground Truth 3D point cloud labeling jobs, all the annotations are generated using the coordinate system of your input data. For some labeling job task types and features, you must provide data in a world coordinate system.

In this topic, you'll learn the following:

- When you are required to provide input data in a world coordinate system or global frame of reference.
- What a world coordinate is and how you can convert point cloud data to a world coordinate system.
- How you can use your sensor and camera extrinsic matrices to provide pose data when using sensor fusion.

Coordinate System Requirements for Labeling Jobs

If your point cloud data was collected in a local coordinate system, you can use an extrinsic matrix of the sensor used to collect the data to convert it to a world coordinate system or a global frame of reference. If you cannot obtain an extrinsic for your point cloud data and, as a result, cannot obtain point clouds in a world coordinate system, you can provide point cloud data in a local coordinate system for 3D point cloud object detection and semantic segmentation task types.

For object tracking, you must provide point cloud data in a world coordinate system. This is because when you are tracking objects across multiple frames, the ego vehicle itself is moving in the world and so all of the frames need a point of reference.

If you include camera data for sensor fusion, it is recommended that you provide camera poses in the same world coordinate system as the 3D sensor (such as a LiDAR sensor).

Using Point Cloud Data in a World Coordinate System

This section explains what a world coordinate system (WCS), also referred to as a global frame of reference, is and explains how you can provide point cloud data in a world coordinate system.
What is a World Coordinate System?

A WCS or global frame of reference is a fixed universal coordinate system in which vehicle and sensor coordinate systems are placed. For example, if multiple point cloud frames are located in different coordinate systems because they were collected from two sensors, a WCS can be used to translate all of the coordinates in these point cloud frames into a single coordinate system, where all frames have the same origin, (0,0,0). This transformation is done by translating the origin of each frame to the origin of the WCS using a translation vector, and rotating the three axes (typically x, y, and z) to the right orientation using a rotation matrix. This rigid body transformation is called a homogeneous transformation.

A world coordinate system is important in global path planning, localization, mapping, and driving scenario simulations. Ground Truth uses the right-handed Cartesian world coordinate system such as the one defined in ISO 8855, where the x axis is forward toward the car’s movement, y axis is left, and the z axis points up from the ground.

The global frame of reference depends on the data. Some datasets use the LiDAR position in the first frame as the origin. In this scenario, all the frames use the first frame as a reference and device heading and position will be near the origin in the first frame. For example, KITTI datasets have the first frame as a reference for world coordinates. Other datasets use a device position that is different from the origin.

Note that this is not the GPS/IMU coordinate system, which is typically rotated by 90 degrees along the z-axis. If your point cloud data is in a GPS/IMU coordinate system (such as OxTS in the open source AV KITTI dataset), then you need to transform the origin to a world coordinate system (typically the vehicle's reference coordinate system). You apply this transformation by multiplying your data with transformation metrics (the rotation matrix and translation vector). This will transform the data from its original coordinate system to a global reference coordinate system. Learn more about this transformation in the next section.

Convert 3D Point Cloud Data to a WCS

Ground Truth assumes that your point cloud data has already been transformed into a reference coordinate system of your choice. For example, you can choose the reference coordinate system of the sensor (such as LiDAR) as your global reference coordinate system. You can also take point clouds from various sensors and transform them from the sensor's view to the vehicle's reference coordinate system view. You use the a sensor's extrinsic matrix, made up of a rotation matrix and translation vector, to convert your point cloud data to a WCS or global frame of reference.
Collectively, the translation vector and rotation matrix can be used to make up an *extrinsic matrix*, which can be used to convert data from a local coordinate system to a WCS. For example, your LiDAR extrinsic matrix may be composed as follows, where $R$ is the rotation matrix and $T$ is the translation vector:

$$\text{LiDAR} \text{ extrinsic} = [R \ T;0 \ 0 \ 0 \ 1]$$

For example, the autonomous driving KITTI dataset includes a rotation matrix and translation vector for the LiDAR extrinsic transformation matrix for each frame. The `pykitti` python module can be used for loading the KITTI data, and in the dataset dataset.oxts[i].T_w_imu gives the LiDAR extrinsic transform for the $i^{th}$ frame with can be multiplied with points in that frame to convert them to a world frame - `np.matmul(lidar_transform_matrix, points)`.

Multiplying a point in LiDAR frame with a LiDAR extrinsic matrix transforms it into world coordinates. Multiplying a point in the world frame with the camera extrinsic matrix gives the point coordinates in the camera's frame of reference.

The following code example demonstrates how you can convert point cloud frames from the KITTI dataset into a WCS.

```python
import pykitti
import numpy as np

basedir = '/Users/nameofuser/kitti-data'
date = '2011_09_26'
drive = '0079'

# The 'frames' argument is optional - default: None, which loads the whole dataset.
# Calibration, timestamps, and IMU data are read automatically.
# Camera and velodyne data are available via properties that create generators
# when accessed, or through getter methods that provide random access.
data = pykitti.raw(basedir, date, drive, frames=range(0, 50, 5))

# i is frame number
i = 0

# lidar extrinsic for the ith frame
lidar_extrinsic_matrix = data.oxts[i].T_w_imu

# velodyne raw point cloud in lidar scanners own coordinate system
points = data.get_velo(i)
```
# transform points from lidar to global frame using lidar_extrinsic_matrix

def generate_transformed_pcd_from_point_cloud(points, lidar_extrinsic_matrix):
    tps = []
    for point in points:
        transformed_points = np.matmul(lidar_extrinsic_matrix, np.array([point[0],
            point[1], point[2], 1], dtype=np.float32).reshape(4,1)).tolist()
        if len(point) > 3 and point[3] is not None:
            tps.append([transformed_points[0][0], transformed_points[1][0],
                transformed_points[2][0], point[3]])

    return tps

# customer transforms points from lidar to global frame using lidar_extrinsic_matrix
transformed_pcl = generate_transformed_pcd_from_point_cloud(points, lidar_extrinsic_matrix)

---

**Sensor Fusion**

Ground Truth supports sensor fusion of point cloud data with up to 8 video camera inputs. This feature allows human labellers to view the 3D point cloud frame side-by-side with the synchronized video frame. In addition to providing more visual context for labeling, sensor fusion allows workers to adjust annotations in the 3D scene and in 2D images and the adjustment are projected into the other view. The following video demonstrates a 3D point cloud labeling job with LiDAR and camera sensor fusion.
For best results, when using sensor fusion, your point cloud should be in a WCS. Ground Truth uses your sensor (such as LiDAR), camera, and ego vehicle pose information to compute extrinsic and intrinsic matrices for sensor fusion.

**Extrinsic Matrix**

Ground Truth uses sensor (such as LiDAR) extrinsic and camera extrinsic and intrinsic matrices to project objects to and from the point cloud data's frame of reference to the camera's frame of reference.

For example, in order to project a label from the 3D point cloud to camera image plane, Ground Truth transforms 3D points from LiDAR’s own coordinate system to the camera's coordinate system. This is typically done by first transforming 3D points from LiDAR’s own coordinate system to a world coordinate system (or a global reference frame) using the LiDAR extrinsic matrix. Ground Truth then uses the camera inverse extrinsic (which converts points from a global frame of reference to the camera's frame of reference) to transform the 3D points from world coordinate system obtained in previous step into the camera image plane. The LiDAR extrinsic matrix can also be used to transform 3D data into a world coordinate system. If your 3D data is already transformed into world coordinate system then the first transformation doesn’t have any impact on label translation, and label translation only depends on the camera inverse extrinsic. A view matrix is used to visualize projected labels. To learn more about these transformations and the view matrix, see [Ground Truth Sensor Fusion Transformations](#).

Ground Truth computes these extrinsic matrices by using LiDAR and camera pose data that you provide: heading (in quaternions: qx, qy, qz, and qw) and position (x, y, z). For the vehicle, typically the heading and position are described in vehicle's reference frame in a world coordinate system and are called a *ego vehicle pose*. For each camera extrinsic, you can add pose information for that camera. For more information, see [Pose](#).

**Intrinsic Matrix**

Ground Truth use the camera extrinsic and intrinsic matrices to compute view metrics to transform labels to and from the 3D scene to camera images. Ground Truth computes the camera intrinsic matrix using camera focal length (fx, fy) and optical center coordinates (cx, cy) that you provide. For more information, see [Intrinsic and Distortion](#).

**Image Distortion**

Image distortion can occur for a variety of reasons. For example, images may be distorted due to barrel or fish-eye effects. Ground Truth uses intrinsic parameters along with distortion co-efficient
to undistort images you provide when creating 3D point cloud labeling jobs. If a camera image is already been undistorted, all distortion coefficients should be set to 0.

For more information about the transformations Ground Truth performs to undistort images, see Camera Calibrations: Extrinsic, Intrinsic and Distortion.

**Ego Vehicle**

To collect data for autonomous driving applications, the measurements used to generate point cloud data and are taken from sensors mounted on a vehicle, or the *ego vehicle*. To project label adjustments to and from the 3D scene and 2D images, Ground Truth needs your ego vehicle pose in a world coordinate system. The ego vehicle pose is comprised of position coordinates and orientation quaternion.

Ground Truth uses your ego vehicle pose to compute rotation and transformations matrices. Rotations in 3 dimensions can be represented by a sequence of 3 rotations around a sequence of axes. In theory, any three axes spanning the 3D Euclidean space are enough. In practice, the axes of rotation are chosen to be the basis vectors. The three rotations are expected to be in a global frame of reference (extrinsic). Ground Truth does not a support body centered frame of reference (intrinsic) which is attached to, and moves with, the object under rotation. To track objects, Ground Truth needs to measure from a global reference where all vehicles are moving. When using Ground Truth 3D point cloud labeling jobs, z specifies the axis of rotation (extrinsic rotation) and yaw Euler angles are in radians (rotation angle).

**Pose**

Ground Truth uses pose information for 3D visualizations and sensor fusion. Pose information you input through your manifest file is used to compute extrinsic matrices. If you already have an extrinsic matrix, you can use it to extract sensor and camera pose data.

For example in the autonomous driving KITTI dataset, the pykitti python module can be used for loading the KITTI data. In the dataset dataset.oxts[i].T_w_imu gives the LiDAR extrinsic transform for the i th frame and it can be multiplied with the points to get them in a world frame - matmul(lidar_transform_matrix, points). This transform can be converted into position (translation vector) and heading (in quaternion) of LiDAR for the input manifest file JSON format. Camera extrinsic transform for cam0 in i th frame can be calculated by inv(matmul(dataset.calib.T_cam0_velo, inv(dataset.oxts[i].T_w_imu))) and this can be converted into heading and position for cam0.

```python
import numpy
```
rotation = [[ 9.96714314e-01, -8.09890350e-02,  1.16333982e-03],
[ 8.09967396e-02,  9.96661051e-01, -1.03090934e-02],
[-3.24531964e-04,  1.03694477e-02,  9.99946183e-01]]

origin= [1.71104606e+00,
9.43144935e-01,
5.80000039e-01]

from scipy.spatial.transform import Rotation as R

# position is the origin
position = origin
r = R.from_matrix(np.asarray(rotation))

# heading in WCS using scipy
heading = r.as_quat()
print(f"pose:{position}\nheading: {heading}"

**Position**

In the input manifest file, `position` refers to the position of the sensor with respect to a world frame. If you are unable to put the device position in a world coordinate system, you can use LiDAR data with local coordinates. Similarly, for mounted video cameras you can specify the position and heading in a world coordinate system. For camera, if you do not have position information, please use (0, 0, 0).

The following are the fields in the position object:

1. **x** (float) – x coordinate of ego vehicle, sensor, or camera position in meters.
2. **y** (float) – y coordinate of ego vehicle, sensor, or camera position in meters.
3. **z** (float) – z coordinate of ego vehicle, sensor, or camera position in meters.

The following is an example of a `position` JSON object:

```json
{
  "position": {
    "y": -152.77584902657554,
    "x": 311.21505956090624,
    "z": -10.854137529636024
```
Heading

In the input manifest file, heading is an object that represents the orientation of a device with respect to world frame. Heading values should be in quaternion. A quaternion is a representation of the orientation consistent with geodesic spherical properties. If you are unable to put the sensor heading in world coordinates, please use the identity quaternion \((q_x = 0, q_y = 0, q_z = 0, q_w = 1)\). Similarly, for cameras, specify the heading in quaternions. If you are unable to obtain extrinsic camera calibration parameters, please also use the identity quaternion.

Fields in heading object are as follows:

1. \(q_x\) (float) - x component of ego vehicle, sensor, or camera orientation.
2. \(q_y\) (float) - y component of ego vehicle, sensor, or camera orientation.
3. \(q_z\) (float) - z component of ego vehicle, sensor, or camera orientation.
4. \(q_w\) (float) - w component of ego vehicle, sensor, or camera orientation.

The following is an example of a heading JSON object:

```
{
    "heading": {
        "qy": -0.7046155108831117,
        "qx": 0.034278837280808494,
        "qz": 0.7070617895701465,
        "qw": -0.04904659893885366
    }
}
```

To learn more, see Compute Orientation Quaternions and Position.

Compute Orientation Quaternions and Position

Ground Truth requires that all orientation, or heading, data be given in quaternions. A quaternion is a representation of the orientation consistent with geodesic spherical properties that can be used to approximate of rotation. Compared to Euler angles they are simpler to compose and avoid the problem of gimbal lock. Compared to rotation matrices they are more compact, more numerically stable, and more efficient.
You can compute quaternions from a rotation matrix or a transformation matrix.

If you have a rotation matrix (made up of the axis rotations) and translation vector (or origin) in world coordinate system instead of a single 4x4 rigid transformation matrix, then you can directly use the rotation matrix and translation vector to compute quaternions. Libraries like scipy and pyquaternion can help. The following code-block shows an example using these libraries to compute quaternion from a rotation matrix.

```python
import numpy

rotation = [[9.96714314e-01, -8.09890350e-02, 1.16333982e-03],
            [8.09967396e-02, 9.96661051e-01, -1.03090934e-02],
            [-3.24531964e-04, 1.03694477e-02, 9.99946183e-01]]

data = 1.71104606e+00,
       5.80000039e-01,
       9.43144935e-01

from scipy.spatial.transform import Rotation as R
# position is the origin
position = origin
r = R.from_matrix(np.asarray(rotation))
# heading in WCS using scipy
heading = r.as_quat()
print(f"position:{position}\nheading: {heading}"

A UI tool like 3D Rotation Converter can also be useful.

If you have a 4x4 extrinsic transformation matrix, note that the transformation matrix is in the form \([R \ T; 0 \ 0 \ 0 \ 1]\) where \(R\) is the rotation matrix and \(T\) is the origin translation vector. That means you can extract rotation matrix and translation vector from the transformation matrix as follows.

```python
import numpy as np

transformation = [[9.96714314e-01, -8.09890350e-02, 1.16333982e-03, 1.71104606e+00],
                  [8.09967396e-02, 9.96661051e-01, -1.03090934e-02, 5.80000039e-01],
                  [-3.24531964e-04, 1.03694477e-02, 9.99946183e-01, 9.43144935e-01],
                  [0, 0, 0, 1]]
```
transformation = np.array(transformation)
rotation = transformation[0:3][0:3]
translation = transformation[0:3][3]

from scipy.spatial.transform import Rotation as R

# position is the origin translation
position = translation
r = R.from_matrix(np.asarray(rotation))

# heading in WCS using scipy
heading = r.as_quat()
print(f"position: {position}
heading: {heading}"

With your own setup, you can compute an extrinsic transformation matrix using the GPS/IMU position and orientation (latitude, longitude, altitude and roll, pitch, yaw) with respect to the LiDAR sensor on the ego vehicle. For example, you can compute pose from KITTI raw data using pose = convertOxtsToPose(oxts) to transform the oxts data into a local euclidean poses, specified by 4x4 rigid transformation matrices. You can then transform this pose transformation matrix to a global reference frame using the reference frames transformation matrix in the world coordinate system.

struct Quaternion
{
    double w, x, y, z;
};

Quaternion ToQuaternion(double yaw, double pitch, double roll) // yaw (Z), pitch (Y), roll (X)
{
    // Abbreviations for the various angular functions
    double cy = cos(yaw * 0.5);
    double sy = sin(yaw * 0.5);
    double cp = cos(pitch * 0.5);
    double sp = sin(pitch * 0.5);
    double cr = cos(roll * 0.5);
    double sr = sin(roll * 0.5);
    Quaternion q;
    q.w = cr * cp * cy + sr * sp * sy;
    q.x = sr * cp * cy - cr * sp * sy;
    q.y = cr * sp * cy + sr * cp * sy;
    q.z = cr * cp * sy - sr * sp * cy;
Ground Truth Sensor Fusion Transformations

The following sections go into greater detail about the Ground Truth sensor fusion transformations that are performed using the pose data you provide.

LiDAR Extrinsic

In order to project to and from a 3D LiDAR scene to a 2D camera image, Ground Truth computes the rigid transformation projection metrics using the ego vehicle pose and heading. Ground Truth computes rotation and translation of a world coordinates into the 3D plane by doing a simple sequence of rotations and translation.

Ground Truth computes rotation metrics using the heading quaternions as follows:

\[
M = \begin{pmatrix}
1 - 2y^2 - 2z^2 & 2xy + 2zw & 2xz - 2yw \\
2xy - 2zw & 1 - 2x^2 - 2z^2 & 2yz + 2xw \\
2xz + 2yw & 2yz - 2xw & 1 - 2x^2 - 2y^2
\end{pmatrix}
\]

Here, \([x, y, z, w]\) corresponds to parameters in the heading JSON object, \([qx, qy, qz, qw]\). Ground Truth computes the translation column vector as \(T = [\text{poseX, poseY, poseZ}]\). Then the extrinsic metrics is simply as follows:

\[
\text{LiDAR\_extrinsic} = [R \; T; 0 \; 0 \; 1]
\]

Camera Calibrations: Extrinsic, Intrinsic and Distortion

Geometric camera calibration, also referred to as camera resectioning, estimates the parameters of a lens and image sensor of an image or video camera. You can use these parameters to correct for lens distortion, measure the size of an object in world units, or determine the location of the camera in the scene. Camera parameters include intrinsics and distortion coefficients.

Camera Extrinsic

If the camera pose is given, then Ground Truth computes the camera extrinsic based on a rigid transformation from the 3D plane into the camera plane. The calculation is the same as the one
used for the LiDAR Extrinsic, except that Ground Truth uses camera pose (position and heading) and computes the inverse extrinsic.

\[
camera_{\text{inverse\_extrinsic}} = \text{inv}([Rc \ Tc; 0 \ 0 \ 0 \ 1]) \quad \# \text{where } Rc \text{ and } Tc \text{ are camera pose components}
\]

**Intrinsic and Distortion**

Some cameras, such as pinhole or fisheye cameras, may introduce significant distortion in photos. This distortion can be corrected using distortion coefficients and the camera focal length. To learn more, see [Camera calibration With OpenCV](#) in the OpenCV documentation.

There are two types of distortion Ground Truth can correct for: radial distortion and tangential distortion.

*Radial distortion* occurs when light rays bend more near the edges of a lens than they do at its optical center. The smaller the lens, the greater the distortion. The presence of the radial distortion manifests in form of the *barrel* or *fish-eye* effect and Ground Truth uses Formula 1 to undistort it.

**Formula 1:**

\[
x_{\text{corrected}} = x(1 + k_1 r^2 + k_2 r^4 + k_3 r^6) \\
y_{\text{corrected}} = y(1 + k_1 r^2 + k_2 r^4 + k_3 r^6)
\]

*Tangential distortion* occurs because the lenses used to take the images are not perfectly parallel to the imaging plane. This can be corrected with Formula 2.

**Formula 2:**

\[
x_{\text{corrected}} = x + [2p_1 xy + p_2 (r^2 + 2x^2)] \\
y_{\text{corrected}} = y + [p_1 (r^2 + 2y^2) + 2p_2 xy]
\]

In the input manifest file, you can provide distortion coefficients and Ground Truth will undistort your images. All distortion coefficients are floats.
• k1, k2, k3, k4 – Radial distortion coefficients. Supported for both fisheye and pinhole camera models.

• p1, p2 – Tangential distortion coefficients. Supported for pinhole camera models.

If images are already undistorted, all distortion coefficients should be 0 in your input manifest.

In order to correctly reconstruct the corrected image, Ground Truth does a unit conversion of the images based on focal lengths. If a common focal length is used with a given aspect ratio for both axes, such as 1, in the upper formula we will have a single focal length. The matrix containing these four parameters is referred to as the *in camera intrinsic calibration matrix*.

\[
\begin{bmatrix}
  x \\
  y \\
  w
\end{bmatrix} =
\begin{bmatrix}
  f_x & 0 & c_x \\
  0 & f_y & c_y \\
  0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
  X \\
  Y \\
  Z
\end{bmatrix}
\]

While the distortion coefficients are the same regardless of the camera resolutions used, these should be scaled with the current resolution from the calibrated resolution.

The following are float values.

• f_x – focal length in x direction.
• f_y – focal length in y direction.
• c_x – x coordinate of principal point.
• c_y – y coordinate of principal point.

Ground Truth use the camera extrinsic and camera intrinsic to compute view metrics as shown in the following code block to transform labels between the 3D scene and 2D images.

```python
def generate_view_matrix(intrinsic_matrix, extrinsic_matrix):
    intrinsic_matrix = np.c_[intrinsic_matrix, np.zeros(3)]
```
When you create a video frame object detection or object tracking labeling job, you can choose video files (MP4 files) or video frames for input data. All worker tasks are created using video frames, so if you choose video files, use the Ground Truth frame extraction tool to extract video frames (images) from your video files.

For both of these options, you can use the **Automated data setup** option in the Ground Truth section of the Amazon SageMaker console to set up a connection between Ground Truth and your input data in Amazon S3 so that Ground Truth knows where to look for your input data when creating your labeling tasks. This creates and stores an input manifest file in your Amazon S3 input dataset location. To learn more, see [Automated Video Frame Input Data Setup](#).

Alternatively, you can manually create sequence files for each sequence of video frames that you want labeled and provide the Amazon S3 location of an input manifest file that references each of these sequences files using the `source-ref` key. To learn more, see [Create a Video Frame Input Manifest File](#).

**Topics**

- [Choose Video Files or Video Frames for Input Data](#)
- [Input Data Setup](#)

**Choose Video Files or Video Frames for Input Data**

When you create a video frame object detection or object tracking labeling job, you can provide a sequence of video frames (images) or you can use the Amazon SageMaker console to have Ground Truth automatically extract video frames from your video files. Use the following sections to learn more about these options.

**Provide Video Frames**

Video frames are sequences of images extracted from a video file. You can create a Ground Truth labeling job to have workers label multiple sequences of video frames. Each sequence is made up of images extracted from a single video.
To create a labeling job using video frame sequences, you must store each sequence using a unique key name prefix in Amazon S3. In the Amazon S3 console, key name prefixes are folders. So in the Amazon S3 console, each sequence of video frames must be located in its own folder in Amazon S3.

For example, if you have two sequences of video frames, you might use the key name prefixes sequence1/ and sequence2/ to identify your sequences. In this example, your sequences may be located in s3://DOC-EXAMPLE-BUCKET/video-frames/sequence1/ and s3://DOC-EXAMPLE-BUCKET/video-frames/sequence2/.

If you are using the Ground Truth console to create an input manifest file, all of the sequence key name prefixes should be in the same location in Amazon S3. For example, in the Amazon S3 console, each sequence could be in a folder in s3://DOC-EXAMPLE-BUCKET/video-frames/. In this example, your first sequence of video frames (images) may be located in s3://DOC-EXAMPLE-BUCKET/video-frames/sequence1/ and your second sequence may be located in s3://DOC-EXAMPLE-BUCKET/video-frames/sequence2/.

**Important**

Even if you only have a single sequence of video frames that you want workers to label, that sequence must have a key name prefix in Amazon S3. If you are using the Amazon S3 console, this means that your sequence is located in a folder. It cannot be located in the root of your S3 bucket.

When creating worker tasks using sequences of video frames, Ground Truth uses one sequence per task. In each task, Ground Truth orders your video frames using UTF-8 binary order.

For example, video frames might be in the following order in Amazon S3:

```
[0001.jpg, 0002.jpg, 0003.jpg, ..., 0011.jpg]
```

They are arranged in the same order in the worker’s task: 0001.jpg, 0002.jpg, 0003.jpg, ..., 0011.jpg.

Frames might also be ordered using a naming convention like the following:

```
[frame1.jpg, frame2.jpg, ..., frame11.jpg]
```
In this case, frame10.jpg and frame11.jpg come before frame2.jpg in the worker task. Your worker sees your video frames in the following order: frame1.jpg, frame10.jpg, frame11.jpg, frame2.jpg, ..., frame9.jpg.

**Provide Video Files**

You can use the Ground Truth frame splitting feature when creating a new labeling job in the console to extract video frames from video files (MP4 files). A series of video frames extracted from a single video file is referred to as a *sequence of video frames*.

You can either have Ground Truth automatically extract all frames, up to 2,000, from the video, or you can specify a frequency for frame extraction. For example, you can have Ground Truth extract every 10\(^{th}\) frame from your videos.

You can provide up to 50 videos when you use automated data setup to extract frames, however your input manifest file cannot reference more than 10 video frame sequence files when you create a video frame object tracking and video frame object detection labeling job. If you use the automated data setup console tool to extract video frames from more than 10 video files, you will need to modify the manifest file the tool generates or create a new one to include 10 video frame sequence files or less. To learn more about these quotas, see [3D Point Cloud and Video Frame Labeling Job Quotas](#).

To use the video frame extraction tool, see [Automated Video Frame Input Data Setup](#).

When all of your video frames have been successfully extracted from your videos, you will see the following in your S3 input dataset location:

- A key name prefix (a folder in the Amazon S3 console) named after each video. Each of these prefixes leads to:
  - A sequence of video frames extracted from the video used to name that prefix.
  - A sequence file used to identify all of the images that make up that sequence.
- An input manifest file with a .manifest extension. This identifies all of the sequence files that will be used to create your labeling job.

All of the frames extracted from a single video file are used for a labeling task. If you extract video frames from multiple video files, multiple tasks are created for your labeling job, one for each sequence of video frames.
Ground Truth stores each sequence of video frames that it extracts in your Amazon S3 location for input datasets using a unique key name prefix. In the Amazon S3 console, key name prefixes are folders.

**Input Data Setup**

When you create a video frame labeling job, you need to let Ground Truth know where to look for your input data. You can do this in one of two ways:

- You can store your input data in Amazon S3 and have Ground Truth automatically detect the input dataset used for your labeling job. See Automated Video Frame Input Data Setup to learn more about this option.

- You can create an input manifest file and sequence files and upload them to Amazon S3. See Manual Input Data Setup to learn more about this option.

**Topics**

- Automated Video Frame Input Data Setup
- Manual Input Data Setup

**Automated Video Frame Input Data Setup**

You can use the Ground Truth automated data setup to automatically detect video files in your Amazon S3 bucket and extract video frames from those files. To learn how, see Provide Video Files.

If you already have video frames in Amazon S3, you can use the automated data setup to use these video frames in your labeling job. For this option, all video frames from a single video must be stored using a unique prefix. To learn about the requirements to use this option, see Provide Video Frames.

Select one of the following sections to learn how to set up your automatic input dataset connection with Ground Truth.

**Provide Video Files and Extract Frames**

Use the following procedure to connect your video files with Ground Truth and automatically extract video frames from those files for video frame object detection and object tracking labeling jobs.
Note

If you use the automated data setup console tool to extract video frames from more than 10 video files, you will need to modify the manifest file the tool generates or create a new one to include 10 video frame sequence files or less. To learn more, see Provide Video Files.

Make sure your video files are stored in an Amazon S3 bucket in the same AWS Region that you perform the automated data setup in.

Automatically connect your video files in Amazon S3 with Ground Truth and extract video frames:


   Your input and output S3 buckets must be located in the same AWS Region that you create your labeling job in. This link puts you in the North Virginia (us-east-1) AWS Region. If your input data is in an Amazon S3 bucket in another Region, switch to that Region. To change your AWS Region, on the navigation bar, choose the name of the currently displayed Region.

2. Select Create labeling job.

3. Enter a Job name.

4. In the section Input data setup, select Automated data setup.

5. Enter an Amazon S3 URI for S3 location for input datasets. An S3 URI looks like the following: s3://DOC-EXAMPLE-BUCKET/path-to-files/. This URI should point to the Amazon S3 location where your video files are stored.

6. Specify your S3 location for output datasets. This is where your output data is stored. You can choose to store your output data in the Same location as input dataset or Specify a new location and entering the S3 URI of the location that you want to store your output data.

7. Choose Video Files for your Data type using the dropdown list.

8. Choose Yes, extract frames for object tracking and detection tasks.


   - When you choose Use all frames extracted from the video to create a labeling task, Ground Truth extracts all frames from each video in your S3 location for input datasets, up
to 2,000 frames. If a video in your input dataset contains more than 2,000 frames, the first 2,000 are extracted and used for that labeling task.

- When you choose **Use every x frame from a video to create a labeling task**, Ground Truth extracts every \( x \) frame from each video in your S3 location for input datasets.

For example, if your video is 2 seconds long, and has a **frame rate** of 30 frames per second, there are 60 frames in your video. If you specify 10 here, Ground Truth extracts every 10\(^{th}\) frame from your video. This means the 1\(^{st}\), 10\(^{th}\), 20\(^{th}\), 30\(^{th}\), 40\(^{th}\), 50\(^{th}\), and 60\(^{th}\) frames are extracted.

10. Choose or create an IAM execution role. Make sure that this role has permission to access your Amazon S3 locations for input and output data specified in steps 5 and 6.

11. Select **Complete data setup**.

**Provide Video Frames**

Use the following procedure to connect your sequences of video frames with Ground Truth for video frame object detection and object tracking labeling jobs.

Make sure your video frames are stored in an Amazon S3 bucket in the same AWS Region that you perform the automated data setup in. Each sequence of video frames should have a unique prefix. For example, if you have two sequences stored in s3://DOC-EXAMPLE-BUCKET/video-frames/sequences/, each should have a unique prefix like sequence1 and sequence2 and should both be located directly under the /sequences/ prefix. In the example above, the locations of these two sequences is: s3://DOC-EXAMPLE-BUCKET/video-frames/sequences/sequence1/ and s3://DOC-EXAMPLE-BUCKET/video-frames/sequences/sequence2/.

**Automatically connect your video frame in Amazon S3 with Ground Truth:**

1. Navigate to the **Create labeling job** page in the Amazon SageMaker console: [https://console.aws.amazon.com/sagemaker/groundtruth](https://console.aws.amazon.com/sagemaker/groundtruth).

   Your input and output S3 buckets must be located in the same AWS Region that you create your labeling job in. This link puts you in the North Virginia (us-east-1) AWS Region. If your input data is in an Amazon S3 bucket in another Region, switch to that Region. To change your AWS Region, on the navigation bar, choose the name of the currently displayed Region.

2. Select **Create labeling job**.

3. Enter a **Job name**.
4. In the section **Input data setup**, select **Automated data setup**.

5. Enter an Amazon S3 URI for **S3 location for input datasets**.
   
   This should be the Amazon S3 location where your sequences are stored. For example, if you have two sequences stored in s3://DOC-EXAMPLE-BUCKET/video-frames/sequences/sequence1/, s3://DOC-EXAMPLE-BUCKET/video-frames/sequences/sequence2/, enter s3://DOC-EXAMPLE-BUCKET/video-frames/sequences/ here.

6. Specify your **S3 location for output datasets**. This is where your output data is stored. You can choose to store your output data in the **Same location as input dataset** or **Specify a new location** and entering the S3 URI of the location that you want to store your output data.

7. Choose **Video frames** for your **Data type** using the dropdown list.

8. Choose or create an IAM execution role. Make sure that this role has permission to access your Amazon S3 locations for input and output data specified in steps 5 and 6.

9. Select **Complete data setup**.

These procedures will create an input manifest in the Amazon S3 location for input datasets that you specified in step 5. If you are creating a labeling job using the SageMaker API or, AWS CLI, or an AWS SDK, use the Amazon S3 URI for this input manifest file as input to the parameter ManifestS3Uri.

**Manual Input Data Setup**

Choose the manual data setup option if you have created sequence files for each of your video frame sequences, and a manifest file listing references to those sequences files.

**Create a Video Frame Input Manifest File**

Ground Truth uses the input manifest file to identify the location of your input dataset when creating labeling tasks. For video frame object detection and object tracking labeling jobs, each line in the input manifest file identifies the location of a video frame sequence file. Each sequence file identifies the images included in a single sequence of video frames.

Use this page to learn how to create a video frame sequence file and an input manifest file for video frame object tracking and object detection labeling jobs.

If you want Ground Truth to automatically generate your sequence files and input manifest file, see [Automated Video Frame Input Data Setup](use-input-and-output-data-1698).
Create a Video Frame Sequence Input Manifest

In the video frame sequence input manifest file, each line in the manifest is a JSON object, with a "source-ref" key that references a sequence file. Each sequence file identifies the location of a sequence of video frames. This is the manifest file formatting required for all video frame labeling jobs.

The following example demonstrates the syntax used for an input manifest file:

```json
{"source-ref": "s3://DOC-EXAMPLE-BUCKET/example-folder/seq1.json"}
{"source-ref": "s3://DOC-EXAMPLE-BUCKET/example-folder/seq2.json"}
```

Create a Video Frame Sequence File

The data for each sequence of video frames needs to be stored in a JSON data object. The following is an example of the format you use for a sequence file. Information about each frame is included as a JSON object and is listed in the frames list. The following JSON has been expanded for readability.

```json
{
    "seq-no": 1,
    "prefix": "s3://mybucket/prefix/video1/",
    "number-of-frames": 3,
    "frames": [
        {"frame-no": 1, "unix-timestamp": 1566861644, "frame": "frame0001.jpg" },
        {"frame-no": 2, "unix-timestamp": 1566861644, "frame": "frame0002.jpg" },
        {"frame-no": 3, "unix-timestamp": 1566861644, "frame": "frame0003.jpg" }
    ]
}
```

The following table provides details about the parameters shown in the this code example.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Required</th>
<th>Accepted Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>seq-no</td>
<td>Yes</td>
<td>Integer</td>
<td>The ordered number of the sequence.</td>
</tr>
<tr>
<td>prefix</td>
<td>Yes</td>
<td>String</td>
<td>The Amazon S3 location where the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accepted Values</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Required</td>
<td>Accepted Values</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>----------</td>
<td>-----------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>s3://&lt;bucket-name&gt; /&lt;prefix&gt;/</td>
<td></td>
<td></td>
<td>sequence files are located. The prefix must end with a forward slash: /.</td>
</tr>
<tr>
<td>number-of-frames</td>
<td>Yes</td>
<td>Integer</td>
<td>The total number of frames included in the sequence file. This number must match the total number of frames listed in the frames parameter in the next row.</td>
</tr>
</tbody>
</table>
| frames                 | Yes      | List of JSON objects  
Required: frame-no, frame  
Optional: unix-timestamp | A list of frame data. The length of the list must equal number-of-frames. In the worker UI, frames in a sequence are ordered in UTF-8 binary order. To learn more about this ordering, see Provide Video Frames. |
| frame-no               | Yes      | Integer         | The frame order number. This will determine the order of a frame in the sequence.                                                                                                                                 |
### Output Data

The output from a labeling job is placed in the Amazon S3 location that you specified in the console or in the call to the `CreateLabelingJob` operation. Output data appears in this location when the workers have submitted one or more tasks, or when tasks expire. Note that it may take a few minutes for output data to appear in Amazon S3 after the worker submits the task or the task expires.

Each line in the output data file is identical to the manifest file with the addition of an attribute and value for the label assigned to the input object. The attribute name for the value is defined in the console or in the call to the `CreateLabelingJob` operation. You can't use `-metadata` in the label attribute name. If you are running an image semantic segmentation, 3D point cloud semantic segmentation, or 3D point cloud object tracking job, the label attribute must end with `-ref`. For any other type of job, the attribute name can't end with `-ref`.

The output of the labeling job is the value of the key-value pair with the label. The label and the value overwrites any existing JSON data in the input file with the new value.

For example, the following is the output from an image classification labeling job where the input data files were stored in an Amazon S3 `AWSDOC-EXAMPLE-BUCKET` and the label attribute name was defined as `sport`. In this example the JSON object is formatted for readability, in the actual output file the JSON object is on a single line. For more information about the data format, see JSON Lines.

```json
{
}
```
The value of the label can be any valid JSON. In this case the label's value is the index of the class in the classification list. Other job types, such as bounding box, have more complex values.

Any key-value pair in the input manifest file other than the label attribute is unchanged in the output file. You can use this to pass data to your application.

The output from a labeling job can be used as the input to another labeling job. You can use this when you are chaining together labeling jobs. For example, you can send one labeling job to determine the sport that is being played. Then you send another using the same data to determine if the sport is being played indoors or outdoors. By using the output data from the first job as the manifest for the second job, you can consolidate the results of the two jobs into one output file for easier processing by your applications.

The output data file is written to the output location periodically while the job is in progress. These intermediate files contain one line for each line in the manifest file. If an object is labeled, the label is included. If the object hasn't been labeled, it is written to the intermediate output file identically to the manifest file.

**Output Directories**

Ground Truth creates several directories in your Amazon S3 output path. These directories contain the results of your labeling job and other artifacts of the job. The top-level directory for a labeling job is given the same name as your labeling job; the output directories are placed beneath it. For example, if you named your labeling job `find-people`, your output would be in the following directories:

```
s3://AWSDOC-EXAMPLE-BUCKET/find-people/activelearning
```
Each directory contains the following output:

**Active Learning Directory**

The `activelearning` directory is only present when you are using automated data labeling. It contains the input and output validation set for automated data labeling, and the input and output folder for automatically labeled data.

**Annotations Directory**

The `annotations` directory contains all of the annotations made by the workforce. These are the responses from individual workers that have not been consolidated into a single label for the data object.

There are three subdirectories in the `annotations` directory.

- The first, `worker-response`, contains the responses from individual workers. This contains a subdirectory for each iteration, which in turn contains a subdirectory for each data object in that iteration. The worker response data for each data object is stored in a timestamped JSON file that contains the answers submitted by each worker for that data object, and if you use a private workforce, metadata about those workers. To learn more about this metadata, see [Worker Metadata](#).

- The second, `consolidated-annotation`, contains information required to consolidate the annotations in the current batch into labels for your data objects.

- The third, `intermediate`, contains the output manifest for the current batch with any completed labels. This file is updated as the label for each data object is completed.

**Note**

We recommend that you do not use files that are not mentioned in the documentation.
Inference Directory

The inference directory is only present when you are using automated data labeling. This directory contains the input and output files for the SageMaker batch transform used while labeling data objects.

Manifest Directory

The manifest directory contains the output manifest from your labeling job. There is one subdirectory in the manifest directory, output. The output directory contains the output manifest file for your labeling job. The file is named output.manifest.

Training Directory

The training directory is only present when you are using automated data labeling. This directory contains the input and output files used to train the automated data labeling model.

Confidence Score

When you have more than one worker annotate a single task, your label results from annotation consolidation. Ground Truth calculates a confidence score for each label. A confidence score is a number between 0 and 1 that indicates how confident Ground Truth is in the label. You can use the confidence score to compare labeled data objects to each other, and to identify the least or most confident labels.

You should not interpret the value of a confidence score as an absolute value, or compare confidence scores across labeling jobs. For example, if all of the confidence scores are between 0.98 and 0.998, you should only compare the data objects with each other and not rely on the high confidence scores.

You should not compare the confidence scores of human-labeled data objects and auto-labeled data objects. The confidence scores for humans are calculated using the annotation consolidation function for the task, while the confidence scores for automated labeling are calculated using a model that incorporates object features. The two models generally have different scales and average confidence.

For a bounding box labeling job, Ground Truth calculates a confidence score per box. You can compare confidence scores within one image or across images for the same labeling type (human or auto). You can't compare confidence scores across labeling jobs.
If a single worker annotates a task (NumberOfHumanWorkersPerDataObject is set to 1 or in the console, you enter 1 for **Number of workers per dataset object**), the confidence score is set to 0.00.

**Worker Metadata**

Ground Truth provides information that you can use to track individual workers in task output data. The following data is located in the directories under the worker-response located in the **Annotations Directory**:

- The acceptanceTime is the time that the worker accepted the task. The format of this date and time stamp is `YYYY-MM-DDTHH:MM:SS.mmmZ` for the year (`YYYY`), month (`MM`), day (`DD`), hour (`HH`), minute (`MM`), second (`SS`) and millisecond (`mmm`). The date and time are separated by a **T**.

- The submissionTime is the time that the worker submitted their annotations using the **Submit** button. The format of this date and time stamp is `YYYY-MM-DDTHH:MM:SS.mmmZ` for the year (`YYYY`), month (`MM`), day (`DD`), hour (`HH`), minute (`MM`), second (`SS`) and millisecond (`mmm`). The date and time are separated by a **T**.

- timeSpentInSeconds reports the total time, in seconds, that a worker actively worked on that task. This metric does not include time when a worker paused or took a break.

- The workerId is unique to each worker.

- If you use a **private workforce**, in workerMetadata, you see the following.
  - The identityProviderType is the service used to manage the private workforce.
  - The issuer is the Cognito user pool or OIDC Identity Provider (IdP) issuer associated with the work team assigned to this human review task.
  - A unique sub identifier refers to the worker. If you create a workforce using Amazon Cognito, you can retrieve details about this worker (such as the name or user name) using this ID using Amazon Cognito. To learn how, see **Managing and Searching for User Accounts** in Amazon Cognito Developer Guide.

The following is an example of the output you may see if you use Amazon Cognito to create a private workforce. This is identified in the identityProviderType.

```json
"submissionTime": "2020-12-28T18:59:58.321Z",
"acceptanceTime": "2020-12-28T18:59:15.191Z",
"timeSpentInSeconds": 40.543,
"workerId": "a12b3cdefg4h5i67",
"workerMetadata": {
```
The following is an example of the workerMetadata you may see if you use your own OIDC IdP to create a private workforce:

```json
"workerMetadata": {
    "identityData": {
        "identityProviderType": "Oidc",
        "issuer": "https://example-oidc-ipd.com/adfs",
        "sub": "aaaaaaaa-bbbb-cccc-dddd-eeeeeeeeeeee"
    }
}
```

To learn more about using private workforces, see Use a Private Workforce.

**Output Metadata**

The output from each job contains metadata about the label assigned to data objects. These elements are the same for all jobs with minor variations. The following example shows the metadata elements:

```json
"confidence": 0.00,
"type": "groundtruth/image-classification",
"job-name": "identify-animal-species",
"human-annotated": "yes",
"creation-date": "2020-10-18T22:18:13.527256"
```

The elements have the following meaning:

- **confidence** – The confidence that Ground Truth has that the label is correct. For more information, see Confidence Score.
- **type** – The type of classification job. For job types, see Built-in Task Types.
- **job-name** – The name assigned to the job when it was created.
- **human-annotated** – Whether the data object was labeled by a human or by automated data labeling. For more information, see Automate Data Labeling.
• creation-date – The date and time that the label was created.

**Classification Job Output**

The following are sample outputs (output manifest files) from an image classification job and a text classification job. They include the label that Ground Truth assigned to the data object, the value for the label, and metadata that describes the label.

In addition to the standard metadata elements, the metadata for a classification job includes the text value of the label's class. For more information, see [Image Classification - MXNet](#).

The red, italicized text in the examples below depends on labeling job specifications and output data.

```
{
    "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/example_image.jpg",
    "species": "0",
    "species-metadata":
    {
        "class-name": "dog",
        "confidence": 0.00,
        "type": "groundtruth/image-classification",
        "job-name": "identify-animal-species",
        "human-annotated": "yes",
        "creation-date": "2018-10-18T22:18:13.527256"
    }
}

{
    "source": "The food was delicious",
    "mood": "1",
    "mood-metadata":
    {
        "class-name": "positive",
        "confidence": 0.8,
        "type": "groundtruth/text-classification",
        "job-name": "label-sentiment",
        "human-annotated": "yes",
        "creation-date": "2020-10-18T22:18:13.527256"
    }
}
```
Multi-label Classification Job Output

The following are example output manifest files from a multi-label image classification job and a multi-label text classification job. They include the labels that Ground Truth assigned to the data object (for example, the image or piece of text) and metadata that describes the labels the worker saw when completing the labeling task.

The label attribute name parameter (for example, `image-label-attribute-name`) contains an array of all of the labels selected by at least one of the workers who completed this task. This array contains integer keys (for example, `[1, 0, 8]`) that correspond to the labels found in `class-map`. In the multi-label image classification example, bicycle, person, and clothing were selected by at least one of the workers who completed the labeling task for the image, `exampleimage.jpg`.

The confidence-map shows the confidence score that Ground Truth assigned to each label selected by a worker. To learn more about Ground Truth confidence scores, see [Confidence Score](#).

The red, italicized text in the examples below depends on labeling job specifications and output data.

The following is an example of a multi-label image classification output manifest file.

```json
{
    "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/example_image.jpg",
    "image-label-attribute-name":[1,0,8],
    "image-label-attribute-name-metadata":
    {
        "job-name":"labeling-job/image-label-attribute-name",
        "class-map":
        {
            "1":"bicycle","0":"person","8":"clothing"
        },
        "human-annotated":"yes",
        "creation-date":"2020-02-27T21:36:25.000201",
        "confidence-map":
        {
            "1":0.95,"0":0.77,"8":0.2
        },
        "type":"groundtruth/image-classification-multilabel"
    }
}
```
The following is an example of a multi-label text classification output manifest file. In this example, approving, sad and critical were selected by at least one of the workers who completed the labeling task for the object exampletext.txt found in AWSDOC-EXAMPLE-BUCKET.

```json
{
    "source-ref": "AWSDOC-EXAMPLE-BUCKET/text_file.txt",
    "text-label-attribute-name":[1,0,4],
    "text-label-attribute-name-metadata":
    {
        "job-name":"labeling-job/text-label-attribute-name",
        "class-map":
        {
            "1":"approving","0":"sad","4":"critical"
        },
        "human-annotated":"yes",
        "creation-date":"2020-02-20T21:36:25.000201",
        "confidence-map":
        {
            "1":0.95,"0":0.77,"4":0.2
        },
        "type":"groundtruth/text-classification-multilabel"
    }
}
```

Bounding Box Job Output

The following is sample output (output manifest file) from a bounding box job. For this task, three bounding boxes are returned. The label value contains information about the size of the image, and the location of the bounding boxes.

The class_id element is the index of the box's class in the list of available classes for the task. The class-map metadata element contains the text of the class.

The metadata has a separate confidence score for each bounding box. The metadata also includes the class-map element that maps the class_id to the text value of the class. For more information, see Object Detection - MXNet.

The red, italicized text in the examples below depends on labeling job specifications and output data.

```json
{
    "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/example_image.png",
    "class-map":
    {
        "1": Approving,"0": Sad,"4": Critical"
    },
    "human-annotated": true,
    "creation-date": "2020-02-20T21:36:25.000201",
    "confidence-map":
    {
        "1": 0.95,"0": 0.77,"4": 0.2
    },
    "type": "groundtruth/bounding-box"
}
```
The output of a bounding box adjustment job looks like the following JSON. Note that the original JSON is kept intact and two new jobs are listed, each with “adjust-” prepended to the original attribute's name.

```json
{
  "source-ref": "S3 bucket location",
  "bounding-box-attribute-name": {
    "image_size": [{ "width": 500, "height": 400, "depth": 3 }],
    "annotations": [
      { "class_id": 0, "left": 111, "top": 134, "width": 61, "height": 128 },
      { "class_id": 5, "left": 161, "top": 250, "width": 30, "height": 30 },
      { "class_id": 5, "left": 20, "top": 20, "width": 30, "height": 30 }
    ]
  },
  "bounding-box-attribute-name-metadata": {
    "objects": [{ "confidence": 0.8 }, { "confidence": 0.9 }, { "confidence": 0.9 }],
    "class-map": {
      "0": "dog",
      "5": "bone"
    },
    "type": "groundtruth/object-detection",
    "human-annotated": "yes",
    "creation-date": "2018-10-18T22:18:13.527256",
    "job-name": "identify-dogs-and-toys"
  }
}
```
Use Input and Output Data
In this output, the job's type doesn't change, but an adjustment-status field is added. This field has the value of adjusted or unadjusted. If multiple workers have reviewed the object and at least one adjusted the label, the status is adjusted.

**Named Entity Recognition**

The following is an example output manifest file from a named entity recognition (NER) labeling task. For this task, seven entities are returned.

In the output manifest, the JSON object, annotations, includes a list of the labels (label categories) that you provided.

Worker responses are in a list named entities. Each entity in this list is a JSON object that contains a label value that matches one in the labels list, an integer startOffset value for labeled span's starting Unicode offset, and an integer endOffset value for the ending Unicode offset.

The metadata has a separate confidence score for each entity. If a single worker labeled each data object, the confidence value for each entity will be zero.

The red, italicized text in the examples below depends on labeling job inputs and worker responses.

```json
{
  "source": "Amazon SageMaker is a cloud machine-learning platform that was launched in November 2017. SageMaker enables developers to create, train, and deploy machine-
```
learning (ML) models in the cloud. SageMaker also enables developers to deploy ML models on embedded systems and edge-devices",
"
"ner-labeling-job-attribute-name": {
    "annotations": {
        "labels": [
            {
                "label": "Date",
                "shortDisplayName": "dt"
            },
            {
                "label": "Verb",
                "shortDisplayName": "vb"
            },
            {
                "label": "Thing",
                "shortDisplayName": "tng"
            },
            {
                "label": "People",
                "shortDisplayName": "ppl"
            }
        ],
        "entities": [
            {
                "label": "Thing",
                "startOffset": 22,
                "endOffset": 53
            },
            {
                "label": "Thing",
                "startOffset": 269,
                "endOffset": 281
            },
            {
                "label": "Verb",
                "startOffset": 63,
                "endOffset": 71
            },
            {
                "label": "Verb",
                "startOffset": 228,
                "endOffset": 234
            },
            {
                "label": "Verb",
                "startOffset": 293,
                "endOffset": 301
            }
        ]
    }
}
Use Input and Output Data

```
"ner-labeling-job-attribute-name-metadata": {
  "job-name": "labeling-job/example-ner-labeling-job",
  "type": "groundtruth/text-span",
  "creation-date": "2020-10-29T00:40:39.398470",
  "human-annotated": "yes",
  "entities": [
    {
      "confidence": 0
    },
    {
      "confidence": 0
    },
    {
      "confidence": 0
    },
    {
      "confidence": 0
    },
    {
      "confidence": 0
    },
    {
      "confidence": 0
    }
  ]
}
```
Label Verification Job Output

The output (output manifest file) of a bounding box verification job looks different than the output of a bounding box annotation job. That's because the workers have a different type of task. They're not labeling objects, but evaluating the accuracy of prior labeling, making a judgment, and then providing that judgment and perhaps some comments.

If human workers are verifying or adjusting prior bounding box labels, the output of a verification job would look like the following JSON. The red, italicized text in the examples below depends on labeling job specifications and output data.

```
{
    "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/image_example.png",
    "bounding-box-attribute-name": {
        "image_size": [{ "width": 500, "height": 400, "depth": 3 }],
        "annotations": [
            { "class_id": 0, "left": 111, "top": 134,
              "width": 61, "height": 128},
            { "class_id": 5, "left": 161, "top": 250,
              "width": 30, "height": 30},
            { "class_id": 5, "left": 20, "top": 20,
              "width": 30, "height": 30}
        ]
    },
    "bounding-box-attribute-name-metadata": {
        "objects": [
            { "confidence": 0.8},
            { "confidence": 0.9},
            { "confidence": 0.9}
        ],
        "class-map": {
            "0": "dog",
            "5": "bone"
        }
    }
}
```
Although the type on the original bounding box output was `groundtruth/object-detection`, the new type is `groundtruth/label-verification`. Also note that the `worker-feedback` array provides worker comments. If the worker doesn't provide comments, the empty fields are excluded during consolidation.

**Semantic Segmentation Job Output**

The following is the output manifest file from a semantic segmentation labeling job. The value of the label for this job is a reference to a PNG file in an Amazon S3 bucket.

In addition to the standard elements, the metadata for the label includes a color map that defines which color is used to label the image, the class name associated with the color, and the confidence score for each color. For more information, see [Semantic Segmentation Algorithm](#).

The red, italicized text in the examples below depends on labeling job specifications and output data.

```json
{
  "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/example_city_image.png",
  "city-streets-ref": "S3 bucket location",
  "city-streets-ref-metadata": {
    "internal-color-map": {
```

Use Input and Output Data
Confidence is scored on a per-image basis. Confidence scores are the same across all classes within an image.

The output of a semantic segmentation adjustment job looks similar to the following JSON.
Use Input and Output Data
Video Frame Object Detection Output

The following is the output manifest file from a video frame object detection labeling job. The red, italicized text in the examples below depends on labeling job specifications and output data.

In addition to the standard elements, the metadata includes a class map that lists each class that has at least one label in the sequence. The metadata also includes job-name which is the name you assigned to the labeling job. For adjustment tasks, if one or more bounding boxes were modified, there is an adjustment-status parameter in the metadata for audit workflows that is set to adjusted.

```json
{
    "source-ref": "s3://DOC-EXAMPLE-BUCKET/example-path/input-manifest.json",
    "CarObjectDetection-ref": "s3://AWSDOC-EXAMPLE-BUCKET/output/labeling-job-name/
annotations/consolidated-annotation/output/0/SeqLabel.json",
    "CarObjectDetection-ref-metadata": {
        "class-map": {
            "0": "car",
            "1": "bus"
        },
        "job-name": "labeling-job/labeling-job-name",
        "human-annotated": "yes",
        "creation-date": "2021-09-29T05:50:35.566000",
        "type": "groundtruth/video-object-detection"
    }
}
```

Ground Truth creates one output sequence file for each sequence of video frames that was labeled. Each output sequence file contains the following:
- All annotations for all frames in a sequence in the detection-annotations list of JSON objects.
- For each frame that was annotated by a worker, the frame file name (frame), number (frame-no), a list of JSON objects containing annotations (annotations), and if applicable, frame-attributes. The name of this list is defined by the task type you use: polylines, polygons, keypoints, and for bounding boxes, annotations.

Each JSON object contains information about a single annotation and associated label. The following table outlines the parameters you'll see for each video frame task type.

<table>
<thead>
<tr>
<th>Task Type</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bounding Box</td>
<td>Box dimensions: height and width</td>
</tr>
<tr>
<td></td>
<td>Box top, left corner pixel location: top and left</td>
</tr>
<tr>
<td>Keypoint</td>
<td>Keypoint vertices: { &quot;x&quot;: int, &quot;y&quot;: int }</td>
</tr>
<tr>
<td>Polygon</td>
<td>A list of polygon vertices: vertices { &quot;x&quot;: int, &quot;y&quot;: int }</td>
</tr>
<tr>
<td></td>
<td>A polygon is a closed shape and so the first point will also represent the last point.</td>
</tr>
<tr>
<td>Polyline</td>
<td>A list of polyline vertices: vertices { &quot;x&quot;: int, &quot;y&quot;: int }</td>
</tr>
</tbody>
</table>

In addition to task type specific values, you will see the following in each JSON object:
- Values of any label-category-attributes that were specified for that label.
- The class-id of the box. Use the class-map in the output manifest file to see which label category this ID maps to.
The following is an example of a SeqLabel.json file from a bounding box video frame object detection labeling job. This file will be located under s3://your-output-bucket/output-prefix/annotations/consolidated-annotation/output/annotation-number/

```json
{
    "detection-annotations": [
        {
            "annotations": [
                {
                    "height": 41,
                    "width": 53,
                    "top": 152,
                    "left": 339,
                    "class-id": "1",
                    "label-category-attributes": {
                        "occluded": "no",
                        "size": "medium"
                    }
                },
                {
                    "height": 24,
                    "width": 37,
                    "top": 148,
                    "left": 183,
                    "class-id": "0",
                    "label-category-attributes": {
                        "occluded": "no"
                    }
                }
            ],
            "frame-no": 0,
            "frame": "frame_0000.jpeg",
            "frame-attributes": {"name": value, "name": value}
        },
        {
            "annotations": [
                {
                    "height": 41,
                    "width": 53,
                    "top": 152,
                    "left": 341,
                    "class-id": "0",
                    "label-category-attributes": {}
                }
            ]
        }
    ]
}
```
Video Frame Object Tracking Output

The following is the output manifest file from a video frame object tracking labeling job. The red, italicized text in the examples below depends on labeling job specifications and output data.

In addition to the standard elements, the metadata includes a class map that lists each class that has at least one label in the sequence of frames. The metadata also includes job-name which is the name you assigned to the labeling job. For adjustment tasks, if one or more bounding boxes were modified, there is an adjustment-status parameter in the metadata for audit workflows that is set to adjusted.

```json
{
    "source-ref": "s3://DOC-EXAMPLE-BUCKET/example-path/input-manifest.json",
    "CarObjectTracking-ref": "s3://AWSDOC-EXAMPLE-BUCKET/output/labeling-job-name/
annotations/consolidated-annotation/output/0/SeqLabel.json",
    "CarObjectTracking-ref-metadata": {
        "class-map": {
            "0": "car",
            "1": "bus"
        },
        "job-name": "labeling-job/labeling-job-name",
        "human-annotated": "yes",
        "creation-date": "2021-09-29T05:50:35.566000",
        "type": "groundtruth/video-object-tracking"
    }
}
```
Ground Truth creates one output sequence file for each sequence of video frames that was labeled. Each output sequence file contains the following:

- All annotations for all frames in a sequence in the `tracking-annotations` list of JSON objects.
- For each frame that was annotated by a worker, the frame (`frame`), number (`frame-no`), a list of JSON objects containing annotations (`annotations`), and if applicable, frame attributes (`frame-attributes`). The name of this list is defined by the task type you use: polylines, polygons, keypoints, and for bounding boxes, annotations.

Each JSON object contains information about a single annotation and associated label. The following table outlines the parameters you'll see for each video frame task type.

<table>
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</tr>
<tr>
<td></td>
<td>Box top, left corner pixel location: top and left</td>
</tr>
<tr>
<td>Keypoint</td>
<td>Keypoint vertices: <code>{ &quot;x&quot;: int, &quot;y&quot;: int }</code></td>
</tr>
<tr>
<td>Polygon</td>
<td>A list of polygon vertices: <code>vertices</code></td>
</tr>
<tr>
<td></td>
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<td>Polyline</td>
<td>A list of polyline vertices: <code>vertices</code></td>
</tr>
<tr>
<td></td>
<td>Polyline vertices: <code>{ &quot;x&quot;: int, &quot;y&quot;: int }</code></td>
</tr>
</tbody>
</table>

In addition to task type specific values, you will see the following in each JSON object:
- Values of any label-category-attributes that were specified for that label.
- The class-id of the box. Use the class-map in the output manifest file to see which label category this ID maps to.
- An object-id which identifies an instance of a label. This ID will be the same across frames if a worker identified the same instance of an object in multiple frames. For example, if a car appeared in multiple frames, all bounding boxes uses to identify that car would have the same object-id.
- The object-name which is the instance ID of that annotation.

The following is an example of a SeqLabel.json file from a bounding box video frame object tracking labeling job. This file will be located under s3://your-output-bucket/output-prefix/annotations/consolidated-annotation/output/annotation-number/

```json
{
    "tracking-annotations": [
        {
            "annotations": [
                {
                    "height": 36,
                    "width": 46,
                    "top": 178,
                    "left": 315,
                    "class-id": "0",
                    "label-category-attributes": {
                        "occluded": "no"
                    },
                    "object-id": "480dc450-c0ca-11ea-961f-a9b1c5c97972",
                    "object-name": "car:1"
                }
            ],
            "frame-no": 0,
            "frame": "frame_0001.jpeg",
            "frame-attributes": {}
        },
        {
            "annotations": [
                {
                    "height": 30,
                    "width": 47,
                    "top": 163,
                    "left": 315,
                    "class-id": "0",
                    "label-category-attributes": {
                        "occluded": "no"
                    },
                    "object-id": "480dc450-c0ca-11ea-961f-a9b1c5c97972",
                    "object-name": "car:1"
                }
            ],
            "frame-no": 1,
            "frame": "frame_0002.jpeg",
            "frame-attributes": {}
        }
    ]
}
```
3D Point Cloud Semantic Segmentation Output

The following is the output manifest file from a 3D point cloud semantic segmentation labeling job.

In addition to the standard elements, the metadata for the label includes a color map that defines which color is used to label the image, the class name associated with the color, and the confidence score for each color. Additionally, there is an adjustment-status parameter in the metadata for audit workflows that is set to adjusted if the color mask is modified. If you added one or more frameAttributes to your label category configuration file, worker responses for frame attributes are in the JSON object, dataset-object-attributes.
The **your-label-attribute-ref** parameter contains the location of a compressed file with a .zlib extension. When you uncompress this file, it contains an array. Each index in the array corresponds to the index of an annotated point in the input point cloud. The value of the array at a given index gives the class of the point at the same index in the point cloud, based on the semantic color map found in the **color-map** parameter of the metadata.

You can use Python code similar to the following to decompress a .zlib file:

```python
import zlib
from array import array

# read the label file
compressed_binary_file = open(zlib_file_path/file.zlib, 'rb').read()

# uncompress the label file
binary_content = zlib.decompress(compressed_binary_file)

# load labels to an array
my_int_array_data = array('B', binary_content);

print(my_int_array_data)
```

The code block above will produce an output similar to the following. Each element of the printed array contains the class of a point at that index in the point cloud. For example, `my_int_array_data[0] = 1` means point[0] in the input point cloud has a class 1. In the following output manifest file example, class 0 corresponds with "Background", 1 with Car, and 2 with Pedestrian.

```python
>> array('B', [1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 2, 2, 2, 2, 2, 2, 2, 2, 2, 2, 2])
```

The following is an example of a semantic segmentation 3D point cloud labeling job output manifest file. The red, italicized text in the examples below depends on labeling job specifications and output data.

```json
{
    "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/examplefolder/frame1.bin",
    "source-ref-metadata": {
        "format": "binary/xyz",
        "unix-timestamp": 1566861644.759115,
        "ego-vehicle-pose": {...}
    }
}
```
3D Point Cloud Object Detection Output

The following is sample output from a 3D point cloud objected detection job. For this task type, the data about 3D cuboids is returned in the 3d-bounding-box parameter, in a list named annotations. In this list, each 3D cuboid is described using the following information.
• Each class, or label category, that you specify in your input manifest is associated with a class-id. Use the class-map to identify the class associated with each class ID.

• These classes are used to give each 3D cuboid an object-name in the format `<class>:<integer>` where integer is a unique number to identify that cuboid in the frame.

• center-x, center-y, and center-z are the coordinates of the center of the cuboid, in the same coordinate system as the 3D point cloud input data used in your labeling job.

• length, width, and height describe the dimensions of the cuboid.

• yaw is used to describe the orientation (heading) of the cuboid in radians.

**Note**

yaw is now in the right-handed Cartesian system. Since this feature was added on September 02, 2022 19:02:17 UTC, you can convert the yaw measurement in the output data prior to that using the following (all units are in radians):

```
old_yaw_in_output = pi - yaw
```

• In our definition, +x is to the right, +y is to the forward, and +z is up from the ground plane. The rotation order is x - y - z. The roll, pitch and yaw are represented in the right-handed Cartesian system. In 3D space, roll is along the x-axis, pitch is along the y-axis and yaw is along the z-axis. All three are counterclockwise.

• If you included label attributes in your input manifest file for a given class, a label-category-attributes parameter is included for all cuboids for which workers selected label attributes.

If one or more cuboids were modified, there is an adjustment-status parameter in the metadata for audit workflows that is set to adjusted. If you added one or more frameAttributes to your label category configuration file, worker responses for frame attributes are in the JSON object, dataset-object-attributes.

The *red, italicized text* in the examples below depends on labeling job specifications and output data. The ellipses (...) denote a continuation of that list, where additional objects with the same format as the proceeding object can appear.

```json
{
    "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/examplefolder/frame1.txt",
    "source-ref-metadata":{
```
"format": "text/xyz",
"unix-timestamp": 1566861644.759115,
"prefix": "s3://AWSDOC-EXAMPLE-BUCKET/lidar_singleframe_dataset/prefix",
"ego-vehicle-pose": {
  "heading": {
    "qx": -0.02111296123795955,
    "qy": -0.006495469416730261,
    "qz": -0.008024565904865688,
    "qw": 0.9997181192298087
  },
  "position": {
    "x": -2.7161461413869947,
    "y": 116.25822288149078,
    "z": 1.8348751887989483
  }
},
"images": [
  {
    "fx": 847.7962624528487,
    "fy": 850.0340893791985,
    "cx": 576.2129134707038,
    "cy": 317.2423573573745,
    "k1": 0,
    "k2": 0,
    "k3": 0,
    "k4": 0,
    "p1": 0,
    "p2": 0,
    "skew": 0,
    "unix-timestamp": 1566861644.759115,
    "image-path": "images/frame_0_camera_0.jpg",
    "position": {
      "x": -2.2722515189268138,
      "y": 116.86003310568965,
      "z": 1.454614668542299
    },
    "heading": {
      "qx": 0.7594754093069037,
      "qy": 0.02181790885672969,
      "qz": -0.02461725233103356,
      "qw": -0.6496916273040025
    },
    "camera_model": "pinhole"
  }
]
"3d-bounding-box":
{
  "annotations": [
    {
      "label-category-attributes": {
        "Occlusion": "Partial",
        "Type": "Sedan"
      },
      "object-name": "Car:1",
      "class-id": 0,
      "center-x": -2.616382013657516,
      "center-y": 125.04149850484193,
      "center-z": 0.311272296465834,
      "length": 2.993000265181146,
      "width": 1.8355260519692056,
      "height": 1.3233490884304047,
      "roll": 0,
      "pitch": 0,
      "yaw": 1.6479308313703527
    },
    {
      "label-category-attributes": {
        "Occlusion": "Partial",
        "Type": "Sedan"
      },
      "object-name": "Car:2",
      "class-id": 0,
      "center-x": -5.188984560617168,
      "center-y": 99.7954483288783,
      "center-z": 0.2226435567445657,
      "length": 4,
      "width": 2,
      "height": 2,
      "roll": 0,
      "pitch": 0,
      "yaw": 1.6243170732068055
    }
  ]
},
"3d-bounding-box-metadata":
{
  "objects": []
}
"class_map":
{
   "0": "Car",
},
"type": "groundtruth/point_cloud_object_detection",
"human-annotated": "yes",
"creation-date": "2018-10-18T22:18:13.527256",
"job-name": "identify-3d-objects",
"adjustment-status": "adjusted",
"dataset-object-attributes": {name: value, name: value}
}

3D Point Cloud Object Tracking Output

The following is an example of an output manifest file from a 3D point cloud object tracking labeling job. The red, italicized text in the examples below depends on labeling job specifications and output data. The ellipses (...) denote a continuation of that list, where additional objects with the same format as the proceeding object can appear.

In addition to the standard elements, the metadata includes a class map that lists each class that has at least one label in the sequence. If one or more cuboids were modified, there is an adjustment-status parameter in the metadata for audit workflows that is set to adjusted.

{ 
   "source-ref": "s3://AWSDOC-EXAMPLE-BUCKET/myfolder/seq1.json",
   "lidar-label-attribute-ref": "s3://<CustomerOutputLocation>/<labelingJobName>/annotations/consolidated-annotation/output/<datasetObjectId>/SeqLabel.json",
   "lidar-label-attribute-ref-metadata": {
      "objects": [
         {
            "frame-no": 300,
            "confidence": []
         },
         {
            "frame-no": 301,
            "confidence": []
         },
         ...
      ],
      'class-map': {'0': 'Car', '1': 'Person'},
      'type': 'groundtruth/point_cloud_object_tracking',
   }
}
In the above example, the cuboid data for each frame in seq1.json is in SeqLabel.json in the Amazon S3 location, s3://<customerOutputLocation>/<labelingJobName>/annotations/consolidated-annotation/output/<datasetObjectId>/SeqLabel.json. The following is an example of this label sequence file.

For each frame in the sequence, you see the frame-number, frame-name, if applicable, frame-attributes, and a list of annotations. This list contains 3D cuboids that were drawn for that frame. Each annotation includes the following information:

- An object-name in the format <class>:<integer> where class identifies the label category and integer is a unique ID across the dataset.
- When workers draw a cuboid, it is associated with a unique object-id which is associated with all cuboids that identify the same object across multiple frames.
- Each class, or label category, that you specified in your input manifest is associated with a class-id. Use the class-map to identify the class associated with each class ID.
- center-x, center-y, and center-z are the coordinates of the center of the cuboid, in the same coordinate system as the 3D point cloud input data used in your labeling job.
- length, width, and height describe the dimensions of the cuboid.
- yaw is used to describe the orientation (heading) of the cuboid in radians.

**Note**

yaw is now in the right-handed Cartesian system. Since this feature was added on September 02, 2022 19:02:17 UTC, you can convert the yaw measurement in the output data prior to that using the following (all units are in radians):

```plaintext
old_yaw_in_output = pi - yaw
```

- In our definition, +x is to the right, +y is to the forward, and +z is up from the ground plane. The rotation order is x - y - z. The roll, pitch and yaw are represented in the right-handed
Cartesian system. In 3D space, roll is along the x-axis, pitch is along the y-axis and yaw is along the z-axis. All three are counterclockwise.

- If you included label attributes in your input manifest file for a given class, a label-category-attributes parameter is included for all cuboids for which workers selected label attributes.

```json
{
    "tracking-annotations": [
        {
            "frame-number": 0,
            "frame-name": "0.txt.pcd",
            "frame-attributes": {"name": value, "name": value},
            "annotations": [
                {
                    "label-category-attributes": {},
                    "object-name": "Car:4",
                    "class-id": 0,
                    "center-x": -2.2906369208300674,
                    "center-y": 103.73924823843463,
                    "center-z": 0.37634114027023313,
                    "length": 4,
                    "width": 2,
                    "height": 2,
                    "roll": 0,
                    "pitch": 0,
                    "yaw": 1.5827222214406014,
                    "object-id": "ae5dc770-a782-11ea-b57d-67c51a0561a1"
                },
                {
                    "label-category-attributes": {
                        "Occlusion": "Partial",
                        "Type": "Sedan"
                    },
                    "object-name": "Car:1",
                    "class-id": 0,
                    "center-x": -2.6451293634707413,
                    "center-y": 124.9534455706848,
                    "center-z": 0.5020834081743839,
                    "length": 4,
                    "width": 2,
                    "height": 2.080488827301309,
                    "roll": 0,
                    "pitch": 0,
```
"yaw": -1.5963335581398077,
"object-id": "06efb020-a782-11ea-b57d-67c51a0561a1"
],
{
"label-category-attributes": {
"Occlusion": "Partial",
"Type": "Sedan"
},
"object-name": "Car:2",
"class-id": 0,
"center-x": -5.205611313118477,
"center-y": 99.91731932137061,
"center-z": 0.22917217081212138,
"length": 3.8747142207671956,
"width": 1.9999999999999918,
"height": 2,
"roll": 0,
"pitch": 0,
"yaw": 1.5672228760316775,
"object-id": "26fad020-a782-11ea-b57d-67c51a0561a1"
}
],
{
"frame-number": 1,
"frame-name": "1.txt.pcd",
"frame-attributes": {},
"annotations": [
{
"label-category-attributes": {},
"object-name": "Car:4",
"class-id": 0,
"center-x": -2.2906369208300674,
"center-y": 103.73924823843463,
"center-z": 0.37634114027023313,
"length": 4,
"width": 2,
"height": 2,
"roll": 0,
"pitch": 0,
"yaw": 1.582722214406014,
"object-id": "ae5dc770-a782-11ea-b57d-67c51a0561a1"
}
]
"label-category-attributes": {
"Occlusion": "Partial",
"Type": "Sedan"
},
"object-name": "Car:1",
"class-id": 0,
"center-x": -2.6451293634707413,
"center-y": 124.9534455706848,
"center-z": 0.5020834081743839,
"length": 4,
"width": 2,
"height": 2.080488827301309,
"roll": 0,
"pitch": 0,
"yaw": -1.5963335581398077,
"object-id": "06efb020-a782-11ea-b57d-67c51a0561a1"
},
{
"label-category-attributes": {
"Occlusion": "Partial",
"Type": "Sedan"
},
"object-name": "Car:2",
"class-id": 0,
"center-x": -5.221311072916759,
"center-y": 100.4639841045424,
"center-z": 0.22917217081212138,
"length": 3.8747142207671956,
"width": 1.9999999999999918,
"height": 2,
"roll": 0,
"pitch": 0,
"yaw": 1.5672228760316775,
"object-id": "26fad020-a782-11ea-b57d-67c51a0561a1"
}
3D-2D Object Tracking Point Cloud Object Tracking Output

The following is an example of an output manifest file from a 3D point cloud object tracking labeling job. The *red, italicized text* in the examples below depends on labeling job specifications and output data. The ellipses (...) denote a continuation of that list, where additional objects with the same format as the proceeding object can appear.

In addition to the standard elements, the metadata includes a class map that lists each class that has at least one label in the sequence. If one or more cuboids were modified, there is an adjustment-status parameter in the metadata for audit workflows that is set to adjusted.

```json
{
    "source-ref": "s3://iad-groundtruth-lidar-test-bucket/artifacts/gt-point-cloud-demos/sequences/seq2.json",
    "source-ref-metadata": {
        "json-paths": [
            "number-of-frames",
            "prefix",
            "frames[frame-no, frame]"
        ]
    },
    "3D2D-linking-ref": "s3://iad-groundtruth-lidar-test-bucket/xyz/3D2D-linking/annotations/consolidated-annotation/output/0/SeqLabel.json",
    "3D2D-linking-ref-metadata": {
        "objects": [
            {
                "frame-no": 0,
                "confidence": []
            },
            {
                "frame-no": 1,
                "confidence": []
            },
            {
                "frame-no": 2,
                "confidence": []
            },
            {
                "frame-no": 3,
                "confidence": []
            },
            {
                "frame-no": 4,
                "confidence": []
            }
        ]
    }
}
```
"confidence": []
},
{
  "frame-no": 5,
  "confidence": []
},
{
  "frame-no": 6,
  "confidence": []
},
{
  "frame-no": 7,
  "confidence": []
},
{
  "frame-no": 8,
  "confidence": []
},
{
  "frame-no": 9,
  "confidence": []
}
],
"class-map": {
  "0": "Car"
},
"type": "groundtruth/point_cloud_object_tracking",
"human-annotated": "yes",
"creation-date": "2023-01-19T02:55:10.206508",
"job-name": "mcm-linking"
},
"3D2D-linking-chain-ref": "s3://iad-groundtruth-lidar-test-bucket/xyz/3D2D-linking-chain/annotations/consolidated-annotation/output/0/SeqLabel.json",
"3D2D-linking-chain-ref-metadata": {
  "objects": [
    {
      "frame-no": 0,
      "confidence": []
    },
    {
      "frame-no": 1,
      "confidence": []
    }
  ]
Use Input and Output Data
In the above example, the cuboid data for each frame in seq2.json is in SeqLabel.json in the Amazon S3 location, s3://<customerOutputLocation>/<labelingJobName>/annotations/consolidated-annotation/output/<datasetObjectId>/SeqLabel.json. The following is an example of this label sequence file.

For each frame in the sequence, you see the frame-number, frame-name, if applicable, frame-attributes, and a list of annotations. This list contains 3D cuboids that were drawn for that frame. Each annotation includes the following information:

• An object-name in the format <class>:<integer> where class identifies the label category and integer is a unique ID across the dataset.

• When workers draw a cuboid, it is associated with a unique object-id which is associated with all cuboids that identify the same object across multiple frames.

• Each class, or label category, that you specified in your input manifest is associated with a class-id. Use the class-map to identify the class associated with each class ID.

• center-x, center-y, and center-z are the coordinates of the center of the cuboid, in the same coordinate system as the 3D point cloud input data used in your labeling job.

• length, width, and height describe the dimensions of the cuboid.

• yaw is used to describe the orientation (heading) of the cuboid in radians.

Note

yaw is now in the right-handed Cartesian system. Since this feature was added on September 02, 2022 19:02:17 UTC, you can convert the yaw measurement in the output data prior to that using the following (all units are in radians):

\[ \text{old}_yaw\text{in}_output = \pi - yaw \]

• In our definition, +x is to the right, +y is to the forward, and +z is up from the ground plane. The rotation order is x - y - z. The roll, pitch and yaw are represented in the right-handed Cartesian system. In 3D space, roll is along the x-axis, pitch is along the y-axis and yaw is along the z-axis. All three are counterclockwise.

• If you included label attributes in your input manifest file for a given class, a label-category-attributes parameter is included for all cuboids for which workers selected label attributes.
"lidar": {
    "tracking-annotations": [
        {
            "frame-number": 0,
            "frame-name": "0.txt.pcd",
            "annotations": [
                {
                    "label-category-attributes": {
                        "Type": "Sedan"
                    },
                    "object-name": "Car:1",
                    "class-id": 0,
                    "center-x": 12.172361721602815,
                    "center-y": 120.23067521992364,
                    "center-z": 1.590525771183712,
                    "length": 4,
                    "width": 2,
                    "height": 2,
                    "roll": 0,
                    "pitch": 0,
                    "yaw": 0,
                    "object-id": "505b39e0-97a4-11ed-8903-dd5b8b903715"
                },
                {
                    "label-category-attributes": {},
                    "object-name": "Car:4",
                    "class-id": 0,
                    "center-x": 17.192725195301094,
                    "center-y": 114.55705365827872,
                    "center-z": 1.590525771183712,
                    "length": 4,
                    "width": 2,
                    "height": 2,
                    "roll": 0,
                    "pitch": 0,
                    "yaw": 0,
                    "object-id": "1afcb670-97a9-11ed-9a84-ff627d099e16"
                }
            ]
        }
    ],
    "frame-attributes": {}
}
"frame-number": 1,
"frame-name": "1.txt.pcd",
"annotations": [
    {
        "label-category-attributes": {
            "Type": "Sedan"
        },
        "object-name": "Car:1",
        "class-id": 0,
        "center-x": -1.6841480600695489,
        "center-y": 126.20198882749516,
        "center-z": 1.590525771183712,
        "length": 4,
        "width": 2,
        "height": 2,
        "roll": 0,
        "pitch": 0,
        "yaw": 0,
        "object-id": "505b39e0-97a4-11ed-8903-dd5b8b903715"
    },
    {
        "label-category-attributes": {},
        "object-name": "Car:4",
        "class-id": 0,
        "center-x": 17.192725195301094,
        "center-y": 114.55705365827872,
        "center-z": 1.590525771183712,
        "length": 4,
        "width": 2,
        "height": 2,
        "roll": 0,
        "pitch": 0,
        "yaw": 0,
        "object-id": "1afcb670-97a9-11ed-9a84-ff627d099e16"
    }
],
"frame-attributes": {}]
}
"Type": "Sedan",
"object-name": "Car:1",
"class-id": 0,
"center-x": -1.6841480600695489,
"center-y": 126.2019882749516,
"center-z": 1.590525771183712,
"length": 4,
"width": 2,
"height": 2,
"roll": 0,
"pitch": 0,
"yaw": 0,
"object-id": "505b39e0-97a4-11ed-8903-dd5b8b903715"
},
{

"label-category-attributes": {},
"object-name": "Car:4",
"class-id": 0,
"center-x": 17.192725195301094,
"center-y": 114.55705365827872,
"center-z": 1.590525771183712,
"length": 4,
"width": 2,
"height": 2,
"roll": 0,
"pitch": 0,
"yaw": 0,
"object-id": "1afcb670-97a9-11ed-9a84-ff627d099e16"
}
],
"frame-attributes": {}
}

"camera-0": {
"tracking-annotations": [
{

"frame-no": 0,
"frame": "0.txt.pcd",
"annotations": [
{

"label-category-attributes": {
"Occlusion": "Partial"
The cuboid and bounding box for an object are linked through a common object-id.

**Enhanced Data Labeling**

Amazon SageMaker Ground Truth manages sending your data objects to workers to be labeled. Labeling each data object is a task. Workers complete each task until the entire labeling job is complete. Ground Truth divides the total number of tasks into smaller batches that are sent to workers. A new batch is sent to workers when the previous one is finished.
Ground Truth provides two features that help improve the accuracy of your data labels and reduce the total cost of labeling your data:

- *Annotation consolidation* helps to improve the accuracy of your data object labels. It combines the results of multiple workers' annotation tasks into one high-fidelity label.

- *Automated data labeling* uses machine learning to label portions of your data automatically without having to send them to human workers.

Topics

- [Control the Flow of Data Objects Sent to Workers](#)
- [Consolidate Annotations](#)
- [Automate Data Labeling](#)
- [Chaining Labeling Jobs](#)

**Control the Flow of Data Objects Sent to Workers**

Depending on the type of labeling job you create, Amazon SageMaker Ground Truth sends data objects to workers in batches or in a streaming fashion. You can control the flow of data objects to workers in the following ways:

- For both types of labeling jobs, you can use `MaxConcurrentTaskCount` to control the total number of data objects available to all workers at a given point in time when the labeling job is running.

- For streaming labeling jobs, you can control the flow of data objects to workers by monitoring and controlling the number of data objects sent to the Amazon SQS associated with your labeling job.

Use the following sections to learn more about these options. To learn more about streaming labeling jobs, see [Ground Truth Streaming Labeling Jobs](#).

Topics

- [Use MaxConcurrentTaskCount to Control the Flow of Data Objects](#)
- [Use Amazon SQS to Control the Flow of Data Objects to Streaming Labeling Jobs](#)
Use MaxConcurrentTaskCount to Control the Flow of Data Objects

MaxConcurrentTaskCount defines the maximum number of data objects that can be labeled by human workers at the same time. If you use the console, this parameter is set to 1,000. If you use CreateLabelingJob, you can set this parameter to any integer between 1 and 1,000, inclusive.

When you start a labeling job using an input manifest file, Ground Truth does the following:

1. For each data object listed in your input manifest file, one or more tasks are created, depending on the value you specify for NumberOfHumanWorkersPerDataObject. For example, if you set the number of workers per data object to 3, 3 tasks will be created for each dataset object. To be marked as successfully labeled, at least one worker must label the object. Alternatively, the tasks can expire or be declined.

2. If you are using the Mechanical Turk workforce, Ground Truth first sends a batch of 10 dataset objects to your workers. It uses this small batch to set up the labeling job and to make sure that the job is correctly configured.

3. Next, Ground Truth sends MaxConcurrentTaskCount number of dataset objects to workers. For example, if you have 2,000 input data objects in your input manifest file and have set the number of workers per data object to 3 and set MaxConcurrentTaskCount to 900, the first 900 data objects in your input manifest are sent to workers, corresponding to 2,700 tasks (900 x 3). This is the first full-sized set of objects sent to workers.

4. What happens next depends on the type of labeling job you create. This step assumes one or more dataset objects in your input manifest file, or sent using an Amazon SNS input data source (in a streaming labeling job) were not include in the set sent to workers in step 3.

   - **Streaming labeling job:** As long as the total number of objects available to workers is equal to MaxConcurrentTaskCount, all remaining dataset objects on your input manifest file and that you send in real time using Amazon SNS are placed on an Amazon SQS queue. When the total number of objects available to workers falls below MaxConcurrentTaskCount minus NumberOfHumanWorkersPerDataObject, a new data object from the queue is used to create NumberOfHumanWorkersPerDataObject-tasks, which are sent to workers in real time.

   - **Non-streaming labeling job:** As workers finish labeling one set of objects, up to MaxConcurrentTaskCount times NumberOfHumanWorkersPerDataObject number of new tasks will be sent to workers. This process is repeated until all data objects in the input manifest file are labeled.
Use Amazon SQS to Control the Flow of Data Objects to Streaming Labeling Jobs

When you create a streaming labeling job, an Amazon SQS queue is automatically created in your account. Data objects are only added to the Amazon SQS queue when the total number of objects sent to workers is above `MaxConcurrentTaskCount`. Otherwise, objects are sent directly to workers.

You can use this queue to manage the flow of data objects to your labeling job. To learn more, see [Manage Labeling Requests with an Amazon SQS Queue](#).

**Consolidate Annotations**

An *annotation* is the result of a single worker's labeling task. *Annotation consolidation* combines the annotations of two or more workers into a single label for your data objects. A label, which is assigned to each object in the dataset, is a probabilistic estimate of what the true label should be. Each object in the dataset typically has multiple annotations, but only one label or set of labels.

You decide how many workers annotate each object in your dataset. Using more workers can increase the accuracy of your labels, but also increases the cost of labeling. To learn more about Ground Truth pricing, see [Amazon SageMaker Ground Truth pricing](#).

If you use the Amazon SageMaker console to create a labeling job, the following are the defaults for the number of workers who can annotate objects:

- Text classification—3 workers
- Image classification—3 workers
- Bounding boxes—5 workers
- Semantic segmentation—3 workers
- Named entity recognition—3 workers

When you use the `CreateLabelingJob` operation, you set the number of workers to annotate each data object with the `NumberOfHumanWorkersPerDataObject` parameter. You can override the default number of workers that annotate a data object using the console or the `CreateLabelingJob` operation.

Ground Truth provides an annotation consolidation function for each of its predefined labeling tasks: bounding box, image classification, name entity recognition, semantic segmentation, and text classification. These are the functions:
Multi-class annotation consolidation for image and text classification uses a variant of the Expectation Maximization approach to annotations. It estimates parameters for each worker and uses Bayesian inference to estimate the true class based on the class annotations from individual workers.

Bounding box annotation consolidates bounding boxes from multiple workers. This function finds the most similar boxes from different workers based on the Jaccard index, or intersection over union, of the boxes and averages them.

Semantic segmentation annotation consolidation treats each pixel in a single image as a multi-class classification. This function treats the pixel annotations from workers as "votes," with more information from surrounding pixels incorporated by applying a smoothing function to the image.

Named entity recognition clusters text selections by Jaccard similarity and calculates selection boundaries based on the mode, or the median if the mode isn't clear. The label resolves to the most assigned entity label in the cluster, breaking ties by random selection.

You can use other algorithms to consolidate annotations. For information, see Create Your Own Annotation Consolidation Function.

Create Your Own Annotation Consolidation Function

You can choose to use your own annotation consolidation function to determine the final labels for your labeled objects. There are many possible approaches for writing a function and the approach that you take depends on the nature of the annotations to consolidate. Broadly, consolidation functions look at the annotations from workers, measure the similarity between them, and then use some form of probabilistic judgment to determine what the most probable label should be.

If you want to use other algorithms to create annotation consolidations functions, you can find the worker responses in the [project-name]/annotations/worker-response folder of the Amazon S3 bucket where you direct the job output.

Assess Similarity

To assess the similarity between labels, you can use one of the following strategies, or you can use one that meets your data labeling needs:

- For label spaces that consist of discrete, mutually exclusive categories, such as multi-class classification, assessing similarity can be straightforward. Discrete labels either match or do not match.
• For label spaces that don’t have discrete values, such as bounding box annotations, find a broad measure of similarity. For bounding boxes, one such measure is the Jaccard index. This measures the ratio of the intersection of two boxes with the union of the boxes to assess how similar they are. For example, if there are three annotations, then there can be a function that determines which annotations represent the same object and should be consolidated.

Assess the Most Probable Label

With one of the strategies detailed in the previous sections in mind, make some sort of probabilistic judgment on what the consolidated label should be. In the case of discrete, mutually exclusive categories, this can be straightforward. One of the most common ways to do this is to take the results of a majority vote between the annotations. This weights the annotations equally.

Some approaches attempt to estimate the accuracy of different annotators and weight their annotations in proportion to the probability of correctness. An example of this is the Expectation Maximization method, which is used in the default Ground Truth consolidation function for multi-class annotations.

For more information about creating an annotation consolidation function, see Step 3: Processing with AWS Lambda.

Automate Data Labeling

If you choose, Amazon SageMaker Ground Truth can use active learning to automate the labeling of your input data for certain built-in task types. Active learning is a machine learning technique that identifies data that should be labeled by your workers. In Ground Truth, this functionality is called automated data labeling. Automated data labeling helps to reduce the cost and time that it takes to label your dataset compared to using only humans. When you use automated labeling, you incur SageMaker training and inference costs.

We recommend using automated data labeling on large datasets because the neural networks used with active learning require a significant amount of data for every new dataset. Typically, as you provide more data, the potential for high accuracy predictions goes up. Data will only be auto-labeled if the neural network used in the auto-labeling model can achieve an acceptably high level of accuracy. Therefore, with larger datasets, there is more potential to automatically label the data because the neural network can achieve high enough accuracy for auto-labeling. Automated data labeling is most appropriate when you have thousands of data objects. The minimum number of objects allowed for automated data labeling is 1,250, but we strongly suggest providing a minimum of 5,000 objects.
Automated data labeling is available only for the following Ground Truth built-in task types:

- Image Classification (Single Label)
- Image Semantic Segmentation
- Object detection (Bounding Box)
- Text Classification (Single Label)

Streaming labeling jobs do not support automated data labeling.

To learn how to create a custom active learning workflow using your own model, see Set up an active learning workflow with your own model.

Input data quotas apply for automated data labeling jobs. See Input Data Quotas for information about dataset size, input data size and resolution limits.

ℹ️ Note

Before you use an the automated-labeling model in production, you need to fine-tune or test it, or both. You might fine-tune the model (or create and tune another supervised model of your choice) on the dataset produced by your labeling job to optimize the model's architecture and hyperparameters. If you decide to use the model for inference without fine-tuning it, we strongly recommend making sure that you evaluate its accuracy on a representative (for example, randomly selected) subset of the dataset labeled with Ground Truth and that it matches your expectations.

How it Works

You enable automated data labeling when you create a labeling job. This is how it works:

1. When Ground Truth starts an automated data labeling job, it selects a random sample of input data objects and sends them to human workers. If more than 10% of these data objects fail, the labeling job will fail. If the labeling job fails, in addition to reviewing any error message Ground Truth returns, check that your input data is displaying correctly in the worker UI, instructions are clear, and that you have given workers enough time to complete tasks.

2. When the labeled data is returned, it is used to create a training set and a validation set. Ground Truth uses these datasets to train and validate the model used for auto-labeling.
3. Ground Truth runs a batch transform job, using the validated model for inference on the validation data. Batch inference produces a confidence score and quality metric for each object in the validation data.

4. The auto labeling component will use these quality metrics and confidence scores to create a confidence score threshold that ensures quality labels.

5. Ground Truth runs a batch transform job on the unlabeled data in the dataset, using the same validated model for inference. This produces a confidence score for each object.

6. The Ground Truth auto labeling component determines if the confidence score produced in step 5 for each object meets the required threshold determined in step 4. If the confidence score meets the threshold, the expected quality of automatically labeling exceeds the requested level of accuracy and that object is considered auto-labeled.

7. Step 6 produces a dataset of unlabeled data with confidence scores. Ground Truth selects data points with low confidence scores from this dataset and sends them to human workers.

8. Ground Truth uses the existing human-labeled data and this additional labeled data from human workers to update the model.

9. The process is repeated until the dataset is fully labeled or until another stopping condition is met. For example, auto-labeling stops if your human annotation budget is reached.

The preceding steps happen in iterations. Select each tab in the following table to see an example of the processes that happen in each iteration for an object detection automated labeling job. The number of data objects used in a given step in these images (for example, 200) is specific to this example. If there are fewer than 5,000 objects to label, the validation set size is 20% of the whole dataset. If there are more than 5,000 objects in your input dataset, the validation set size is 10% of the whole dataset. You can control the number of human labels collected per active learning iteration by changing the value for `MaxConcurrentTaskCount` when using the API operation `CreateLabelingJob`. This value is set to 1,000 when you create a labeling job using the console. In the active learning flow illustrated under the **Active Learning** tab, this value is set to 200.
Model Training

Iterations 1, 2, and 3: Model training

Unlabeled images → Send randomly selected images to annotators → Iterations 1, 2: 200 human-annotated training labels → Iteration 3: Train an object detection model → Iteration 3: 200 human-annotated validation labels
Automated Labeling

Iteration 4: Automated labeling

200 human-annotated validation labels

Compare human-annotated and automated labels

Confidence threshold

200 automated validation labels with confidence scores

Trained model

Automated labels: Unlabeled image labels with confidence > θ

Unlabeled images

Trained model

Active Learning

Iteration 4: Active learning

Unlabeled images

Trained model

Automated labels with confidence scores

Send 200 images with lowest label confidence to annotators

400 human-annotated training labels
Accuracy of Automated Labels

The definition of accuracy depends on the built-in task type that you use with automated labeling. For all task types, these accuracy requirements are pre-determined by Ground Truth and cannot be manually configured.

- For image classification and text classification, Ground Truth uses logic to find a label-prediction confidence level that corresponds to at least 95% label accuracy. This means Ground Truth expects the accuracy of the automated labels to be at least 95% when compared to the labels that human labelers would provide for those examples.
- For bounding boxes, the expected mean Intersection Over Union (IoU) of the auto-labeled images is 0.6. To find the mean IoU, Ground Truth calculates the mean IoU of all the predicted and missed boxes on the image for every class, and then averages these values across classes.
- For semantic segmentation, the expected mean IoU of the auto-labeled images is 0.7. To find the mean IoU, Ground Truth takes the mean of the IoU values of all the classes in the image (excluding the background).

At every iteration of Active Learning (steps 3-6 in the list above), the confidence threshold is found using the human-annotated validation set so that the expected accuracy of the auto-labeled objects satisfies certain predefined accuracy requirements.

Create an Automated Data Labeling Job (Console)

To create a labeling job that uses automated labeling in the SageMaker console, use the following procedure.

**To create an automated data labeling job (console)**

2. Using Create a Labeling Job (Console) as a guide, complete the Job overview and Task type sections. Note that auto labeling is not supported for custom task types.
3. Under Workers, choose your workforce type.
4. In the same section, choose Enable automated data labeling.
5. Using Step 4: Configure the Bounding Box Tool as a guide, create worker instructions in the section Task Type labeling tool. For example, if you chose Semantic segmentation as your labeling job type, this section is called Semantic segmentation labeling tool.
6. To preview your worker instructions and dashboard, choose Preview.
7. Choose Create. This creates and starts your labeling job and the auto labeling process.

You can see your labeling job appear in the Labeling jobs section of the SageMaker console. Your output data appears in the Amazon S3 bucket that you specified when creating the labeling job. For more information about the format and file structure of your labeling job output data, see Output Data.

Create an Automated Data Labeling Job (API)

To create an automated data labeling job using the SageMaker API, use the LabelingJobAlgorithmsConfig parameter of the CreateLabelingJob operation. To learn how to start a labeling job using the CreateLabelingJob operation, see Create a Labeling Job (API).

Specify the Amazon Resource Name (ARN) of the algorithm that you are using for automated data labeling in the LabelingJobAlgorithmSpecificationArn parameter. Choose from one of the four Ground Truth built-in algorithms that are supported with automated labeling:

- **Image Classification (Single Label)**
- **Image Semantic Segmentation**
- **Object detection (Bounding Box)**
- **Text Classification (Single Label)**

When an automated data labeling job finishes, Ground Truth returns the ARN of the model it used for the automated data labeling job. Use this model as the starting model for similar auto-labeling job types by providing the ARN, in string format, in the InitialActiveLearningModelArn parameter. To retrieve the model's ARN, use an AWS Command Line Interface (AWS CLI) command similar to the following.

```
# Fetch the mARN of the model trained in the final iteration of the previous labeling job.Ground Truth
pretrained_model_arn = sagemaker_client.describe_labeling_job(LabelingJobName=job_name)['LabelingJobOutput']['FinalActiveLearningModelArn']
```

To encrypt data on the storage volume attached to the ML compute instance(s) that are used in automated labeling, include an AWS Key Management Service (AWS KMS) key in the
VolumeKmsKeyId parameter. For information about AWS KMS keys, see What is AWS Key Management Service? in the AWS Key Management Service Developer Guide.

For an example that uses the CreateLabelingJob operation to create an automated data labeling job, see the object_detection_tutorial example in the SageMaker Examples, Ground Truth Labeling Jobs section of a SageMaker notebook instance. To learn how to create and open a notebook instance, see Create a Notebook Instance. To learn how to access SageMaker example notebooks, see Example Notebooks.

Amazon EC2 Instances Required for Automated Data Labeling

The following table lists the Amazon Elastic Compute Cloud (Amazon EC2) instances that you need to run automated data labeling for training and batch inference jobs.

<table>
<thead>
<tr>
<th>Automated Data Labeling Job Type</th>
<th>Training Instance Type</th>
<th>Inference Instance Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image classification</td>
<td>ml.p3.2xlarge*</td>
<td>ml.c5.xlarge</td>
</tr>
<tr>
<td>Object detection (bounding box)</td>
<td>ml.p3.2xlarge*</td>
<td>ml.c5.4xlarge</td>
</tr>
<tr>
<td>Text classification</td>
<td>ml.c5.2xlarge</td>
<td>ml.m4.xlarge</td>
</tr>
<tr>
<td>Semantic segmentation</td>
<td>ml.p3.2xlarge*</td>
<td>ml.p3.2xlarge*</td>
</tr>
</tbody>
</table>

* In the Asia Pacific (Mumbai) Region (ap-south-1) use ml.p2.8xlarge instead.

Ground Truth manages the instances that you use for automated data labeling jobs. It creates, configures, and terminates the instances as needed to perform your job. These instances don’t appear in your Amazon EC2 instance dashboard.

Set up an active learning workflow with your own model

You can create an active learning workflow with your own algorithm to run training and inferencess in that workflow to auto-label your data. The notebook bring_your_own_model_for_sagemaker_labeling_workflows_with_active_learning.ipynb demonstrates this using the SageMaker built-in algorithm, BlazingText. This notebook provides an AWS CloudFormation stack that you can use to execute this workflow using AWS Step Functions. You can find the notebook and supporting files in this GitHub repository.
Chaining Labeling Jobs

Amazon SageMaker Ground Truth can reuse datasets from prior jobs in two ways: cloning and chaining.

Cloning copies the setup of a prior labeling job and allows you to make additional changes before setting it to run.

Chaining uses not only the setup of the prior job, but also the results. This allows you to continue an incomplete job and add labels or data objects to a completed job. Chaining is a more complex operation.

For data processing:

- Cloning uses the prior job's input manifest, with optional modifications, as the new job's input manifest.
- Chaining uses the prior job's output manifest as the new job's input manifest.

Chaining is useful when you need to:

- Continue a labeling job that was manually stopped.
- Continue a labeling job that failed mid-job, after fixing issues.
- Switch to automated data labeling after manually labeling part of a job (or the other way around).
- Add more data objects to a completed job and start the job from there.
- Add another annotation to a completed job. For example, you have a collection of phrases labeled for topic, then want to run the set again, categorizing them by the topic's implied audience.

In Amazon SageMaker Ground Truth you can configure a chained labeling job with either the console or the API.
Key Term: Label Attribute Name

The *label attribute name* (LabelAttributeName in the API) is a string used as the key for the key-value pair formed with the label that a worker assigns to the data object.

The following rules apply for the label attribute name:

- It can't end with `-metadata`.
- The names source and source-ref are reserved and can't be used.
- For semantic segmentation labeling jobs, it must end with `-ref`. For all other labeling jobs, it can't end with `-ref`. If you use the console to create the job, Amazon SageMaker Ground Truth automatically appends `-ref` to all label attribute names except for semantic segmentation jobs.
- For a chained labeling job, if you're using the same label attribute name from the originating job and you configure the chained job to use auto-labeling, then if it had been in auto-labeling mode at any point, Ground Truth uses the model from the originating job.

In an output manifest, the label attribute name appears similar to the following.

```json
"source-ref": "<S3 URI>",
"<label attribute name>": {
  "annotations": [ {
    "class_id": 0,
    "width": 99,
    "top": 87,
    "height": 62,
    "left": 175
  }],
  "image_size": [ {
    "width": 344,
    "depth": 3,
    "height": 234
  }]
},
"<label attribute name>-metadata": {
  "job-name": "<job name>",
  "class-map": {
    "0": "<label attribute name>"
  },
  "human-annotated": "yes",
  "objects": [ {
    "confidence": 0.09
  }
},
```

Enhanced Data Labeling
If you're creating a job in the console and don't explicitly set the label attribute name value, Ground Truth uses the job name as the label attribute name for the job.

**Start a Chained Job (Console)**

Choose a stopped, failed, or completed labeling job from the list of your existing jobs. This enables the *Actions* menu.

From the *Actions* menu, choose *Chain*.

**Job Overview Panel**

In the *Job overview* panel, a new *Job name* is set based on the title of the job from which you are chaining this one. You can change it.

You may also specify a label attribute name different from the labeling job name.

If you're chaining from a completed job, the label attribute name uses the name of the new job you're configuring. To change the name, select the check box.

If you're chaining from a stopped or failed job, the label attribute name uses the name of the job from which you're chaining. It's easy to see and edit the value because the name check box is checked.

**Attribute label naming considerations**

- **The default** uses the label attribute name Ground Truth has selected. All data objects without data connected to that label attribute name are labeled.
- **Using a label attribute name** not present in the manifest causes the job to process *all* the objects in the dataset.

The *input dataset location* in this case is automatically selected as the output manifest of the chained job. The input field is not available, so you cannot change it.
Adding data objects to a labeling job

You cannot specify an alternate manifest file. Manually edit the output manifest from the previous job to add new items before starting a chained job. The Amazon S3 URI helps you locate where you are storing the manifest in your Amazon S3 bucket. Download the manifest file from there, edit it locally on your computer, and then upload the new version to replace it. Make sure you are not introducing errors during editing. We recommend you use JSON linter to check your JSON. Many popular text editors and IDEs have linter plugins available.

Start a Chained Job (API)

The procedure is almost the same as setting up a new labeling job with CreateLabelingJob, except for two primary differences:

- **Manifest location**: Rather than use your original manifest from the prior job, the value for the ManifestS3Uri in the DataSource should point to the Amazon S3 URI of the output manifest from the prior labeling job.

- **Label attribute name**: Setting the correct LabelAttributeName value is important here. This is the key portion of a key-value pair where labeling data is the value. Sample use cases include:
  - **Adding new or more specific labels to a completed job** — Set a new label attribute name.
  - **Labeling the unlabeled items from a prior job** — Use the label attribute name from the prior job.

Use a Partially Labeled Dataset

You can get some chaining benefits if you use an augmented manifest that has already been partially labeled. Check the **Label attribute name** check box and set the name so that it matches the name in your manifest.

If you're using the API, the instructions are the same as those for starting a chained job. However, be sure to upload your manifest to an Amazon S3 bucket and use it instead of using the output manifest from a prior job.

The **Label attribute name** value in the manifest has to conform to the naming considerations discussed earlier.
Ground Truth Security and Permissions

Use the topics on this page to learn about Ground Truth security features and how to configure AWS Identity and Access Management (IAM) permissions to allow a user or role to create a labeling job. Additionally, learn how to create an execution role. An execution role is the role that you specify when you create a labeling job. This role is used to start your labeling job.

If you are a new user and want to get started quickly, or if you do not require granular permissions, see Use IAM Managed Policies with Ground Truth.

For more information about IAM users and roles, see Identities (Users, Groups, and Roles) in the IAM User Guide.

To learn more about using IAM with SageMaker, see Identity and Access Management for Amazon SageMaker.

Topics

- CORS Permission Requirement
- Assign IAM Permissions to Use Ground Truth
- Using Amazon SageMaker Ground Truth in an Amazon Virtual Private Cloud
- Output Data and Storage Volume Encryption
- Workforce Authentication and Restrictions

CORS Permission Requirement

Earlier in 2020, widely used browsers like Chrome and Firefox changed their default behavior for rotating images based on image metadata, referred to as EXIF data. Previously, browsers would always display images in exactly the manner in which they are stored on disk, which is typically unrotated. After the change, images now rotate according to a piece of image metadata called orientation value. This has important implications for the entire machine learning (ML) community. For example, if applications that annotate images do not consider the EXIF orientation, they may display images in unexpected orientations, resulting in incorrect labels.

Starting with Chrome 89, AWS can no longer automatically prevent the rotation of images because the web standards group W3C has decided that the ability to control rotation of images violates the web's Same-origin Policy. Therefore, to ensure human workers annotate your input images in a predictable orientation when you submit requests to create a labeling job, you must add a CORS header policy to the Amazon S3 buckets that contain your input images.
Important

If you do not add a CORS configuration to the Amazon S3 buckets that contain your input data, labeling tasks for those input data objects will fail.

If you create a job through the Ground Truth console, CORS is enabled by default. If all of your input data is not located in the same Amazon S3 bucket as your input manifest file, you must add a CORS configuration to all Amazon S3 buckets that contain input data using the following instructions.

If you are using the CreateLabelingJob API to create a Ground Truth labeling job, you can add a CORS policy to an Amazon S3 bucket that contains input data in the S3 console. To set the required CORS headers on the Amazon S3 bucket that contain your input images in the Amazon S3 console, follow the directions detailed in How do I add cross-domain resource sharing with CORS?. Use the following CORS configuration code for the buckets that host your images. If you use the Amazon S3 console to add the policy to your bucket, you must use the JSON format.

Important

If you create a 3D point cloud or video frame labeling job, you must add additional rules to your CORS configuration. To learn more, see 3D Point Cloud Labeling Job Permission Requirements and Video Frame Job Permission Requirements respectively.

### JSON

```json
[
  
  "AllowedHeaders": [],
  "AllowedMethods": ["GET"],
  "AllowedOrigins": ["*"],
  "ExposeHeaders": ["Access-Control-Allow-Origin"]
]
```

### XML

```xml
<CORSConfiguration>
  <CORSRule>
    <AllowedOrigin>*</AllowedOrigin>
    <AllowedMethod>GET</AllowedMethod>
  </CORSRule>
</CORSConfiguration>
```
Assign IAM Permissions to Use Ground Truth

Use the topics in this section to learn how to use AWS Identity and Access Management (IAM) managed and custom policies to manage access to Ground Truth and associated resources.

You can use the sections on this page to learn the following:

- How to create IAM policies that grant a user or role permission to create a labeling job. Administrators can use IAM policies to restrict access to Amazon SageMaker and other AWS services that are specific to Ground Truth.
- How to create a SageMaker execution role. An execution role is the role that you specify when you create a labeling job. The role is used to start and manage your labeling job.

The following is an overview of the topics you'll find on this page:

- If you are getting started using Ground Truth, or you do not require granular permissions for your use case, it is recommended that you use the IAM managed policies described in Use IAM Managed Policies with Ground Truth.
- Learn about the permissions required to use the Ground Truth console in Grant IAM Permission to Use the Amazon SageMaker Ground Truth Console. This section includes policy examples that grant an IAM entity permission to create and modify private work teams, subscribe to vendor work teams, and create custom labeling workflows.
- When you create a labeling job, you must provide an execution role. Use Create a SageMaker Execution Role for a Ground Truth Labeling Job to learn about the permissions required for this role.

Use IAM Managed Policies with Ground Truth

SageMaker and Ground Truth provide AWS managed policies that you can use to create a labeling job. If you are getting started using Ground Truth and you do not require granular permissions for your use case, it is recommended that you use the following policies:

- AmazonSageMakerFullAccess – Use this policy to give a user or role permission to create a labeling job. This is a broad policy that grants a entity permission to use SageMaker features,
as well as features of necessary AWS services through the console and API. This policy gives the entity permission to create a labeling job and to create and manage workforces using Amazon Cognito. To learn more, see AmazonSageMakerFullAccess Policy.

- **AmazonSageMakerGroundTruthExecution** – To create an execution role, you can attach the policy AmazonSageMakerGroundTruthExecution to a role. An execution role is the role that you specify when you create a labeling job and it is used to start your labeling job. This policy allows you to create both streaming and non-streaming labeling jobs, and to create a labeling job using any task type. Note the following limits of this managed policy.

  - **Amazon S3 permissions**: This policy grants an execution role permission to access Amazon S3 buckets with the following strings in the name: GroundTruth, Groundtruth, groundtruth, SageMaker, Sagemaker, and sagemaker or a bucket with an object tag that includes SageMaker in the name (case insensitive). Make sure your input and output bucket names include these strings, or add additional permissions to your execution role to grant it permission to access your Amazon S3 buckets. You must give this role permission to perform the following actions on your Amazon S3 buckets: AbortMultipartUpload, GetObject, and PutObject.

  - **Custom Workflows**: When you create a custom labeling workflow, this execution role is restricted to invoking AWS Lambda functions with one of the following strings as part of the function name: GtRecipe, SageMaker, Sagemaker, sagemaker, or LabelingFunction. This applies to both your pre-annotation and post-annotation Lambda functions. If you choose to use names without those strings, you must explicitly provide lambda:InvokeFunction permission to the execution role used to create the labeling job.

To learn how to attach an AWS managed policy to a user or role, refer to Adding and removing IAM identity permissions in the IAM User Guide.

### Grant IAM Permission to Use the Amazon SageMaker Ground Truth Console

To use the Ground Truth area of the SageMaker console, you need to grant permission to an entity to access SageMaker and other AWS services that Ground Truth interacts with. Required permissions to access other AWS services depends on your use-case:

- Amazon S3 permissions are required for all use cases. These permissions must grant access to the Amazon S3 buckets that contain input and output data.

- AWS Marketplace permissions are required to use a vendor workforce.

- Amazon Cognito permission are required for private work team setup.
AWS KMS permissions are required to view available AWS KMS keys that can be used for output data encryption.

IAM permissions are required to either list pre-existing execution roles, or to create a new one. Additionally, you must use a PassRole permission to allow SageMaker to use the execution role chosen to start the labeling job.

The following sections list policies you may want to grant to a role to use one or more functions of Ground Truth.

Topics

- Ground Truth Console Permissions
- Custom Labeling Workflow Permissions
- Private Workforce Permissions
- Vendor Workforce Permissions

Ground Truth Console Permissions

To grant permission to a user or role to use the Ground Truth area of the SageMaker console to create a labeling job, attach the following policy to the user or role. The following policy will give an IAM role permission to create a labeling job using a built-in task type task type. If you want to create a custom labeling workflow, add the policy in Custom Labeling Workflow Permissions to the following policy. Each Statement included in the following policy is described below this code block.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "SageMakerApis",
            "Effect": "Allow",
            "Action": ["sagemaker:*"],
            "Resource": "*"
        },
        {
            "Sid": "KmsKeysForCreateForms",
            "Effect": "Allow",
            "Action": ["kms:*"],
            "Resource": "*"
        }
    ]
}
```
"Action": [
    "kms:DescribeKey",
    "kms:ListAliases"
],
"Resource": "*"
},
{
    "Sid": "AccessAwsMarketplaceSubscriptions",
    "Effect": "Allow",
    "Action": [
        "aws-marketplace:ViewSubscriptions"
    ],
    "Resource": "*"
},
{
    "Sid": "SecretsManager",
    "Effect": "Allow",
    "Action": [
        "secretsmanager:CreateSecret",
        "secretsmanager:DescribeSecret",
        "secretsmanager:ListSecrets"
    ],
    "Resource": "*"
},
{
    "Sid": "ListAndCreateExecutionRoles",
    "Effect": "Allow",
    "Action": [
        "iam:ListRoles",
        "iam:CreateRole",
        "iam:CreatePolicy",
        "iam:AttachRolePolicy"
    ],
    "Resource": "*"
},
{
    "Sid": "PassRoleForExecutionRoles",
    "Effect": "Allow",
    "Action": [
        "iam:PassRole"
    ],
    "Resource": "*",
    "Condition": {
        "StringEquals": {
            ...
        }
    }
}
"iam:PassedToService": "sagemaker.amazonaws.com"

},
],
{
"Sid": "GroundTruthConsole",
"Effect": "Allow",
"Action": [
"groundtruthlabeling:*",
"lambda:InvokeFunction",
"lambda:ListFunctions",
"s3:GetObject",
"s3:PutObject",
"s3:ListBucket",
"s3:GetBucketCors",
"s3:PutBucketCors",
"s3:ListAllMyBuckets",
"cognito-idp:AdminAddUserToGroup",
"cognito-idp:AdminCreateUser",
"cognito-idp:AdminDeleteUser",
"cognito-idp:AdminDisableUser",
"cognito-idp:AdminEnableUser",
"cognito-idp:AdminRemoveUserFromGroup",
"cognito-idp:CreateGroup",
"cognito-idp:CreateUserPool",
"cognito-idp:CreateUserPoolClient",
"cognito-idp:CreateUserPoolDomain",
"cognito-idp:DescribeUserPool",
"cognito-idp:DescribeUserPoolClient",
"cognito-idp:ListGroups",
"cognito-idp:ListIdentityProviders",
"cognito-idp:ListUsers",
"cognito-idp:ListUsersInGroup",
"cognito-idp:ListUserPoolClients",
"cognito-idp:ListUserPools",
"cognito-idp:UpdateUserPool",
"cognito-idp:UpdateUserPoolClient"
],
"Resource": "**"
This policy includes the following statements. You can scope down any of these statements by adding specific resources to the Resource list for that statement.

**SageMakerApis**

This statement includes `sagemaker:*`, which allows the user to perform all SageMaker API actions. You can reduce the scope of this policy by restricting users from performing actions that are not used to create and monitoring a labeling job.

**KmsKeysForCreateForms**

You only need to include this statement if you want to grant a user permission to list and select AWS KMS keys in the Ground Truth console to use for output data encryption. The policy above grants a user permission to list and select any key in the account in AWS KMS. To restrict the keys that a user can list and select, specify those key ARNs in Resource.

**SecretsManager**

This statement gives the user permission to describe, list, and create resources in AWS Secrets Manager required to create the labeling job.

**ListAndCreateExecutionRoles**

This statement gives a user permission to list (ListRoles) and create (CreateRole) IAM roles in your account. It also grants the user permission to create (CreatePolicy) policies and attach (AttachRolePolicy) policies to entities. These are required to list, select, and if required, create an execution role in the console.

If you have already created an execution role, and want to narrow the scope of this statement so that users can only select that role in the console, specify the ARNs of the roles you want the user to have permission to view in Resource and remove the actions CreateRole, CreatePolicy, and AttachRolePolicy.

**AccessAwsMarketplaceSubscriptions**

These permissions are required to view and choose vendor work teams that you are already subscribed to when creating a labeling job. To give the user permission to subscribe to vendor work teams, add the statement in Vendor Workforce Permissions to the policy above.

**PassRoleForExecutionRoles**
This is required to give the labeling job creator permission to preview the worker UI and verify that input data, labels, and instructions display correctly. This statement gives an entity permissions to pass the IAM execution role used to create the labeling job to SageMaker to render and preview the worker UI. To narrow the scope of this policy, add the role ARN of the execution role used to create the labeling job under Resource.

**GroundTruthConsole**

- **groundtruthlabeling** – This allows a user to perform actions required to use certain features of the Ground Truth console. These include permissions to describe the labeling job status (DescribeConsoleJob), list all dataset objects in the input manifest file (ListDatasetObjects), filter the dataset if dataset sampling is selected (RunFilterOrSampleDatasetJob), and to generate input manifest files if automated data labeling is used (RunGenerateManifestByCrawlingJob). These actions are only available when using the Ground Truth console and cannot be called directly using an API.

- **lambda:InvokeFunction and lambda:ListFunctions** – these actions give users permission to list and invoke Lambda functions that are used to run a custom labeling workflow.

- **s3:** – All Amazon S3 permissions included in this statement are used to view Amazon S3 buckets for **automated data setup** (ListAllMyBuckets), access input data in Amazon S3 (ListBucket, GetObject), check for and create a CORS policy in Amazon S3 if needed (GetBucketCors and PutBucketCors), and write labeling job output files to S3 (PutObject).

- **cognito-idp** – These permissions are used to create, view and manage and private workforce using Amazon Cognito. To learn more about these actions, refer to the Amazon Cognito API References.

**Custom Labeling Workflow Permissions**

Add the following statement to a policy similar to the one in Ground Truth Console Permissions to give a user permission to select pre-existing pre-annotation and post-annotation Lambda functions while creating a custom labeling workflow.

```json
{
    "Sid": "GroundTruthConsoleCustomWorkflow",
    "Effect": "Allow",
    "Action": [
        "lambda:InvokeFunction",
        "lambda:ListFunctions"
    ],
}
```
To learn how to give an entity permission to create and test pre-annotation and post-annotation Lambda functions, see Required Permissions To Use Lambda With Ground Truth.

Private Workforce Permissions

When added to a permissions policy, the following permission grants access to create and manage a private workforce and work team using Amazon Cognito. These permissions are not required to use an OIDC IdP workforce.

```
{
    "Effect": "Allow",
    "Action": [
        "cognito-idp:AdminAddUserToGroup",
        "cognito-idp:AdminCreateUser",
        "cognito-idp:AdminDeleteUser",
        "cognito-idp:AdminDisableUser",
        "cognito-idp:AdminEnableUser",
        "cognito-idp:AdminRemoveUserFromGroup",
        "cognito-idp:CreateGroup",
        "cognito-idp:CreateUserPool",
        "cognito-idp:CreateUserPoolClient",
        "cognito-idp:CreateUserPoolDomain",
        "cognito-idp:DescribeUserPool",
        "cognito-idp:DescribeUserPoolClient",
        "cognito-idp:ListGroups",
        "cognito-idp:ListIdentityProviders",
        "cognito-idp:ListUsers",
        "cognito-idp:ListUsersInGroup",
        "cognito-idp:ListUserPoolClients",
        "cognito-idp:ListUserPools",
        "cognito-idp:UpdateUserPool",
        "cognito-idp:UpdateUserPoolClient"
    ],
    "Resource": "*"
}
```

To learn more about creating private workforce using Amazon Cognito, see Create and Manage Amazon Cognito Workforce.
Vendor Workforce Permissions

You can add the following statement to the policy in Grant IAM Permission to Use the Amazon SageMaker Ground Truth Console to grant an entity permission to subscribe to a vendor workforce.

```
{
    "Sid": "AccessAwsMarketplaceSubscriptions",
    "Effect": "Allow",
    "Action": [
        "aws-marketplace:Subscribe",
        "aws-marketplace:Unsubscribe",
        "aws-marketplace:ViewSubscriptions"
    ],
    "Resource": "*"
}
```

Create a SageMaker Execution Role for a Ground Truth Labeling Job

When you configure your labeling job, you need to provide an execution role, which is a role that SageMaker has permission to assume to start and run your labeling job.

This role must give Ground Truth permission to access the following:

- Amazon S3 to retrieve your input data and write output data to an Amazon S3 bucket. You can either grant permission for an IAM role to access an entire bucket by providing the bucket ARN, or you can grant access to the role to access specific resources in a bucket. For example, the ARN for a bucket may look similar to `arn:aws:s3:::awsexamplebucket1` and the ARN of a resource in an Amazon S3 bucket may look similar to `arn:aws:s3:::awsexamplebucket1/prefix/file-name.png`. To apply an action to all resources in an Amazon S3 bucket, you can use the wild card: `*`. For example, `arn:aws:s3:::awsexamplebucket1/prefix/*`. For more information, see Amazon Amazon S3 Resources in the Amazon Simple Storage Service User Guide.

- CloudWatch to log worker metrics and labeling job statuses.

- AWS KMS for data encryption. (Optional)

- AWS Lambda for processing input and output data when you create a custom workflow.

Additionally, if you create a streaming labeling job, this role must have permission to access:

- Amazon SQS to create an interact with an SQS queue used to manage labeling requests.
Amazon SNS to subscribe to and retrieve messages from your Amazon SNS input topic and to send messages to your Amazon SNS output topic.

All of these permissions can be granted with the AmazonSageMakerGroundTruthExecution managed policy except:

- Data and storage volume encryption of your Amazon S3 buckets. To learn how to configure these permissions, see Encrypt Output Data and Storage Volume with AWS KMS.

- Permission to select and invoke Lambda functions that do not include GtRecipe, SageMaker, Sagemaker, sagemaker, or LabelingFunction in the function name.

- Amazon S3 buckets that do not include either GroundTruth, Groundtruth, groundtruth, SageMaker, Sagemaker, and sagemaker in the prefix or bucket name or an object tag that includes SageMaker in the name (case insensitive).

If you require more granular permissions than the ones provided in AmazonSageMakerGroundTruthExecution, use the following policy examples to create an execution role that fits your specific use case.

Topics

- Built-In Task Types (Non-streaming) Execution Role Requirements
- Built-In Task Types (Streaming) Execution Role Requirements
- Execution Role Requirements for Custom Task Types
- Automated Data Labeling Permission Requirements

Built-In Task Types (Non-streaming) Execution Role Requirements

The following policy grants permission to create a labeling job for a built-in task type. This execution policy does not include permissions for AWS KMS data encryption or decryption. Replace each red, italicized ARN with your own Amazon S3 ARNs.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "S3ViewBuckets",
         "Effect": "Allow",
         "Action": "s3:ListBucket",
         "Resource": "arn:aws:s3:::<bucket_name>"
      }
   ]
}
```
Built-In Task Types (Streaming) Execution Role Requirements

If you create a streaming labeling job, you must add a policy similar to the following to the execution role you use to create the labeling job. To narrow the scope of the policy, replace the * in Resource with specific AWS resources that you want to grant the IAM role permission to access and use.
{ "Version": "2012-10-17", "Statement": [ {
  "Effect": "Allow",
  "Action": [ "s3:AbortMultipartUpload", "s3:GetObject", "s3:PutObject"
  ],
  "Resource": [ "arn:aws:s3::<input-bucket-name>/*", "arn:aws:s3::<output-bucket-name>/*" ]
}, {
  "Effect": "Allow",
  "Action": [ "s3:GetObject" ],
  "Resource": "*",
  "Condition": {
    "StringEqualsIgnoreCase": { "s3:ExistingObjectTag/SageMaker": "true" }
  }
}, {
  "Effect": "Allow",
  "Action": [ "s3:GetBucketLocation", "s3:ListBucket" ],
  "Resource": [ "arn:aws:s3::<input-bucket-name>", "arn:aws:s3::<output-bucket-name>" ]
}, {
  "Sid": "CloudWatch",
  "Effect": "Allow",
  "Action": [ "cloudwatch:PutMetricData", "cloudwatch:GetMetricData" ],
  "Resource": "*"
} ]}
"logs:DescribeLogStreams",
"logs:PutLogEvents"
],
"Resource": "*"
},
{
"Sid": "StreamingQueue",
"Effect": "Allow",
"Action": [
"sqs:CreateQueue",
"sqs:DeleteMessage",
"sqs:GetQueueAttributes",
"sqs:GetQueueUrl",
"sqs:ReceiveMessage",
"sqs:SendMessage",
"sqs:SendMessageBatch",
"sqs:SetQueueAttributes"
],
"Resource": "arn:aws:sqs:*:*:*GroundTruth*"
},
{
"Sid": "StreamingTopicSubscribe",
"Effect": "Allow",
"Action": "sns:Subscribe",
"Resource": [  
"arn:aws:sns:<aws-region>:<aws-account-number>:<output-topic-name>"
],
"Condition": {
"StringEquals": {
  "sns:Protocol": "sqs"
},
"StringLike": {
}
}
},
{
"Sid": "StreamingTopic",
"Effect": "Allow",
"Action": [
"sqs:CreateQueue",
"sqs:DeleteMessage",
"sqs:GetQueueAttributes",
"sqs:GetQueueUrl",
"sqs:ReceiveMessage",
"sqs:SendMessage",
"sqs:SendMessageBatch",
"sqs:SetQueueAttributes"
],
"Resource": "arn:aws:sqs:*:*:*GroundTruth*"
]
Execution Role Requirements for Custom Task Types

If you want to create a custom labeling workflow, add the following statement to an execution role policy like the ones found in Built-In Task Types (Non-streaming) Execution Role Requirements or Built-In Task Types (Streaming) Execution Role Requirements.

This policy gives the execution role permission to Invoke your pre-annotation and post-annotation Lambda functions.

```json
{
    "Sid": "LambdaFunctions",
    "Effect": "Allow",
    "Action": [
        "lambda:InvokeFunction"
    ],
    "Resource": [
        "arn:aws:lambda:<region>:<account-id>:function:<post-annotation-lambda-name>"
    ]
}
```
Automated Data Labeling Permission Requirements

If you want to create a labeling job with automated data labeling enabled, you must 1) add one policy to the IAM policy attached to the execution role and 2) update the trust policy of the execution role.

The following statement allows the IAM execution role to be passed to SageMaker so that it can be used to run the training and inference jobs used for active learning and automated data labeling respectively. Add this statement to an execution role policy like the ones found in Built-In Task Types (Non-streaming) Execution Role Requirements or Built-In Task Types (Streaming) Execution Role Requirements. Replace arn:aws:iam::<account-number>:role/<role-name> with the execution role ARN. You can find your IAM role ARN in the IAM console under Roles.

```
{
   "Effect": "Allow",
   "Action": [ "iam:PassRole"
   ],
   "Resource": "arn:aws:iam::<account-number>:role/<execution-role-name>",
   "Condition": {
      "StringEquals": { "iam:PassedToService": [ "sagemaker.amazonaws.com" ]
   }
}
```

The following statement allows SageMaker to assume the execution role to create and manage the SageMaker training and inference jobs. This policy must be added to the trust relationship of the execution role. To learn how to add or modify an IAM role trust policy, see Modifying a role in the IAM User Guide.

```
{
   "Version": "2012-10-17",
   "Statement": {
      "Effect": "Allow",
      "Principal": {"Service": "sagemaker.amazonaws.com" },
      "Action": "sts:AssumeRole"
   }
}
```
Encrypt Output Data and Storage Volume with AWS KMS

You can use AWS Key Management Service (AWS KMS) to encrypt output data from a labeling job by specifying a customer managed key when you create the labeling job. If you use the API operation CreateLabelingJob to create a labeling job that uses automated data labeling, you can also use a customer managed key to encrypt the storage volume attached to the ML compute instances to run the training and inference jobs.

This section describes the IAM policies you must attach to your customer managed key to enable output data encryption and the policies you must attach to your customer managed key and execution role to use storage volume encryption. To learn more about these options, see Output Data and Storage Volume Encryption.

Encrypt Output Data using KMS

If you specify an AWS KMS customer managed key to encrypt output data, you must add an IAM policy similar to the following to that key. This policy gives the IAM execution role that you use to create your labeling job permission to use this key to perform all of the actions listed in "Action". To learn more about these actions, see AWS KMS permissions in the AWS Key Management Service Developer Guide.

To use this policy, replace the IAM service-role ARN in "Principal" with the ARN of the execution role you use to create the labeling job. When you create a labeling job in the console, this is the role you specify for IAM Role under the Job overview section. When you create a labeling job using CreateLabelingJob, this is ARN you specify for RoleArn.

```json
{
    "Sid": "AllowUseOfKmsKey",
    "Effect": "Allow",
    "Principal": {
        "AWS": "arn:aws:iam::111122223333:role/service-role/example-role"
    },
    "Action": [
        "kms:Encrypt",
        "kms:Decrypt",
        "kms:ReEncrypt*",
        "kms:GenerateDataKey*",
        "kms:DescribeKey"
    ],
    "Resource": "*
    
```
Encrypt Automated Data Labeling ML Compute Instance Storage Volume

If you specify a `VolumeKmsKeyId` to encrypt the storage volume attached to the ML compute instance used for automated data labeling training and inference, you must do the following:

- Attach permissions described in [Encrypt Output Data using KMS](#) to the customer managed key.
- Attach a policy similar to the following to the IAM execution role you use to create your labeling job. This is the IAM role you specify for `RoleArn` in `CreateLabelingJob`. To learn more about the "kms:CreateGrant" action that this policy permits, see [CreateGrant](#) in the AWS Key Management Service API Reference.

```json
{
  "Version": "2012-10-17",
  "Statement":
  [
    {
      "Effect": "Allow",
      "Action": [
        "kms:CreateGrant"
      ],
      "Resource": "*"
    }
  ]
}
```

To learn more about Ground Truth storage volume encryption, see [Use Your KMS Key to Encrypt Automated Data Labeling Storage Volume (API Only)](#).

Using Amazon SageMaker Ground Truth in an Amazon Virtual Private Cloud

[Amazon Virtual Private Cloud](#) (Amazon VPC) is a service with which you can launch AWS resources in a logically isolated virtual network that you define. You can create and run a Ground Truth labeling job inside of an Amazon VPC instead of connecting over the internet. When you launch a labeling job in an Amazon VPC, communication between your VPC and Ground Truth is conducted entirely and securely within the AWS network.

A Ground Truth labeling job in Amazon VPC follows the same behavior as PrivateLink Endpoints. For more information, see [Shared subnets](#).
This guide shows how you can use Ground Truth in an Amazon VPC in the following ways:

1. Run an Amazon SageMaker Ground Truth Labeling Job in an Amazon Virtual Private Cloud
2. Use Amazon VPC Mode from a Private Worker Portal

Run an Amazon SageMaker Ground Truth Labeling Job in an Amazon Virtual Private Cloud

Amazon SageMaker Ground Truth supports the following functionalities.

- You can use Amazon S3 bucket policies to control access to buckets from specific Amazon VPC endpoints, or specific VPCs. If you launch a labeling job and your input data is located in an Amazon S3 bucket with access restricted to users in your VPC, you can add a bucket policy to also grant a Ground Truth endpoint permission to access the bucket. To learn more, see Allow Ground Truth to Access VPC Restricted Amazon S3 Buckets.

- You can launch an automated data labeling job in your VPC. You use a VPC configuration to specify VPC subnets and security groups. SageMaker uses this configuration to launch the training and inference jobs used for automated data labeling in your VPC. To learn more, see Create an Automated Data Labeling Job in a VPC.

You may want to use these options in any of the following ways.

- You can use both of these methods to launch a labeling job using a VPC-protected Amazon S3 bucket with automated data labeling enabled.
- You can launch a labeling job using any built-in task type using a VPC-protected bucket.
- You can launch a custom labeling workflow using a VPC-protected bucket. Ground Truth interacts with your pre-annotation and post-annotation Lambda functions using an AWS PrivateLink endpoint.

We recommend that you review Prerequisites to Run a Ground Truth Labeling Job in a VPC before you create a labeling job in an Amazon VPC.

Prerequisites to Run a Ground Truth Labeling Job in a VPC

Review the following prerequisites before you create a Ground Truth labeling job in an Amazon VPC.
• If you are a new user of Ground Truth, review Getting started to learn how to create a labeling job.

• If your input data is located in a VPC-protected Amazon S3 bucket, your workers must access the worker portal from your VPC.

⚠️ **Note**

When you launch a labeling job in your VPC, you must use a private work team. To learn more about creating a private work team, see Use a Private Workforce.

• If you want to launch an automated data labeling job in your VPC, review the following prerequisites.
  
  • Use the instructions in Create an Amazon S3 VPC Endpoint. Training and inference containers used in the automated data labeling workflow use this endpoint to communicate with your buckets in Amazon S3.
  
  • Review Automate Data Labeling to learn more about this feature. Note that automated data labeling is supported for the following built-in task types: Image Classification (Single Label), Image Semantic Segmentation, Bounding Box, and Text Classification (Single Label). Streaming labeling jobs do not support automated data labeling.

• Review the Ground Truth Security and Permissions section and ensure that you have met the following conditions.
  
  • The user creating the labeling job has all necessary permissions
  
  • You have created an IAM execution role with required permissions. If you do not require fine-tuned permissions for your use case, we recommend you use the IAM managed policies described in Grant General Permissions To Get Started Using Ground Truth.
  
  • Allow your VPC to have access to the sagemaker-labeling-data-region and sm-bxcb-region-saved-task-states S3 buckets. These are system owned regionalized S3 buckets that are accessed from worker portal when worker is working on a task. We use these buckets to interact with system managed data.

**Allow Ground Truth to Access VPC Restricted Amazon S3 Buckets**

The following sections provide details about the permissions Ground Truth requires to launch labeling jobs using Amazon S3 buckets that have access restricted to your VPC and VPC endpoints. To learn how to restrict access to an Amazon S3 bucket to a VPC, see Controlling access from VPC.
endpoints with bucket policies in the Amazon Simple Storage Service User Guide guide. To learn how to add a policy to an S3 bucket, see Adding a bucket policy using the Amazon S3 console.

Note
Modifying policies on existing buckets can cause IN_PROGRESS Ground Truth jobs to fail. We recommend you start new jobs using a new bucket. If you want to continue using the same bucket, you can do one of the following.

- Wait for an IN_PROGRESS job to finish.
- Terminate the job using the console or the AWS CLI.

You can restrict Amazon S3 bucket access to users in your VPC using an AWS PrivateLink endpoint. For example, the following S3 bucket policy allows access to a specific bucket, <bucket-name>, from <vpc> and the endpoint <vpc-endpoint> only. When you modify this policy, you must replace all red-italized text with your resources and specifications.

Note
The following policy denies all entities other than users within a VPC to perform the actions listed in Action. If you do not include actions in this list, they are still accessible to any entity that has access to this bucket and permission to perform those actions. For example, if a user has permission to perform GetBucketLocation on your Amazon S3 bucket, the policy below does not restrict the user from performing this action outside of your VPC.

```json
{
  "Version": "2012-10-17",
  "Id": "Policy1415115909152",
  "Statement": [
    {
      "Sid": "Access-to-specific-VPCE-only",
      "Principal": "*",
      "Action": [
        "s3:GetObject",
        "s3:PutObject"
      ],
      "Effect": "Deny",
    }
  ]
}
```
Ground Truth must be able to perform the following Amazon S3 actions on the S3 buckets you use to configure the labeling job.

```
"s3:AbortMultipartUpload",
"s3:GetObject",
"s3:PutObject",
"s3:ListBucket",
"s3:GetBucketLocation"
```

You can do this by adding a Ground Truth endpoint to the bucket policy like the one previously mentioned. The following table includes Ground Truth service endpoints for each AWS Region. Add an endpoint in the same **AWS Region** you use to run your labeling job to your bucket policy.

<table>
<thead>
<tr>
<th>AWS Region</th>
<th>Ground Truth endpoint</th>
</tr>
</thead>
<tbody>
<tr>
<td>us-east-2</td>
<td>vpce-02569ba1c40aad0bc</td>
</tr>
<tr>
<td>us-east-1</td>
<td>vpce-08408e335ebf95b40</td>
</tr>
<tr>
<td>us-west-2</td>
<td>vpce-0ea07aa498eb78469</td>
</tr>
<tr>
<td>ca-central-1</td>
<td>vpce-0d46ea4c9ff55e1b7</td>
</tr>
<tr>
<td>eu-central-1</td>
<td>vpce-0865e7194a099183d</td>
</tr>
</tbody>
</table>
For example, the following policy restricts GetObject and PutObject actions on:

- An Amazon S3 bucket to users in a VPC (<vpc>)
- A VPC endpoint (<vpc-endpoint>)
- A Ground Truth service endpoint (<ground-truth-endpoint>)

```json
{
    "Version": "2012-10-17",
    "Id": "1",
    "Statement": [
        {
            "Sid": "DenyAccessFromNonGTandCustomerVPC",
            "Effect": "Deny",
            "Principal": "*",
            "Action": [
                "s3:GetObject",
                "s3:PutObject"
            ],
            "Resource": [
                "arn:aws:s3:::<bucket-name>",
                "arn:aws:s3:::<bucket-name>/*
            ],
            "Condition": {
                "ForAllValues:StringNotEquals": {
```
If you want a user to have permission to launch a labeling job using the Ground Truth console, you must also add the user's ARN to the bucket policy using the \texttt{aws:PrincipalArn} condition. This user must also have permission to perform the following Amazon S3 actions on the bucket you use to launch the labeling job.

\begin{verbatim}
"s3:GetObject",
"s3:PutObject",
"s3:ListBucket",
"s3:GetBucketCors",
"s3:PutBucketCors",
"s3:ListAllMyBuckets",
\end{verbatim}

The following code is an example of a bucket policy that restricts permission to perform the actions listed in \texttt{Action} on the S3 bucket \texttt{<bucket-name>} to the following.

- \texttt{<role-name>}
- The VPC endpoints listed in \texttt{aws:sourceVpce}
- Users within the VPC named \texttt{<vpc>}

\begin{verbatim}
{
    "Version": "2012-10-17",
    "Id": "1",
    "Statement": [
        {
            "Sid": "DenyAccessFromNonGTandCustomerVPC",
            "Effect": "Deny",
            "Principal": "*",
            "Action": ["s3:GetObject",
                        "s3:PutObject",
                        "s3:ListBucket",
                        "s3:GetBucketCors",
                        "s3:PutBucketCors",
                        "s3:ListAllMyBuckets"]
        }
    ]
}
\end{verbatim}
"s3:PutObject",
  "Resource": [
    "arn:aws:s3::<bucket-name>/*",
    "arn:aws:s3::<bucket-name>"
  ],
  "Condition": {
    "ForAllValues:StringNotEquals": {
      "aws:sourceVpce": [
        "<vpc-endpoint>",
        "<ground-truth-endpoint>"
      ],
      "aws:PrincipalArn": "arn:aws:iam::<aws-account-id>:role/<role-name>"
    }"},
  "aws:SourceVpc": "<vpc>"}
]
}

⚠️ Note

The Amazon VPC interface endpoints and the protected Amazon S3 buckets you use for input and output data must be located in the same AWS Region that you use to create the labeling job.

After you have granted Ground Truth permission to access your Amazon S3 buckets, you can use one of the topics in Create a Labeling Job to launch a labeling job. Specify the VPC-restricted Amazon S3 buckets for your input and output data buckets.

Create an Automated Data Labeling Job in a VPC

To create an automated data labeling job using an Amazon VPC, you provide a VPC configuration using the Ground Truth console or CreateLabelingJob API operation. SageMaker uses the subnets and security groups you provide to launch the training and inferences jobs used for automated labeling.
## Important

Before you launch an automated data labeling job with a VPC configuration, make sure you have created an Amazon S3 VPC endpoint using the VPC you want to use for the labeling job. To learn how, see [Create an Amazon S3 VPC Endpoint](#).

Additionally, if you create an automated data labeling job using a VPC-restricted Amazon S3 bucket, you must follow the instructions in [Allow Ground Truth to Access VPC Restricted Amazon S3 Buckets](#) to give Ground Truth permission to access the bucket.

Use the following procedures to learn how to add a VPC configuration to your labeling job request.

### Add a VPC configuration to an automated data labeling job (console):

1. Follow the instructions in [Create a Labeling Job (console)](#) and complete each step in the procedure, up to step 15.

2. In the **Workers** section, select the checkbox next to **Enable automated data labeling**.

3. Maximize the **VPC configuration** section of the console by selecting the arrow.

4. Specify the **Virtual private cloud (VPC)** that you want to use for your automated data labeling job.

5. Choose the dropdown list under **Subnets** and select one or more subnets.

6. Choose the dropdown list under **Security groups** and select one or more groups.

7. Complete all remaining steps of the procedure in [Create a Labeling Job (Console)](#).

### Add a VPC configuration to an automated data labeling job (API):

To configure a labeling job using the Ground Truth API operation, `CreateLabelingJob`, follow the instructions in [Create an Automated Data Labeling Job (API)](#) to configure your request. In addition to the parameters described in this documentation, you must include a `VpcConfig` parameter in `LabelingJobResourceConfig` to specify one or more subnets and security groups using the following schema.

```json
"LabelingJobAlgorithmsConfig": {  
  "InitialActiveLearningModelArn": "string",  
  "LabelingJobAlgorithmSpecificationArn": "string",  
  "LabelingJobResourceConfig": {  
```

---

**Security and Permissions**

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1786
The following is an example of an AWS Python SDK (Boto3) request to create an automated data labeling job in the US East (N. Virginia) Region using a private workforce. Replace all red-italicized text with your labeling job resources and specifications. To learn more about the CreateLabelingJob operation, see the Create a Labeling Job (API) tutorial and CreateLabelingJob API documentation.

```python
import boto3
client = boto3.client(service_name='sagemaker')

response = client.create_labeling_job(
    LabelingJobName="example-labeling-job",
    LabelAttributeName="label",
    InputConfig={
        'DataSource': {
            'S3DataSource': {
                'ManifestS3Uri': "s3://bucket/path/manifest-with-input-data.json"
            }
        }
    },
    "LabelingJobAlgorithmsConfig": {
        "LabelingJobResourceConfig": {
            "VpcConfig": {
                "SecurityGroupIds": [ "sg-0123456789", "sg-987654321" ],
                "Subnets": [ "subnet-e0123456", "subnet-e7891011" ]
            }
        }
    },
    OutputConfig={
        'S3OutputPath': "s3://bucket/path/file-to-store-output-data",
        'KmsKeyId': "string"
    },
    RoleArn="arn:aws:iam::*:role/*",
    LabelCategoryConfigS3Uri="s3://bucket/path/label-categories.json",
)```
Use Amazon VPC Mode from a Private Worker Portal

To restrict worker portal access to labelers working inside of your Amazon VPC, you can add a VPC configuration when you create a Ground Truth private workforce. You can also add a VPC configuration to an existing private workforce. Ground Truth automatically creates VPC interface endpoints in your VPC and sets up AWS PrivateLink between your VPC endpoint and the Ground Truth services. The worker portal URL associated with the workforce can be accessed from your VPC. The worker portal URL can also be accessed from public internet until you set the restriction
on the public internet. When you delete the workforce or remove the VPC configuration from your workforce, Ground Truth automatically deletes the VPC endpoints associated with the workforce.

Note
There can be only one VPC supported for a workforce.

Point Cloud and video tasks do not support loading through a VPC.

The guide demonstrates how to complete the necessary steps to add and delete an Amazon VPC configuration to your workforce, and satisfy the prerequisites.

Prerequisites

To run a Ground Truth labeling job in Amazon VPC, review the following prerequisites.

- You have an Amazon VPC configured that you can use. If you have not configured a VPC, follow these instructions for creating a VPC.
- Depending on how a Worker Task Template is written, labeling data stored in an Amazon S3 bucket may be accessed directly from Amazon S3 during labeling tasks. In these cases, the VPC network must be configured to allow traffic from the device used by the human labeler to the S3 bucket containing labeling data.
- Follow View and update DNS attributes for your VPC to enable DNS hostnames and DNS resolution for your VPC.

Note
There are two ways to configure your VPC for your workforce. You can do this through the console or the AWS SageMaker CLI.

Using the SageMaker console to manage a VPC config

You can use the SageMaker console to add or remove a VPC configuration. You can also delete an existing workforce.
Adding a VPC configuration to your workforce

Create a private workforce

- [Create a private workforce using Amazon Cognito](#)
- [Create a private workforce using OpenID Connect (OIDC) Identity Provider(IdP)](#)

After you have created your private workforce, add a VPC configuration to it.

1. Navigate to [Amazon SageMaker Runtime](#) in your console.
2. Select **Labeling workforces** in the left panel.
3. Select **Private** to access your private workforce. After your **Workforce status** is **Active**, select **Add** next to **VPC**.
4. When you are prompted to configure your VPC, provide the following:
   a. Your **VPC**
   b. **Subnets**
      i. Ensure that your VPC has an existing subnet
   c. **Security groups**
      i. **Note**
         You cannot select more than 5 security groups.
   d. After filling in this information, choose **Confirm**.
5. After you choose **Confirm**, you are redirected back to the **Private** page under **Labeling workforces**. You should see a green banner at the top that reads **Your private workforce update with VPC configuration was successfully initialized**. The workforce status is **Updating**. Next to the **Delete workforce** button is the **Refresh** button, which can be used to retrieve the latest **Workforce status**. After the workforce status has changed to **Active**, the VPC endpoint ID is updated as well.

Removing a VPC configuration from your workforce

Use the following information to remove a VPC configuration from your workforce using the console.

1. Navigate to [Amazon SageMaker Runtime](#) in your console.
2. Select **Labeling workforces** in the left panel.

3. Find and select your workforce.

4. Under **Private workforce summary**, find **VPC** and choose **Remove** next to it.

5. Select **Remove**.

**Deleting a workforce through the console**

If you delete a workforce, you should not have any teams associated with it. You can delete a workforce only if the workforce status is **Active** or **Failed**.

Use the following information to delete a workforce using the console.

1. Navigate to **Amazon SageMaker Runtime** in your console.

2. Select **Labeling workforces** in the left panel.

3. Find and select your workforce.

4. Choose **Delete workforce**.

5. Choose **Delete**.

**Using the SageMaker AWS API to manage a VPC config**

Download the following files to use a new **VpcConfig** parameter into the SageMaker workforce CLI:

**sagemaker-2017-07-24.normal.json**

**sagemaker-2017-07-24.paginators.json**

**sagemaker-2017-07-24.waiters-2.json**

After downloading the files, run the following commands in your CLI:

```
aws configure add-model --service-model file://./sagemaker-2017-07-24.normal.json --service-name sagemaker

```
You can now test your API changes using AWS CLI. You can either create a new workforce with a VPC configuration or update an existing workforce to add a VPC configuration. You can also remove a VPC configuration from an existing workforce.

**Create a workforce with a VPC configuration**

If the account already has a workforce, then you must delete it first. You can also update the workforce with VPC configuration.

```bash
```

```
aws sagemaker create-workforce --cognito-config '{"ClientId": "app-client-id","UserPool": "Pool_ID"}', --workforce-vpc-config "{"VpcId": "vpc-id", "SecurityGroupIds": ["sg-0123456789abcdef0"], "Subnets": ["subnet-0123456789abcdef0"]}" --workforce-name workforce-name
```

Describe the workforce and make sure the status is Initializing.

```
aws sagemaker describe-workforce --workforce-name workforce-name
```

```json
{
    "Workforce": {
        "WorkforceName": "workforce-name",
        "LastUpdatedDate": 1622151252.451,
        "SourceIpConfig": {
            "Cidrs": []
        },
        "SubDomain": "subdomain.us-west-2.sagemaker.aws.com",
        "CognitoConfig": {
            "UserPool": "Pool_ID",
            "ClientId": "app-client-id"
        },
        "CreateDate": 1622151252.451,
        "WorkforceVpcConfig": {
```
Navigate to the Amazon VPC console. Select **Endpoints** from the left panel. There should be two VPC endpoints created in your account.

**Adding a VPC configuration your workforce**

Update a non-VPC private workforce with a VPC configuration using the following command.

```bash
aws sagemaker update-workforce --workforce-name workforce-name
--workforce-vpc-config "{"VpcId": "vpc-id", "SecurityGroupIds": ["sg-0123456789abcdef0"], "Subnets": ["subnet-0123456789abcdef0"]}"
```

Describe the workforce and make sure the status is **Updating**.

```bash
aws sagemaker describe-workforce --workforce-name workforce-name
{
   "Workforce": {
      "WorkforceName": "workforce-name",
      "LastUpdatedDate": 1622151252.451,
      "SourceIpConfig": {
         "Cidrs": []
      },
      "SubDomain": "subdomain.us-west-2.sagemaker.aws.com",
      "CognitoConfig": {
         "UserPool": "Pool_ID",
         "ClientId": "app-client-id"
      }
   }
}
```
Navigate to your Amazon VPC console. Select **Endpoints** from the left panel. There should be two VPC endpoints created in your account.

**Removing a VPC configuration from your workforce**

Update a VPC private workforce with an empty VPC configuration to remove VPC resources.

```
aws sagemaker update-workforce --workforce-name \n--workforce-vpc-config "{}"
```

Describe the workforce and make sure the status is **Updating**.

```
aws sagemaker describe-workforce --workforce-name \nworkforce-name
{
  "Workforce": {
    "WorkforceName": "workforce-name",
    "LastUpdatedDate": 1622151252.451,
    "SourceIpConfig": {
      "Cidrs": []
    },
    "SubDomain": "subdomain.us-west-2.sagemaker.aws.com",
    "CognitoConfig": {
```
Navigate to your Amazon VPC console. Select **Endpoints** from the left panel. The two VPC endpoints should be deleted.

**Restrict public access to the worker portal while maintaining access through a VPC**

The workers in a VPC or non-VPC worker portal are be able to see the labeling job tasks assigned to them. The assignment comes from assigning workers in a work team through OIDC groups. It is the customer's responsibility to restrict the access to their public worker portal by setting the `sourceIpConfig` in their workforce.

⚠️ **Note**

You can restrict access to the worker portal only through the SageMaker API. This cannot be done through the console.

Use the following command to restrict public access to the worker portal.

```bash
aws sagemaker update-workforce --region us-west-2 \
--workforce-name workforce-demo --source-ip-config '{"Cidrs":["10.0.0.0/16"]}'
```

After the `sourceIpConfig` is set on the workforce, the workers can access the worker portal in VPC but not through public internet.

⚠️ **Note**

You can not set the `sourceIP` restriction for worker portal in VPC.
Output Data and Storage Volume Encryption

With Amazon SageMaker Ground Truth, you can label highly sensitive data, stay in control of your data, and employ security best practices. While your labeling job is running, Ground Truth encrypts data in transit and at rest. Additionally, you can use AWS Key Management Service (AWS KMS) with Ground Truth to do the following:

- Use a [customer managed key](#) to encrypt your output data.
- Use AWS KMS customer managed key with your automated data labeling job to encrypt the storage volume attached to the compute instance used for model training and inference.

Use the topics on this page to learn more about these Ground Truth security features.

Use Your KMS Key to Encrypt Output Data

Optionally, you can provide an AWS KMS customer managed key when you create a labeling job, which Ground Truth uses to encrypt your output data.

If you don't provide a customer managed key, Amazon SageMaker uses the default AWS managed key for Amazon S3 for your role's account to encrypt your output data.

If you provide a customer managed key, you must add the required permissions to the key described in [Encrypt Output Data and Storage Volume with AWS KMS](#). When you use the API operation CreateLabelingJob, you can specify your customer managed key ID using the parameter `KmsKeyId`. See the following procedure to learn how to add a customer managed key when you create a labeling job using the console.

To add an AWS KMS key to encrypt output data (console):

1. Complete the first 7 steps in [Create a Labeling Job (Console)](#).
2. In step 8, select the arrow next to **Additional configuration** to expand this section.
3. For **Encryption key**, select the AWS KMS key that you want to use to encrypt output data.
4. Complete the rest of steps in [Create a Labeling Job (Console)](#) to create a labeling job.

Use Your KMS Key to Encrypt Automated Data Labeling Storage Volume (API Only)

When you create a labeling job with automated data labeling using the CreateLabelingJob API operation, you have the option to encrypt the storage volume attached to the ML compute instances that run the training and inference jobs. To add encryption to your storage volume,
use the parameter `VolumeKmsKeyId` to input an AWS KMS customer managed key. For more information about this parameter, see `LabelingJobResourceConfig`.

If you specify a key ID or ARN for `VolumeKmsKeyId`, your SageMaker execution role must include permissions to call `kms:CreateGrant`. To learn how to add this permission to an execution role, see `Create a SageMaker Execution Role for a Ground Truth Labeling Job`.

**Note**

If you specify an AWS KMS customer managed key when you create a labeling job in the console, that key is *only* used to encrypt your output data. It is not used to encrypt the storage volume attached to the ML compute instances used for automated data labeling.

---

**Workforce Authentication and Restrictions**

Ground Truth enables you to use your own private workforce to work on labeling jobs. A *private workforce* is an abstract concept which refers to a set of people who work for you. Each labeling job is created using a work team, composed of workers in your workforce. Ground Truth supports private workforce creation using Amazon Cognito.

A Ground Truth workforce maps to a Amazon Cognito user pool. A Ground Truth work team maps to a Amazon Cognito user group. Amazon Cognito manages the worker authentication. Amazon Cognito supports Open ID connection (OIDC) and customers can set up Amazon Cognito federation with their own identity provider (IdP).

Ground Truth only allows one workforce per account per AWS Region. Each workforce has a dedicated Ground Truth work portal login URL.

You can also restrict workers to a Classless Inter-Domain Routing (CIDR) block/IP address range. This means annotators must be on a specific network to access the annotation site. You can add up to ten CIDR blocks for one workforce. To learn more, see `Manage Private Workforce Using the Amazon SageMaker API`.

To learn how you can create a private workforce, see `Create a Private Workforce (Amazon Cognito)`.

**Restrict Access to Workforce Types**

Amazon SageMaker Ground Truth work teams fall into one of three *workforce types*: public (with Amazon Mechanical Turk), private, and vendor. To restrict user access to a specific work team using one of these types or the work team ARN, use the `sagemaker:WorkteamType` and/or the
sagemaker:WorkteamArn condition keys. For the sagemaker:WorkteamType condition key, use string condition operators. For the sagemaker:WorkteamArn condition key, use Amazon Resource Name (ARN) condition operators. If the user attempts to create a labeling job with a restricted work team, SageMaker returns an access denied error.

The policies below demonstrate different ways to use the sagemaker:WorkteamType and sagemaker:WorkteamArn condition keys with appropriate condition operators and valid condition values.

The following example uses the sagemaker:WorkteamType condition key with the StringEquals condition operator to restrict access to a public work team. It accepts condition values in the following format: workforcetype-crowd, where workforcetype can equal public, private, or vendor.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "RestrictWorkteamType",
         "Effect": "Deny",
         "Action": "sagemaker:CreateLabelingJob",
         "Resource": "*",
         "Condition": {
            "StringEquals": {
               "sagemaker:WorkteamType": "public-crowd"
            }
         }
      }
   ]
}
```

The following policies show how to restrict access to a public work team using the sagemaker:WorkteamArn condition key. The first shows how to use it with a valid IAM regex-variant of the work team ARN and the ArnLike condition operator. The second shows how to use it with the ArnEquals condition operator and the work team ARN.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "RestrictWorkteamType",
         "Effect": "Deny",
         "Action": "sagemaker:CreateLabelingJob",
         "Resource": "*",
         "Condition": {
            "ArnLike": {
            }
         }
      }
   ]
}
```
Monitor Labeling Job Status

To monitor the status of your labeling jobs, you can set up an Amazon CloudWatch Events (CloudWatch Events) rule for Amazon SageMaker Ground Truth (Ground Truth) to send an event to CloudWatch Events when a labeling job status changes to Completed, Failed, or Stopped or when a worker accepts, declines, submits, or returns a task.

Once you create a rule, you can add a target to it. CloudWatch Events uses this target to invoke another AWS service to process the event. For example, you can create a target using a Amazon Simple Notification Service (Amazon SNS) topic to send a notification to your email when a labeling job status changes.
Prerequisites:

To create a CloudWatch Events rule, you will need an AWS Identity and Access Management (IAM) role with an events.amazonaws.com trust policy attached. The following is an example of an events.amazonaws.com trust policy.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "",
            "Effect": "Allow",
            "Principal": {
                "Service": ["events.amazonaws.com"
            ],
            "Action": "sts:AssumeRole"
        }
    ]
}
```

Topics

- Send Events to CloudWatch Events
- Set Up a Target to Process Events
- Labeling Job Expiration
- Declining Tasks

Send Events to CloudWatch Events

To configure a CloudWatch Events rule to get status updates, or events, for your Ground Truth labeling jobs, use the AWS Command Line Interface (AWS CLI) `put-rule` command. You can filter events that are sent to your rule by status change. For example, you can create a rule that notifies you only if a labeling job status changes to Completed. When using the `put-rule` command, specify the following to receive labeling job statuses:

- "source": "aws.sagemaker"
- "detail-type": "SageMaker Ground Truth Labeling Job State Change"
To configure a CloudWatch Events rule to watch for all status changes, use the following command and replace the placeholder text. For example, replace "GTLabelingJobStateChanges" with a unique CloudWatch Events rule name and "arn:aws:iam::111122223333:role/MyRoleForThisRule" with the Amazon Resource Number (ARN) of an IAM role with an events.amazonaws.com trust policy attached.

```
aws events put-rule --name "GTLabelingJobStateChanges"
    --event-pattern "{"source":["aws.sagemaker"],"detail-type":["SageMaker Ground Truth Labeling Job State Change"]}
    --role-arn "arn:aws:iam::111122223333:role/MyRoleForThisRule"
    --region "region"
```

To filter by job status, use the "detail":{"LabelingJobStatus":["Status"]} syntax. Valid values for Status are Completed, Failed, and Stopped.

The following example creates a CloudWatch Events rule that notifies you when a labeling job in us-west-2 (Oregon) changes to Completed.

```
aws events put-rule --name "LabelingJobCompleted"
    --event-pattern "{"source":["aws.sagemaker"],"detail-type":["SageMaker Ground Truth Labeling Job State Change"], "detail":{"LabelingJobStatus": ["Completed"]}
    --role-arn "arn:aws:iam::111122223333:role/MyRoleForThisRule"
    --region us-west-2
```

The following example creates a CloudWatch Events rule that notifies you when a labeling job in us-east-1 (Virginia) changes to Completed or Failed.

```
aws events put-rule --name "LabelingJobCompletedOrFailed"
    --event-pattern "{"source":["aws.sagemaker"],"detail-type":["SageMaker Ground Truth Labeling Job State Change"], "detail":{"LabelingJobStatus": ["Completed", "Failed"]}
    --role-arn "arn:aws:iam::111122223333:role/MyRoleForThisRule"
    --region us-east-1
```

To learn more about the put-rule request, see Event Patterns in CloudWatch Events in the Amazon CloudWatch Events User Guide.
Set Up a Target to Process Events

After you have created a rule, events similar to the following are sent to CloudWatch Events. In this example, the labeling job test-labeling-job's status changed to Completed.

```
{
    "version": "0",
    "id": "11e1111-11d1-11f-b111-1111b11dcb11",
    "detail-type": "SageMaker Ground Truth Labeling Job State Change",
    "source": "aws.sagemaker",
    "account": "111122223333",
    "time": "2018-10-06T12:26:13Z",
    "region": "us-east-1",
    "resources": [
    ],
    "detail": {
        "LabelingJobStatus": "Completed"
    }
}
```

To process events, you need to set up a target. For example, if you want to receive an email when your labeling job status changes, use a procedure in Setting Up Amazon SNS Notifications in the Amazon CloudWatch User Guide to set up an Amazon SNS topic and subscribe your email to it. Once you have create a topic, you can use it to create a target.

**To add a target to your CloudWatch Events rule**

1. Open the CloudWatch console: [https://console.aws.amazon.com/cloudwatch/home](https://console.aws.amazon.com/cloudwatch/home)
2. In the navigation pane, choose Rules.
3. Choose the rule that you want to add a target to.
4. Choose Actions, and then choose Edit.
5. Under Targets, choose Add Target and choose the AWS service you want to act when a labeling job status change event is detected.
6. Configure your target. For instructions, see the topic for configuring a target in the AWS documentation for that service.
7. Choose Configure details.
8. For Name, enter a name and, optionally, provide details about the purpose of the rule in Description.
9. Make sure that the check box next to **State** is selected so that your rule is listed as **Enabled**.

10. Choose **Update rule**.

**Labeling Job Expiration**

If your labeling job is not completed after 30 days, it will expire. If your labeling job expires, you can chain the job to create a new labeling job that will only send unlabeled data to workers. For more information, and to learn how to create a labeling job using chaining, see [Chaining Labeling Jobs](#).

**Declining Tasks**

Workers are able to decline tasks.

Workers decline a task if the instructions are not clear, input data is not displaying correctly, or if they encounter some other issue with the task. If the number of workers per dataset object (NumberofHumanWorkersPerDataObject) decline the task, the data object is marked as expired and will not be sent to additional workers.

**Use Amazon SageMaker Ground Truth Plus to Label Data**

Amazon SageMaker Ground Truth Plus is a turnkey data labeling service that uses an expert workforce to deliver high-quality annotations quickly and reduces costs by up to 40%. Using SageMaker Ground Truth Plus, data scientists and business managers, such as data operations managers and program managers, can create high-quality training datasets without having to build labeling applications and manage labeling workforces on their own. You can get started with Amazon SageMaker Ground Truth Plus by uploading data along with the labeling requirements in Amazon S3.

**Why use SageMaker Ground Truth Plus?**

To train a machine learning (ML) model, data scientists need large, high-quality, labeled datasets. As ML adoption grows, labeling needs increase. This forces data scientists to spend weeks on building data labeling workflows and managing a data labeling workforce. Unfortunately, this slows down innovation and increases cost. To ensure data scientists can spend their time building, training, and deploying ML models, data scientists typically task other in-house teams consisting of data operations managers and program managers to produce high-quality training datasets.
However, these teams typically don’t have access to skills required to deliver high-quality training datasets, which affects ML results. As a result, you look for a data labeling partner that can help them create high-quality training datasets at scale without consuming their in-house resources.

When you upload the data, SageMaker Ground Truth Plus sets up the data labeling workflows and operates them on your behalf. From there, an expert workforce trained on a variety of machine learning (ML) tasks performs data labeling. SageMaker Ground Truth Plus currently offers two types of expert workforce: an Amazon employed workforce and a curated list of third-party vendors. SageMaker Ground Truth Plus provides you with the flexibility to choose the labeling workforce. AWS experts select the best labeling workforce based on your project requirements. For example, if you need people proficient in labeling audio files, specify that in the guidelines provided to SageMaker Ground Truth Plus, and the service automatically selects labelers with those skills.

⚠️ **Important**

SageMaker Ground Truth Plus does not support PHI, PCI or FedRAMP certified data, and you should not provide this data to SageMaker Ground Truth Plus.

**How does SageMaker Ground Truth Plus work?**

There are five main components to a workflow.

- Requesting a project
- Creating a project team
- Accessing the project portal to monitor progress of training datasets and review labeled data
- Creating a batch
- Receiving the labeled data

**How do I use SageMaker Ground Truth Plus?**

If you are a first-time user of SageMaker Ground Truth Plus, use [Getting Started with Amazon SageMaker Ground Truth Plus](#) to get started. To access SageMaker Ground Truth Plus using the SageMaker console, you must be in US East (N. Virginia) (us-east-1).
Getting Started with Amazon SageMaker Ground Truth Plus.

The guide demonstrates how to complete the necessary steps to start an Amazon SageMaker Ground Truth Plus project, review labels, and satisfy SageMaker Ground Truth Plus prerequisites.

To get started using SageMaker Ground Truth Plus, review [Set Up Amazon SageMaker Ground Truth Plus Prerequisites](#) and [Core Components of Amazon SageMaker Ground Truth Plus](#).

Set Up Amazon SageMaker Ground Truth Plus Prerequisites

Use the following information to sign up for an AWS account. If you already have an AWS account, skip this step.

Sign up for an AWS account

If you do not have an AWS account, complete the following steps to create one.

To sign up for an AWS account

2. Follow the online instructions.

   Part of the sign-up procedure involves receiving a phone call and entering a verification code on the phone keypad.

   When you sign up for an AWS account, an *AWS account root user* is created. The root user has access to all AWS services and resources in the account. As a security best practice, assign administrative access to an administrative user, and use only the root user to perform tasks that require root user access.

AWS sends you a confirmation email after the sign-up process is complete. At any time, you can view your current account activity and manage your account by going to [https://aws.amazon.com/](https://aws.amazon.com/) and choosing My Account.

Create an administrative user

After you sign up for an AWS account, secure your AWS account root user, enable AWS IAM Identity Center, and create an administrative user so that you don't use the root user for everyday tasks.
Secure your AWS account root user

1. Sign in to the AWS Management Console as the account owner by choosing Root user and entering your AWS account email address. On the next page, enter your password.

   For help signing in by using root user, see Signing in as the root user in the AWS Sign-In User Guide.

2. Turn on multi-factor authentication (MFA) for your root user.

   For instructions, see Enable a virtual MFA device for your AWS account root user (console) in the IAM User Guide.

Create an administrative user

1. Enable IAM Identity Center.

   For instructions, see Enabling AWS IAM Identity Center in the AWS IAM Identity Center User Guide.

2. In IAM Identity Center, grant administrative access to an administrative user.

   For a tutorial about using the IAM Identity Center directory as your identity source, see Configure user access with the default IAM Identity Center directory in the AWS IAM Identity Center User Guide.

Sign in as the administrative user

- To sign in with your IAM Identity Center user, use the sign-in URL that was sent to your email address when you created the IAM Identity Center user.

   For help signing in using an IAM Identity Center user, see Signing in to the AWS access portal in the AWS Sign-In User Guide.

Core Components of Amazon SageMaker Ground Truth Plus

The following terms are key to understanding the capabilities of SageMaker Ground Truth Plus:

- **Project**: Each qualified engagement with an AWS expert results in a SageMaker Ground Truth Plus project. A project can be in the pilot or production stage.
• **Batch**: A batch is a collection of similar recurring data objects such as images, video frames and text to be labeled. A project can have multiple batches.

• **Metrics**: Metrics are data about your SageMaker Ground Truth Plus project for a specific date or over a date range.

• **Task type**: SageMaker Ground Truth Plus supports five task types for data labeling. You can also have a custom task type. These include text, image, video, audio, and 3D point cloud.

• **Data objects**: Individual items that are to be labeled.

---

## Request a Project

To use Amazon SageMaker Ground Truth Plus, get started by requesting a project.

1. Under the Ground Truth tab of Amazon SageMaker, choose **Plus**.
2. On the **SageMaker Ground Truth Plus** page, choose **Request project**.
3. A page titled **Request a project** opens. The page includes fields for **General information** and **Project overview**. Enter the following information

   a. Under **General information**, enter your **First name**, **Last name** and **Business email address**. An AWS expert uses this information for contacting you to discuss the project after you submit the request.

   b. Under **Project overview**, enter your **Project name** and **Project description**. Choose the **Task type** based on your data and use case. You can also indicate if your data contains personally identifiable information (PII).

   c. Create or select an IAM role that grants SageMaker Ground Truth Plus permissions to perform a labeling job by choosing one of the options below.

      i. You can **Create an IAM role** that provides access to any S3 bucket you specify.

      ii. You can **Enter a custom IAM role ARN**.

      iii. You can choose an existing role.

      iv. If you use an existing role or a custom IAM role ARN, make sure you have the following IAM role and trust policy.

         ```json
         {
           "Version": "2012-10-17",
         }
         ```
"Statement": [ 
    { 
        "Effect": "Allow",
        "Action": [ 
            "s3:GetObject",
            "s3:GetBucketLocation",
            "s3:ListBucket",
            "s3:PutObject"
        ],
        "Resource": [ 
            "arn:aws:s3:::your-bucket-name",
            "arn:aws:s3:::your-bucket-name/**
        ]
    }
]

Trust policy

{
    "Version": "2012-10-17",
    "Statement": [ 
        { 
            "Effect": "Allow",
            "Principal": { 
                "Service": "sagemaker-ground-truth-plus.amazonaws.com"
            },
            "Action": "sts:AssumeRole"
        }
    ]
}

4. Choose **Request a project**.

Once you create a project, you can find it on the **SageMaker Ground Truth Plus** page, under the Projects section. The project status should be **Review in-progress**.

⚠️ **Note**

You cannot have more than 5 projects with the **Review in progress** status.
Create a Project Team

A project team provides access to the members from your organization or team to track projects, view metrics, and review annotations. You can create a SageMaker Ground Truth Plus project team once you have shared your data in an Amazon S3 bucket.

To add team members using Amazon Cognito, you have two options:

1. Create a new Amazon Cognito user group
   a. Enter an Amazon Cognito user group name. This name cannot be changed.
   b. Enter the email addresses of up to 50 team members in the Email addresses field. The addresses must be separated by a comma.
   c. Choose Create project team.
Your team members receive an email inviting them to join the SageMaker Ground Truth Plus project team as shown in the following image.
2. Import team members from existing Amazon Cognito user groups.
   a. Choose a user pool that you have created. User pools require a domain and an existing user group. If you get an error that the domain is missing, set it in the Domain name options on the App integration page of the Amazon Cognito console for your group.
   b. Choose an app client. We recommend using a client generated by Amazon SageMaker.
   c. Choose a user group from your pool to import its members.
   d. Choose Create project team.

You can view and manage the list of team members through the AWS console.

To add team members after creating the project team:

1. Choose Invite new members in the Members section.
2. Enter the email addresses of up to 50 team members in the Email addresses field. The addresses must be separated by a comma.

3. Choose **Invite new members**

**To delete existing team members:**

1. Choose the team member to be deleted in the Members section.
2. Choose **Delete**.

Once you have added members to your project team, you can open the project portal to access your projects.

**Open the Project Portal**

Once you have successfully submitted the intake form and created a project team, you can access the SageMaker Ground Truth Plus project by choosing the **Open project portal** on the AWS console.

Each project consists of one or more batches. A *batch* is a collection of recurring similar data objects (text, image, video frame, and point cloud) to be labeled. The project portal provides you with transparency into the data labeling process. You can stay updated about a project, create batches within a project, review the progress of the datasets across multiple projects, and analyze project metrics. The project portal also allows you to review a subset of the labeled data and provide feedback. You can configure the columns displayed in your project and batch table.

You can use the SageMaker Ground Truth Plus project portal to track the following details about your project.

**Project name:** Each project is identified using a unique name.

**Status:** A SageMaker Ground Truth Plus project has one of the following status types:
1. **Review in progress**: You have successfully submitted the project request form. An AWS expert is currently reviewing your request.

2. **Request approved**: Your project request is approved. You can now share your data by creating a new batch from the project portal.

3. **Workflow design and setup progress**: An AWS expert is setting up your project.

4. **Pilot in-progress**: Object labeling for the project in the pilot stage is currently in progress.

5. **Pilot complete**: Object labeling is complete and the labeled data is stored in your Amazon S3 bucket.

6. **Pricing complete**: An AWS expert shares the pricing for the production project with you.

7. **Contract executed**: The contract is complete.

8. **Production in-progress**: Labeling for the project in the production stage is in progress.

9. **Production complete**: Object labeling is complete and the labeled data is stored in your Amazon S3 bucket.

10. **Paused**: Project is currently paused at your request.

**Task type**: SageMaker Ground Truth Plus lets you label five types of tasks that include text, image, video, audio, and point cloud.

**Batches**: Total number of batches within a project.

**Project creation date**: Starting date of a project.

**Total objects**: Total number of objects to be labeled across all batches.

**Objects completed**: Number of labeled objects.

**Remaining objects**: Number of objects left to be labeled.

**Failed objects**: Number of objects that cannot be labeled due to an issue with the input data.

### Create a Batch

You can use the project portal to create batches for a project after the project status is changed to **Request approved**.
Create a batch

A batch is a collection of similar recurring data objects such as images, video frames and text to be labeled. A project can have multiple batches. Create a batch by following the steps below.

Basic Information

**Batch name**
Enter the name of your batch.

**Batch description - optional**
Provide a brief description of the batch.

**Data setup**

*S3 location for input datasets info*
This is the location in S3 where your dataset objects are stored. Ground Truth Plus will use all data objects in this location for your labeling job.

*S3 location for output datasets info*
This is the location in S3 where your labeling job output data is stored.

To create a batch, do the following.

1. Select a project by choosing the project name.
2. A page titled with the project name opens. Under the Batches section, choose Create batch.
3. Enter the Batch name, Batch description, S3 location for input datasets, and S3 location for output datasets.
4. Choose Submit.
To create a batch successfully, make sure you meet the following criteria:

- Your data is in the US East (N. Virginia) Region.
- The maximum size for each file is no more than 2 gigabytes.
- The maximum number of files in a batch is 10,000.
- The total size of a batch is less than 100 gigabytes.
- You have no more than 5 batches with the Data transfer in-progress status.

Note
You cannot create a batch before the project status changes to Request approved.

Review Metrics

Metrics are data about your SageMaker Ground Truth Plus project for a specific date or over a date range.

You can review metrics for all batches or choose a batch of your choice as shown in the following image.
You can review the following metrics about the batch:

**Total objects**: Total number of objects in a batch or across all batches.

**Objects completed by day**: Total numbers of objects labeled on a specific date or over a date range.

**Labels completed by day**: Total numbers of labels completed on a specific date or over a date range. An object can have more than one label.
Review Batches

Every Amazon SageMaker Ground Truth Plus project consists of one or more batches. Each batch is made up of data objects to be labeled. You can view all the batches for your project using the project portal as shown in the following image.

You can use the SageMaker Ground Truth Plus project portal to track the following details about every batch:

**Batch name**: Each batch is identified with a unique batch name.

**Status**: A SageMaker Ground Truth Plus batch has one of the following status types:

1. **Request submitted**: You have successfully submitted a new batch.
2. **Data transfer failed**: Data transfer failed with errors. Check the error reason and create a new batch after fixing the error.
3. **Data received**: We have received your unlabeled input data.
4. **In-progress**: Data labeling is in progress.
5. **Ready for review**: Data labeling is completed. A subset of labeled objects from the batch are ready for you to review. This is an optional step.
6. **Review submission in-progress**: Review feedback is currently being processed.
7. **Review complete**: You have successfully reviewed the batch. Next, you have to accept or reject it. This action can not be undone.
8. **Accepted**: You have accepted the labeled data and will receive it in your Amazon S3 bucket shortly.

9. **Rejected**: Labeled data needs to be reworked.

10. **Sent for rework**: Labeled data is sent for rework. You can review the batch after its status changes to **Ready for review**.

11. **Ready for delivery**: Labeled data is ready to be transferred to your Amazon S3 bucket.

12. **Data delivered**: Object labeling is complete and the labeled data is stored in your Amazon S3 bucket.

13. **Paused**: Batch is paused at your request.

**Task type**: SageMaker Ground Truth Plus lets you label five types of tasks that include text, image, video, audio, and point cloud.

**Batch creation date**: Date when the batch was created.

**Total objects**: Total number of objects to be labeled across a batch.

**Completed objects**: Number of labeled objects.

**Remaining objects**: Number of objects left to be labeled.

**Failed objects**: Number of objects that cannot be labeled due to an issue with the input data.

**Objects to review**: Number of objects that are ready for your review.

**Objects with feedback**: Number of objects that have gotten feedback from the team members.

SageMaker Ground Truth Plus lets you review a sample set of your labeled data (determined during the initial consultation call) through the review UI shown in the following image.
The portal allows your project team members and you to review a small sample set of the labeled objects for each batch. You can provide feedback for each labeled object within that subset through this UI. The review UI allows you to navigate across the subset of labeled objects and provide feedback for those labeled objects.

You can perform the following actions using the review UI.

- Use the arrow controls on the bottom left to navigate through the data objects.
- You can provide feedback for each object. The Feedback section is in the right panel. Choose Submit to submit feedback for all images.
- Use the image controls in the bottom tray to zoom, pan, and control contrast.
- If you plan on returning to finish up your review, choose Stop and resume later on the top right.
- Choose Save to save your progress. Your progress is also autosaved every 15 minutes.
- To exit the review UI, choose Close on the upper right corner of the review UI.
- You can verify the Label attributes and Frame attributes on each frame using the panel on the right. You cannot create new objects or modify existing objects in this task.
Accept or Reject Batches

After you have reviewed a batch, you must choose to accept or reject it.

If you accept a batch, the output from that labeling job is placed in the Amazon S3 bucket that you specify. Once the data is delivered to your S3 bucket, the status of your batch changes from Accepted to Data delivered.

If you reject a batch, you can provide feedback and explain your reasons for rejecting the batch.

SageMaker Ground Truth Plus allows you to provide feedback at the data object level as well as the batch level. You can provide feedback for data objects through the review UI. You can use the project portal to provide feedback for each batch. When you reject a batch, an AWS expert contacts you to determine the rework process and the next steps for the batch.

Note

Accepting or rejecting a batch is a one-time action and cannot be undone. It is necessary to either accept or reject every batch of the project.

Create and Manage Workforces

A workforce is the group of workers that you have selected to label your dataset. You can choose either the Amazon Mechanical Turk workforce, a vendor-managed workforce, or you can create your own private workforce to label or review your dataset. Whichever workforce type you choose, Amazon SageMaker takes care of sending tasks to workers.

When you use a private workforce, you also create work teams, a group of workers from your workforce that are assigned to specific jobs—Amazon SageMaker Ground Truth labeling jobs or Amazon Augmented AI human review tasks. You can have multiple work teams and can assign one or more work teams to each job.

You can use Amazon Cognito or your own private OpenID Connect (OIDC) Identity Provider (IdP) to manage your private workforce and work teams. For more information about the permissions required to manage your workforce this way, see Permissions Required to Use the Amazon SageMaker Ground Truth Console.

Topics

- Using the Amazon Mechanical Turk Workforce
• **Managing Vendor Workforces**

• **Use a Private Workforce**

**Using the Amazon Mechanical Turk Workforce**

The Amazon Mechanical Turk (Mechanical Turk) workforce provides the most workers for your Amazon SageMaker Ground Truth labeling job and Amazon Augmented AI human review task. The Amazon Mechanical Turk workforce is a world-wide resource. Workers are available 24 hours a day, 7 days a week. You typically get the fastest turnaround for your human review tasks and labeling jobs when you use the Amazon Mechanical Turk workforce.

Any Amazon Mechanical Turk workforce billing is handled as part of your Ground Truth or Amazon Augmented AI billing. You do not need to create a separate Mechanical Turk account to use the Amazon Mechanical Turk workforce.

⚠️ **Important**

You should not share confidential information, personal information, or protected health information with this workforce. You should not use the Amazon Mechanical Turk workforce when you use Amazon A2I in conjunction with AWS HIPAA-eligible services, such as Amazon Textract and Amazon Rekognition, for workloads containing protected health information.

You can choose Mechanical Turk as your workforce when you create a Ground Truth labeling job or Amazon A2I human review workflow (flow definition). You can create a labeling job and a human review workflow using the SageMaker console and API.

When you use an API operation to create a labeling job or human review workflow, you use the following ARN for the Amazon Mechanical Turk workforce for your WorkteamArn. Replace `region` with the AWS Region you are using to create the labeling job or human loops. For example, if you create a labeling job in US West (Oregon), replace `region` with `us-west-2`.

• arn:aws:sagemaker:`region`:394669845002:workteam/public-crowd/default

Ground Truth and Amazon A2I require that your input data is free of personally identifiable information (PII) when you use Mechanical Turk. If you use the Mechanical Turk workforce and do
not specify that your input data is free of PII, your Ground Truth labeling jobs and Augmented AI tasks will fail. You specify that your input data is free of PII when you create a Ground Truth labeling job and when you create a Amazon A2I human loop using a built-in integration or the StartHumanLoop operation.

Use the following sections to learn how to use Mechanical Turk with these services.

**Topics**
- Use Mechanical Turk with Ground Truth
- Use Mechanical Turk with Amazon A2I
- When is Mechanical Turk Not Supported?

**Use Mechanical Turk with Ground Truth**

You can use Mechanical Turk with Ground Truth when you create a labeling job using the console, or the CreateLabelingJob operation.

When you create a labeling job, we recommend you adjust the number of workers that annotate each data object based on the complexity of the job and the quality that you need. Amazon SageMaker Ground Truth uses annotation consolidation to improve the quality of the labels. More workers can make a difference in the quality of the labels for more complex labeling jobs, but might not make a difference for simpler jobs. For more information, see Consolidate Annotations. Note that annotation consolidation is not supported for Amazon A2I human review workflows.

**To use Mechanical Turk when you create a labeling job (console):**

1. Use the following to create a labeling job using the Ground Truth area of the SageMaker console: Create a Labeling Job (Console).
2. When you are selecting Worker types in the Workers section, select Amazon Mechanical Turk.
3. Specify the total amount of time workers have to complete a task using Task timeout.
4. Specify the total amount of time a task remains available to workers in Task expiration. This is how long workers have to pick up a task before it fails.
5. Select the Price per task using the dropdown list. This is the amount of money a worker receives for completing a single task.
6. (Optional) If applicable, select The dataset does not contain adult content. SageMaker may restrict the Mechanical Turk workers that can view your task if it contains adult content.
7. You must read and confirm the following statement by selecting the check box to use the Mechanical Turk workforce. If your input data contains confidential information, personal information, or protected health information, you must select another workforce.

    **You understand and agree that the Mechanical Turk workforce consists of independent contractors located worldwide and that you should not share confidential information, personal information, or protected health information with this workforce.**

8. (Optional) Select the check box next to **Enable automated data labeling** if you want to enable automated data labeling. To learn more about this feature, see [Automate Data Labeling](#).

9. You can specify the **Number of workers per dataset object** under **Additional configuration**. For example, if you enter 3 in this field, each data object will be labeled by 3 workers.

When you create your labeling job by selecting **Create**, your labeling tasks are sent to Mechanical Turk workers.

**To use Mechanical Turk when you create a labeling job (API):**

1. Use the following to create a labeling job using the `CreateLabelingJob` operation: [Create a Labeling Job (API)](#).

2. Use the following for the `WorkteamArn`. Replace **region** with the AWS Region you are using to create the labeling job.

   ```
   arn:aws:sagemaker:region:394669845002:workteam/public-crowd/default
   ```

3. Use `TaskTimeLimitInSeconds` to specify the total amount of time workers have to complete a task.

4. Use `TaskAvailabilityLifetimeInSeconds` to specify the total amount of time a task remains available to workers. This is how long workers have to pick up a task before it fails.

5. Use `NumberOfHumanWorkersPerDataObject` to specify the number of workers per dataset object.

6. Use `PublicWorkforceTaskPrice` to set the price per task. This is the amount of money a worker receives for completing a single task.

7. Use `DataAttributes` to specify that your input data is free of confidential information, personal information, or protected health information.

    Ground Truth **requires** that your input data is free of personally identifiable information (PII) if you use the Mechanical Turk workforce. If you use Mechanical Turk and do not specify that
your input data is free of PII using the FreeOfPersonallyIdentifiableInformation flag, your labeling job will fail.

Use the FreeOfAdultContent flag to declare that your input data is free of adult content. SageMaker may restrict the Mechanical Turk workers that can view your task if it contains adult content.

You can see examples of how to use this API in the following notebooks, found on GitHub: [Ground Truth Jupyter Notebook Examples](#). You can access these notebooks under the SageMaker [Example Notebooks](#) in a [notebook instance](#).

### Use Mechanical Turk with Amazon A2I

You can specify that you want to use Mechanical Turk with Amazon A2I when you create a human review workflow, also referred to as a flow definition, in the console, or with the CreateFlowDefinition API operation. When you use this human review workflow to configure human loops, you must specify that your input data is free of PII.

#### To use Mechanical Turk when you create a human review workflow (console):

1. Use the following to create a human review workflow in the Augmented AI section of the SageMaker console: [Create a Human Review Workflow (Console)](#).
2. When you are selecting **Worker types** in the **Workers** section, select Amazon Mechanical Turk.
3. Select the **Price per task** using the dropdown list. This is the amount of money a worker receives for completing a single task.
4. (Optional) You can specify the **Number of workers per dataset object** under **Additional configuration**. For example, if you enter 3 in this field, each data object will be labeled by 3 workers.
5. (Optional) Specify the total amount of time workers have to complete a task using **Task timeout**.
6. (Optional) Specify the total amount of time a task remains available to workers in **Task expiration**. This is how long workers have to pick up a task before it fails.
7. Once you have created your human review workflow, you can use it to configure a human loop by providing its Amazon Resource Name (ARN) in the parameter FlowDefinitionArn. You configure a human loop using one of the API operations of a built-in task type, or the Amazon A2I runtime API operation, StartHumanLoop. To learn more, see [Create and Start a Human Loop](#).
When you configure your human loop, you must specify that your input data is free of personally identifiable information (PII) using the FreeOfPersonallyIdentifiableInformation content classifier in DataAttributes. If you use Mechanical Turk and do not specify that your input data is free of PII, your human review tasks will fail.

Use the FreeOfAdultContent flag to declare that your input data is free of adult content. SageMaker may restrict the Mechanical Turk workers that can view your task if it contains adult content.

To use Mechanical Turk when you create a human review workflow (API):

1. Use the following to create a human review workflow using the CreateFlowDefinition operation: Create a Human Review Workflow (API).
2. Use the following for the WorkteamArn. Replace region with the AWS Region you are using to create the labeling job.

```
arn:aws:sagemaker:region:394669845002:workteam/public-crowd/default
```
3. Use TaskTimeLimitInSeconds to specify the total amount of time workers have to complete a task.
4. Use TaskAvailabilityLifetimeInSeconds to specify the total amount of time a task remains available to workers. This is how long workers have to pick up a task before it fails.
5. Use TaskCount to specify the number of workers per dataset object. For example, if you specify 3 for this parameter, each data object will be labeled by 3 workers.
6. Use PublicWorkforceTaskPrice to set the price per task. This is the amount of money a worker receives for completing a single task.
7. Once you have created your human review workflow, you can use it to configure a human loop by providing its Amazon Resource Name (ARN) in the parameter FlowDefinitionArn. You configure a human loop using one of the API operations of a built-in task type, or the Amazon A2I runtime API operation, StartHumanLoop. To learn more, see Create and Start a Human Loop.

When you configure your human loop, you must specify that your input data is free of personally identifiable information (PII) using the FreeOfPersonallyIdentifiableInformation content classifier in DataAttributes.
If you use Mechanical Turk and do not specify that your input data is free of PII, your human review tasks will fail.

Use the FreeOfAdultContent flag to declare that your input data is free of adult content. SageMaker may restrict the Mechanical Turk workers that can view your task if it contains adult content.

You can see examples of how to use this API in the following notebooks, found on GitHub: Amazon A2I Jupyter Notebook Examples.

When is Mechanical Turk Not Supported?

This workforce is not supported under the following scenarios. In each scenario, you must use a private or vendor workforce.

- This workforce is not supported for Ground Truth video frame labeling jobs and 3D point cloud labeling jobs.
- You cannot use this workforce if your input data contains personally identifiable information (PII).
- Mechanical Turk is not available in some of the AWS special regions. If applicable, refer to the documentation for your special region for more information.

Managing Vendor Workforces

You can use a vendor-managed workforce to label your data using Amazon SageMaker Ground Truth (Ground Truth) and Amazon Augmented AI (Amazon A2I). Vendors have extensive experience in providing data labeling services for the purpose of machine learning. Vendor workforces for these two services must be created and managed separately through the Amazon SageMaker console.

Vendors make their services available via the AWS Marketplace. You can find details of the vendor's services on their detail page, such as the number of workers and the hours that they work. You can use these details to make estimates of how much the labeling job will cost and the amount of time that you can expect the job to take. Once you have chosen a vendor you subscribe to their services using the AWS Marketplace.
A subscription is an agreement between you and the vendor. The agreement spells out the details of the agreement, such as price, schedule, or refund policy. You work directly with the vendor if there are any issues with your labeling job.

You can subscribe to any number of vendors to meet your data annotation needs. When you create a labeling job or human review workflow you can specify that the job be routed to a specific vendor.

⚠️ Important

Before you send sensitive data to a vendor, check the vendor’s security and compliance practices on their detail page and review the end user license agreement (EULA) that is part of your subscription agreement. You are responsible for ensuring that the vendor meets your compliance requirements for personal or confidential information. Do not share protected health information with this workforce.

You must use the console to subscribe to a vendor workforce. Once you have a subscription, you can use the `ListSubscribedWorkteams` operation to list your subscribed vendors.

**To subscribe to a vendor workforce**

2. Choose the appropriate page in the SageMaker console.
   - For Ground Truth labeling jobs, choose **Labeling workforces**, choose **Vendor**, and then choose **Find data labeling services**.
   - For Amazon A2I human review workflows, choose **Human review workforces**, choose **Vendor**, and then choose **Find human review services**.
3. The console opens the AWS Marketplace with:
   - data labeling services category selected for Ground Truth
   - human review services category selected for Amazon A2I

   Here you see a list of the vendor services available for this service.
4. Choose a vendor. The AWS Marketplace shows detailed information about the data labeling or human review service. Use this information to determine if the vendor meets your requirements for your task.
5. If the vendor meets your requirements, choose **Continue to subscribe**.

6. Review the details of the subscription. If you agree to the terms, choose **Subscribe** to complete your subscription to the service.

## Use a Private Workforce

A **private workforce** is a group of workers that you choose. These can be employees of your company or a group of subject matter experts from your industry. For example, if the task is to label medical images, you could create a private workforce of people knowledgeable about the images in question.

Each AWS account has access to a single private workforce per region, and the owner has the ability to create multiple **private work teams** within that workforce. A single private work team is used to complete a labeling job or human review task, or a **job**. You can assign each work team to a separate job or use a single team for multiple jobs. A single worker can be in more than one work team.

Your private workforce can either be created and managed using **Amazon Cognito** or your own private OpenID Connect (OIDC) Identity Provider (IdP).

If you are a new user of **Amazon SageMaker Ground Truth** or **Amazon Augmented AI** and do not require your workers to be managed with your own IdP, it is recommended that you use Amazon Cognito to create and manage your private workforce.

After you create a workforce, in addition to creating and managing work teams, you can do the following:

- **Track worker performance**
- **Create and manage Amazon SNS topics** to notify workers when labeling tasks are available
- **Manage Private Workforce Access to Tasks Using IP Addresses**

**Note**

Your private workforce is shared between Ground Truth and Amazon A2I. To create and manage private work teams used by Augmented AI, use the Ground Truth section of the SageMaker console.
Create and Manage Amazon Cognito Workforce

Create and manage your private workforce using Amazon Cognito when you want to create your workforce using the Amazon SageMaker console or you don't want the overhead of managing worker credentials and authentication. When you create a private workforce with Amazon Cognito, it provides authentication, authorization, and user management for your private workers.

Topics

- Create a Private Workforce (Amazon Cognito)
- Manage a Private Workforce (Amazon Cognito)

Create a Private Workforce (Amazon Cognito)

When you use Amazon Cognito, you can create a private workforce in one of the following ways:

- Create a new workforce while you are creating your labeling job. To learn how, see Create an Amazon Cognito Workforce When Creating a Labeling Job.
- Create a new workforce before you create your labeling job. To learn how, see Create an Amazon Cognito Workforce Using the Labeling Workforces Page.
- Import an existing workforce after creating a user pool in the Amazon Cognito console. To learn how, see Create a Private Workforce (Amazon Cognito Console).

Once you create a private workforce, that workforce and all work teams and workers associated with it are available to use for all Ground Truth labeling job tasks and Amazon Augmented AI human review workflows tasks.

If you are new to Amazon SageMaker and want to test Ground Truth or Amazon A2I, we suggest that you create a private work team consisting of people from your organization using the console.
Use this work team when creating labeling or human review workflows (flow definitions) to test your worker UI and job workflow.

**Topics**

- [Create a Private Workforce (Amazon SageMaker Console)](#)
- [Create a Private Workforce (Amazon Cognito Console)](#)

**Create a Private Workforce (Amazon SageMaker Console)**

You can create a private workforce in the Amazon SageMaker console in one of two ways:

- When creating a labeling job in the **Labeling jobs** page of the Amazon SageMaker Ground Truth section.
- Using the **Labeling workforces** page of the Amazon SageMaker Ground Truth section. If you are creating a private workforce for an Amazon A2I human review workflow, use this method.

Both of these methods also create a default work team containing all of the members of the workforce. This private workforce is available to use for both Ground Truth and Amazon Augmented AI jobs.

When you create a private workforce using the console, SageMaker uses Amazon Cognito as an identity provider for your workforce. If you want to use your own OpenID Connect (OIDC) Identity Provider (IdP) to create and manage your private workforce, you must create a workforce using the SageMaker API operation `CreateWorkforce`. To learn more, see [Create a Private Workforce (OIDC IdP)](#).

**Create an Amazon Cognito Workforce When Creating a Labeling Job**

If you haven’t created a private workforce when you create your labeling job and you choose to use private workers, you are prompted to create a work team. This will create a private workforce using Amazon Cognito.

**To create a workforce while creating a labeling job (console)**

2. In the navigation pane, choose **Labeling jobs** and fill in all required fields. For instructions on how to start a labeling job, see [Getting started]. Choose **Next**.
3. Choose **Private** for the workforce type.
4. In the **Workers** section, enter:

   a. The **Team name**.

   b. Email addresses for up to 100 workforce members. Email addresses are case sensitive. Your workers must log in using the same case used when the address was initially entered. You can add additional workforce members after the job has been created.

   c. The name of your organization. SageMaker uses this to customize the email sent to the workers.

   d. A contact email address for workers to report issues related to the task.

When you create the labeling job, an email is sent to each worker inviting them to join the workforce. After creating the workforce, you can add, delete, and disable workers using the SageMaker console or the Amazon Cognito console.

**Create an Amazon Cognito Workforce Using the Labeling Workforces Page**

To create and manage your private workforce using Amazon Cognito, you can use the **Labeling workforces** page. When following the instructions below, you have the option to create a private workforce by entering worker emails importing a pre-existing workforce from an Amazon Cognito user pool. To import a workforce, see [Create a Private Workforce (Amazon Cognito Console)].

**To create a private workforce using worker emails**

2. In the navigation pane, choose **Labeling workforces**.
3. Choose **Private**, then choose **Create private team**.
4. Choose **Invite new workers by email**.
5. Paste or type a list of up to 50 email addresses, separated by commas, into the email addresses box.
6. Enter an organization name and contact email.
7. Optionally, choose an SNS topic to which to subscribe the team so workers are notified by email when new Ground Truth labeling jobs become available. Amazon SNS notifications are supported by Ground Truth and are not supported by Augmented AI. If you subscribe workers to receive SNS notifications, they only receive notifications about Ground Truth labeling jobs. They do not receive notifications about Augmented AI tasks.
8. Click the **Create private team** button.
After you import your private workforce, refresh the page. On the **Private workforce summary** page, you can see information about the Amazon Cognito user pool for your workforce, a list of work teams for your workforce, and a list of all of the members of your private workforce.

**Note**

If you delete all of your private work teams, you have to repeat this process to use a private workforce in that region.

### Create a Private Workforce (Amazon Cognito Console)

Amazon Cognito is used to define and manage your private workforce and your work teams. It is a service that you can use to create identities for your workers and authenticate these identities with identity providers. A private workforce corresponds to a single **Amazon Cognito user pool**. Private work teams correspond to **Amazon Cognito user groups** within that user pool.

Example identity providers supported by Amazon Cognito:

- Social sign-in providers such as Facebook and Google
- OpenID Connect (OIDC) providers
- Security Assertion Markup Language (SAML) providers such as Active Directory
- The Amazon Cognito built-in identity provider

For more information, see [What Is Amazon Cognito?](#).

To create a private workforce using Amazon Cognito, you must have an existing Amazon Cognito user pool containing at least one user group. See [Tutorial: Creating a User Pool](#) to learn how to create a user pool. See [Adding Groups to a User Pool](#) to learn how to add a user group to a pool.

Once your user pool has been created, follow the steps below to create a private workforce by importing that user pool into Amazon SageMaker.

**To create a private workforce by importing a Amazon Cognito user pool**

2. In the navigation pane, choose **Labeling workforces**.
3. Choose **Private**.
4. Choose **Create private team**. This creates a private workforce and a work team.

5. Choose **Import workers from existing Amazon Cognito user groups**.

6. Choose a user pool that you have created. User pools require a domain and an existing user group. If you get an error that the domain is missing, set it in the **Domain name** options on the **App integration** page of the Amazon Cognito console for your group.

7. Choose an app client. We recommend using a client generated by SageMaker.

8. Choose a user group from your pool to import its members.

9. Optionally choose an Amazon Simple Notification Service (Amazon SNS) topic to which to subscribe the team so that workers are notified by email when new labeling jobs become available. Amazon SNS notifications are supported by Ground Truth and are not supported by Augmented AI. If you subscribe workers to receive SNS notifications, they only receive notifications about Ground Truth labeling jobs. They do not receive notifications about Augmented AI tasks.

10. Choose **Create private team**.

---

⚠️ **Important**

After you create a workforce using an Amazon Cognito user pool, it should not be deleted without first deleting all work teams associated with that pool in the SageMaker console.

---

After you import your private workforce, refresh the page to see the **Private workforce summary** page. On this page, you can see information about the Amazon Cognito user pool for your workforce, a list of work teams for your workforce, and a list of all of the members of your private workforce. This workforce is now available to use in both Amazon Augmented AI and Amazon SageMaker Ground Truth for human review tasks and data labeling jobs respectively.

**Manage a Private Workforce (Amazon Cognito)**

After you have created a private workforce using Amazon Cognito, you can create and manage work teams using the Amazon SageMaker console and API operations.

You can do the following using either the [SageMaker console](https://console.aws.amazon.com/sagemaker) or [Amazon Cognito console](https://console.aws.amazon.com/cognito).  

- Add and delete work teams.
- Add workers to your workforce and one or more work teams.
• Disable or remove workers from your workforce and one or more workteams. If you add workers to a workforce using the Amazon Cognito console, you must use the same console to remove the worker from the workforce.

You can restrict access to tasks to workers at specific IP addresses using the SageMaker API. For more information, see Manage Private Workforce Using the Amazon SageMaker API.

Topics

• Manage a Workforce (Amazon SageMaker Console)
• Manage a Private Workforce (Amazon Cognito Console)

Manage a Workforce (Amazon SageMaker Console)

You can use the Amazon SageMaker console to create and manage the work teams and individual workers that make up a private workforce.

Use a work team to assign members of your private workforce to a labeling or human review job. When you create your workforce using the SageMaker console, there is a work team called Everyone-in-private-workforce that enables you to assign your entire workforce to a job. Because an imported Amazon Cognito user pool may contain members that you don't want to include in your work teams, a similar work team is not created for Amazon Cognito user pools.

You have two choices to create a new work team:

• You can create a work team in the SageMaker console and add members from your workforce to the team.
• You can create a user group by using the Amazon Cognito console and then create a work team by importing the user group. You can import more than one user group into each work team. You manage the members of the work team by updating the user group in the Amazon Cognito console. See Manage a Private Workforce (Amazon Cognito Console) for more information.

Create a Work Team Using the SageMaker Console

You can create a new Amazon Cognito user group or import an existing user group using the SageMaker console, on the Labeling workforces page. For more information on creating a user group in the Amazon Cognito console, see Manage a Private Workforce (Amazon Cognito Console).
To create a work team using the SageMaker console

2. Choose **Labeling workforces** from the left menu.
3. Under **Private**, choose **Create private team**.
4. Under **Team details**, enter a **Team name**. The name must be unique in your account in an AWS Region.
5. Under **Add workers**, choose a method to add workers to the team using a user group.
   - If you chose **Create a team by adding workers to a new Amazon Cognito user group**, select the workers to add to the team.
   - If you chose **Create a team by importing existing Amazon Cognito user groups**, choose the user groups that are part of the new team.
6. If you select an **SNS topic**, all workers added to the team are subscribed to the Amazon SNS topic and notified when new work items are available to the team. Select from a list of your existing Ground Truth related Amazon SNS topics or select **Create new topic** to open a topic-creation dialog.

   Amazon SNS notifications are supported by Ground Truth and are not supported by Augmented AI. If you subscribe workers to receive SNS notifications, they only receive notifications about Ground Truth labeling jobs. They do not receive notifications about Augmented AI tasks.

Workers in a workteam subscribed to a topic receive notifications when a new Ground Truth labeling job for that team becomes available and when one is about to expire.

Read [Create and manage Amazon SNS topics for your work teams](#) for more information about using Amazon SNS topic.

**Subscriptions**

After you have created a work team, you can see more information about the team and change or set the Amazon SNS topic to which its members are subscribed by visiting the Amazon Cognito console. If you added any team members before you subscribed the team to a topic, you need to manually subscribe those members to that topic. Read [Create and manage Amazon SNS topics for your work teams](#) for more information on creating and managing the Amazon SNS topic.
Add or Remove Workers

A work team is a group of workers within your workforce to whom you can assign jobs. A worker can be added to more than one work team. Once a worker has been added to a work team, that worker can be disabled or removed.

Add Workers to the Workforce

Adding a worker to the workforce enables you to add that worker to any work team within that workforce.

To add workers using the private workforce summary page

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Labeling workforces to navigate to your private workforce summary page.
3. Choose Private.
5. Paste or type a list of email addresses, separated by commas, into the email addresses box. You can have up to 50 email addresses in this list.

Add a Worker to a Work Team

A worker must be added to the workforce before being added to a work team. To add a worker to a work team, first navigate to the Private workforce summary page using the steps above.

To add a worker to a work team from the private workforce summary page

1. In the Private teams section, choose the team to which you want to add the workers.
2. Choose the Workers tab.
3. Choose Add workers to team and choose the boxes next to the workers that you want to add.
4. Click Add workers to team.

Disable and Remove a Worker from the Workforce

Disabling a worker stops the worker from receiving a job. This action does not remove the worker from the workforce, or from any work team with which the worker is associated. To disable or remove a worker from a work team, first navigate to the private workforce summary page using the steps above.
To deactivate a worker using the private workforce summary page

1. In the **Workers** section, choose the worker that you would like to disable.
2. Choose **Disable**.

If desired, you can subsequently **Enable** a worker after they have been disabled.

You can remove workers from your private workforce directly in the SageMaker console if that worker was added in this console. If you added the worker (user) in the Amazon Cognito console, see [Manage a Private Workforce (Amazon Cognito Console)](#) to learn how to remove the worker in the Amazon Cognito console.

To remove a worker using the private workforce summary page

1. In the **Workers** section, choose the worker that you would like to delete.
2. If the worker has not been disabled, choose **Disable**.
3. Select the worker and choose **Delete**.

**Manage a Private Workforce (Amazon Cognito Console)**

A private workforce corresponds to a single **Amazon Cognito user pool**. Private work teams correspond to **Amazon Cognito user groups** within that user pool. Workers correspond to **Amazon Cognito users** within those groups.

After your workforce has been created, you can add work teams and individual workers through the Amazon Cognito console. You can also delete workers from your private workforce or remove them from individual teams in the Amazon Cognito console.

⚠️ **Important**

You can’t delete work teams from the Amazon Cognito console. Deleting an Amazon Cognito user group that is associated with an Amazon SageMaker work team will result in an error. To remove work teams, use the SageMaker console.
Create Work Teams (Amazon Cognito Console)

You can create a new work team to complete a job by adding a Amazon Cognito user group to the user pool associated with your private workforce. To add a Amazon Cognito user group to an existing worker pool, see Adding groups to a User Pool.

To create a work team using an existing Amazon Cognito user group

2. In the navigation pane, choose Workforces.
3. For Private teams, choose Create private team.
4. Under Team details, give the team a name. The name must be unique in your account in an AWS Region.
5. For Add workers, choose Import existing Amazon Cognito user groups, and choose one or more user groups that are part of the new team.
6. If you choose an SNS topic, all workers added to the team are subscribed to the Amazon Simple Notification Service (Amazon SNS) topic and notified when new work items are available to the team. Choose from a list of your existing SNS topics related to SageMaker Ground Truth or Amazon Augmented AI or choose Create new topic to create one.

Note

Amazon SNS notifications are supported by Ground Truth and are not supported by Augmented AI. If you subscribe workers to receive SNS notifications, they only receive notifications about Ground Truth labeling jobs. They do not receive notifications about Augmented AI tasks.

Subscriptions

After you have created a work team, you can see more information about the team and change or set the SNS topic to which its members are subscribed using the Amazon Cognito console. If you added any team members before you subscribed the team to a topic, you need to manually subscribe those members to that topic. For more information, see Create and manage Amazon SNS topics for your work teams.
Add and Remove Workers (Amazon Cognito Console)

When using the Amazon Cognito console to add workers to a work team, you must add a user to the user pool associated with the workforce before adding that user to a user group. Users can be added to a user pool in various ways. For more information, see Signing Up and Confirming User Accounts.

Add a Worker to a Work Team

After a user has been added to a pool, the user can be associated with user groups inside of that pool. After a user has been added to a user group, that user becomes a worker on any work team created using that user group.

To add a user to a user group

1. Open the Amazon Cognito console: https://console.aws.amazon.com/cognito/.
2. Choose Manage User Pools.
3. Choose the user pool associated with your SageMaker workforce.
4. Under General Settings, choose Users and Groups and do one of the following:
   - Choose Groups, choose the group that you want to add the user to, and choose Add users. Choose the users that you want to add by choosing the plus-icon to the right of the user's name.
   - Choose Users, choose the user that you want to add to the user group, and choose Add to group. From the dropdown menu, choose the group and choose Add to group.

Disable and Remove a Worker From a Work Team

Disabling a worker stops the worker from receiving jobs. This action doesn't remove the worker from the workforce, or from any work team the worker is associated with. To remove a user from a work team in Amazon Cognito, you remove the user from the user group associated with that team.

To deactivate a worker (Amazon Cognito console)

1. Open the Amazon Cognito console: https://console.aws.amazon.com/cognito/.
2. Choose Manage User Pools.
3. Choose the user pool associated with your SageMaker workforce.
4. Under **General Settings**, choose **Users and Groups**.

5. Choose the user that you want to disable.

6. Choose **Disable User**.

You can enable a disabled user by choosing **Enable User**.

**To remove a user from a user group (Amazon Cognito console)**

1. Open the Amazon Cognito console: [https://console.aws.amazon.com/cognito/](https://console.aws.amazon.com/cognito/).

2. Choose **Manage User Pools**.

3. Choose the user pool associated with your SageMaker workforce.

4. Under **General Settings**, choose **Users and Groups**.

5. For **User** tab, choose the X icon to the right of the group from which you want to remove the user.

**Create and Manage OIDC IdP Workforce**

Create a private workforce using an OpenID Connect (OIDC) Identity Provider (IdP) when you want to manage and authenticate your workers using your own OIDC IdP. Individual worker credentials and other data will be kept private. Ground Truth and Amazon A2I will only have visibility into worker information you provide through the claims that you send to these services. To create a workforce using an OIDC IdP, your IdP must support *groups* because Ground Truth and Amazon A2I map one or more groups in your IdP to a work team. To learn more, see [Send Required and Optional Claims to Ground Truth and Amazon A2I](https://docs.aws.amazon.com/groundtruth/latest/dg/oidc-idp-claims.html).

If you are a new user of Ground Truth or Amazon A2I, you can test your worker UI and job workflow by creating a private work team and adding yourself as a worker. Use this work team when you create a labeling job or human review workflow. First, create a private OIDC IdP workforce using the instructions in [Create a Private Workforce (OIDC IdP)](https://docs.aws.amazon.com/groundtruth/latest/dg/oidc-idp-workforce.html). Next, refer to [Manage a Private Workforce (OIDC IdP)](https://docs.aws.amazon.com/groundtruth/latest/dg/oidc-idp-workforce.html) to learn how to create a work team.

**Topics**

- [Create a Private Workforce (OIDC IdP)](https://docs.aws.amazon.com/groundtruth/latest/dg/oidc-idp-workforce.html)
- [Manage a Private Workforce (OIDC IdP)](https://docs.aws.amazon.com/groundtruth/latest/dg/oidc-idp-workforce.html)
Create a Private Workforce (OIDC IdP)

Create a private workforce using an OpenID Connect (OIDC) Identity Provider (IdP) when you want to authenticate and manage workers using your own identity provider. Use this page to learn how to configure your IdP to communicate with Amazon SageMaker Ground Truth (Ground Truth) or Amazon Augmented AI (Amazon A2I) and to learn how to create a workforce using your own IdP.

To create a workforce using an OIDC IdP, your IdP must support *groups* because Ground Truth and Amazon A2I use one or more groups that you specify to create work teams. You use work teams to specify workers for your labeling jobs and human review tasks. Because groups are not a standard claim, your IdP may have a different naming convention for a group of users (workers). Therefore, you must identify one or more user groups to which a worker belongs using the custom claim `sagemaker:groups` that is sent to Ground Truth or Amazon A2I from your IdP. To learn more, see *Send Required and Optional Claims to Ground Truth and Amazon A2I*.

You create an OIDC IdP workforce using the SageMaker API operation *CreateWorkforce*. Once you create a private workforce, that workforce and all work teams and workers associated with it are available to use for all Ground Truth labeling job tasks and Amazon A2I human review workflows tasks. To learn more, see *Create an OIDC IdP Workforce*.

Send Required and Optional Claims to Ground Truth and Amazon A2I

When you use your own IdP, Ground Truth and Amazon A2I use your Issuer, ClientId, and ClientSecret to authenticate workers by obtaining an authentication CODE from your AuthorizationEndpoint.

Ground Truth and Amazon A2I will use this CODE to obtain a custom claim from either your IdP's TokenEndpoint or UserInfoEndpoint. You can either configure TokenEndpoint to return a JSON web token (JWT) or UserInfoEndpoint to return a JSON object. The JWT or JSON object must contain required and optional claims that you specify. A *claim* is a key-value pair that contains information about a worker or metadata about the OIDC service. The following table lists the claims that must be included, and that can optionally be included in the JWT or JSON object that your IdP returns.

<table>
<thead>
<tr>
<th>Claim</th>
<th>Required/Optional</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>sub</code></td>
<td>Required</td>
<td>Subject ID (unique identifier)</td>
</tr>
<tr>
<td><code>name</code></td>
<td>Optional</td>
<td>Name of the user (not required if <code>name</code> is included in the custom claim)</td>
</tr>
<tr>
<td><code>email</code></td>
<td>Optional</td>
<td>Email address of the user (not required if <code>email</code> is included in the custom claim)</td>
</tr>
<tr>
<td><code>phone</code></td>
<td>Optional</td>
<td>Phone number of the user (not required if <code>phone</code> is included in the custom claim)</td>
</tr>
<tr>
<td><code>locale</code></td>
<td>Optional</td>
<td>Language of the user (not required if <code>locale</code> is included in the custom claim)</td>
</tr>
<tr>
<td><code>birthdate</code></td>
<td>Optional</td>
<td>Date of birth of the user (not required if <code>birthdate</code> is included in the custom claim)</td>
</tr>
<tr>
<td><code>picture</code></td>
<td>Optional</td>
<td>User's profile picture (not required if <code>picture</code> is included in the custom claim)</td>
</tr>
<tr>
<td><code>groups</code></td>
<td>Required</td>
<td>One or more groups that a worker belongs to</td>
</tr>
<tr>
<td><code>role</code></td>
<td>Optional</td>
<td>Role of the user (not required if <code>role</code> is included in the custom claim)</td>
</tr>
</tbody>
</table>

**Note**

Some of the parameters in the following table can be specified using a : or a -. For example, you can specify the groups a worker belongs to using `sagemaker:groups` or `sagemaker-groups` in your claim.
<table>
<thead>
<tr>
<th>Name</th>
<th>Required</th>
<th>Accepted Format and Values</th>
<th>Description</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>sagemaker:groups</code> or <code>sagemaker-groups</code></td>
<td>Yes</td>
<td>Data type:</td>
<td>Assigns a worker to one or more groups. Groups are used to map the worker into work teams.</td>
<td>Example of worker that belongs to a single group: &quot;work_team1&quot;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If a worker belongs to a single group, identify the group using a string.</td>
<td></td>
<td>Example of a worker that belongs to more than one groups: [&quot;work_team1&quot;, &quot;work_team2&quot;]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If a worker belongs to multiple groups, use a list of up to 10 strings.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Allowable characters:</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Regex: \p{L}\p{M}\p{S}\p{N}\p{P} +</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Quotas:</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>10 groups per worker</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>63 characters per group name</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>sagemaker:sub</code> or <code>sagemaker-sub</code></td>
<td>Yes</td>
<td>Data type:</td>
<td>This is mandatory to track a worker identity inside the Ground Truth platform for auditing and to identify tasks worked on by that worker.</td>
<td>&quot;11101110 1-1234567 89-368705 6437-1111&quot;</td>
</tr>
<tr>
<td>Name</td>
<td>Required</td>
<td>Accepted Format and Values</td>
<td>Description</td>
<td>Example</td>
</tr>
<tr>
<td>------</td>
<td>----------</td>
<td>----------------------------</td>
<td>-------------</td>
<td>---------</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>For ADFS: Customers must use the Primary Security Identifier (SID).</td>
<td></td>
</tr>
<tr>
<td>sagemaker :client_id or sagemaker -client_id</td>
<td>Yes</td>
<td>Data type: String Allowable characters: Regex: `[\w+-]+ Quotes: 128 characters</td>
<td>A client ID. All tokens must be issued for this client ID.</td>
<td>&quot;00b600bb-1f00-05d0-bd00-00be00fbd0e0&quot;</td>
</tr>
<tr>
<td>sagemaker :name or sagemaker-name</td>
<td>Yes</td>
<td>Data type: String</td>
<td>The worker name to be displayed in the worker portal.</td>
<td>&quot;Jane Doe&quot;</td>
</tr>
<tr>
<td>Name</td>
<td>Required</td>
<td>Accepted Format and Values</td>
<td>Description</td>
<td>Example</td>
</tr>
<tr>
<td>---------------</td>
<td>----------</td>
<td>-----------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>--------------------------</td>
</tr>
<tr>
<td>email</td>
<td>No</td>
<td>Data type: String</td>
<td>The worker email. Ground Truth uses this email to notify workers that they have been invited to work on labeling tasks. Ground Truth will also use this email to notify your workers when labeling tasks become available if you set up an Amazon SNS topic for a work team that this worker is on.</td>
<td>&quot;<a href="mailto:example-email@domain.com">example-email@domain.com</a>&quot;</td>
</tr>
<tr>
<td>email_verified</td>
<td>No</td>
<td>Data type: Bool, Accepted Values: True, False</td>
<td>Indicates if the user email was verified or not.</td>
<td>True</td>
</tr>
</tbody>
</table>

The following an example of the JSON object syntax your UserInfoEndpoint can return.

```json
{
    "sub": "122",
    "exp": "10000",
    "sagemaker-groups":["group1","group2"]
    "sagemaker-name":"name",
    "sagemaker-sub": "122",
    "sagemaker-client_id": "123456"
}
```
Ground Truth or Amazon A2I compares the groups listed in `sagemaker:groups` or `sagemaker-groups` to verify that your worker belongs to the work team specified in the labeling job or human review task. After the work team has been verified, labeling or human review tasks are sent to that worker.

**Create an OIDC IdP Workforce**

You can create a workforce using the SageMaker API operation `CreateWorkforce` and associated language-specific SDKs. Specify a `WorkforceName` and information about your OIDC IDP in the parameter `OidcConfig`. It is recommended that you configure your OIDC with a place-holder redirect URI, and then update the URI with the worker portal URL after you create the workforce. To learn more, see [Configure your OIDC IdP](#).

The following shows an example of the request. See [CreateWorkforce](#) to learn more about each parameter in this request.

```java
CreateWorkforceRequest: {
  #required fields
  WorkforceName: "example-oidc-workforce",
  OidcConfig: {
    ClientId: "clientId",
    ClientSecret: "secret",
    Issuer: "https://example-oidc-idp.com/adfs",
    TokenEndpoint: "https://example-oidc-idp.com/adfs/oauth2/token",
    UserInfoEndpoint: "https://example-oidc-idp.com/adfs/oauth2/userInfo",
    JwksUri: "https://example-oidc-idp.com/adfs/discovery/keys"
  },
  SourceIpConfig: {
    Cidrs: ["string", "string"]
  }
}
```

**Configure your OIDC IdP**

How you configure your OIDC IdP depends on the IdP you use, and your business requirements.

When you configure your IdP, you must to specify a callback or redirect URI. After Ground Truth or Amazon A2I authenticates a worker, this URI will redirect the worker to the worker portal where the workers can access labeling or human review tasks. To create a worker portal URL, you need
to create a workforce with your OIDC IdP details using the CreateWorkforce API operation. Specifically, you must configure your OIDC IdP with required custom sagemaker claims (see the next section for more details). Therefore, it is recommended that you configure your OIDC with a place-holder redirect URI, and then update the URI after you create the workforce. See Create an OIDC IdP Workforce to learn how to create a workforce using this API.

You can view your worker portal URL in the SageMaker Ground Truth console, or using the SageMaker API operation, DescribeWorkforce. The worker portal URL is in the SubDomain parameter in the response.

⚠️ Important

Make sure you add the workforce subdomain to your OIDC IdP allow list. When you add the subdomain to your allow list, it must end with /oauth2/idpresponse.

To view your worker portal URL after creating a private workforce (Console):

2. In the navigation pane, choose Labeling workforces.
3. Select the Private tab.
4. In Private workforce summary you will see Labeling portal sign-in URL. This is your worker portal URL.

To view your worker portal URL after creating a private workforce (API):

When you create a private workforce using CreateWorkforce, you specify a WorkforceName. Use this name to call DescribeWorkforce. The following table includes examples of requests using the AWS CLI and AWS SDK for Python (Boto3).

SDK for Python (Boto3)

```python
response = client.describe_workforce(WorkforceName='string')
print(f'The workforce subdomain is: {response['SubDomain']}'
```

AWS CLI

```
$ C:\> describe-workforce --workforce-name 'string'
```
Validate Your OIDC IdP Workforce Authentication Response

After you have created your OIDC IdP workforce, you can use the following procedure to validate its authentication workflow using cURL. This procedure assumes you have access to a terminal, and that you have cURL installed.

To validate your OIDC IdP authorization response:

1. Get an authorization code using a URI configured as follows:

   `{AUTHORIZE ENDPOINT}?client_id={CLIENT ID}&redirect_uri={REDIRECT URI}&scope={SCOPE}&response_type=code`

   a. Replace `{AUTHORIZE ENDPOINT}` with the authorize endpoint for your OIDC IdP.
   b. Replace `{CLIENT ID}` with the Client ID from your OAuth client.
   c. Replace `{REDIRECT URI}` with the worker portal URL. If it is not already present, you must add /oauth2/idpresponse to the end of the URL.
   d. If you have a custom scope, use it to replace `{SCOPE}`. If you do not have a custom scope, replace `{SCOPE}` with openid.

   The following is an example of a URI after the modifications above are made:

   ```
   https://example.com/authorize?
   client_id=f490a907-9bf1-4471-97aa-6bfd159f81ac&redirect_uri=https%3A%2F%2Fexample.labeling.sagemaker.aws%2Foauth2%2Fidpresponse&response_type=code&scope=openid
   ```

2. Copy and paste the modified URI from step 1 into your browser and press Enter on your keyboard.

3. Authenticate using your IdP.

4. Copy the authentication code query parameter in the URI. This parameter begins with code=.
   The following is an example of what the response might look like. In this example, copy code=MCNYDB...
   and everything thereafter.

   ```
   https://example.labeling.sagemaker.aws/oauth2/idpresponse?code=MCNYDB....
   ```

5. Open a terminal and enter the following command after making required modifications listed below:
curl --request POST
   --url '{TOKEN ENDPOINT}'
   --header 'content-type: application/x-www-form-urlencoded'
   --data grant_type=authorization_code
   --data client_id={CLIENT ID}
   --data client_secret={CLIENT SECRET}
   --data code={CODE}
   --data 'redirect_uri={REDIRECT URI}'

a. Replace {TOKEN ENDPOINT} with the token endpoint for your OIDC IdP.

b. Replace {CLIENT ID} with the Client ID from your OAuth client.

c. Replace {CLIENT SECRET} with the Client Secret from your OAuth client.

d. Replace {CODE} with the authentication code query parameter you copied in step 4.

e. Replace {REDIRECT URI} with the worker portal URL.

The following is an example of the cURL request after making the modifications described above:

curl --request POST
   --url 'https://example.com/token'
   --header 'content-type: application/x-www-form-urlencoded'
   --data grant_type=authorization_code
   --data 'client_id=f490a907-9bf1-4471-97aa-6bfd159f81ac'
   --data client_secret=client-secret
   --data code=MCNYDB...
   --data 'redirect_uri=https://example.labeling.sagemaker.aws/oauth2/idpresponse'

6. This step depends on the type of access_token your IdP returns, a plain text access token or a JWT access token.

   • If your IdP does not support JWT access tokens, access_token may be plain text (for example, a UUID). The response you see may look similar to the following. In this case, move to step 7.

   ```json
   {
       "access_token":"179c144b-fccb-4d96-a28f-eea060f39c13",
       "token_type":"Bearer",
       "expires_in":3600,
   }```
If your IdP supports JWT access tokens, step 5 should generate an access token in JWT format. For example, the response may look similar to the following:

```json
{
    "access_token":"eyJh...JV_adQssw5c",
    "refresh_token":"i6mapTIAVSp2oJkgUnCACKKfZxt_H5MBLiqcybBBd04",
    "refresh_token_expires_in":6327,
    "scope":"openid",
    "id_token":"eyJ0eXAiOiJK9...-rDaQzUHl6cQQWNiDpWO1_1xXjQEwQ"
}
```

Copy the JWT and decode it. You can use python script or a third party website to decode it. For example, you can go to the website [https://jwt.io/](https://jwt.io/) and paste the JWT into the Encoded box to decode it.

Make sure the decoded response contains the following:

- **The Required** SageMaker claims in the table found in [Send Required and Optional Claims to Ground Truth and Amazon A2I](https://aws.amazon.com). If it does not, you must reconfigure your OIDC IdP to contain these claims.

- **The Issuer** you specified when you set up the IdP workforce.

7. In a terminal and enter the following command after making required modifications listed below:

   ```bash
curl -X POST -H 'Authorization: Bearer {ACCESS_TOKEN}' -d '' -k -v {USERINFO ENDPOINT}
```

   a. Replace `{USERINFO ENDPOINT}` with the user info endpoint for your OIDC IdP.

   b. Replace `{ACCESS_TOKEN}` with the access token in the response you received in step 7. This is the entry for the "access_token" parameter.

   The following is an example of the cURL request after making the modifications described above:
8. The response to the final step in the procedure above may look similar to the following code block.

   If the access_token returned in step 6 was plain text, you must verify that this response contains required information. In this case, the response must contain the **Required** SageMaker claims in the table found in Send Required and Optional Claims to Ground Truth and Amazon A2I. For example, sagemaker-groups, sagemaker-name.

   ```json
   {
     "sub":"122",
     "exp":10000,
     "sagemaker-groups":["group1","group2"],
     "sagemaker-name":"name",
     "sagemaker-sub":"122",
     "sagemaker-client_id":"123456"
   }
   ```

**Next Steps**

Once you've created a private workforce using your IdP and verified your IdP authentication response, you can create work teams using your IdP groups. To learn more, see [Manage a Private Workforce (OIDC IdP)](#).

You can restrict worker access to tasks to specific IP addresses, and update or delete your workforce using the SageMaker API. To learn more, see [Manage Private Workforce Using the Amazon SageMaker API](#).

**Manage a Private Workforce (OIDC IdP)**

Once you've created a private workforce using your OpenID Connect (OIDC) Identity Provider (IdP), you can manage your workers using your IdP. For example, you can add, remove, and group workers directly through your IdP.

To add workers to an Amazon SageMaker Ground Truth (Ground Truth) labeling job or Amazon Augmented AI (Amazon A2I) human review task, you create work teams using 1-10 IdP groups and assign that work team to the job or task. You assign a work team to a job or task by specifying that
work team when you create a labeling job (Ground Truth) or a human review workflow (Amazon A2I).

You can only assign one team to each labeling job or human review workflow. You can use the same team to create multiple labeling jobs or human review tasks. You can also create multiple work teams to work on different labeling jobs or human review tasks.

Prerequisites

To create and manage private work teams using your OIDC IdP groups, first you must create a workforce using the SageMaker API operation `CreateWorkforce`. To learn more, see Create a Private Workforce (OIDC IdP).

Add work teams

You can use the SageMaker console to create a private work team using your OIDC IdP workforce on the Labeling workforces page under Ground Truth. If you are creating a Ground Truth labeling job, you can also create a private work team while creating a labeling job.

Note

You create and manage work teams for Amazon A2I in the Ground Truth area of the SageMaker console.

You can also use the SageMaker API and associated language-specific SDKs to create a private work team.

Use the following procedures to learn how to create a private work team using the SageMaker console and API.

To create a private work team on the Labeling workforces page (console)

2. Select Labeling workforces.
3. Select Private.
4. In the Private teams section, select Create private team.
5. In the **Team details** section, enter a **Team name**.

6. In the **Add workers** section, enter the name of a single user group. All workers associated with this group in your IdP are added to this work team.

7. To add more than one user group, select **Add new user group** and enter the names of the user groups you want to add to this work team. Enter one user group per line.

8. (Optional) For Ground Truth labeling jobs, if you provide an email for workers in your JWT, Ground Truth notifies workers when a new labeling task is available if you select an SNS topic.

9. Select **Create private team**.

---

**To create a private work team while creating a Ground Truth labeling job (console)**


2. Select **Labeling jobs**.

3. Use the instructions in [Create a Labeling Job (Console)](https://console.aws.amazon.com/sagemaker/groundtruth) to create a labeling job. Stop when you get to the **Workers** section on the second page.

4. Select **Private** for your worker type.

5. Enter a **Team name**.

6. In the **Add workers** section, enter the name of a single user group under **User groups**. All workers associated with this group in your IdP are added to this work team.

   **Important**

   The group names you specify for **User groups** must match the group names specified in your OIDC IdP.

7. To add more than one user group, select **Add new user group** and enter the names of the user groups you want to add to this work team. Enter one user group per line.

8. Complete all remaining steps to create your labeling job.

The private team that you create is used for this labeling job, and is listed in the **Labeling workforces** section of the SageMaker console.

---

**To create a private work team using the SageMaker API**
You can create a private work team using the SageMaker API operation `CreateWorkteam`.

When you use this operation, list all user groups that you want included in the work team in the `OidcMemberDefinition` parameter `Groups`.

⚠️ **Important**

The group names you specify for `Groups` must match the group names specified in your OIDC IdP.

For example, if your user group names are `group1`, `group2`, and `group3` in your OIDC IdP, configure `OidcMemberDefinition` as follows:

```json
"OidcMemberDefinition": {
  "Groups": ["group1", "group2", "group3"]
}
```

Additionally, you must give the work team a name using the `WorkteamName` parameter.

### Add or remove IdP groups from work teams

After you've created a work team, you can use the SageMaker API to manage that work team. Use the `UpdateWorkteam` operation to update the IdP user groups included in that work team.

- Use the `WorkteamName` parameter to identify the work team that you want to update.
- When you use this operation, list all user groups that you want included in the work team in the `OidcMemberDefinition` parameter `Groups`. If a user group is associated with a work team and you do *not* include it in this list, that user group is no longer associated with this work team.

### Delete a work team

You can delete a work team using the SageMaker console and SageMaker API.

**To delete a private work team in the SageMaker console**

2. Select **Labeling workforces**.
3. Select **Private**.

4. In the **Private teams** section, select the work team that you want to delete.

5. Select **Delete**.

**To delete a private work team (API)**

You can delete a private work team using the SageMaker API operation **DeleteWorkteam**.

**Manage Individual Workers**

When you create a workforce using your own OIDC IdP, you cannot use Ground Truth or Amazon A2I to manage individual workers.

- To add a worker to a work team, add that worker to a group associated with that work team.
- To remove a worker from a work team, remove that worker from all user groups associated with that work team.

**Update, Delete, and Describe Your Workforce**

You can update, delete, and describe your OIDC IdP workforce using the SageMaker API. The following is a list of API operations that you can use to manage your workforce. For additional details, including how you can locate your workforce name, see **Manage Private Workforce Using the Amazon SageMaker API**.

- **UpdateWorkforce** – You may want to update a workforce created using your own OIDC IdP to specify a different authorization endpoint, token endpoint, or issuer. You can update any parameter found in **OidcConfig** using this operation.

  You can only update your OIDC IdP configuration when there are no work teams associated with your workforce. To learn how to delete work teams, see **Delete a work team**.

- **DeleteWorkforce** – Use this operation to delete your private workforce. If you have any work teams associated with your workforce, you must delete those work teams before you delete your work force. For more information, see **Delete a work team**.

- **DescribeWorkforce** – Use this operation to list private workforce information, including workforce name, Amazon Resource Name (ARN), and, if applicable, allowed IP address ranges (CIDRs).
Manage Private Workforce Using the Amazon SageMaker API

You can use Amazon SageMaker API operations to manage, update, and delete your private workforce. For each API operation linked on this page, you can find a list of supported language-specific SDKs and their documentation in the See Also section of the API documentation.

Find Your Workforce Name

Some of the SageMaker workforce-related API operations require your workforce name as input. You can see your Amazon Cognito or OIDC IdP private and vendor workforce names in an AWS Region using the ListWorkforces API operation in that AWS Region.

If you created your workforce using your own OIDC IdP, you can find your workforce name in the Ground Truth area of the SageMaker console.

To find your workforce name in the SageMaker console

2. Select Labeling workforces.
3. Select Private.
4. In the Private workforce summary section, locate your workforce ARN. Your workforce name is located at the end of this ARN. For example, if the ARN is arn:aws:sagemaker:us-east-2:111122223333:workforce/example-workforce, the workforce name is example-workforce.

Restrict Worker Access to Tasks to Allowable IP Addresses

By default, a workforce isn't restricted to specific IP addresses. You can use the UpdateWorkforce operation to require that workers use a specific range of IP addresses (CIDRs) to access tasks. If you specify one or more CIDRs, workers who attempt to access tasks using any IP address outside the specified ranges are denied and will get a HTTP 204 No Content error message on the worker portal. You can specify up to 10 CIDR values using UpdateWorkforce.

After you have restricted your workforce to one or more CIDRs, the output of UpdateWorkforce lists all allowable CIDRs. You can also use the DescribeWorkforce operation to view all allowable CIDRs for a workforce.
Update OIDC Identity Provider Workforce Configuration

You may want to update a workforce created using your own OIDC IdP to specify a different authorization endpoint, token endpoint, or issuer. You can update any parameter found in `OidcConfig` using the `UpdateWorkforce` operation.

⚠️ Important

You can only update your OIDC IdP configuration when there are no work teams associated with your workforce. You can delete a private work team using the `DeleteWorkteam` operation.

Delete a Private Workforce

You can only have one private workforce in each AWS Region. You may want to delete your private workforce in an AWS Region when:

- You want to create a workforce using a new Amazon Cognito user pool.
- You have already created a private workforce using Amazon Cognito and you want to create a workforce using your own OpenID Connect (OIDC) Identity Provider (IdP).

To delete a private workforce, use the `DeleteWorkforce` API operation. If you have any work teams associated with your workforce, you must delete those work teams before you delete your workforce. You can delete a private work team using the `DeleteWorkteam` operation.

Track Worker Performance

Amazon SageMaker Ground Truth logs worker events to Amazon CloudWatch, such as when a worker starts or submits a task. Use Amazon CloudWatch metrics to measure and track throughput across a team or for individual workers.

⚠️ Important

Worker event tracking is not available for Amazon Augmented AI human review workflows.
Enable Tracking

During the set-up process for a new work team, the permissions for Amazon CloudWatch logging of worker events are created. Since this feature was added in August 2019, work teams created prior to that may not have the correct permissions. If all of your work teams were created before August 2019, create a new work team. It does not need any members and may be deleted after creation, but by creating it, you establish the permissions and apply them to all of your work teams, regardless of when they were created.

Examine Logs

After tracking is enabled, the activity of your workers is logged. Open the Amazon CloudWatch console and choose Logs in the navigation pane. You should see a log group named /aws/sagemaker/groundtruth/WorkerActivity.

Each completed task is represented by a log entry, which contains information about the worker, their team, the job, when the task was accepted, and when it was submitted.

Example Log entry

```
{
    "worker_id": "cd449a289e129409",
    "cognito_user_pool_id": "us-east-2_IpicJXXXX",
    "cognito_sub_id": "d6947aeb-0650-447a-ab5d-894db61017fd",
    "task_accepted_time": "Wed Aug 14 16:00:59 UTC 2019",
    "task_submitted_time": "Wed Aug 14 16:01:04 UTC 2019",
    "task_returned_time": "",
    "task_declined_time": "",
    "workteam_arn": "arn:aws:sagemaker:us-east-2:############:workteam/private-crowd/Sample-labeling-team",
    "labeling_job_arn": "arn:aws:sagemaker:us-east-2:############:labeling-job/metrics-demo",
    "work_requester_account_id": "############",
    "job_reference_code": "############",
    "job_type": "Private",
    "event_type": "TasksSubmitted",
    "event_timestamp": "1565798464"
}
```

A useful data point in each event is the cognito_sub_id. You can match that to an individual worker.
1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.

2. Under the **Ground Truth** section, choose **Workforces**.

3. Choose **Private**.

4. Choose the name of a team in the **Private teams** section.

5. In the **Team summary** section, choose the user group identified under **Amazon Cognito user group**. That will take you to the group in the Amazon Cognito console.

6. The **Group** page lists the users in the group. Choose any user's link in the **Username** column to see more information about the user, including a unique **sub** ID.

To get information about all of the team's members, use the ListUsers action (examples) in the Amazon Cognito API.

**Use Log Metrics**

If you don't want to write your own scripts to process and visualize the raw log information, Amazon CloudWatch metrics provide insights into worker activity for you.

**To view metrics**


2. In the navigation pane, choose **Metrics**.

3. Choose the AWS/SageMaker/Workteam name space, then explore the available metrics. For example, selecting the **Workteam** and **Workforce** metrics lets you calculate the average time per submitted task for a specific labeling job.

For more information, see **Using Amazon CloudWatch Metrics**.

**Create and manage Amazon SNS topics for your work teams**

Use the procedures in this topic when you want to:

- Create a topic to which you want an existing work team to subscribe.
- Create a topic before you've created a work team.
- Create or modify the work team with an API call, and specify a topic Amazon Resource Name (ARN).
If you create a work team using the console, the console provides an option to create a new topic for the team so that you don't have to perform these steps.

⚠️ **Important**
The Amazon SNS feature is not supported by Amazon A2I. If you subscribe your work team to an Amazon SNS topic, workers will only receive notifications about Ground Truth labeling jobs. Workers will not receive notifications about new Amazon A2I human review tasks.

Create the Amazon SNS topic

The steps for creating Amazon SNS topics for work team notifications are similar to the steps in Getting Started in the Amazon SNS Developer Guide, with one significant addition—you must add an access policy so that Amazon SageMaker can publish messages to the topic on your behalf.

To add the policy when you create the topic

2. In Create topic, enter the name of your topic and then choose Next steps.
4. In the JSON editor, find the Resource property, which displays the topic's ARN.
5. Copy the Resource ARN value.
6. Before the final closing brace (]), add the following policy.

```
,
   
   "Sid": "AwsSagemaker_SnsAccessPolicy",
   "Effect": "Allow",
   "Principal": {
      "Service": "sagemaker.amazonaws.com"
   },
   "Action": "sns:Publish",
   "Resource": "arn:partition:sns:region:111122223333:MyTopic", # ARN of the topic you copied in the previous step
   "Condition": {
      "ArnLike": {
         "aws:SourceArn":
         "arn:partition:sagemaker:region:111122223333:workteam/**" # Workteam ARN
   ```
7. Create the topic.

After you create the topic, it appears in your Topics summary screen. For more information about creating topics, see Creating a Topic in the Amazon SNS Developer Guide.

Manage worker subscriptions

If you subscribe a work team to a topic after you've already created the work team, the individual work team members who were added to the team when the work team was created are not automatically subscribed to the topic. For information about subscribing workers' email addresses to the topic, see Subscribing an Endpoint to an Amazon SNS Topic in the Amazon SNS Developer Guide.

The only situation in which workers are automatically subscribed to your topic is when you create or import an Amazon Cognito user group at the time that you create a work team and you set up the topic subscription when you create that work team. For more information about creating and managing your workteams with Amazon Cognito, see Create Work Teams (Amazon Cognito Console).

Crowd HTML Elements Reference

Crowd HTML Elements are web components, a web standard that abstracts HTML markup, CSS, and JavaScript functionality into an HTML tag or set of tags. Amazon SageMaker provides customers with the ability to design their own custom task templates in HTML.

As a starting point, you can use a template built using Crowd HTML Elements from one of the following GitHub repositories:

- Example task UIs for Amazon SageMaker Ground Truth
- Over 60 example task UIs for Amazon Augmented AI (A2I)
These repositories include templates designed for audio, image, text, video, and other types of data labeling and annotation tasks.

For more information about how to implement custom templates in Amazon SageMaker Ground Truth, see Creating Custom Labeling Workflows. To learn more about custom templates in Amazon Augmented AI, see Create Custom Worker Task Templates.

**SageMaker Crowd HTML Elements**

The following is a list of Crowd HTML Elements that make building a custom template easier and provide a familiar UI for workers. These elements are supported in Ground Truth, Augmented AI, and Mechanical Turk.

**Topics**
- crowd-alert
- crowd-badge
- crowd-button
- crowd-bounding-box
- crowd-card
- crowd-checkbox
- crowd-classifier
- crowd-classifier-multi-select
- crowd-entity-annotation
- crowd-fab
- crowd-form
- crowd-icon-button
- crowd-image-classifier
- crowd-image-classifier-multi-select
- crowd-input
- crowd-instance-segmentation
- crowd-instructions
- crowd-keypoint
- crowd-line
- crowd-modal
crowd-polygon

crowd-polyline

crowd-radio-button

crowd-radio-group

crowd-semantic-segmentation

crowd-slider

crowd-tab

crowd-tabs

crowd-text-area

crowd-toast

crowd-toggle-button

crowd-alert

A message that alerts the worker to a current situation.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of a Liquid template that uses the `<crowd-alert>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<template src="https://assets.crowd.aws/crowd-html-elements.js"></template>

<template>
  <div id="errorBox"></div>

  <template>
    <template src="{{ task.input.taskObject | grant_read_access }}">
      <template labels="["Item A", "Item B", "Item C"]">
        <template header="Please locate the centers of each item."">
          <template name="annotatedResult">
            <template short-instructions>
              Describe your task briefly here and give examples
            </template>
            <template full-instructions>
              Give additional instructions and good/bad examples here
            </template>
          </template>
        </template>
      </template>
    </template>
  </template>
</template>
```
```javascript
var num_obj = 1;

document.querySelector('crowd-form').onsubmit = function(e) {
    const keypoints = document.querySelector('crowd-keypoint').value.keypoints || document.querySelector('crowd-keypoint')._submittableValue.keypoints;
    const labels = keypoints.map(function(p) {
        return p.label;
    });

    // 1. Make sure total number of keypoints is correct.
    var original_num_labels = document.getElementsByTagName("crowd-keypoint") [0].getAttribute("labels");
    original_num_labels = original_num_labels.substring(2, original_num_labels.length - 2).split("",",");
    var goalNumKeypoints = num_obj*original_num_labels.length;
    if (keypoints.length != goalNumKeypoints) {
        e.preventDefault();
        errorBox.innerHTML = '<crowd-alert type="error" dismissible>You must add all keypoint annotations and use each label only once.</crowd-alert>';  
        errorBox.scrollIntoView();
        return;
    }

    // 2. Make sure all labels are unique.
    labelCounts = {};
    for (var i = 0; i < labels.length; i++) {
        if (!labelCounts[labels[i]]) {
            labelCounts[labels[i]] = 0;
        }
        labelCounts[labels[i]]++;
    }
    const goalNumSingleLabel = num_obj;
    const numLabels = Object.keys(labelCounts).length;
    Object.entries(labelCounts).forEach(entry => {
        if (entry[1] != goalNumSingleLabel) {
            e.preventDefault();
        }else{
            labelCounts[labels[i]]++;
        }
    })
    const goalNumKeypoints = num_obj*original_num_labels.length;
    if (keypoints.length != goalNumKeypoints) {
        e.preventDefault();
        errorBox.innerHTML = '<crowd-alert type="error" dismissible>You must add all keypoint annotations and use each label only once.</crowd-alert>';  
        errorBox.scrollIntoView();
        return;
    }
    // 1. Make sure total number of keypoints is correct.
    var original_num_labels = document.getElementsByTagName("crowd-keypoint") [0].getAttribute("labels");
    original_num_labels = original_num_labels.substring(2, original_num_labels.length - 2).split("",",");
    var goalNumKeypoints = num_obj*original_num_labels.length;
    if (keypoints.length != goalNumKeypoints) {
        e.preventDefault();
        errorBox.innerHTML = '<crowd-alert type="error" dismissible>You must add all keypoint annotations and use each label only once.</crowd-alert>';  
        errorBox.scrollIntoView();
        return;
    }

    // 2. Make sure all labels are unique.
    labelCounts = {};
    for (var i = 0; i < labels.length; i++) {
        if (!labelCounts[labels[i]]) {
            labelCounts[labels[i]] = 0;
        }
        labelCounts[labels[i]]++;
    }
    const goalNumSingleLabel = num_obj;
    const numLabels = Object.keys(labelCounts).length;
    Object.entries(labelCounts).forEach(entry => {
        if (entry[1] != goalNumSingleLabel) {
            e.preventDefault();
        }
    })
```
The following attributes are supported by this element.

**dismissible**

A Boolean switch that, if present, allows the message to be closed by the worker.

**type**

A string that specifies the type of message to be displayed. The possible values are "info" (the default), "success", "error", and "warning".

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements:** crowd-form
- **Child elements:** none

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

**crowd-badge**

An icon that floats over the top right corner of another element to which it is attached.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).
Amazon SageMaker

Developer Guide

The following is an example of a template that uses the <crowd-badge> element. Copy the
following code and save it in a ﬁle with the extension .html. Open the ﬁle in any browser to
preview and interact with this template.

<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<crowd-image-classifier
name="crowd-image-classifier"
src="https://unsplash.com/photos/NLUkAA-nDdE"
header="Choose the correct category for this image."
categories="['Person', 'Umbrella', 'Chair', 'Dolphin']"
>
<full-instructions header="Classification Instructions">
<p>Read the task carefully and inspect the image.</p>
<p>Choose the appropriate label that best suits the image.</p>
</full-instructions>
<short-instructions id="short-instructions">
<p>Read the task carefully and inspect the image.</p>
<p>Choose the appropriate label that best suits the image.</p>
<crowd-badge icon="star" for="short-instructions"/>
</short-instructions>
</crowd-image-classifier>
</crowd-form>

Attributes
The following attributes are supported by this element.
for
A string that speciﬁes the ID of the element to which the badge is attached.
icon
A string that speciﬁes the icon to be displayed in the badge. The string must be either the name of
an icon from the open-source iron-icons set, which is pre-loaded, or the URL to a custom icon.
This attribute overrides the label attribute.
SageMaker Crowd HTML Elements

1865


The following is an example of the syntax that you can use to add an iron-icon to a `<crowd-badge>` HTML element. Replace `icon-name` with the name of the icon you'd like to use from this Icons set.

```
<crowd-badge icon="icon-name" for="short-instructions"/>
```

**label**

The text to display in the badge. Three characters or less is recommended because text that is too large will overflow the badge area. An icon can be displayed instead of text by setting the `icon` attribute.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: none

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

**crowd-button**

A styled button that represents some action.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).

The following is an example of a template that uses the `<crowd-button>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
```
Attributes

The following attributes are supported by this element.

`disabled`

A Boolean switch that, if present, displays the button as disabled and prevents clicks.

`form-action`

A switch that either submits its parent `<crowd-form>` element, if set to "submit", or resets its parent `<crowd-form>` element, if set to "reset".

`href`

The URL to an online resource. Use this property if you need a link styled as a button.

`icon`

A string that specifies the icon to be displayed next to the button's text. The string must be the name of an icon from the open-source `iron-icons` set, which is pre-loaded. For example, to insert the `search` iron-icon, use the following:

```html
<iron-icon icon="question-answer"/>
```
The icon is positioned to either the left or the right of the text, as specified by the `icon-align` attribute.

To use a custom icon see `icon-url`.

**icon-align**

The left or right position of the icon relative to the button's text. The default is "left".

**icon-url**

A URL to a custom image for the icon. A custom image can be used in place of a standard icon that is specified by the `icon` attribute.

**loading**

A Boolean switch that, if present, displays the button as being in a loading state. This attribute has precedence over the `disabled` attribute if both attributes are present.

**target**

When you use the `href` attribute to make the button act as a hyperlink to a specific URL, the `target` attribute optionally targets a frame or window where the linked URL should load.

**variant**

The general style of the button. Use "primary" for primary buttons, "normal" for secondary buttons, "link" for tertiary buttons, or "icon" to display only the icon without text.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements:** `crowd-form`
- **Child elements:** none

**See Also**

For more information, see the following.
crowd-bounding-box

A widget for drawing rectangles on an image and assigning a label to the portion of the image that is enclosed in each rectangle.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](https://codepen.io).

The following is an example of a Liquid template that uses the `<crowd-bounding-box>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template. For more examples, see this [GitHub repository](https://github.com).

```html
<crowd-form>
  <crowd-bounding-box
    name="annotatedResult"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="Draw bounding boxes around all the cats and dogs in this image"
    labels="['Cat', 'Dog']"
  >
    <full-instructions header="Bounding Box Instructions">
      <p>Use the bounding box tool to draw boxes around the requested target of interest:</p>
      <ol>
        <li>Draw a rectangle using your mouse over each instance of the target.</li>
        <li>Make sure the box does not cut into the target, leave a 2 - 3 pixel margin</li>
        <li>When targets are overlapping, draw a box around each object, include all contiguous parts of the target in the box. Do not include parts that are completely overlapped by another object.</li>
        <li>Do not include parts of the target that cannot be seen, even though you think you can interpolate the whole shape of the target.</li>
        <li>Avoid shadows, they're not considered as a part of the target.</li>
        <li>If the target goes off the screen, label up to the edge of the image.</li>
      </ol>
    </full-instructions>
  </crowd-bounding-box>
</crowd-form>
```
Attributes

The following attributes are supported by this element.

**header**

The text to display above the image. This is typically a question or simple instruction for the worker.

**initial-value**

An array of JSON objects, each of which sets a bounding box when the component is loaded. Each JSON object in the array contains the following properties. Bounding boxes set via the `initial-value` property can be adjusted and whether or not a worker answer was adjusted is tracked via an `initialValueModified` boolean in the worker answer output.

- **height** – The height of the box in pixels.
- **label** – The text assigned to the box as part of the labeling task. This text must match one of the labels defined in the `labels` attribute of the `<crowd-bounding-box>` element.
- **left** – Distance of the top-left corner of the box from the left side of the image, measured in pixels.
- **top** – Distance of the top-left corner of the box from the top of the image, measured in pixels.
- **width** – The width of the box in pixels.

You can extract the bounding box initial value from a manifest file of a previous job in a custom template using the Liquid templating language:

```liquid
initial-value="[
    {% for box in task.input.manifestLine.label-attribute-name-from-prior-job.annotations %}
        {% capture class_id %}{{ box.class_id }}{% endcapture %}
        {% assign label = task.input.manifestLine.label-attribute-name-from-prior-job-metadata.class-map[class_id] %}
    {% endfor %}
]
```
labels

A JSON formatted array of strings, each of which is a label that a worker can assign to the image portion enclosed by a rectangle. **Limit:** 10 labels.

name

The name of this widget. It's used as a key for the widget's input in the form output.

src

The URL of the image on which to draw bounding boxes.

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements:** crowd-form
- **Child elements:** full-instructions, short-instructions

Regions

The following regions are required by this element.

full-instructions

General instructions about how to draw bounding boxes.

short-instructions

Important task-specific instructions that are displayed in a prominent place.
Output

The following output is supported by this element.

**boundingBoxes**

An array of JSON objects, each of which specifies a bounding box that has been created by the worker. Each JSON object in the array contains the following properties.

- **height** – The height of the box in pixels.
- **label** – The text assigned to the box as part of the labeling task. This text must match one of the labels defined in the `labels` attribute of the `<crowd-bounding-box>` element.
- **left** – Distance of the top-left corner of the box from the left side of the image, measured in pixels.
- **top** – Distance of the top-left corner of the box from the top of the image, measured in pixels.
- **width** – The width of the box in pixels.

**inputImageProperties**

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.

**Example: Sample Element Outputs**

The following are samples of outputs from common use scenarios for this element.

**Single Label, Single Box / Multiple Label, Single Box**

```json
[
  {
    "annotatedResult": {
      "boundingBoxes": [
        {
          "height": 401,
          "label": "Dog",
```
Single Label, Multiple Box

[ {
   "annotatedResult": {
      "boundingBoxes": [
      {
         "height": 401,
         "label": "Dog",
         "left": 243,
         "top": 117,
         "width": 187
      },
      {
         "height": 283,
         "label": "Dog",
         "left": 684,
         "top": 120,
         "width": 116
      }
      ],
   "inputImageProperties": {
      "height": 533,
      "width": 800
   }
   }
} ]

Multiple Label, Multiple Box
[{
  "annotatedResult": {
    "boundingBoxes": [
      {
        "height": 395,
        "label": "Dog",
        "left": 241,
        "top": 125,
        "width": 158
      },
      {
        "height": 298,
        "label": "Cat",
        "left": 699,
        "top": 116,
        "width": 101
      }
    ],
    "inputImageProperties": {
      "height": 533,
      "width": 800
    }
  }
}]

You could have many labels available, but only the ones that are used appear in the output.

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

**crowd-card**

A box with an elevated appearance for displaying information.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.
The following is an example of a template designed for sentiment analysis tasks that uses the `<crowd-card>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<style>
  h3 {
    margin-top: 0;
  }

  crowd-card {
    width: 100%;
  }

  .card {
    margin: 10px;
  }

  .left {
    width: 70%;
    margin-right: 10px;
    display: inline-block;
    height: 200px;
  }

  .right {
    width: 20%;
    height: 200px;
    display: inline-block;
  }
</style>

<crowd-form>
  <short-instructions>
    Your short instructions here.
  </short-instructions>

  <full-instructions>
    Your full instructions here.
  </full-instructions>

  <div class="left">
    SageMaker Crowd HTML Elements
  </div>
</crowd-form>
```
<h3>What sentiment does this text convey?</h3>
<crowd-card>
  <div class="card">
    Nothing is great.
  </div>
</crowd-card>
</div>

<div class="right">
<h3>Select an option</h3>
<select name="sentiment1" style="font-size: large" required>
  <option value="">(Please select)</option>
  <option>Negative</option>
  <option>Neutral</option>
  <option>Positive</option>
  <option>Text is empty</option>
</select>
</div>

<div class="left">
<h3>What sentiment does this text convey?</h3>
<crowd-card>
  <div class="card">
    Everything is great!
  </div>
</crowd-card>
</div>

<div class="right">
<h3>Select an option</h3>
<select name="sentiment2" style="font-size: large" required>
  <option value="">(Please select)</option>
  <option>Negative</option>
  <option>Neutral</option>
  <option>Positive</option>
  <option>Text is empty</option>
</select>
</div>

</crowd-form>
Attributes

The following attributes are supported by this element.

**heading**

The text displayed at the top of the box.

**image**

A URL to an image to be displayed within the box.

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: `crowd-form`
- **Child elements**: none

See Also

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

crowd-checkbox

A UI component that can be checked or unchecked allowing a user to select multiple options from a set.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).

The following is an example of a Liquid template that uses the `<crowd-checkbox>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
Attributes

The following attributes are supported by this element.

checked

A Boolean switch that, if present, displays the check box as checked.

The following is an example of the syntax used to check a checkbox by default.

```html
<crowd-checkbox name="checkedBox" value="checked" checked>This box is checked</crowd-checkbox>
```

disabled

A Boolean switch that, if present, displays the check box as disabled and prevents it from being checked.

The following is an example of the syntax used to disable a checkbox.

```html
<crowd-checkbox name="disabledCheckBox" value="Disabled" disabled>Cannot be selected</crowd-checkbox>
```

name

A string that is used to identify the answer submitted by the worker. This value will match a key in the JSON object that specifies the answer.

required

A Boolean switch that, if present, requires the worker to provide input.
The following is an example of the syntax used to require a checkbox be selected.

```html
<crowd-checkbox name="work_verified" required>Instructions were clear</crowd-checkbox>
```

**value**

A string used as the name for the check box state in the output. Defaults to "on" if not specified.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: `crowd-form`
- **Child elements**: none

**Output**

Provides a JSON object. The name string is the object name and the value string is the property name for a Boolean value based on the check box state; true if checked, false if not checked.

**Example: Sample Element Outputs**

**Using the same name value for multiple boxes.**

```html
<!-- INPUT -->
<div><crowd-checkbox name="image_attributes" value="blurry"> Blurry </crowd-checkbox></div>
<div><crowd-checkbox name="image_attributes" value="dim"> Too Dim </crowd-checkbox></div>
<div><crowd-checkbox name="image_attributes" value="exposed"> Too Bright </crowd-checkbox></div>

//Output with "blurry" and "dim" checked
[
    {
        "image_attributes": {
            "blurry": true,
            "dim": true,
            "exposed": false
        }
    }
]
Note that all three color values are properties of a single object.

**Using different name values for each box.**

```html
<!-- INPUT -->
<div><crowd-checkbox name="Stop" value="Red"> Red </crowd-checkbox></div>
<div><crowd-checkbox name="Slow" value="Yellow"> Yellow </crowd-checkbox></div>
<div><crowd-checkbox name="Go" value="Green"> Green </crowd-checkbox></div>

//Output with "Red" checked
[
  {
    "Go": {
      "Green": false
    },
    "Slow": {
      "Yellow": false
    },
    "Stop": {
      "Red": true
    }
  }
]
```

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

**crowd-classifier**

A widget for classifying non-image content, such as audio, video, or text.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).
Amazon SageMaker

Developer Guide

The following is an example of an HTML worker task template built using crowd-classifier.
This example uses the Liquid template language to automate:
• Label categories in the categories parameter
• The objects that are being classiﬁed in the classification-target parameter.
Copy the following code and save it in a ﬁle with the extension .html. Open the ﬁle in any browser
to preview and interact with this template.
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<crowd-classifier
name="category"
categories="{{ task.input.labels | to_json | escape }}"
header="What type of a document is this?"
>
<classification-target>
<iframe style="width: 100%; height: 600px;" src="{{ task.input.taskObject
grant_read_access }}" type="application/pdf"></iframe>
</classification-target>

|

<full-instructions header="Document Classification Instructions">
<p>Read the task carefully and inspect the document.</p>
<p>Choose the appropriate label that best suits the document.</p>
</full-instructions>
<short-instructions>
Please choose the correct category for the document
</short-instructions>
</crowd-classifier>
</crowd-form>

Attributes
The following attributes are supported by this element.
categories
A JSON formatted array of strings, each of which is a category that a worker can assign to the
text. You should include "other" as a category, otherwise the worker my not be able to provide an
answer.
SageMaker Crowd HTML Elements

1881


header

The text to display above the image. This is typically a question or simple instruction for the worker.

name

The name of this widget. It is used as a key for the widget's input in the form output.

Element Hierarchy

This element has the following parent and child elements.

- Parent elements: crowd-form
- Child elements: classification-target, full-instructions, short-instructions

Regions

The following regions are supported by this element.

classification-target

The content to be classified by the worker. This can be plain text or HTML. Examples of how the HTML can be used include but are not limited to embedding a video or audio player, embedding a PDF, or performing a comparison of two or more images.

full-instructions

General instructions about how to do text classification.

short-instructions

Important task-specific instructions that are displayed in a prominent place.

Output

The output of this element is an object using the specified name value as a property name, and a string from the categories as the property's value.

Example: Sample Element Outputs

The following is a sample of output from this element.

[ ]
crowd-classifier-multi-select

A widget for classifying various forms of content—such as audio, video, or text—into one or more categories. The content to classify is referred to as an object.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of an HTML worker task template built using this element. Copy the following code and save it in a file with the extension .html. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-classifier-multi-select
    name="category"
    categories="['Positive', 'Negative', 'Neutral']"
    header="Select the relevant categories"
    exclusion-category={ "text": 'None of the above' }"
  >
    <classification-target>
      {{ task.input.taskObject }}
    </classification-target>
  </crowd-classifier-multi-select>

  <full-instructions header="Text Categorization Instructions">
    <p><strong>Positive</strong> sentiment include: joy, excitement, delight</p>
    <p><strong>Negative</strong> sentiment include: anger, sarcasm, anxiety</p>
  </full-instructions>
</crowd-form>
```
Attributes

The following attributes are supported by the `crowd-classifier-multi-select` element. Each attribute accepts a string value or string values.

**categories**

Required. A JSON-formatted array of strings, each of which is a category that a worker can assign to the object.

**header**

Required. The text to display above the image. This is typically a question or simple instruction for workers.

**name**

Required. The name of this widget. In the form output, the name is used as a key for the widget's input.

**exclusion-category**

Optional. A JSON-formatted string with the following format: `"{ text: 'default-value' }"`. This attribute sets a default value that workers can choose if none of the labels applies to the object shown in the worker UI.

**Element Hierarchy**

This element has the following parent and child elements:

- **Parent elements:** crowd-form
• **Child elements:** classification-target, full-instructions, short-instructions

**Regions**

This element uses the following regions.

**classification-target**

The content to be classified by the worker. Content can be plain text or an object that you specify in the template using HTML. For example, you can use HTML elements to include a video or audio player, embedding a PDF file, or include a comparison of two or more images.

**full-instructions**

General instructions about how to classify text.

**short-instructions**

Important task-specific instructions. These instructions are displayed prominently.

**Output**

The output of this element is an object that uses the specified name value as a property name, and a string from categories as the property's value.

**Example : Sample Element Outputs**

The following is a sample of output from this element.

```
[
  {
    "<name>": {
      "labels": ["label_a", "label_b"]
    }
  }
]
```

**See Also**

For more information, see the following:

- [Text Classification (Multi-label)]
- [Use Amazon SageMaker Ground Truth to Label Data]
**crowd-entity-annotation**

A widget for labeling words, phrases, or character strings within a longer text. Workers select a label, and highlight the text that the label applies to.

---

**Important: Self-contained Widget**

Do not use `<crowd-entity-annotation>` element with the `<crowd-form>` element. It contains its own form submission logic and **Submit** button.

---

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](https://codepen.io).

The following is an example of a template that uses the `<crowd-entity-annotation>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-entity-annotation
  name="crowd-entity-annotation"
  header="Highlight parts of the text below"
  labels="[\{\'label\': \'person\', \'shortDisplayName\': \'per\', \'fullDisplayName\': \'Person\'},\n  \{\'label\': \'date\', \'shortDisplayName\': \'dat\', \'fullDisplayName\': \'Date\'},\n  \{\'label\': \'company\', \'shortDisplayName\': \'com\', \'fullDisplayName\': \'Company\'}]\"
  text="Amazon SageMaker Ground Truth helps you build highly accurate training datasets for machine learning quickly."
>
  <full-instructions header="Named entity recognition instructions">
    <ol>
      <li><strong>Read</strong> the text carefully.</li>
      <li><strong>Highlight</strong> words, phrases, or sections of the text.</li>
      <li><strong>Choose</strong> the label that best matches what you have highlighted.</li>
      <li>To <strong>change</strong> a label, choose highlighted text and select a new label.</li>
      <li>To <strong>remove</strong> a label from highlighted text, choose the X next to the abbreviated label name on the highlighted text.</li>
    </ol>
  </full-instructions>

</crowd-entity-annotation>
```
You can select all of a previously highlighted text, but not a portion of it.

Apply labels to words or phrases.

What is the overall subject of this text?

Technology

Politics

The following attributes are supported by this element.

header

The text to display above the image. This is typically a question or simple instruction for the worker.
initial-value

A JSON formatted array of objects, each of which defines an annotation to apply to the text at initialization. Objects contain a label value that matches one in the labels attribute, an integer startOffset value for labeled span's starting unicode offset, and an integer endOffset value for the ending unicode offset.

Example

```json
[
  {
    label: 'person',
    startOffset: 0,
    endOffset: 16
  },
  ...
]
```

labels

A JSON formatted array of objects, each of which contains:

- **label** (required): The name used to identify entities.
- **fullDisplayName** (optional): Used for the label list in the task widget. Defaults to the label value if not specified.
- **shortDisplayName** (optional): An abbreviation of 3-4 letters to display above selected entities. Defaults to the label value if not specified.

Information:

ℹ️ **shortDisplayName is highly recommended**

Values displayed above the selections can overlap and create difficulty managing labeled entities in the workspace. Providing a 3-4 character shortDisplayName for each label is highly recommended to prevent overlap and keep the workspace manageable for your workers.

Example

```json
[
  {

```
name

Serves as the widget's name in the DOM. It is also used as the label attribute name in form output and the output manifest.

text

The text to be annotated. The templating system escapes quotes and HTML strings by default. If your code is already escaped or partially escaped, see Variable filters for more ways to control escaping.

Element Hierarchy

This element has the following parent and child elements.

- **Child elements**: full-instructions, short-instructions

Regions

The following regions are supported by this element.

full-instructions

General instructions about how to work with the widget.

short-instructions

Important task-specific instructions that are displayed in a prominent place.

Output

The following output is supported by this element.

entities

A JSON object that specifies the start, end, and label of an annotation. This object contains the following properties.
- **label** – The assigned label.
- **startOffset** – The Unicode offset of the beginning of the selected text.
- **endOffset** – The Unicode offset of the first character after the selection.

**Example: Sample Element Outputs**

The following is a sample of the output from this element.

```json
{
  "myAnnotatedResult": {
    "entities": [
      {
        "endOffset": 54,
        "label": "person",
        "startOffset": 47
      },
      {
        "endOffset": 97,
        "label": "event",
        "startOffset": 93
      },
      {
        "endOffset": 219,
        "label": "date",
        "startOffset": 212
      },
      {
        "endOffset": 271,
        "label": "location",
        "startOffset": 260
      }
    ]
  }
}
```

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)
crowd-fab

A floating button with an image in its center.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of a Liquid template designed for image classification that uses the <crowd-fab> element. This template uses JavaScript to enable workers to report issues with the worker UI. Copy the following code and save it in a file with the extension .html. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
  <crowd-image-classifier
    src="${image_url}"
    categories="['Cat', 'Dog', 'Bird', 'None of the Above']"
    header="Choose the correct category for the image"
    name="category">
    <short-instructions>
      <p>Read the task carefully and inspect the image.</p>
      <p>Choose the appropriate label that best suits the image.</p>
      <p>If there is an issue with the image or tools, please select</p>
      <b>None of the Above</b>, describe the issue in the text box and click
      the button below.</p>
    </short-instructions>
  
  <crowd-input label="Report an Issue" name="template-issues"></crowd-input>
  <crowd-fab id="button1" icon="report-problem" title="Issue"/>
    
  </short-instructions>
</crowd-image-classifier>
</crowd-form>
```
button1,
].forEach(function(button) {
    button.addEventListener('click', function() {
        document.querySelector('crowd-form').submit();
    });
});
</script>

Attributes

The following attributes are supported by this element.

**disabled**

A Boolean switch that, if present, displays the floating button as disabled and prevents clicks.

**icon**

A string that specifies the icon to be displayed in the center of the button. The string must be either the name of an icon from the open-source *iron-icons* set, which is pre-loaded, or the URL to a custom icon.

The following is an example of the syntax that you can use to add an iron-icon to a `<crowd-fab>` HTML element. Replace `icon-name` with the name of the icon you'd like to use from this [Icons set](#)

```html
<crowd-fab "id="button1" icon="icon-name" title="Issue"/>
```

**label**

A string consisting of a single character that can be used instead of an icon. Emojis or multiple characters may result in the button displaying an ellipsis instead.

**title**

A string that will display as a tool tip when the mouse hovers over the button.

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-form
• **Child elements**: none

See Also

For more information, see the following.

• [Use Amazon SageMaker Ground Truth to Label Data](#)
• [Crowd HTML Elements Reference](#)

**crowd-form**

The form wrapper for all custom tasks. Sets and implements important actions for the proper submission of your form data.

If a `crowd-button` of type "submit" is not included inside the `<crowd-form>` element, it will automatically be appended within the `<crowd-form>` element.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).

The following is an example of an image classification template that uses the `<crowd-form>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-image-classifier
    src="${image_url}"
    categories="['Cat', 'Dog', 'Bird', 'None of the Above']"
    header="Choose the correct category for the image"
    name="category">

    <short-instructions>
      <p>Read the task carefully and inspect the image.</p>
      <p>Choose the appropriate label that best suits the image.</p>
    </short-instructions>

    <full-instructions header="Classification Instructions">

  </full-instructions>
</crowd-form>
```
Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: none
- **Child elements**: Any of the [UI Template] elements

Element Events

The `<crowd-form>` element extends the [standard HTML form element] and inherits its events, such as `onclick` and `onsubmit`.

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data]
- [Crowd HTML Elements Reference]

**crowd-icon-button**

A button with an image placed in the center. When the user touches the button, a ripple effect emanates from the center of the button.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen].

The following is an example of a Liquid template designed for image classification that uses the `<crowd-icon-button>` element. This template uses JavaScript to enable workers to report issues with the worker UI. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.
Attributes

The following attributes are supported by this element.
**disabled**

A Boolean switch that, if present, displays the button as disabled and prevents clicks.

**icon**

A string that specifies the icon to be displayed in the center of the button. The string must be either the name of an icon from the open-source iron-icons set, which is pre-loaded, or the URL to a custom icon.

The following is an example of the syntax that you can use to add an iron-icon to a `<crowd-icon-button>` HTML element. Replace `icon-name` with the name of the icon you'd like to use from this Icons set.

```html
<crowd-icon-button id="button1" icon="icon-name" title="Issue"/>
```

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: none

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

**crowd-image-classifier**

A widget for classifying an image. Use one of the following supported image formats: APNG, BMP, GIF, ICO, JPEG, PNG, SVG. Images do not have a size limit.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).

The following is an example of an image classification template that uses the `<crowd-image-classifier>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.
Attributes

The following attributes are required by this element.

**categories**

A JSON formatted array of strings, each of which is a category that a worker can assign to the image. You should include "other" as a category, so that the worker can provide an answer. You can specify up to 10 categories.

**header**

The text to display above the image. This is typically a question or simple instruction for the worker.

**name**

The name of this widget. It is used as a key for the widget's input in the form output.
overlay

Information to be overlaid on the source image. This is for verification workflows of bounding-box, semantic-segmentation, and instance-segmentation tasks.

It is a JSON object containing an object with the name of the task-type in camelCase as the key. That key's value is an object that contains the labels and other necessary information from the previous task.

An example of a crowd-image-classifier element with attributes for verifying a bounding-box task follows:

```xml
<crowd-image-classifier
   name="boundingBoxClassification"
   header="Rate the quality of the annotations based on the background section in the instructions on the left hand side."
   src="https://i.imgur.com/CIPKVJo.jpg"
   categories="["good", "bad", "okay"]"
   overlay='{
       "boundingBox": {
           labels: ["bird", "cat"],
           value: [
               {
                   height: 284,
                   label: "bird",
                   left: 230,
                   top: 974,
                   width: 223
               },
               {
                   height: 69,
                   label: "bird",
                   left: 79,
                   top: 889,
                   width: 247
               }
           ]
       }
   }
>

A semantic segmentation verification task would use the overlay value as follows:
An instance-segmentation task would use the overlay value as follows:

```
<crowd-image-classifier
name='crowd-image-classifier'
categories='["good", "bad"]'
src='URL of image to be classified'
header='Please classify instances of each category'
overlay='{
  "instanceSegmentation": {
    "labels": ["Cat", "Dog", "Bird", "Cow"],
    "instances": [
      {
        "color": "#2ca02c",
        "label": "Cat"
      },
      {
        "color": "#2acf59",
        "label": "Dog"
      }
    ],
    "src": "URL of overlay image",
  }
}

> ... </crowd-image-classifier>
```
The URL of the image to be classified.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: full-instructions, short-instructions, worker-comment

**Regions**

The following regions are used by this element.

**full-instructions**

General instructions for the worker on how to classify an image.

**short-instructions**

Important task-specific instructions that are displayed in a prominent place.

**worker-comment**

Use this in verification workflows when you need workers to explain why they made the choice they did. Use the text between the opening and closing tags to provide instructions for workers on what information should be included in the comment.

It uses the following attributes:
header

A phrase with a call to action for leaving a comment. Used as the title text for a modal window where the comment is added.

Optional. Defaults to "Add a comment."

link-text

This text appears below the categories in the widget. When clicked, it opens a modal window where the worker may add a comment.

Optional. Defaults to "Add a comment."

placeholder

An example text in the comment text area that is overwritten when worker begins to type. This does not appear in output if the worker leaves the field blank.

Optional. Defaults to blank.

Output

The output of this element is a string that specifies one of the values defined in the categories attribute of the <crowd-image-classifier> element.

Example : Sample Element Outputs

The following is a sample of output from this element.

```json
[
  {
    "<name>": {
      "label": "<value>"
      "workerComment": "Comment - if no comment is provided, this field will not be present"
    }
  }
]
```

See Also

For more information, see the following.
crowd-image-classifier-multi-select

A widget for classifying an image into one or more categories. Use one of the following supported image formats: APNG, BMP, GIF, ICO, JPEG, PNG, SVG. Images do not have a size limit.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of an HTML worker task template built using this crowd element. Copy the following code and save it in a file with the extension .html. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-image-classifier-multi-select
      name="animals"
      categories="['Cat', 'Dog', 'Horse', 'Pig', 'Bird']"
      src="https://images.unsplash.com/photo-1509205477838-a534e43a849f?
      ixlib=rb-1.2.1&ixid=eyJhcHBfaWQiOjEyMDd9&auto=format&fit=crop&w=1998&q=80"
      header="Please identify the animals in this image"
      exclusion-category="{ text: 'None of the above' }"
  >
    <full-instructions header="Classification Instructions">
      <p>If more than one label applies to the image, select multiple labels.</p>
      <p>If no labels apply, select <b>None of the above</b></p>
    </full-instructions>

    <short-instructions>
      <p>Read the task carefully and inspect the image.</p>
      <p>Choose the appropriate label(s) that best suit the image.</p>
    </short-instructions>

  </crowd-image-classifier-multi-select>
</crowd-form>
```

Attributes

The following attributes are supported by the crowd-image-classifier-multi-select element. Each attribute accepts a string value or string values.
categories

Required. A JSON-formatted array of strings, each of which is a category that a worker can assign to the image. A worker must choose at least one category and can choose all categories.

header

Required. The text to display above the image. This is typically a question or simple instruction for workers.

name

Required. The name of this widget. In the form output, the name is used as a key for the widget's input.

src

Required. The URL of the image to be classified.

exclusion-category

Optional. A JSON-formatted string with the following format: "\{ text: 'default-value' \}". This attribute sets a default value that workers can choose if none of the labels applies to the image shown in the worker UI.

Element Hierarchy

This element has the following parent and child elements:

- **Parent elements**: crowd-form
- **Child elements**: full-instructions, short-instructions, worker-comment

Regions

This element uses the following regions

full-instructions

General instructions for the worker on how to classify an image.

short-instructions

Important task-specific instructions. These instructions are displayed prominently.
Output

The output of this element is a string that specifies one or more of the values defined in the categories attribute of the `<crowd-image-classifier-multi-select>` element.

Example: Sample Element Outputs

The following is a sample of output from this element.

```json
[
  {
    "name": {
      labels: ["label_a", "label_b"]
    }
  }
]
```

See Also

For more information, see the following:

- [Image Classification (Multi-label)]
- [Use Amazon SageMaker Ground Truth to Label Data]
- [Crowd HTML Elements Reference]

**crowd-input**

A box that accepts input data.

⚠️ Cannot be self-closing

Unlike the `input` element in the HTML standard, this element cannot be self-closed by putting a slash before the ending bracket, e.g. `<crowd-input ... />`. It must be followed with a `<crowd-input>` to close the element.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](https://codepen.io).

The following is an example of a Liquid template that uses the `<crowd-input>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.
Attributes

The following attributes are supported by this element.

**allowed-pattern**

A regular expression that is used with the `auto-validate` attribute to ignore non-matching characters as the worker types.

**auto-focus**

When the value is set to true, the browser places focus inside the input area after loading. This way, the worker can start typing without having to select it first.

**auto-validate**

A Boolean switch that, if present, turns on input validation. The behavior of the validator can be modified by the `error-message` and `allowed-pattern` attributes.

**disabled**

A Boolean switch that, if present, displays the input area as disabled.

**error-message**

The text to be displayed below the input field, on the left side, if validation fails.
**label**

A string that is displayed inside a text field. This text shrinks and rises up above a text field when the worker starts typing in the field or when the `value` attribute is set.

**max-length**

A maximum number of characters the input will accept. Input beyond this limit is ignored.

**min-length**

A minimum length for the input in the field

**name**

Sets the name of the input to be used in the DOM and the output of the form.

**placeholder**

A string value that is used as placeholder text, displayed until the worker starts entering data into the input. It is not used as a default value.

**required**

A Boolean switch that, if present, requires the worker to provide input.

**type**

Takes a string to set the HTML5 `input-type` behavior for the input. Examples include `file` and `date`.

**value**

A preset that becomes the default if the worker does not provide input. The preset appears in a text field.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: `crowd-form`
- **Child elements**: none
Output

Provides a name string as the property name, and the text that was entered in the field as its value.

Example: Sample JSON Output

The values for multiple elements are output in the same object, with their name attribute value as their property name. Elements with no input do not appear in the output. For example, let's use three inputs:

```html
<crowd-input name="tag1" label="Word/phrase 1"></crowd-input>
<crowd-input name="tag2" label="Word/phrase 2"></crowd-input>
<crowd-input name="tag3" label="Word/phrase 3"></crowd-input>
```

This is the output if only two have input:

```json
[
  {
    "tag1": "blue",
    "tag2": "red"
  }
]
```

This means any code built to parse these results should be able to handle the presence or absence of each input in the answers.

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

**crowd-instance-segmentation**

A widget for identifying individual instances of specific objects within an image and creating a colored overlay for each labeled instance.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.
The following is an example of a Liquid template that uses the `<crowd-instance-segmentation>`. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-instance-segmentation
    name="annotatedResult"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="Please label each of the requested objects in this image"
    labels="['Cat', 'Dog', 'Bird']"
  >
    <full-instructions header="Segmentation Instructions">
      <ol>
        <li><strong>Read</strong> the task carefully and inspect the image.</li>
        <li><strong>Read</strong> the options and review the examples provided to understand more about the labels.</li>
        <li><strong>Choose</strong> the appropriate label that best suits the image.</li>
      </ol>
    </full-instructions>

    <short-instructions>
      <p>Use the tools to label all instances of the requested items in the image</p>
    </short-instructions>
  </crowd-instance-segmentation>
</crowd-form>

Use a template similar to the following to allow workers to add their own categories (labels).

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-instance-segmentation
    id="annotator"
    name="myTexts"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="Click Instructions to add new labels."
    labels="['placeholder']"
  >
    <short-instructions>
      <h3>Add a label to describe each type of object in this image.</h3>
      <h3>Cover each instance of each object with a segmentation mask.</h3>
    </short-instructions>
  </crowd-instance-segmentation>
</crowd-form>
```
<br>
<h3>Add new label</h3>
<crowd-input name="_customLabel" id="customLabel"></crowd-input>
<crowd-button id="addLabel">Add</crowd-button>

<br><br><br>
<h3>Manage labels</h3>
<div id="labelsSection"></div>
</crowd-instance-segmentation>
</crowd-form>

<script>
document.addEventListener('all-crowd-elements-ready', function(event) {
    document.querySelector('crowd-instance-segmentation').labels = [];
});

function populateLabelsSection() {
    labelsSection.innerHTML = '';  
    annotator.labels.forEach(function(label) {
        const labelContainer = document.createElement('div');
        labelContainer.innerHTML = label + ' <a href="javascript:void(0)">(Delete)</a>';
        labelContainer.querySelector('a').onclick = function() {
            return 1 !== label;
        }
        populateLabelsSection();
    });
    labelsSection.appendChild(labelContainer);
});

addLabel.onclick = function() {
    annotator.labels = annotator.labels.concat([customLabel.value]);
    customLabel.value = null;
</script>
Attributes

The following attributes are supported by this element.

header

The text to display above the image. This is typically a question or simple instruction for the worker.

labels

A JSON formatted array of strings, each of which is a label that a worker can assign to an instance of an object in the image. Workers can generate different overlay colors for each relevant instance by selecting "add instance" under the label in the tool.

name

The name of this widget. It is used as a key for the labeling data in the form output.

src

The URL of the image that is to be labeled.

initial-value

A JSON object containing the color mappings of a prior instance segmentation job and a link to the overlay image output by the prior job. Include this when you want a human worker to verify the results of a prior labeling job and adjust it if necessary.

The attribute will appear as follows:

```javascript
initial-value="{
  "instances": [
    {
      "color": "#2ca02c",
      "label": "Cat"
    },
    {
```
Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: full-instructions, short-instructions

Regions

The following regions are supported by this element.

**full-instructions**

General instructions about how to do image segmentation.

**short-instructions**

Important task-specific instructions that are displayed in a prominent place.

Output

The following output is supported by this element.

**labeledImage**

A JSON Object containing a Base64 encoded PNG of the labels.

**instances**

A JSON Array containing objects with the instance labels and colors.
- **color** – The hexadecimal value of the label's RGB color in the `labeledImage` PNG.
- **label** – The label given to overlay(s) using that color. This value may repeat, because the different instances of the label are identified by their unique color.

**inputImageProperties**

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.

**Example: Sample Element Outputs**

The following is an example of output from this element.

```
[
  {
    "annotatedResult": {
      "inputImageProperties": {
        "height": 533,
        "width": 800
      },
      "instances": [
        {
          "color": "#1f77b4",
          "label": "<Label 1>"
        },
        {
          "color": "#2ca02c",
          "label": "<Label 1>"
        },
        {
          "color": "#ff7f0e",
          "label": "<Label 3>"
        },
      ],
      "labeledImage": {
        "pngImageData": "<Base-64 Encoded Data>"
      }
    }
  }
]
```
Amazon SageMaker

Developer Guide

}
]

See Also
For more information, see the following.
• Use Amazon SageMaker Ground Truth to Label Data
• Crowd HTML Elements Reference

crowd-instructions
An element that displays instructions on three tabbed pages, Summary, Detailed Instructions, and
Examples, when the worker clicks on a link or button.
See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.
The following is an example of a Liquid template that used the <crowd-instructions> element.
Copy the following code and save it in a ﬁle with the extension .html. Open the ﬁle in any browser
to preview and interact with this template.
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<crowd-instructions link-text="View instructions" link-type="button">
<short-summary>
<p>Given an image, write three words or short phrases that summarize its
contents.</p>
</short-summary>
<detailed-instructions>
<p>Imagine that you are describing an image to a friend or tagging it for a news
website. Provide three specific words or short phrases that describe it.</p>
</detailed-instructions>
<positive-example>
<p>
<ul>
<li>Highway</li>
<li>Cars</li>
<li>Gas station</li>
</ul>
</p>
SageMaker Crowd HTML Elements

1913


These are not specific enough:

- Trees
- Outside
- Daytime

Instructions: Given an image, write three words or short phrases that summarize its contents.

- If someone were to see these three words or phrases, they should understand the subject and context of the image, as well as any important actions.
- View the instructions for detailed instructions and examples.

Attributes

The following attributes are supported by this element.

**link-text**

The text to display for opening the instructions. The default is **Click for instructions**.

**link-type**

A string that specifies the type of trigger for the instructions. The possible values are "link" (default) and "button".

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: none
Regions

The following regions are supported by this element.

detailed-instructions

Content that provides specific instructions for a task. This appears on the page of the "Detailed Instructions" tab.

negative-example

Content that provides examples of inadequate task completion. This appears on the page of the "Examples" tab. More than one example may be provided within this element.

positive-example

Content that provides examples of proper task completion. This appears on the page of the "Examples" tab.

short-summary

A brief statement that summarizes the task to be completed. This appears on the page of the "Summary" tab. More than one example may be provided within this element.

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

crowd-keypoint

Generates a tool to select and annotate key points on an image.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of an Liquid template that uses the <crowd-keypoint> element. Copy the following code and save it in a file with the extension .html. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
```
<crowd-form>
  <div id="errorBox"></div>

  <crowd-keypoint
    src="{{ task.input.taskObject | grant_read_access }}"
    labels="["Item A", "Item B", "Item C"]"
    header="Please locate the centers of each item."
    name="annotatedResult">
    <short-instructions>
      Describe your task briefly here and give examples
    </short-instructions>
    <full-instructions>
      Give additional instructions and good/bad examples here
    </full-instructions>
  </crowd-keypoint>
</crowd-form>

<script>
  var num_obj = 1;

  document.querySelector('crowd-form').onsubmit = function(e) {
    const keypoints = document.querySelector('crowd-keypoint').value.keypoints ||
    document.querySelector('crowd-keypoint')._submittableValue.keypoints;
    const labels = keypoints.map(function(p) {
      return p.label;
    });

    // 1. Make sure total number of keypoints is correct.
    var original_num_labels = document.getElementsByTagName("crowd-keypoint")
    [0].getAttribute("labels");

    original_num_labels = original_num_labels.substring(2, original_num_labels.length - 2).split("", ",");
    var goalNumKeypoints = num_obj*original_num_labels.length;
    if (keypoints.length != goalNumKeypoints) {
      e.preventDefault();
      errorBox.innerHTML = '<crowd-alert type="error" dismissible>You must add all keypoint annotations and use each label only once.</crowd-alert>,'
      errorBox.scrollIntoView();
      return;
    }

    // 2. Make sure all labels are unique.
    labelCounts = {};
  }
</script>
for (var i = 0; i < labels.length; i++) {
    if (!labelCounts[labels[i]]) {
        labelCounts[labels[i]] = 0;
    }
    labelCounts[labels[i]]++;
}
const goalNumSingleLabel = num_obj;

const numLabels = Object.keys(labelCounts).length;

Object.entries(labelCounts).forEach(entry => {
    if (entry[1] != goalNumSingleLabel) {
        e.preventDefault();
        errorBox.innerHTML = '<crowd-alert type="error" dismissible>You must use each label only once.</crowd-alert>';
        errorBox.scrollIntoView();
    }
});
</script>

Attributes

The following attributes are supported by this element.

header

The text to display above the image. This is typically a question or simple instruction for the worker.

initial-value

An array, in JSON format, of keypoints to be applied to the image on start. For example:

```javascript
initial-value="[
  {
    'label': 'Left Eye',
    'x': 1022,
    'y': 429
  },
  {
    'label': 'Beak',
    'x': 941,
```
Note

Please note that label values used in this attribute must have a matching value in the labels attribute or the point will not be rendered.

labels

An array, in JSON format, of strings to be used as keypoint annotation labels.

name

A string used to identify the answer submitted by the worker. This value will match a key in the JSON object that specifies the answer.

src

The source URI of the image to be annotated.

Element Hierarchy

This element has the following parent and child elements.

- Parent elements: crowd-form
- Child elements: full-instructions, short-instructions

Regions

The following regions are required by this element.

full-instructions

General instructions about how to annotate the image.

short-instructions

Important task-specific instructions that are displayed in a prominent place.
Output

The following output is supported by this element.

**inputImageProperties**

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.

**keypoints**

An array of JSON objects containing the coordinates and label of a keypoint. Each object contains the following properties.

- **label** – The assigned label for the keypoint.
- **x** – The X coordinate, in pixels, of the keypoint on the image.
- **y** – The Y coordinate, in pixels, of the keypoint on the image.

**Note**

X and Y coordinates are based on 0,0 being the top left corner of the image.

**Example : Sample Element Outputs**

The following is a sample output from using this element.

```json
[
  {
    "crowdKeypoint": {
      "inputImageProperties": {
        "height": 1314,
        "width": 962
      },
      "keypoints": [
        {
```
You may have many labels available, but only the ones that are used appear in the output.

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference
Amazon SageMaker

Developer Guide

crowd-line
A widget for drawing lines on an image. Each line is associated with a label, and output data will
report the starting and ending points of each line.
See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.
The following is an example of a Liquid template that uses the <crowd-line> element. Copy
the following code and save it in a ﬁle with the extension .html. Open the ﬁle in any browser to
preview and interact with this template. For more examples, see this GitHub repository.
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<crowd-line
name="crowdLine"
src="{{ task.input.taskObject | grant_read_access }}"
header="Add header here to describe the task"
labels="['car','pedestrian','street car']"
>
<short-instructions>
<p>Read the task carefully and inspect the image.</p>
<p>Choose the appropriate label that best suits the image.</p>
<p>Draw a line on each objects that the label applies to.</p>
</short-instructions>
<full-instructions>
<p>Read the task carefully and inspect the image.</p>
<p>Choose the appropriate label that best suits the image.
<p>Draw a line along each object that the image applies to.
Make sure that the line does not extend beyond the boundaries
of the object.
</p>
<p>Each line is defined by a starting and ending point. Carefully
place the starting and ending points on the boundaries of the object.</p>
</full-instructions>
</crowd-line>
</crowd-form>

Attributes
The following attributes are supported by this element.
SageMaker Crowd HTML Elements

1921


**header**

Optional. The text to display above the image. This is typically a question or simple instruction for the worker.

**initial-value**

Optional. An array of JSON objects, each of which sets a line when the component is loaded. Each JSON object in the array contains the following properties:

- **label** – The text assigned to the line as part of the labeling task. This text must match one of the labels defined in the `labels` attribute of the `<crowd-line>` element.
- **vertices** – the x and y pixel coordinates of the start point and end point of the line, relative to the top-left corner of the image.

```json
initial-value="{
  lines: [
    {
      label: 'sideline', // label of this line annotation
      vertices:[         // an array of vertices which decide the position of the line
        { x: 84,
          y: 110
        },
        { x: 60,
          y: 100
        }
      ]
    },
    {
      label: 'yardline',
      vertices:[
        { x: 651,
          y: 498
        },
        { x: 862,
          y: 869
        }
      ]
    }
  ]
}
```
Lines set via the `initial-value` property can be adjusted. Whether or not a worker answer was adjusted is tracked via an `initialValueModified` boolean in the worker answer output.

**labels**

Required. A JSON formatted array of strings, each of which is a label that a worker can assign to the line.

**Limit:** 10 labels

**label-colors**

Optional. An array of strings. Each string is a hexadecimal (hex) code for a label.

**name**

Required. The name of this widget. It's used as a key for the widget's input in the form output.

**src**

Required. The URL of the image on which to draw lines.

**Regions**

The following regions are required by this element.

**full-instructions**

General instructions about how to draw lines.

**short-instructions**

Important task-specific instructions that are displayed in a prominent place.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements:** crowd-form
- **Child elements:** short-instructions, full-instructions
Output

inputImageProperties

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.

lines

A JSON Array containing objects with the line labels and vertices.

- **label** – The label given to a line.
- **vertices** – the x and y pixel coordinates of the start point and end point of the line, relative to the top-left corner of the image.

Example: Sample Element Outputs

The following is an example of output from this element.

```json
{
   "crowdLine": { //This is the name you set for the crowd-line
      "inputImageProperties": {
         "height": 1254,
         "width": 2048
      },
      "lines": [
         {
            "label": "yardline",
            "vertices": [
               {
                  "x": 58,
                  "y": 295
               },
               {
                  "x": 1342,
                  "y": 398
               }
            ]
         }
      ]
   }
}
```
See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

**crowd-modal**

A small window that pops up on the display when it is opened.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](https://codepen.io).

The following is an example of the syntax that you can use with the `<crowd-modal>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-modal
link-text = "See Examples"
link-type = "button">
Example Modal Text</crowd-modal>
```
Attributes

The following attributes are supported by this element.

**link-text**

The text to display for opening the modal. The default is "Click to open modal".

**link-type**

A string that specifies the type of trigger for the modal. The possible values are "link" (default) and "button".

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: none

See Also

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

crowd-polygon

A widget for drawing polygons on an image and assigning a label to the portion of the image that is enclosed in each polygon.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).

The following is an example of a Liquid template that uses the `<crowd-polygon>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
```
<crowd-polygon
    name="annotatedResult"
    src="{{ task.input.taskObject | grant_read_access }}"
    header="Draw a polygon around each of the requested target(s) of interest"
    labels="['Cat', 'Dog', 'Bird']"
> 
    <full-instructions header="Polygon instructions">
        <ul>
            <li>Make the polygon tight around the object</li>
            <li>You need to select a label before starting a polygon</li>
            <li>You will need to select a label again after completing a polygon</li>
            <li>To select a polygon, you can click on its borders</li>
            <li>You can start drawing a polygon from inside another polygon</li>
            <li>You can undo and redo while you're drawing a polygon to go back and forth between points you've placed</li>
            <li>You are prevented from drawing lines that overlap other lines from the same polygon</li>
        </ul>
    </full-instructions>

    <short-instructions>
        <p>Draw a polygon around each of the requested target(s) of interest</p>
        <p>Make the polygon tight around the object</p>
    </short-instructions>
</crowd-polygon>

Attributes

The following attributes are supported by this element.

header

The text to display above the image. This is typically a question or simple instruction for the worker.

labels

A JSON formatted array of strings, each of which is a label that a worker can assign to the image portion enclosed by a polygon.

name

The name of this widget. It's used as a key for the widget's input in the form output.
**src**

The URL of the image on which to draw polygons.

**initial-value**

An array of JSON objects, each of which defines a polygon to be drawn when the component is loaded. Each JSON object in the array contains the following properties.

- **label** – The text assigned to the polygon as part of the labeling task. This text must match one of the labels defined in the `labels` attribute of the `<crowd-polygon>` element.
- **vertices** – An array of JSON objects. Each object contains an x and y coordinate value for a point in the polygon.

**Example**

An `initial-value` attribute might look something like this.

```javascript
initial-value = 
'[
  {
    "label": "dog",
    "vertices":
      [
        {
          "x": 570,
          "y": 239
        },
        ...
        {
          "x": 759,
          "y": 281
        }
      ]
  }
]
```

Because this will be within an HTML element, the JSON array must be enclosed in single or double quotes. The example above uses single quotes to encapsulate the JSON and double quotes within the JSON itself. If you must mix single and double quotes inside your JSON, replace them with their HTML entity codes (" for double quote, ’ for single) to safely escape them.
Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: [crowd-form](#)
- **Child elements**: [full-instructions](#), [short-instructions](#)

Regions

The following regions are required.

**full-instructions**

General instructions about how to draw polygons.

**short-instructions**

Important task-specific instructions that are displayed in a prominent place.

Output

The following output is supported by this element.

**polygons**

An array of JSON objects, each of which describes a polygon that has been created by the worker. Each JSON object in the array contains the following properties.

- **label** – The text assigned to the polygon as part of the labeling task.
- **vertices** – An array of JSON objects. Each object contains an x and y coordinate value for a point in the polygon. The top left corner of the image is 0,0.

**inputImageProperties**

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.
Example: Sample Element Outputs

The following are samples of outputs from common use scenarios for this element.

Single Label, Single Polygon

```json
{
    "annotatedResult": {
        "inputImageProperties": {
            "height": 853,
            "width": 1280
        },
        "polygons": [
            {
                "label": "dog",
                "vertices": [
                    {
                        "x": 570,
                        "y": 239
                    },
                    {
                        "x": 603,
                        "y": 513
                    },
                    {
                        "x": 823,
                        "y": 645
                    },
                    {
                        "x": 901,
                        "y": 417
                    },
                    {
                        "x": 759,
                        "y": 281
                    }
                ]
            }
        ]
    }
}
```
Single Label, Multiple Polygons

[  
  {  
    "annotatedResult": {  
      "inputImageProperties": {  
        "height": 853,  
        "width": 1280  
      },  
      "polygons": [  
        {  
          "label": "dog",  
          "vertices": [  
            {  
              "x": 570,  
              "y": 239  
            },  
            {  
              "x": 603,  
              "y": 513  
            },  
            {  
              "x": 823,  
              "y": 645  
            },  
            {  
              "x": 901,  
              "y": 417  
            },  
            {  
              "x": 759,  
              "y": 281  
            }  
          ]  
        },  
        {  
          "label": "dog",  
          "vertices": [  
            {  
              "x": 870,  
              "y": 278  
            }  
          ]  
        }  
      ]  
    }  
  ]
Multiple Labels, Multiple Polygons

```json
[
  {
    "annotatedResult": {
      "inputImageProperties": {
        "height": 853,
        "width": 1932
      }
    }
  }
]
```
"width": 1280
},
"polygons": [
{
"label": "dog",
"vertices": [
{
"x": 570,
"y": 239
},
{
"x": 603,
"y": 513
},
{
"x": 823,
"y": 645
},
{
"x": 901,
"y": 417
},
{
"x": 759,
"y": 281
}
]
},
{
"label": "cat",
"vertices": [
{
"x": 870,
"y": 278
},
{
"x": 908,
"y": 446
},
{
"x": 1009,
"y": 602
}
]
You could have many labels available, but only the ones that are used appear in the output.

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)

**crowd-polyline**

A widget for drawing polylines or lines on an image. Each polyline is associated with a label and can include two or more vertices. A polyline can intersect itself and its starting and ending points can be placed anywhere on the image.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](#).
The following is an example of a Liquid template that uses the `<crowd-polyline>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template. For more examples, see this [GitHub repository](#).

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-polyline
      name="crowdPolyline"
      src="{{ task.input.taskObject | grant_read_access }}"
      header="Add header here to describe the task"
      labels="['car','pedestrian','street car']"
    >
      <full-instructions>
        <p>Read the task carefully and inspect the image.</p>
        <p>Choose the appropriate label that best suits the image.</p>
        <p>Draw a polyline around the boundaries of all objects that the label applies to.</p>
        <p>Use the `<b>Enter</b>` key to complete a polyline.</p>
        <p>Make sure that the polyline fits tightly around the boundary of the object.</p>
      </full-instructions>

      <short-instructions>
        <p>Read the task carefully and inspect the image.</p>
        <p>Review the tool guide to learn how to use the polyline tool.</p>
        <p>Choose the appropriate label that best suits the image.</p>
        <p>To draw a polyline, select a label that applies to an object of interest and add a single point to the photo by clicking on that point. Continue to draw the polyline around the object by adding additional points around the object boundary.</p>
        <p>After you place the final point on the polyline, press `<b>Enter</b>` on your keyboard to complete the polyline.</p>
      </short-instructions>
  </crowd-polyline>
</crowd-form>

Attributes

The following attributes are supported by this element.
header

Optional. The text to display above the image. This is typically a question or simple instruction for the worker.

initial-value

Optional. An array of JSON objects, each of which sets a polyline when the component is loaded. Each JSON object in the array contains the following properties:

- **label** – The text assigned to the polyline as part of the labeling task. This text must match one of the labels defined in the `labels` attribute of the `<crowd-polyline>` element.
- **vertices** – the x and y pixel coordinates of the vertices of a polyline, relative to the top-left corner of the image.

```javascript
initial-value = "{ "polylines": [ 
    { 
        label: 'sideline', // label of this line annotation
        vertices:[ // an array of vertices which decide the position of the line 
            { 
                x: 84, 
                y: 110 
            }, 
            { 
                x: 60, 
                y: 100 
            } 
        ] 
    }, 
    { 
        label: 'yardline', 
        vertices:[ 
            { 
                x: 651, 
                y: 498 
            }, 
            { 
                x: 862, 
                y: 869 
            } 
        ] 
    } 
] 
"}
```
Polylines set via the `initial-value` property can be adjusted. Whether or not a worker answer was adjusted is tracked via an `initialValueModified` boolean in the worker answer output.

**labels**

Required. A JSON formatted array of strings, each of which is a label that a worker can assign to the line.

**Limit**: 10 labels

**label-colors**

Optional. An array of strings. Each string is a hexadecimal (hex) code for a label.

**name**

Required. The name of this widget. It's used as a key for the widget's input in the form output.

**src**

Required. The URL of the image on which to draw polylines.

**Regions**

The following regions are required by this element.

**full-instructions**

General instructions about how to draw polylines.

**short-instructions**

Important task-specific instructions that are displayed in a prominent place.
**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: short-instructions, full-instructions

**Output**

**inputImageProperties**

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.

**polylines**

A JSON Array containing objects with polylines' labels and vertices.

- **label** – The label given to a line.
- **vertices** – the x and y pixel coordinates of the vertices of a polyline, relative to the top-left corner of the image.

**Example : Sample Element Outputs**

The following is an example of output from this element.

```json
{
    "crowdPolyline": { //This is the name you set for the crowd-polyline
        "inputImageProperties": {
            "height": 1254,
            "width": 2048
        },
        "polylines": [
            {
                "label": "sideline",
            }
        ]
    }
}
```
See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference
Amazon SageMaker

Developer Guide

crowd-radio-button
A button that can be either checked or unchecked. When radio buttons are inside a radio group,
exactly one radio button in the group can be checked at any time. The following is an example of
how to conﬁgure a crowd-radio-button element inside of a crowd-radio-group element.
See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.
The following is an example of the syntax that you can use with the <crowd-radio-button>
element. Copy the following code and save it in a ﬁle with the extension .html. Open the ﬁle in
any browser to preview and interact with this template.
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<crowd-radio-group>
<crowd-radio-button name="tech" value="tech">Technology</crowd-radio-button>
<crowd-radio-button name="politics" value="politics">Politics</crowd-radio-button>
</crowd-radio-group>
</crowd-form>

The previous example can be seen in a custom worker task template in this GitHub example: entity
recognition labeling job custom template.
Crowd HTML Element radio buttons do not support the HTML tag, required. To make a radio
button selection required, use <input type="radio"> elements to create radio buttons and add
the required tag. The name attribute for all <input> elements that belong to the same group of
radio buttons must be the same. For example, the following template requires the user to select a
radio button in the animal-type group before submitting.
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
<p>Select an animal type:</p>
<img src="https://images.unsplash.com/photo-1537151608828-ea2b11777ee8?
ixlib=rb-1.2.1&ixid=eyJhcHBfaWQiOjEyMDd9&auto=format&fit=crop&w=1539&q=80"
style="height: 500; width: 400;"/>
<br><br>
<div>
<input type="radio" id="cat" name="animal-type" value="cat" required>
<label for="cat">Cat</label>
</div>
<div>
SageMaker Crowd HTML Elements

1940


Attributes

The following attributes are supported by this element.

**checked**

A Boolean switch that, if present, displays the radio button as checked.

**disabled**

A Boolean switch that, if present, displays the button as disabled and prevents it from being checked.

**name**

A string that is used to identify the answer submitted by the worker. This value will match a key in the JSON object that specifies the answer.

>Note

If you use the buttons outside of a crowd-radio-group element, but with the same name string and different value strings, the name object in the output will contain a Boolean value for each value string. To ensure that only one button in a group is selected, make them children of a crowd-radio-group element and use different name values.
value

A property name for the element's boolean value. If not specified, it uses "on" as the default, e.g. 
{ "<name>": { "<value>": <true or false> } }.

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-radio-group
- **Child elements**: none

Output

Outputs an object with the following pattern: 
{ "<name>": { "<value>": <true or false> } }.

If you use the buttons outside of a crowd-radio-group element, but with the same name string and different value strings, the name object will contain a Boolean value for each value string. To ensure that only one in a group of buttons is selected, make them children of a crowd-radio-group element and use different name values.

Example Sample output of this element

```
[
  {
    "btn1": {
      "yes": true
    },
    "btn2": {
      "no": false
    }
  }
]
```

See Also

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)
Amazon SageMaker

Developer Guide

crowd-radio-group
A group of radio buttons. Only one radio button within the group can be selected. Choosing one
radio button clears any previously chosen radio button within the same group. For an example
of a custom UI template that uses the crowd-radio-group element, see this entity recognition
labeling job custom template.
See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.
The following is an example of the syntax that you can use with the <crowd-radio-group>
element. Copy the following code and save it in a ﬁle with the extension .html. Open the ﬁle in
any browser to preview and interact with this template.

<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<style>
body {
padding-left: 20px;
margin-bottom: 20px;
}
#outer-container {
display: flex;
justify-content: space-around;
max-width: 900px;
margin-left: 100px;
}
.left-container {
margin-right: auto;
padding-right: 50px;
}
.right-container {
margin-left: auto;
padding-left: 50px;
}
#vertical-separator {
border: solid 1px #d5dbdb;
}
</style>
<crowd-form>
<div>
<h1>Instructions</h1>
SageMaker Crowd HTML Elements

1943


Lorem ipsum...
</div>
</div>

<h2>Background</h2>
<p>Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua. Ut enim ad minim veniam, quis nostrud exercitation ullamco laboris nisi ut aliquip ex ea commodo consequat.</p>

</div>

<div id="outer-container">
  <span class="left-container">
    <h2>Option 1</h2>
    <p>Nulla facilisi morbi tempus iaculis urna. Orci dapibus ultrices in iaculis nunc sed augue lacus.</p>
  </span>
  <span id="vertical-separator"></span>
  <span class="right-container">
    <h2>Option 2</h2>
    <p>Ultrices vitae auctor eu augue ut. Pellentesque massa placerat duis ultricies lacus sed turpis tincidunt id.</p>
  </span>
</div>

<h2>Question</h2>
<p>Which do you agree with?</p>
<crowd-radio-group>
  <crowd-radio-button name="option1" value="Option 1">Option 1</crowd-radio-button>
  <crowd-radio-button name="option2" value="Option 2">Option 2</crowd-radio-button>
</crowd-radio-group>

<p>Why did you choose this answer?</p>
<crowd-text-area name="explanation" placeholder="Explain how you reached your conclusion..."></crowd-text-area>
</div>
</crowd-form>

**Attributes**

No special attributes are supported by this element.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements:** [crowd-form](#)
• **Child elements**: crowd-radio-button

**Output**

Outputs an array of objects representing the crowd-radio-button elements within it.

**Example Sample of Element Output**

```
[
  {
    "btn1": {
      "yes": true
    },
    "btn2": {
      "no": false
    }
  }
]
```

**See Also**

For more information, see the following.

• [Use Amazon SageMaker Ground Truth to Label Data](#)
• [Crowd HTML Elements Reference](#)

**crowd-semantic-segmentation**

A widget for segmenting an image and assigning a label to each image segment.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](https://codepen.io/).

The following is an example of a Liquid template that uses the `<crowd-semantic-segmentation>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-semantic-segmentation
    name="annotatedResult"
```
The following attributes are supported by this element.

**header**

The text to display above the image. This is typically a question or simple instruction for the worker.

**initial-value**

A JSON object containing the color mappings of a prior semantic segmentation job and a link to the overlay image output by the prior job. Include this when you want a human worker to verify the results of a prior labeling job and adjust it if necessary.

The attribute would appear as follows:

```
initial-value='{
  "labelMappings": {
    "Bird": {
      "color": "#ff7f0e"
    },
    "Cat": {
```
When using Ground Truth built in task types with annotation consolidation (where more than one worker labels a single image), label mappings are included in individual worker output records, however the overall result is represented as the internal-color-map in the consolidated results.

You can convert the internal-color-map to label-mappings in a custom template using the Liquid templating language:

```liquid
initial-value="{
    'src' : '{{ task.input.manifestLine.label-attribute-name-from-prior-job|grant_read_access }}',
    'labelMappings': {
        {% for box in task.input.manifestLine.label-attribute-name-from-prior-job.metadata.internal-color-map %}
            {% if box[1]['class-name'] != 'BACKGROUND' %}
                {{ box[1]['class-name'] | to_json }}: {
                    'color': {{ box[1]['hex-color'] | to_json }}
                },
            {% endif %}
        {% endfor %}
    }
}"
```

**labels**

A JSON formatted array of strings, each of which is a label that a worker can assign to a segment of the image.

**name**

The name of this widget. It is used as a key for the widget’s input in the form output.
**src**

The URL of the image that is to be segmented.

**Element Hierarchy**

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: full-instructions, short-instructions

**Regions**

The following regions are supported by this element.

**full-instructions**

General instructions about how to do image segmentation.

**short-instructions**

Important task-specific instructions that are displayed in a prominent place.

**Output**

The following output is supported by this element.

**labeledImage**

A JSON Object containing a Base64 encoded PNG of the labels.

**labelMappings**

A JSON Object containing objects with named with the segmentation labels.

- **color** – The hexadecimal value of the label's RGB color in the labeledImage PNG.

**initialValueModified**

A boolean representing whether the initial values have been modified. This is only included when the output is from an adjustment task.
inputImageProperties

A JSON object that specifies the dimensions of the image that is being annotated by the worker. This object contains the following properties.

- **height** – The height, in pixels, of the image.
- **width** – The width, in pixels, of the image.

**Example: Sample Element Outputs**

The following is a sample of output from this element.

```json
[
  {
    "annotatedResult": {
      "inputImageProperties": {
        "height": 533,
        "width": 800
      },
      "labelMappings": {
        "<Label 2>": {
          "color": "#ff7f0e"
        },
        "<label 3>": {
          "color": "#2ca02c"
        },
        "<label 1>": {
          "color": "#1f77b4"
        }
      },
      "labeledImage": {
        "pngImageData": "<Base-64 Encoded Data>"
      }
    }
  }
]
```

**See Also**

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
• **Crowd HTML Elements Reference**

**crowd-slider**

A bar with a sliding knob that allows a worker to select a value from a range of values by moving the knob. The slider makes it a great choice for settings that reflect intensity levels, such as volume, brightness, or color saturation.

See an interactive example of an HTML template that uses this Crowd HTML Element in [CodePen](http://codepen.io).

The following is an example of a survey template that uses the `<crowd-slider>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

crowd-form

crowd-instructions link-text="View instructions" link-type="button">
  short-summary
    p>Provide a brief instruction here</p>
  </short-summary>

  detailed-instructions
    h3>Provide more detailed instructions here</h3>
    p>Include additional information</p>
  </detailed-instructions>

  positive-example
    p>Provide an example of a good answer here</p>
    p>Explain why it's a good answer</p>
  </positive-example>

  negative-example
    p>Provide an example of a bad answer here</p>
    p>Explain why it's a bad answer</p>
  </negative-example>
</crowd-instructions>

div
  p>What is your favorite color for a bird?</p>
  crowd-input name="favoriteColor" placeholder="example: pink" required>
```
Attributes

The following attributes are supported by this element.

**disabled**

A Boolean switch that, if present, displays the slider as disabled.

**editable**

A Boolean switch that, if present, displays an up/down button that can be chosen to select the value.

Selecting the value via the up/down button is an alternative to selecting the value by moving the knob on the slider. The knob on the slider will move synchronously with the up/down button choices.

**max**

A number that specifies the maximum value on the slider.

**min**

A number that specifies the minimum value on the slider.
name

A string that is used to identify the answer submitted by the worker. This value will match a key in the JSON object that specifies the answer.

pin

A Boolean switch that, if present, displays the current value above the knob as the knob is moved.

required

A Boolean switch that, if present, requires the worker to provide input.

secondary-progress

When used with a crowd-slider-secondary-color CSS attribute, the progress bar is colored to the point represented by the secondary-progress. For example, if this was representing the progress on a streaming video, the value would represent where the viewer was in the video timeline. The secondary-progress value would represent the point on the timeline to which the video had buffered.

step

A number that specifies the difference between selectable values on the slider.

value

A preset that becomes the default if the worker does not provide input.

Element Hierarchy

This element has the following parent and child elements.

- Parent elements: crowd-form
- Child elements: none

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference
crowd-tab

A component styled to look like a tab with information below.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example template that uses the `<crowd-tab>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-tabs>
    <crowd-tab header="Tab 1">
      <h2>Image</h2>
      <img src="https://images.unsplash.com/photo-1478382188900-5bb598fe27d3?ixlib=rb-1.2.1&ixid=eyJhcHBfaWQiOjEyMDd9&auto=format&fit=crop&w=1351&q=80" style="max-width: 40%"/>
      <h2>Text</h2>
      <p>Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.</p>
      <p>Sed risus ultricies tristique nulla aliquet enim tortor at auctor. Tempus egestas sed sed risus.</p>
    </crowd-tab>
    <crowd-tab header="Tab 2">
      <h2>Description</h2>
      <p>Sed risus ultricies tristique nulla aliquet enim tortor at auctor. Tempus egestas sed sed risus.</p>
    </crowd-tab>
    <crowd-tab header="Tab 3">
    </crowd-tab>
  </crowd-tabs>
</crowd-form>
```
Lorem ipsum dolor sit amet, lorem a wisi nibh, in pulvinar, consequat praesent vestibulum tellus ante felis auctor, vitae lobortis dictumst mauris. Pellentesque nulla ipsum ante quisque quam augue. Class lacus id euismod, blandit tempor mauris quisque tortor mauris, urna gravida ullam pede libero, ut suscipit orci faucibus lacus varius ornare, pellentesque ipsum. At etiam suspendisse est elementum luctus netus, vel sem nulla sodales, potenti magna enim ipsum diam tortor rutrum, quam donec massa elit ac, nam adipiscing sed at leo ipsum consectetuer. Ac turpis amet wisi, porttitor sint lacus ante, turpis accusantium, ac maecenas deleniti, nisl leo sem integer ac dignissim. Lobortis etiam luctus lectus odio auctor. Justo vitae, felis integer id, bibendum accumsan turpis eu est mus eros, ante id eros.

<p>Sed risus ultricies tristique nulla aliquet enim tortor at auctor. Tempus egestas sed sed risus.</p>

<p>Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.</p>
Attributes

The following attributes are supported by this element.

header

The text appearing on the tab. This is usually some short descriptive name indicative of the information contained below the tab.

Element Hierarchy

This element has the following parent and child elements.

- Parent elements: crowd-tabs
- Child elements: none

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

crowd-tabs

A container for tabbed information.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example template that uses the <crowd-tabs> element. Copy the following code and save it in a file with the extension .html. Open the file in any browser to preview and interact with this template.

```html
<scrip src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-tabs>
    <crowd-tab header="Tab 1">
      <h2>Image</h2>
      <h2>Image</h2>
    </crowd-tab>
  </crowd-tabs>
</crowd-form>
```
Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

Sed risus ultricies tristique nulla aliquet enim tortor at auctor. Tempus egestas sed sed risus.

Sed risus ultricies tristique nulla aliquet enim tortor at auctor. Tempus egestas sed sed risus.

Lorem ipsum dolor sit amet, lorem a wisi nibh, in pulvinar, consequat praesent vestibulum tellus ante felis auctor, vitae lobortis dictumst mauris. Pellentesque nulla ipsum ante quisque quam augue.
Class lacus id euismod, blandit tempor mauris quisque tortor mauris, urna gravida nullam pede libero, ut suscipit orci faucibus lacus varius ornare, pellentesque ipsum.

At etiam suspendisse est elementum luctus netus, vel sem nulla sodales, potenti magna enim ipsum diam tortor rutrum,
qua donec massa elit ac, nam adipiscing sed at leo ipsum consectetur.
Ac turpis amet wisi, porttitor sint lacus ante, turpis accusantium, ac maecenas deleniti,

nisl leo sem integer ac dignissim. Lobortis etiam luctus lectus odio auctor. Justo vitae, felis integer id, bibendum accumsan turpis eu est mus eros, ante id eros.

</div>
</crowd-tab>
</crowd-tabs>

<crowd-input label="Input outside tabs" name="inputOutsideTab"></crowd-input>

<short-instructions>
<p>Sed risus ultricies tristique nulla aliquet enim tortor at auctor. Tempus egestas sed sed risus.</p>
</short-inSTRUCTIONS>

<full-instructions header="Classification Instructions">
<p>Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.</p>
<p>Tempus egestas sed sed risus.</p>
</full-instructions>

</crowd-form>

Attributes

This element has no attributes.

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: crowd-tab
See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

**crowd-text-area**

A field for text input.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of a Liquid template designed to transcribe audio clips that uses the `<crowd-text-area>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <audio controls>
    <source src="{{ task.input.taskObject | grant_read_access }}" type="audio/mpeg">
    Your browser does not support the audio element.
  </audio>
  <h3>Instructions</h3>
  <p>Transcribe the audio</p>
  <p>Ignore "umms", "hmms", "uhs" and other non-textual phrases</p>
  <crowd-text-area name="transcription" rows="4"></crowd-text-area>
</crowd-form>
```

**Attributes**

The following attributes are supported by this element.

**allowed-pattern**

A regular expression that is used with the `auto-validate` attribute to ignore non-matching characters as the worker types.
auto-focus

A Boolean switch that, if present, puts the cursor in this element on-load so that users can immediately begin typing without having to click inside the element.

auto-validate

A Boolean switch that, if present, turns on input validation. The behavior of the validator can be modified by the error-message and allowed-pattern attributes.

char-counter

A Boolean switch that, if present, puts a small text field beneath the lower-right corner of the element, displaying the number of characters inside the element.

disabled

A Boolean switch that, if present, displays the input area as disabled.

error-message

The text to be displayed below the input field, on the left side, if validation fails.

label

A string that is displayed inside a text field.

This text shrinks and rises up above a text field when the worker starts typing in the field or when the value attribute is set.

max-length

An integer that specifies the maximum number of characters allowed by the element. Characters typed or pasted beyond the maximum are ignored.

max-rows

An integer that specifies the maximum number of rows of text that are allowed within a crowd-text-area. Normally the element expands to accommodate new rows. If this is set, after the number of rows exceeds it, content scrolls upward out of view and a scrollbar control appears.

name

A string used to represent the element's data in the output.
placeholder

A string presented to the user as placeholder text. It disappears after the user puts something in the input area.

rows

An integer that specifies the height of the element in rows of text.

data

A preset that becomes the default if the worker does not provide input. The preset appears in a text field.

Element Hierarchy

This element has the following parent and child elements.

- **Parent elements**: crowd-form
- **Child elements**: none

Output

This element outputs the name as a property name and the element's text contents as the value. Carriage returns in the text are represented as \n.

Example Sample output for this element

```
[
  {
    "textInput1": "This is the text; the text that\nmakes the crowd go wild."
  }
]
```

See Also

For more information, see the following.

- [Use Amazon SageMaker Ground Truth to Label Data](#)
- [Crowd HTML Elements Reference](#)
crowd-toast

A subtle notification that temporarily appears on the display. Only one crowd-toast is visible.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following is an example of a Liquid template that uses the `<crowd-toast>` element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <p>Find the official website for: <strong>{{ task.input.company }}</strong></p>
  <p>Do not give Yelp pages, LinkedIn pages, etc.</p>
  <p>Include the http:// prefix from the website</p>
  <crowd-input name="website" placeholder="http://example.com"></crowd-input>

  <crowd-toast duration="10000" opened>
    This is a message that you want users to see when opening the template. This message will disappear in 10 seconds.
  </crowd-toast>
</crowd-form>
```

**Attributes**

The following attributes are supported by this element.

**duration**

A number that specifies the duration, in milliseconds, that the notification appears on the screen.

**text**

The text to display in the notification.

**Element Hierarchy**

This element has the following parent and child elements.
• **Parent elements**: crowd-form

• **Child elements**: none

### See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

### crowd-toggle-button

A button that acts as an ON/OFF switch, toggling a state.

See an interactive example of an HTML template that uses this Crowd HTML Element in CodePen.

The following example shows different ways you can use to use the `<crowd-toggle-button>` HTML element. Copy the following code and save it in a file with the extension `.html`. Open the file in any browser to preview and interact with this template.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <!--Toggle button without value-->
  <crowd-toggle-button name="toggleButtonWithoutValue"></crowd-toggle-button>

  <!--Toggle button with value-->
  <crowd-toggle-button name="toggleButtonWithValue" value="someValue"></crowd-toggle-button>

  <!--Toggle button disabled-->
  <crowd-toggle-button name="toggleButtonDisabled" disabled></crowd-toggle-button>

  <!--Toggle button marked invalid-->
  <crowd-toggle-button name="toggleButtonInvalid" invalid></crowd-toggle-button>

  <!--Toggle button marked required-->
  <crowd-toggle-button name="toggleButtonRequired" required></crowd-toggle-button>
</crowd-form>
```
Attributes

The following attributes are supported by this element.

checked

A Boolean switch that, if present, displays the button switched to the ON position.

disabled

A Boolean switch that, if present, displays the button as disabled and prevents toggling.

invalid

When in an off position, a button using this attribute, will display in an alert color. The standard is red, but may be changed in CSS. When toggled on, the button will display in the same color as other buttons in the on position.

name

A string that is used to identify the answer submitted by the worker. This value matches a key in the JSON object that specifies the answer.

required

A Boolean switch that, if present, requires the worker to provide input.

value

A value used in the output as the property name for the element's Boolean state. Defaults to "on" if not provided.

Element Hierarchy

This element has the following parent and child elements.

- Parent elements: crowd-form
- Child elements: none

Output

This element outputs the name as the name of an object, containing the value as a property name and the element's state as Boolean value for the property. If no value for the element is specified, the property name defaults to "on."
Example Sample output for this element

```
[
  {
    "theToggler": {
      "on": true
    }
  }
]
```

See Also

For more information, see the following.

- Use Amazon SageMaker Ground Truth to Label Data
- Crowd HTML Elements Reference

Augmented AI Crowd HTML Elements

The following Crowd HTML Elements are only available for Amazon Augmented AI human workflow tasks.

Topics

- crowd-textract-analyze-document
- crowd-rekognition-detect-moderation-labels

**crowd-textract-analyze-document**

A widget to enable human review of a Amazon Textract document analysis result.

**Attributes**

The following attributes are supported by this element.

**header**

This is the text that is displayed as the header.

**src**

This is a link to the image to be analyzed by the worker.
initialValue

This sets initial values for attributes found in the worker UI.

The following is an example of an initialValue input:

```json
[
  {
    "blockType": "KEY_VALUE_SET",
    "confidence": 38.43309020996094,
    "geometry": {
      "boundingBox": {
        "width": 0.32613086700439453,
        "weight": 0.0942094624042511,
        "left": 0.4833833575248718,
        "top": 0.5227988958358765
      },
      "polygon": [
        {"x": 0.123, "y": 0.345}, ...
      ]
    },
    "id": "8c97b240-0969-4678-834a-646c95da9cf4",
    "relationships": [
      {
        "type": "CHILD",
        "ids": [
          "7ee7b7da-ee1b-428d-a567-55a3e3affa56",
          "4d6da730-ba43-467c-a9a5-c6137ba0c472"
        ]
      },
      {
        "type": "VALUE",
        "ids": [
          "6ee7b7da-ee1b-428d-a567-55a3e3affa54"
        ]
      }
    ],
    "entityTypes": [
      "KEY"
    ],
    "text": "Foo bar"
  }
]
```
**blockTypes**

This determines the kind of analysis the workers can do. Only KEY_VALUE_SET is currently supported.

**keys**

This specifies new keys and the associated text value the worker can add. The input values for keys can include the following elements:

- **importantFormKey** accepts strings, and is used to specify a single key.
- **importantFormKeyAliases** can be used to specify aliases that are acceptable alternatives to the keys supplied. Use this element to identify alternative spellings or presentations of your keys. This parameter accepts a list of one or more strings.

The following is an example of an input for keys.

```json
[
  {
    importantFormKey: 'Address',
    importantFormKeyAliases: [
      'address',
      'Addr.',
      'Add.'
    ]
  },
  {
    importantFormKey: 'Last name',
    importantFormKeyAliases: ['Surname']
  }
]
```

**no-key-edit**

This prevents the workers from editing the keys of annotations passed through initialValue. This prevents workers from editing the keys that have been detected on your documents. This is required.
no-geometry-edit

This prevents workers from editing the polygons of annotations passed through initialValue. For example, this would prevent the worker from editing the bounding box around a given key. This is required.

Element Hierarchy

This element has the following parent and child elements.

- Parent elements – crowd-form
- Child elements – full-instructions, short-instructions

Regions

The following regions are supported by this element. You can use custom HTML and CSS code within these regions to format your instructions to workers. For example, use the short-instructions section to provide good and bad examples of how to complete a task.

full-instructions

General instructions about how to work with the widget.

short-instructions

Important task-specific instructions that are displayed in a prominent place.

Example of a Worker Template Using the crowd Element

An example of a worker template using this crowd element would look like the following.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
{% capture s3_uri %}http://s3.amazonaws.com/
{{ task.input.aiServiceRequest.document.s3Object.bucket }}/\n{{ task.input.aiServiceRequest.document.s3Object.name }}{% endcapture %}

<crowd-form>
    <crowd-textract-analyze-document
        src="{{ s3_uri | grant_read_access }}"
        initial-value="{{ task.input.selectedAiServiceResponse.blocks }}"
        header="Review the key-value pairs listed on the right and correct them if they don't match the following document."
        no-key-edit
```
Click on a key-value block to highlight the corresponding key-value pair in the document.

If it is a valid key-value pair, review the content for the value. If the content is incorrect, correct it.

The text of the value is incorrect, correct it.

A wrong value is identified, correct it.

If it is not a valid key-value relationship, choose No.

If you can’t find the key in the document, choose Key not found.

If the content of a field is empty, choose Value is blank.

Examples
Key and value are often displayed next or below to each other.

Key and value displayed in one line.
Output

The following is a sample of the output from this element. You can find a detailed explanation of this output in the Amazon Textract AnalyzeDocument API documentation.

```json
{
    "AWS/Textract/AnalyzeDocument/Forms/V1": {
        "blocks": [
            {
                "blockType": "KEY_VALUE_SET",
                "id": "8c97b240-0969-4678-834a-646c95da9cf4",
                "relationships": [
                    {
                        "type": "CHILD",
                        "ids": ["7ee7b7da-ee1b-428d-a567-55a3e3affa56", "4d6da730-ba43-467c-a9a5-c6137ba0c472"]
                    },
                    {
                        "type": "VALUE",
                        "ids": ["6ee7b7da-ee1b-428d-a567-55a3e3affa54"]
                    }
                ],
                "entityTypes": ["KEY"],
                "text": "Foo bar baz"
            }
        ]
    }
}
```
crowd-rekognition-detect-moderation-labels

A widget to enable human review of an Amazon Rekognition image moderation result.

Attributes

The following attributes are supported by this element.

header

This is the text that is displayed as the header.

src

This is a link to the image to be analyzed by the worker.

categories

This supports categories as an array of strings or an array of objects where each object has a name field.

If the categories come in as objects, the following applies:

- The displayed categories are the value of the name field.
- The returned answer contains the full objects of any selected categories.

If the categories come in as strings, the following applies:

- The returned answer is an array of all the strings that were selected.

exclusion-category

By setting this attribute you create a button underneath the categories in the UI.

- When a user chooses the button, all categories are deselected and disabled.
- Choosing the button again re-enables the categories so that users can choose them.
- If you submit after choosing the button, it returns an empty array.
Element Hierarchy

This element has the following parent and child elements.

- Parent elements – crowd-form
- Child elements – full-instructions, short-instructions

AWS Regions

The following AWS Regions are supported by this element. You can use custom HTML and CSS code within these Regions to format your instructions to workers. For example, use the short-instructions section to provide good and bad examples of how to complete a task.

full-instructions

General instructions about how to work with the widget.

short-instructions

Important task-specific instructions that are displayed in a prominent place.

Example Worker Template with the crowd Element

An example of a worker template using the crowd element would look like the following.

```
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
{% capture s3_uri %}http://s3.amazonaws.com/
{{ task.input.aiServiceRequest.image.s3Object.bucket }}/ {{ task.input.aiServiceRequest.image.s3Object.name }}{% endcapture %}

<crowd-form>
  <crowd-rekognition-detect-moderation-labels
categories='[
  {% for label in task.input.selectedAiServiceResponse.moderationLabels %}
    {
      name: "{{ label.name }}",
      parentName: "{{ label.parentName }}",
    },
  {% endfor %}
]
src="{{ s3_uri | grant_read_access }}"
header="Review the image and choose all applicable categories."
>
```
Review the image and choose all applicable categories. If no categories apply, choose None.

<b>Nudity</b>
Visuals depicting nude male or female person or persons

<b>Graphic Male Nudity</b>
Visuals depicting full frontal male nudity, often close ups

<b>Graphic Female Nudity</b>
Visuals depicting full frontal female nudity, often close ups

<b>Sexual Activity</b>
Visuals depicting various types of explicit sexual activities and pornography

<b>Illustrated Nudity or Sexual Activity</b>
Visuals depicting animated or drawn sexual activity, nudity or pornography

<b>Adult Toys</b>
Visuals depicting adult toys, often in a marketing context

<b>Female Swimwear or Underwear</b>
Visuals depicting female person wearing only swimwear or underwear

<b>Male Swimwear Or Underwear</b>
Visuals depicting male person wearing only swimwear or underwear

<b>Partial Nudity</b>
Visuals depicting covered up nudity, for example using hands or pose

<b>Revealing Clothes</b>
Visuals depicting revealing clothes and poses, such as deep cut dresses

<b>Graphic Violence or Gore</b>
Visuals depicting prominent blood or bloody injuries

<b>Physical Violence</b>
Visuals depicting violent physical assault, such as kicking or punching
<b>Weapon Violence</b>
Visuals depicting violence using weapons like firearms or blades, such as shooting

<b>Weapons</b>
Visuals depicting weapons like firearms and blades

<b>Self Injury</b>
Visuals depicting self-inflicted cutting on the body, typically in distinctive patterns using sharp objects

<b>Emaciated Bodies</b>
Visuals depicting extremely malnourished human bodies

<b>Corpses</b>
Visuals depicting human dead bodies

<b>Hanging</b>
Visuals depicting death by hanging

Output
The following is a sample of the output from this element. For details about this output, see Amazon Rekognition <a>DetectModerationLabels</a> API documentation.

```json
{
   "AWS/Rekognition/DetectModerationLabels/Image/V3": {
      "ModerationLabels": [
         { name: 'Gore', parentName: 'Violence' },
         { name: 'Corpses', parentName: 'Violence' }
      ]
   }
}
```
Using Amazon Augmented AI for Human Review

When you use AI applications such as Amazon Rekognition, Amazon Textract, or your custom machine learning (ML) models, you can use Amazon Augmented AI to get human review of low-confidence predictions or random prediction samples.

What is Amazon Augmented AI?

Amazon Augmented AI (Amazon A2I) is a service that brings human review of ML predictions to all developers by removing the heavy lifting associated with building human review systems or managing large numbers of human reviewers.

Many ML applications require humans to review low-confidence predictions to ensure the results are correct. For example, extracting information from scanned mortgage application forms can require human review due to low-quality scans or poor handwriting. Building human review systems can be time-consuming and expensive because it involves implementing complex processes or workflows, writing custom software to manage review tasks and results, and managing large groups of reviewers.

Amazon A2I streamlines building and managing human reviews for ML applications. Amazon A2I provides built-in human review workflows for common ML use cases, such as content moderation and text extraction from documents. You can also create your own workflows for ML models built on SageMaker or any other tools. Using Amazon A2I, you can allow human reviewers to step in when a model is unable to make a high-confidence prediction or to audit its predictions on an ongoing basis.

Amazon A2I Use Case Examples

The following examples demonstrate how you can use Amazon A2I to integrate a human review loop into your ML application. For each of these examples, you can find a Jupyter Notebook that demonstrates that workflow in Use Cases and Examples Using Amazon A2I.

- **Use Amazon A2I with Amazon Textract** – Have humans review important key-value pairs in single-page documents or have Amazon Textract randomly sample and send documents from your dataset to humans for review.

- **Use Amazon A2I with Amazon Rekognition** – Have humans review unsafe images for explicit adult or violent content if Amazon Rekognition returns a low-confidence score, or have Amazon Rekognition randomly sample and send images from your dataset to humans for review.
• **Use Amazon A2I to review real-time ML inferences** – Use Amazon A2I to review real-time, low-confidence inferences made by a model deployed to a SageMaker hosted endpoint and incrementally train your model using Amazon A2I output data.

• **Use Amazon A2I with Amazon Comprehend** – Have humans review Amazon Comprehend inferences about text data such as sentiment analysis, text syntax, and entity detection.

• **Use Amazon A2I with Amazon Transcribe** – Have humans review Amazon Transcribe transcriptions of video or audio files. Use the results of transcription human review loops to create a custom vocabulary and improve future transcriptions of similar video or audio content.

• **Use Amazon A2I with Amazon Translate** – Have humans review low-confidence translations returned from Amazon Translate.

• **Use Amazon A2I to review tabular data** – Use Amazon A2I to integrate a human review loop into an ML application that uses tabular data.

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**Topics**

- [Get Started with Amazon Augmented AI](#)
- [Use Cases and Examples Using Amazon A2I](#)
- [Create a Human Review Workflow](#)
- [Delete a Human Review Workflow](#)
- [Create and Start a Human Loop](#)
- [Delete a Human Loop](#)
- [Create and Manage Worker Task Templates](#)
- [Monitor and Manage Your Human Loop](#)
Get Started with Amazon Augmented AI

To get started using Amazon Augmented AI, review the Core Components of Amazon A2I and Prerequisites to Using Augmented AI. Then, use the following documentation to learn how to use the Amazon A2I console and API.

- Tutorial: Get Started in the Amazon A2I Console
- Tutorial: Get Started Using the Amazon A2I API

You can also get started using the Amazon A2I API by following a Jupyter Notebook tutorial. See Use Cases and Examples Using Amazon A2I for a list of notebooks and use cases.

Core Components of Amazon A2I

Review the following terms to familiarize yourself with the core components of Amazon A2I.

Task Types

The AI/ML workflow into which you integrate Amazon A2I defines an Amazon A2I task type.

Amazon A2I supports:

- Two built-in task types: Amazon Textract key-value pair extraction and Amazon Rekognition image moderation.
- A custom task type: Use a custom task type to integrate a human review loop into any machine learning workflow. You can use a custom task type to integrate Amazon A2I with other AWS services like Amazon Comprehend, Amazon Transcribe, and Amazon Translate, as well as your own custom machine learning workflows. To learn more, see Use Cases and Examples Using Amazon A2I.
Select a tab in the following table to see diagrams that illustrate how Amazon A2I works with each task type. Select the task type page using the links in the preceding list to learn more about that task type.

Amazon Textract – Key-value pair extraction

This image depicts the Amazon A2I built-in workflow with Amazon Textract. On the left, the resources that are required to create an Amazon Textract human review workflow are depicted: an Amazon S3 bucket, activation conditions, a worker task template, and a work team. These resources are used to create a human review workflow, or flow definition. An arrow points right to the next step in the workflow: using Amazon Textract to configure a human loop with the human review workflow. A second arrow points right from this step to the step in which activation conditions specified in the human review workflow are met. This initiates the creation of a human loop. On the right of the image, the human loop is depicted in three steps: 1) the worker UI and tools are generated and the task is made available to workers, 2) workers review input data, and finally, 3) results are saved in Amazon S3.

Amazon Rekognition – Image moderation

This image depicts the Amazon A2I built-in workflow with Amazon Rekognition. On the left, the resources that are required to create an Amazon Rekognition human review workflow are depicted: an Amazon S3 bucket, activation conditions, a worker task template, and a work team. These resources are used to create a human review workflow, or flow definition. An arrow points right to the next step in the workflow: using Amazon Rekognition to configure a human loop.
with the human review workflow. A second arrow points right from this step to the step in which activation conditions specified in the human review workflow are met. This initiates the creation of a human loop. On the right of the image, the human loop is depicted in three steps: 1) the worker UI and tools are generated and the task is made available to workers, 2) workers review input data, and finally, 3) results are saved in Amazon S3.

Custom Task Type

The following image depicts the Amazon A2I custom workflow. A custom ML model is used to generate predictions. The client application filters these predictions using user-defined criteria and determines if a human review is required. If so, these predictions are sent to Amazon A2I for human review. Amazon A2I collects the results of human review in Amazon S3, which can access by the client application. If the filter determines that no human review is needed, predictions can be fed directly to the client application.
Human Review Workflow (Flow Definition)

You use a human review workflow to specify your human work team, to set up your worker UI using a worker task template, and to provide information about how workers should complete the review task.

For built-in task types, you also use the human review workflow to identify the conditions under which a human loop is initiated. For example, Amazon Rekognition can perform image content moderation using machine learning. You can use the human review workflow to specify that an image is sent to a human for content moderation review if Amazon Rekognition's confidence is too low.

You can use a human review workflow to create multiple human loops.

You can create a flow definition in the SageMaker console or with the SageMaker API. To learn more about both of these options, see Create a Human Review Workflow.

Work Team

A work team is a group of human workers to whom you send your human review tasks.

When you create a human review workflow, you specify a single work team.

Your work team can come from the Amazon Mechanical Turk workforce, a vendor-managed workforce, or your own private workforce. When you use the private workforce, you can create
multiple work teams. Each work team can be used in multiple human review workflows. To learn how to create a workforce and work teams, see Create and Manage Workforces.

Worker Task Template and Human Task UI

You use a worker task template to create a worker UI (a human task UI) for your human review tasks.

The human task UI displays your input data, such as documents or images, and instructions to workers. It also provides interactive tools that the worker uses to complete your tasks.

For built-in task types, you must use the Amazon A2I worker task template provided for that task type.

Human Loops

A human loop is used to create a single human review job. For each human review job, you can choose the number of workers that are sent a task to review a single data object. For example, if you set the number of workers per object to 3 for an image classification labeling job, three workers classify each input image. Increasing the number of workers per object can improve label accuracy.

A human loop is created using a human review workflow as follows:

- For built-in task types, the conditions specified in the human review workflow determine when the human loop is created.
- Human review tasks are sent to the work team specified in the human review workflow.
- The worker task template specified in the human review workflow is used to render the human task UI.

When do human loops get created?

When you use one of the built-in task types, the corresponding AWS service creates and starts a human loop on your behalf when the conditions specified in your human review workflow are met. For example:

- When you use Augmented AI with Amazon Textract, you can integrate Amazon A2I into a document review task using the API operation `AnalyzeDocument`. A human loop is created every time Amazon Textract returns inferences about key-value pairs that meet the conditions you specify in your human review workflow.
When you use Augmented AI with Amazon Rekognition, you can integrate Amazon A2I into an image moderation task using the API operation `DetectModerationLabels`. A human loop is created every time Amazon Rekognition returns inferences about image content that meet the conditions you specify in your human review workflow.

When using a *custom task type*, you start a human loop using the [Amazon Augmented AI Runtime API](https://docs.aws.amazon.com/a2i/latest/dg/a2i-runtime-api.html). When you call `StartHumanLoop` in your custom application, a task is sent to human reviewers.

To learn how to create and start a human loop, see [Create and Start a Human Loop](https://docs.aws.amazon.com/a2i/latest/dg/a2i-console-tutorial-human-loop.html).

To generate these resources and create a human review workflow, Amazon A2I integrates multiple APIs, including the Amazon Augmented AI Runtime Model, the SageMaker APIs, and APIs associated with your task type. To learn more, see [Use APIs in Amazon Augmented AI](https://docs.aws.amazon.com/a2i/latest/dg/a2i-console-api-reference.html).

**Note**

AWS Region availability may differ when you use Augmented AI with other AWS services, such as Amazon Textract. Create Augmented AI resources in the same AWS Region that you use to interact with those AWS services. For AWS Region availability for all services, see the [Region Table](https://docs.aws.amazon.com/region-availability/index.html).

**Prerequisites to Using Augmented AI**

Amazon A2I uses resources in IAM, SageMaker, and Amazon S3 to create and run your human review workflows. You can create some of these resources in the Amazon A2I console when you create a human review workflow. To learn how, see [Tutorial: Get Started in the Amazon A2I Console](https://docs.aws.amazon.com/a2i/latest/dg/a2i-console-tutorial-get-started.html).

To use Amazon A2I, you need the following resources:

- One or more Amazon S3 buckets in the same AWS Region as the workflow for your input and output data. To create a bucket, follow the instructions in [Create a Bucket](https://docs.aws.amazon.com/AmazonS3/latest/userguide/creating-bucket.html) in the *Amazon Simple Storage Service Console User Guide*.
- An IAM role with required permissions to create a human review workflow and an IAM user or role with permission to access Augmented AI. For more information, see [Permissions and Security in Amazon Augmented AI](https://docs.aws.amazon.com/a2i/latest/dg/a2i-console-role-permissions.html).
• A public, private, or vendor workforce for your human review workflows. If you plan to use a private workforce, you need to set one up ahead of time in the same AWS Region as your Amazon A2I workflow. To learn more about these workforce types, see Create and Manage Workforces.

**Important**

To learn about the compliance programs that cover Amazon Augmented AI at this time, see AWS Services in Scope by Compliance Program. If you use Amazon Augmented AI in conjunction with other AWS services (such as Amazon Rekognition and Amazon Textract), note that Amazon Augmented AI may not be in scope for the same compliance programs as those other services. You are responsible for how you use Amazon Augmented AI, including understanding how the service processes or stores customer data and any impact on the compliance of your data environment. You should discuss your workload objectives and goals with your AWS account team; they can help you evaluate whether the service is a good fit for your proposed use case and architecture.

**Tutorial: Get Started in the Amazon A2I Console**

The following tutorial shows you how to get started using Amazon A2I in the Amazon A2I console.

The tutorial gives you the option to use Augmented AI with Amazon Textract for document review or Amazon Rekognition for image content review.

**Prerequisites**

To get started using Amazon A2I, complete the following prerequisites.

• Create an Amazon S3 bucket in the same AWS Region as the workflow for your input and output data. For example, if you are using Amazon A2I with Amazon Textract in us-east-1, create your bucket in us-east-1. To create a bucket, follow the instructions in Create a Bucket in the Amazon Simple Storage Service Console User Guide.

• Do one of the following:
  • If you want to complete the tutorial using Amazon Textract, download this sample document and place it in your Amazon S3 bucket.
  • If you want to complete the tutorial using Amazon Rekognition, download this image and place it in your Amazon S3 bucket.
Step 1: Create a Work Team

First, create a work team in the Amazon A2I console and add yourself as a worker so that you can preview the worker review task.

Important

This tutorial uses a private work team. The Amazon A2I private workforce is configured in the Ground Truth area of the SageMaker console and is shared between Amazon A2I and Ground Truth.

To create a private workforce using worker emails

2. In the navigation pane, choose Labeling workforces under Ground Truth.
3. Choose Private, then choose Create private team.
4. Choose Invite new workers by email.
5. For this tutorial, enter your email and any others that you want to be able to preview the human task UI. You can paste or type a list of up to 50 email addresses, separated by commas, into the email addresses box.
6. Enter an organization name and contact email.
7. Optionally, choose an Amazon SNS topic to which to subscribe the team so workers are notified by email when new Ground Truth labeling jobs become available. Amazon SNS notifications are supported by Ground Truth and are not supported by Augmented AI. If you subscribe workers to Amazon SNS notifications, they only receive notifications about Ground Truth labeling jobs. They do not receive notifications about Augmented AI tasks.
8. Choose Create private team.

If you add yourself to a private work team, you receive an email from no-reply@verificationemail.com with login information. Use the link in this email to reset your
password and log in to your worker portal. This is where your human review tasks appear when you create a human loop.

Step 2: Create a Human Review Workflow

In this step, you create a human review workflow. Each human review workflow is created for a specific task type. This tutorial allows you to choose between the built-in task types: Amazon Rekognition and Amazon Textract.

To create a human review workflow:

1. Open the Augmented AI console at https://console.aws.amazon.com/a2i to access the Human review workflows page.
2. Select Create human review workflow.
3. In Workflow settings, enter a workflow Name, S3 bucket, and the IAM role that you created for this tutorial, with the AWS managed policy AmazonAugmentedAIIntegratedAPIAccess attached.
4. For Task type, select Textract – Key-value pair extraction or Rekognition – Image moderation.
5. Select the task type that you chose from the following table for instructions for that task type.

Amazon Textract – Key-value pair extraction

1. Select Trigger a human review for specific form keys based on the form key confidence score or when specific form keys are missing.
2. For Key name, enter Mail Address.
3. Set the identification confidence threshold between 0 and 99.
4. Set the qualification confidence threshold between 0 and 99.
5. Select Trigger a human review for all form keys identified by Amazon Textract with confidence scores in a specific range.
6. Set the identification confidence threshold between 0 and 90.
7. Set the qualification confidence threshold between 0 and 90.
This initiates a human review if Amazon Textract returns a confidence score that is less than 99 for Mail Address and its key, or if it returns a confidence score less than 90 for any key value pair detected in the document.

The following image shows the Amazon Textract form extraction - Conditions for invoking human review section of the Amazon A2I console. In the image, the check boxes for the two types of triggers explained in the proceeding paragraph are checked, and Mail Address is used as a Key name for the first trigger. The identification confidence threshold is defined using confidence scores for key-value pairs detect within the form and is set between 0 and 99. The qualification confidence threshold is defined using confidence scores for text contained within keys and values in a form and is set between 0 and 99.
Amazon Rekognition – Image moderation

1. Select Trigger human review for labels identified by Amazon Rekognition based on label confidence score.

2. Set the Threshold between 0 and 98.

This initiates a human review if Amazon Rekognition returns a confidence score that is less than 98 for an image moderation job.
The following image shows how you can select the Trigger human review for labels identified by Amazon Rekognition based on label confidence score option and enter a Threshold between 0 and 98 in the Amazon A2I console.

6. Under Worker task template creation, select Create from a default template.

7. Enter a Template name.

8. In Task description field, enter the following text:

   Read the instructions carefully and complete the task.


10. Select the private team that you created.

11. Choose Create.

Once your human review workflow is created, it appears in the table on the Human review workflows page. When the Status is Active, copy and save the Workflow ARN. You need it for the next step.

**Step 3: Start a Human Loop**

You must use an API operation to start a human loop. There are a variety of language-specific SDKs that you can use to interact with these API operations. To see documentation for each of these SDKs, refer to the See Also section in the API documentation, as shown in the following image.
For this tutorial, you use one of the following APIs:

- If you chose the Amazon Textract task type, you use the `AnalyzeDocument` operation.
- If you chose the Amazon Rekognition task type, you use the `DetectModerationLabels` operation.

You can interact with these APIs using a SageMaker notebook instance (recommended for new users) or the AWS Command Line Interface (AWS CLI). Choose one of the following to learn more about these options:

- To learn more about and set up a notebook instance, see [Amazon SageMaker Notebook Instances](#).
- To learn more about and get started using the AWS CLI, see [What Is the AWS Command Line Interface?](#) in the [AWS Command Line Interface User Guide](#).
Select your task type in the following table to see example requests for Amazon Textract and Amazon Rekognition using the AWS SDK for Python (Boto3).

Amazon Textract – Key-value pair extraction

The following example uses the AWS SDK for Python (Boto3) to call `analyze_document` in `us-west-2`. Replace the italicized red text with your resources. Include the `DataAttributes` parameter if you are using the Amazon Mechanical Turk workforce. For more information, see the [analyze_document](#) documentation in the [AWS SDK for Python (Boto) API Reference](#).

```python
response = client.analyze_document(
    Document={
        "S3Object": {
            "Bucket": "AWSDOC-EXAMPLE-BUCKET",
            "Name": "document-name.pdf"
        }
    },
    HumanLoopConfig={
        "HumanLoopName": "human-loop-name",
        "DataAttributes": {
            "ContentClassifiers": [
                "FreeOfPersonallyIdentifiableInformation",
                "FreeOfAdultContent"
            ]
        },
        FeatureTypes=["TABLES", "FORMS"]
    })
```

Amazon Rekognition – Image moderation

The following example uses the AWS SDK for Python (Boto3) to call `detect_moderation_labels` in `us-west-2`. Replace the italicized red text with your resources. Include the `DataAttributes` parameter if you are using the Amazon Mechanical Turk workforce. For more information, see the [detect_moderation_labels](#) documentation in the [AWS SDK for Python (Boto) API Reference](#).

```python
response = client.detect_moderation_labels(
    Image={
        "S3Object":{
```
Step 4: View Human Loop Status in Console

When you start a human loop, you can view its status in the Amazon A2I console.

To view your human loop status

1. Open the Augmented AI console at https://console.aws.amazon.com/a2i to access the Human review workflows page.
2. Select the human review workflow that you used to start your human loop.
3. In the Human loops section, you can see your human loop. View its status in the Status column.

Step 5: Download Output Data

Your output data is stored in the Amazon S3 bucket you specified when you created a human review workflow.

To view your Amazon A2I output data

1. Open the Amazon S3 console.
2. Select the Amazon S3 bucket you specified when you created your human review workflow in step 2 of this example.
3. Starting with the folder that is named after your human review workflow, navigate to your output data by selecting the folder with the following naming convention:
4. Select `output.json` and choose **Download**.

**Tutorial: Get Started Using the Amazon A2I API**

This tutorial explains the API operations you can use to get started using Amazon A2I.

To use a Jupyter Notebook to run these operations, select a Jupyter Notebook from **Use Cases and Examples Using Amazon A2I** and use **Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook** to learn how to use it in a SageMaker notebook instance.

To learn more about the API operations you can use with Amazon A2I, see **Use APIs in Amazon Augmented AI**.

**Create a Private Work Team**

You can create a private work team and add yourself as a worker so that you can preview Amazon A2I.

If you are not familiar with Amazon Cognito, we recommend that you use the SageMaker console to create a private workforce and add yourself as a private worker. For instructions, see **Step 1: Create a Work Team**.

If you are familiar with Amazon Cognito, you can use the following instructions to create a private work team using the SageMaker API. After you create a work team, note the work team ARN (WorkteamArn).

To learn more about the private workforce and other available configurations, see **Use a Private Workforce**.

**Create a private workforce**

If you have not created a private workforce, you can do so using an **Amazon Cognito user pool**. Make sure that you have added yourself to this user pool. You can create a private work team using the AWS SDK for Python (Boto3) **create_workforce** function. For other language-specific SDKs, refer to the list in **CreateWorkforce**.
Create a private work team

After you have created a private workforce in the AWS Region to configure and start your human loop, you can create a private work team using the AWS SDK for Python (Boto3) `create_workteam` function. For other language-specific SDKs, refer to the list in CreateWorkteam.

```python
response = client.create_workteam(
    WorkteamName="work-team-name",
    WorkforceName="workforce-name",
    MemberDefinitions=[
        {
            "CognitoMemberDefinition": {
                "UserPool": "<aws-region>_ID",
                "UserGroup": "user-group",
                "ClientId": "app-client-id"
            }
        }
    ]
)
```

Access your work team ARN as follows:

```python
workteamArn = response["WorkteamArn"]
```

List private work teams in your account

If you have already created a private work team, you can list all work teams in a given AWS Region in your account using the AWS SDK for Python (Boto3) `list_workteams` function. For other language-specific SDKs, refer to the list in ListWorkteams.
response = client.list_workteams()

If you have numerous work teams in your account, you may want to use MaxResults, SortBy, and NameContains to filter your results.

Create a Human Review Workflow

You can create a human review workflow using the Amazon A2I CreateFlowDefinition operation. Before you create your human review workflow, you need to create a human task UI. You can do this with the CreateHumanTaskUi operation.

If you are using Amazon A2I with the Amazon Textract or Amazon Rekognition integrations, you can specify activation conditions using a JSON.

Create a Human Task UI

If you are creating a human review workflow to be used with Amazon Textract or Amazon Rekognition integrations, you need to use and modify pre-made worker task template. For all custom integrations, you can use your own custom worker task template. Use the following table to learn how to create a human task UI using a worker task template for the two built-in integrations. Replace the template with your own to customize this request.

Amazon Textract – Key-value pair extraction

To learn more about this template, see Custom Template Example for Amazon Textract.

```template = r""
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
{% capture s3_uri %}http://s3.amazonaws.com/{{ task.input.aiServiceRequest.document.s3Object.bucket }}/\n{{ task.input.aiServiceRequest.document.s3Object.name }}{% endcapture %}
<crowd-form>
  <crowd-textract-analyze-document
    src="{{ s3_uri | grant_read_access }}"
    initial-value="{{ task.input.selectedAiServiceResponse.blocks }}"
    header="Review the key-value pairs listed on the right and correct them if they don't match the following document."
    no-key-edit=""
    no-geometry-edit=""
    keys="{{ task.input.humanLoopContext.importantFormKeys }}"
    block-types='["KEY_VALUE_SET"]'
  >
  <short-instructions header="Instructions">
```
Click on a key-value block to highlight the corresponding key-value pair in the document.

If it is a valid key-value pair, review the content for the value. If the content is incorrect, correct it.

The text of the value is incorrect, correct it.

A wrong value is identified, correct it.

If it is not a valid key-value relationship, choose No.

If you can’t find the key in the document, choose Key not found.

If the content of a field is empty, choose Value is blank.

Examples

Key and value are often displayed next or below to each other.

Key and value displayed in one line.

If the content of the value has multiple lines, enter all the text without line break.

Include all value text even if it extends beyond the highlight box.

If it is a valid key-value pair, review the content for the value. If the content is incorrect, correct it.
Amazon Rekognition – Image moderation

To learn more about this template, see Custom Template Example for Amazon Rekognition.

```template
<template
<template
<template
<template
<template
<template
<template
<template
<template
<template
<template
<template
<template
<template
```
<b>Physical Violence</b>
Visuals depicting violent physical assault, such as kicking or punching

<b>Weapon Violence</b>
Visuals depicting violence using weapons like firearms or blades, such as shooting

<b>Weapons</b>
Visuals depicting weapons like firearms and blades

Custom Integration

The following is an example template that can be used in a custom integration. This template is used in this notebook, demonstrating a custom integration with Amazon Comprehend.

```
template = r'""
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>

<crowd-form>
  <crowd-classifier
    name="sentiment"
    categories=['"Positive", "Negative", "Neutral", "Mixed"]'
    initial-value="\{ task.input.initialValue \}"
    header="What sentiment does this text convey?"
  >
    <classification-target>
      \{ task.input.taskObject \}
    </classification-target>

    <full-instructions header="Sentiment Analysis Instructions">
      <p><strong>Positive</strong> sentiment include: joy, excitement, delight</p>
      <p><strong>Negative</strong> sentiment include: anger, sarcasm, anxiety</p>
      <p><strong>Neutral</strong>: neither positive or negative, such as stating a fact</p>
      <p><strong>Mixed</strong>: when the sentiment is mixed</p>
    </full-instructions>

  <short-instructions>
    Choose the primary sentiment that is expressed by the text.
  </short-instructions>
</crowd-form>
```
Using the template specified above, you can create a template using the AWS SDK for Python (Boto3) `create_human_task_ui` function. For other language-specific SDKs, refer to the list in CreateHumanTaskUi.

```python
response = client.create_human_task_ui(
    HumanTaskUiName="human-task-ui-name",
    UiTemplate={
        "Content": template
    }
)
```

This response element contains the human task UI ARN. Save this as follows:

```python
humanTaskUiArn = response["HumanTaskUiArn"]
```

### Create JSON to specify activation conditions

For Amazon Textract and Amazon Rekognition built-in integrations, you can save activation conditions in a JSON object and use this in your CreateFlowDefinition request.

Next, select a tab to see example activation conditions you can use for these built-in integrations. For additional information about activation condition options, see JSON Schema for Human Loop Activation Conditions in Amazon Augmented AI.

**Amazon Textract – Key-value pair extraction**

This example specifies conditions for specific keys (such as `Mail address`) in the document. If Amazon Textract’s confidence falls outside of the thresholds set here, the document is sent to a human for review, with the specific keys that initiated the human loop prompted to the worker.

```python
import json

humanLoopActivationConditions = json.dumps(
    humanLoopActivationConditions)  
```
{ "Conditions": [
    {
        "Or": [
            {
                "ConditionType": "ImportantFormKeyConfidenceCheck",
                "ConditionParameters": {
                    "ImportantFormKey": "Mail address",
                    "ImportantFormKeyAliases": ["Mail Address:","Mail address:","Mailing Add:","Mailing Addresses"],
                    "KeyValueBlockConfidenceLessThan": 100,
                    "WordBlockConfidenceLessThan": 100
                }
            },
            {
                "ConditionType": "MissingImportantFormKey",
                "ConditionParameters": {
                    "ImportantFormKey": "Mail address",
                    "ImportantFormKeyAliases": ["Mail Address:","Mail address:","Mailing Add:","Mailing Addresses"]
                }
            }
        ]
    },
    {
        "ConditionType": "ImportantFormKeyConfidenceCheck",
        "ConditionParameters": {
            "ImportantFormKey": "Phone Number",
            "ImportantFormKeyAliases": ["Phone number:","Phone No.:","Number:"],
            "KeyValueBlockConfidenceLessThan": 100,
            "WordBlockConfidenceLessThan": 100
        }
    },
    {
        "ConditionType": "ImportantFormKeyConfidenceCheck",
        "ConditionParameters": {
            "ImportantFormKey": "*",
            "KeyValueBlockConfidenceLessThan": 100,
            "WordBlockConfidenceLessThan": 100
        }
    }
]}

Amazon Rekognition – Image moderation

The human loop activation conditions used here are tailored towards Amazon Rekognition content moderation; they are based on the confidence thresholds for the Suggestive and Female Swimwear Or Underwear moderation labels.

```python
import json

humanLoopActivationConditions = json.dumps(
    {
        "Conditions": [
            {
                "Or": [
                    {
                        "ConditionType": "ModerationLabelConfidenceCheck",
                        "ConditionParameters": {
                            "ModerationLabelName": "Suggestive",
                            "ConfidenceLessThan": 98
                        }
                    },
                    {
                        "ConditionType": "ModerationLabelConfidenceCheck",
                        "ConditionParameters": {
                            "ModerationLabelName": "Female Swimwear Or Underwear",
                            "ConfidenceGreaterThan": 98
                        }
                    }
                ]
            }
        ]
    }
)
```
Create a human review workflow

This section gives an example of the CreateFlowDefinition AWS SDK for Python (Boto3) request using the resources created in the previous sections. For other language-specific SDKs, refer to the list in CreateFlowDefinition. Use the tabs in the following table to see the requests to create a human review workflow for Amazon Textract and Amazon Rekognition built-in integrations.

Amazon Textract – Key-value pair extraction

If you use the built-in integration with Amazon Textract, you must specify "AWS/Textract/AnalyzeDocument/Forms/V1" for "AwsManagedHumanLoopRequestSource" in HumanLoopRequestSource.

```python
response = client.create_flow_definition(
    FlowDefinitionName="human-review-workflow-name",
    HumanLoopRequestSource={
        "AwsManagedHumanLoopRequestSource": "AWS/Textract/AnalyzeDocument/Forms/V1"
    },
    HumanLoopActivationConfig={
        "HumanLoopActivationConditionsConfig": {
            "HumanLoopActivationConditions": humanLoopActivationConditions
        }
    },
    HumanLoopConfig={
        "WorkteamArn": workteamArn,
        "HumanTaskUiArn": humanTaskUiArn,
        "TaskTitle": "Document entry review",
        "TaskDescription": "Review the document and instructions. Complete the task",
        "TaskCount": 1,
        "TaskAvailabilityLifetimeInSeconds": 43200,
        "TaskTimeLimitInSeconds": 3600,
        "TaskKeywords": [
            "document review",
        ]
    })
```
Amazon Rekognition – Image moderation

If you use the built-in integration with Amazon Rekognition, you must specify "AWS/Rekognition/DetectModerationLabels/Image/V3" for "AwsManagedHumanLoopRequestSource" in HumanLoopRequestSource.

```python
response = client.create_flow_definition(
    FlowDefinitionName="human-review-workflow-name",
    HumanLoopRequestSource={
        "AwsManagedHumanLoopRequestSource": "AWS/Rekognition/DetectModerationLabels/Image/V3"
    },
    HumanLoopActivationConfig={
        "HumanLoopActivationConditionsConfig": {
            "HumanLoopActivationConditions": humanLoopActivationConditions
        }
    },
    HumanLoopConfig={
        "WorkteamArn": workteamArn,
        "HumanTaskUiArn": humanTaskUiArn,
        "TaskTitle": "Image content moderation",
        "TaskDescription": "Review the image and instructions. Complete the task",
        "TaskCount": 1,
        "TaskAvailabilityLifetimeInSeconds": 43200,
        "TaskTimeLimitInSeconds": 3600,
        "TaskKeywords": ["content moderation"],
    }
)
```
Custom Integration

If you use a custom integration, exclude the following parameters:
HumanLoopRequestSource, HumanLoopActivationConfig.

```python
response = client.create_flow_definition(
    FlowDefinitionName="human-review-workflow-name",
    HumanLoopConfig={
        "WorkteamArn": workteamArn,
        "HumanTaskUiArn": humanTaskUiArn,
        "TaskTitle": "Image content moderation",
        "TaskDescription": "Review the image and instructions. Complete the task",
        "TaskCount": 1,
        "TaskAvailabilityLifetimeInSeconds": 43200,
        "TaskTimeLimitInSeconds": 3600,
        "TaskKeywords": [
            "content moderation",
        ],
    },
    OutputConfig={
        "S3OutputPath": "s3://DOC-EXAMPLE-BUCKET/prefix/",
    },
    RoleArn="arn:aws:iam::<account-number>:role/<role-name>",
    Tags=[
        {
            "Key": "string",
            "Value": "string"
        },
    ]
)`
After you create a human review workflow, you can retrieve the flow definition ARN from the response:

```python
humanReviewWorkflowArn = response['FlowDefinitionArn']```

Create a Human Loop

The API operation you use to start a human loop depends on the Amazon A2I integration you use.

- If you use the Amazon Textract built-in integration, you use the `AnalyzeDocument` operation.
- If you use the Amazon Rekognition built-in integration, you use the `DetectModerationLabels` operation.
- If you use a custom integration, you use the `StartHumanLoop` operation.

Select your task type in the following table to see example requests for Amazon Textract and Amazon Rekognition using the AWS SDK for Python (Boto3).

Amazon Textract – Key-value pair extraction

The following example uses the AWS SDK for Python (Boto3) to call `analyze_document` in `us-west-2`. Replace the italicized red text with your resources. Include the `DataAttributes` parameter if you are using the Amazon Mechanical Turk workforce. For more information, see the `analyze_document` documentation in the `AWS SDK for Python (Boto) API Reference`.

```python
response = client.analyze_document(
    Document={"S3Object": {"Bucket": "AWSDOC-EXAMPLE-BUCKET", "Name": "document-name.pdf"},
    HumanLoopConfig={
        "HumanLoopName": "human-loop-name",
        "DataAttributes": {ContentClassifiers: [
            "FreeOfPersonallyIdentifiableInformation", "FreeOfAdultContent"]
    }
    FeatureTypes=['FORMS']
)```
Human loops are only created if Amazon Textract's confidence for document analysis task meets the activation conditions you specified in your human review workflow. You can check the response element to determine if a human loop has been created. To see everything included in this response, see HumanLoopActivationOutput.

```python
if "HumanLoopArn" in analyzeDocumentResponse["HumanLoopActivationOutput"]:
    # A human loop has been started!
    print(f"A human loop has been started with ARN:
    {analyzeDocumentResponse["HumanLoopActivationOutput"]['HumanLoopArn']}")
```

Amazon Rekognition – Image moderation

The following example uses the AWS SDK for Python (Boto3) to call detect_moderation_labels in us-west-2. Replace the italicized red text with your resources. Include the DataAttributes parameter if you are using the Amazon Mechanical Turk workforce. For more information, see the detect_moderation_labels documentation in the AWS SDK for Python (Boto) API Reference.

```python
response = client.detect_moderation_labels(
    Image={"S3Object":{"Bucket": "AWSDOC-EXAMPLE-BUCKET", "Name": "imageName.png"}},
    HumanLoopConfig={
        "HumanLoopName": "human-loop-name",
        "DataAttributes": {"ContentClassifiers": ["FreeOfPersonallyIdentifiableInformation","FreeOfAdultContent"]}
    }
)
```

Human loops are only created if Amazon Rekognition's confidence for an image moderation task meets the activation conditions you specified in your human review workflow. You can check the response element to determine if a human loop has been created. To see everything included in this response, see HumanLoopActivationOutput.
if "HumanLoopArn" in response["HumanLoopActivationOutput"]:
    # A human loop has been started!
    print(f"A human loop has been started with ARN: {response["HumanLoopActivationOutput"]['HumanLoopArn']}")

Custom Integration

The following example uses the AWS SDK for Python (Boto3) to call `start_human_loop` in `us-west-2`. Replace the italicized red text with your resources. Include the `DataAttributes` parameter if you are using the Amazon Mechanical Turk workforce. For more information, see the `start_human_loop` documentation in the `AWS SDK for Python (Boto) API Reference`.

```python
response = client.start_human_loop(
    HumanLoopName= "human-loop-name",
    HumanLoopInput={"InputContent": inputContentJson},
    DataAttributes="ContentClassifiers": [
        "FreeOfPersonallyIdentifiableInformation","FreeOfAdultContent"
    ]
)
```

This example stores input content in the variable `inputContentJson`. Assume that the input content contains two elements: a text blurb and sentiment (such as Positive, Negative, or Neutral), and it is formatted as follows:

```python
inputContent = {
    "initialValue": sentiment,
    "taskObject": blurb
}
```

The keys `initialValue` and `taskObject` must correspond to the keys used in the liquid elements of the worker task template. Refer to the custom template in `Create a Human Task UI` to see an example.

To create `inputContentJson`, do the following:

```python
import json
```
inputContentJson = json.dumps(inputContent)

A human loop starts each time you call start_human_loop. To check the status of your human loop, use describe_human_loop:

human_loop_info = a2i.describe_human_loop(HumanLoopName="human_loop_name")
print(f"HumanLoop Status: {resp["HumanLoopStatus"]}" )
print(f"HumanLoop Output Destination: {resp["HumanLoopOutput"]}")

Use Cases and Examples Using Amazon A2I

You can use Amazon Augmented AI to incorporate a human review into your workflow for built-in task types, Amazon Textract and Amazon Rekognition, or your own custom tasks using a custom task type.

When you create a human review workflow using one of the built-in task types, you can specify conditions, such as confidence thresholds, that initiate a human review. The service (Amazon Rekognition or Amazon Textract) creates a human loop on your behalf when these conditions are met and supplies your input data directly to Amazon A2I to send to human reviewers. To learn more about the built-in task types, use the following:

- Use Amazon Augmented AI with Amazon Textract
- Use Amazon Augmented AI with Amazon Rekognition

When you use a custom task type, you create and start a human loop using the Amazon A2I Runtime API. Use the custom task type to incorporate a human review workflow with other AWS services or your own custom ML application.

- For more details, see Use Amazon Augmented AI with Custom Task Types

The following table outlines a variety of Amazon A2I use cases that you can explore using SageMaker Jupyter Notebooks. To get started with a Jupyter Notebook, use the instructions in Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook. For more examples, see this GitHub repository.
<table>
<thead>
<tr>
<th>Use Case</th>
<th>Description</th>
<th>Task Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use Amazon A2I with Amazon Textract</td>
<td>Have humans review single-page documents to review important form key-value pairs, or have Amazon Textract randomly sample and send documents from your dataset to humans for review.</td>
<td>Built-in</td>
</tr>
<tr>
<td>Use Amazon A2I with Amazon Rekognition</td>
<td>Have humans review unsafe images for explicit adult or violent content if Amazon Rekognition returns a low confidence score, or have Amazon Rekognition randomly sample and send images from your dataset to humans for review.</td>
<td>Built-in</td>
</tr>
<tr>
<td>Use Amazon A2I with Amazon Comprehend</td>
<td>Have humans review Amazon Comprehend inferences about text data such as sentiment analysis, text syntax, and entity detection.</td>
<td>Custom</td>
</tr>
<tr>
<td>Use Amazon A2I with Amazon Transcribe</td>
<td>Have humans review Amazon Transcribe transcriptions of video or audio files. Use the results of transcription human review loops to create a custom vocabulary and improve future transcriptions of similar video or audio content.</td>
<td>Custom</td>
</tr>
<tr>
<td>Use Case</td>
<td>Description</td>
<td>Task Type</td>
</tr>
<tr>
<td>----------</td>
<td>-------------</td>
<td>-----------</td>
</tr>
<tr>
<td>Use Amazon A2I with Amazon Translate</td>
<td>Have humans review low-confidence translations returned from Amazon Translate.</td>
<td>Custom</td>
</tr>
<tr>
<td>Use Amazon A2I to review real time ML inferences</td>
<td>Use Amazon A2I to review real-time, low-confidence inferences made by a model deployed to a SageMaker hosted endpoint and incrementally train your model using Amazon A2I output data.</td>
<td>Custom</td>
</tr>
<tr>
<td>Use Amazon A2I to review tabular data</td>
<td>Use Amazon A2I to integrate a human review loop into an ML application that uses tabular data.</td>
<td>Custom</td>
</tr>
</tbody>
</table>

Topics

- Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook
- Use Amazon Augmented AI with Amazon Textract
- Use Amazon Augmented AI with Amazon Rekognition
- Use Amazon Augmented AI with Custom Task Types

Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook

For an end-to-end example that demonstrates how to integrate an Amazon A2I human review loop into a machine learning workflow, you can use a Jupyter Notebook from this GitHub Repository in a SageMaker notebook instance.
To use an Amazon A2I custom task type sample notebook in an Amazon SageMaker notebook instance:

1. If you do not have an active SageMaker notebook instance, create one by following the instructions in [Step 1: Create an Amazon SageMaker Notebook Instance](#).

2. When your notebook instance is active, choose **Open JupyterLab** to the right of the notebook instance's name. It may take a few moments for JupyterLab to load.

3. Choose the ![clone](#) icon to clone a GitHub repository into your workspace.

4. Enter the [amazon-a2i-sample-jupyter-notebooks](#) repository HTTPS URL.

5. Choose **CLONE**.

6. Open the notebook that you would like to run.

7. Follow the instructions in the notebook to configure your human review workflow and human loop and run the cells.

8. To avoid incurring unnecessary charges, when you are done with the demo, stop and delete your notebook instance in addition to any Amazon S3 buckets, IAM roles, and CloudWatch Events resources created during the walkthrough.

---

Use Amazon Augmented AI with Amazon Textract

Amazon Textract enables you to add document text detection and analysis to your applications. Amazon Augmented AI (Amazon A2I) directly integrates with Amazon Textract's AnalyzeDocument API operation. You can use AnalyzeDocument to analyze a document for relationships between detected items. When you add an Amazon A2I human review loop to an AnalyzeDocument request, Amazon A2I monitors the Amazon Textract results and sends a document to one or more human workers for review when the conditions specified in your flow definition are met. For example, if you want a human to review a specific key like Full name: and their associated input values, you can create an activation condition that starts a human review any time the Full name: key is detected or when the inference confidence for that key falls within a range that you specify.

The following image depicts the Amazon A2I built-in workflow with Amazon Textract. On the left, the resources that are required to create an Amazon Textract human review workflow are depicted: and Amazon S3 bucket, activation conditions, a worker task template, and a work team.
These resources are used to create a human review workflow, or flow definition. An arrow points right to the next step in the workflow: using Amazon Textract to configure a human loop with the human review workflow. A second arrow points right from this step to the step in which activation conditions specified in the human review workflow are met. This initiates the creation of a human loop. On the right of the image, the human loop is depicted in three steps: 1) the worker UI and tools are generated and the task is made available to workers, 2) workers review input data, and finally, 3) results are saved in Amazon S3.

You can specify when Amazon Textract sends a task to a human worker for review when creating a human review workflow or flow definition by specifying *activation conditions*.

You can set the following activation conditions when using the Amazon Textract task type:

- Initiate a human review for specific form keys based on the form key confidence score.
- Initiate a human review when specific form keys are missing.
- Initiate human review for all form keys identified by Amazon Textract with confidence scores in a specified range.
- Randomly send a sample of forms to humans for review.

When your activation condition depends on form key confidence scores, you can use two types of prediction confidence to initiate human loops:

- **Identification confidence** – The confidence score for key-value pairs detected within a form.
• **Qualification confidence** – The confidence score for text contained within key and value in a form.

In the image in the following section, **Full Name: Jane Doe** is the key-value pair, **Full Name** is the key, and **Jane Doe** is the value.

You can set these activation conditions using the Amazon SageMaker console when you create a human review workflow, or by creating a JSON for human loop activation conditions and specifying this as input in the `HumanLoopActivationConditions` parameter of the `CreateFlowDefinition` API operation. To learn how specify activation conditions in JSON format, see [JSON Schema for Human Loop Activation Conditions in Amazon Augmented AI](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-human-loop-activate-conditions-json.html) and [Use Human Loop Activation Conditions JSON Schema with Amazon Textract](https://docs.aws.amazon.com/textract/latest/dg/a2i-human-loop-activate-conditions-json.html).

**Note**
When using Augmented AI with Amazon Textract, create Augmented AI resources in the same AWS Region you use to call `AnalyzeDocument`.

---

**Get Started: Integrate a Human Review into an Amazon Textract Analyze Document Job**

To integrate a human review into an Amazon Textract text detection and analysis job, you need to create a flow definition, and then use the Amazon Textract API to integrate that flow definition into your workflow. To learn how to create a flow definition using the SageMaker console or Augmented AI API, see the following topics:

- [Create a Human Review Workflow (Console)](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-human-loop-activate-conditions-json.html)
- [Create a Human Review Workflow (API)](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-human-loop-activate-conditions-json.html)

After you've created your flow definition, see [Using Augmented AI with Amazon Textract](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-human-loop-activate-conditions-json.html) to learn how to integrate your flow definition into your Amazon Textract task.

**End-to-End Example Using Amazon Textract and Amazon A2I**

For an end-to-end example that demonstrates how to use Amazon Textract with Amazon A2I using the console, see [Tutorial: Get Started in the Amazon A2I Console](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-human-loop-activate-conditions-json.html).

To learn how to use the Amazon A2I API to create and start a human review, you can use [Amazon Augmented AI (Amazon A2I) integration with Amazon Textract's Analyze Document [Example]](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-human-loop-activate-conditions-json.html) in a...
SageMaker Notebook instance. To get started, see Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook.

**A2I Textract Worker Console Preview**

When they're assigned a review task in an Amazon Textract workflow, workers might see a user interface similar to the following:

You can customize this interface in the SageMaker console when you create your human review definition, or by creating and using a custom template. To learn more, see Create and Manage Worker Task Templates.

**Use Amazon Augmented AI with Amazon Rekognition**

Amazon Rekognition makes it easy to add image analysis to your applications. The Amazon Rekognition DetectModerationLabels API operation is directly integrated with Amazon A2I so that you can easily create a human loop to review unsafe images, such as explicit adult or violent content. You can use DetectModerationLabels to configure a human loop using a flow definition ARN. This enables Amazon A2I to analyze predictions made by Amazon Rekognition and send results to a human for review to ensure they meet the conditions set in your flow definition.

The following image depicts the Amazon A2I built-in workflow with Amazon Rekognition. On the left, the resources that are required to create an Amazon Rekognition human review workflow
are depicted: and Amazon S3 bucket, activation conditions, a worker task template, and a work
team. These resources are used to create a human review workflow, or flow definition. An arrow
points right to the next step in the workflow: using Amazon Rekognition to configure a human loop
with the human review workflow. A second arrow points right from this step to the step in which
activation conditions specified in the human review workflow are met. This initiates the creation of
a human loop. On the right of the image, the human loop is depicted in three steps: 1) the worker
UI and tools are generated and the task is made available to workers, 2) workers review input data,
and finally, 3) results are saved in Amazon S3.

You can set the following activation conditions when using the Amazon Rekognition task type:

- Initiate human review for labels identified by Amazon Rekognition based on the label confidence
  score.
- Randomly send a sample of images to humans for review.

You can set these activation conditions using the Amazon SageMaker console when you
create a human review workflow, or by creating a JSON for human loop activation conditions
and specifying this as input in the HumanLoopActivationConditions parameter of the
CreateFlowDefinition API operation. To learn how specify activation conditions in JSON
format, see JSON Schema for Human Loop Activation Conditions in Amazon Augmented AI and
Use Human Loop Activation Conditions JSON Schema with Amazon Rekognition.
Note

When using Augmented AI with Amazon Rekognition, create Augmented AI resources in the same AWS Region you use to call DetectModerationLabels.

Get Started: Integrate a Human Review into an Amazon Rekognition Image Moderation Job

To integrate a human review into an Amazon Rekognition, see the following topics:

- Create a Human Review Workflow (Console)
- Create a Human Review Workflow (API)

After you've created your flow definition, see Using Augmented AI with Amazon Rekognition to learn how to integrate your flow definition into your Amazon Rekognition task.

End-to-end Demo Using Amazon Rekognition and Amazon A2I

For an end-to-end example that demonstrates how to use Amazon Rekognition with Amazon A2I using the console, see Tutorial: Get Started in the Amazon A2I Console.

To learn how to use the Amazon A2I API to create and start a human review, you can use Amazon Augmented AI (Amazon A2I) integration with Amazon Rekognition [Example] in a SageMaker notebook instance. To get started, see Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook.

A2I Rekognition Worker Console Preview

When they're assigned a review task in an Amazon Rekognition workflow, workers might see a user interface similar to the following:
You can customize this interface in the SageMaker console when you create your human review definition, or by creating and using a custom template. To learn more, see Create and Manage Worker Task Templates.

Use Amazon Augmented AI with Custom Task Types

You can use Amazon Augmented AI (Amazon A2I) to incorporate a human review (human loop) into any machine learning workflow using the custom task type. This options gives you the most flexibility to customize the conditions under which your data objects are sent to humans for review, as well as the look and feel of your worker user interface.

When you use a custom task type, you create a custom human review workflow and specify the conditions under which a data object is sent for human review directly in your application.

The following image depicts the Amazon A2I custom workflow. A custom ML model is used to generate predictions. The client application filters these predictions using user-defined criteria and determines if a human review is required. If so, these predictions are sent to Amazon A2I for human review. Amazon A2I collects the results of human review in Amazon S3, which can access by the client application. If the filter determines that no human review is needed, predictions can be fed directly to the client application.
Use the procedures on this page to learn how to integrate Amazon A2I into any machine learning workflow using the custom task type.

Create a human loop using a flow definition, integrate it into your application, and monitor the results

1. Complete the Amazon A2I Prerequisites to Using Augmented AI. Note the following:
   - The path to the Amazon Simple Storage Service (Amazon S3) bucket or buckets where you store your input and output data.
   - The Amazon Resource Name (ARN) of an AWS Identity and Access Management (IAM) role with required permissions attached.
   - (Optional) The ARN of your private workforce, if you plan to use one.

2. Using HTML elements, create a custom worker template which Amazon A2I uses to generate your worker task UI. To learn how to create a custom template, see Create Custom Worker Task Templates.

3. Use the custom worker template from step 2 to generate a worker task template in the Amazon SageMaker console. To learn how, see Create a Worker Task Template.

In the next step, you create a flow definition:
• If you want to create a flow definition using the SageMaker API, note the ARN of this worker task template for the next step.

• If you are creating a flow definition using the console, your template automatically appears in **Worker task template** section when you choose **Create human review workflow**.

4. When creating your flow definition, provide the path to your S3 buckets, your IAM role ARN, and your worker template.

• To learn how to create a flow definition using the SageMaker CreateFlowDefinition API, see **Create a Human Review Workflow (API)**.

• To learn how to create a flow definition using the SageMaker console, see **Create a Human Review Workflow (Console)**.

5. Configure your human loop using the **Amazon A2I Runtime API**. To learn how, see **Create and Start a Human Loop**.

6. To control when human reviews are initiated in your application, specify conditions under which **StartHumanLoop** is called in your application. Human loop activation conditions, such as confidence thresholds that initiate the human loop, are not available when using Amazon A2I with custom task types. Every **StartHumanLoop** invocation results in a human review.

Once you have started a human loop, you can manage and monitor your loops using the Amazon Augmented AI Runtime API and Amazon EventBridge (also known as Amazon CloudWatch Events). To learn more, see **Monitor and Manage Your Human Loop**.

**End-to-end Tutorial Using Amazon A2I Custom Task Types**

For an end-to-end examples that demonstrates how to integrate Amazon A2I into a variety of ML workflows, see the table in **Use Cases and Examples Using Amazon A2I**. To get started using one of these notebooks, see **Use SageMaker Notebook Instance with Amazon A2I Jupyter Notebook**.

**Create a Human Review Workflow**

Use an Amazon Augmented AI (Amazon A2I) **human review workflow**, or **flow definition**, to specify the following:

• For the Amazon Textract and Amazon Rekognition built-in task types, the conditions under which your human loop is called

• The workforce to which your tasks are sent
• The set of instructions that your workforce receives, which is called a worker task template
• The configuration of your worker tasks, including the number of workers that receive a task and time limits to complete tasks
• Where your output data is stored

You can create a human review workflow in the SageMaker console or using the SageMaker CreateFlowDefinition operation. You can build a worker task template using the console for Amazon Textract and Amazon Rekognition task types while creating your flow definition.

⚠️ Important

Human loop activation conditions, which initiate the human loop—for example, confidence thresholds—aren’t available for Amazon A2I custom task types. When using the console to create a flow definition for a custom task type, you can’t specify activation conditions. When using the Amazon A2I API to create a flow definition for a custom task type, you can’t set the HumanLoopActivationConditions attribute of the HumanLoopActivationConditionsConfig parameter. To control when human reviews are initiated, specify conditions under which StartHumanLoop is called in your custom application. In this case, every StartHumanLoop invocation results in a human review. For more information, see Use Amazon Augmented AI with Custom Task Types.

Prerequisites

To create a human review workflow definition, you must have completed the prerequisites described in Prerequisites to Using Augmented AI.

If you use the API to create a flow definition for any task type, or if you use a custom task type when creating a flow definition in the console, first create a worker task template. For more information, see Create and Manage Worker Task Templates.

If you want to preview your worker task template while creating a flow definition for a built-in task type in the console, ensure that you grant the role that you use to create the flow definition permission to access the Amazon S3 bucket that contains your template artifacts using a policy like the one described in Enable Worker Task Template Previews.

Topics
Create a Human Review Workflow (Console)

Use this procedure to create an Amazon Augmented AI (Amazon A2I) human review workflow using the SageMaker console. If you are new to Amazon A2I, we recommend that you create a private work team using people in your organization, and use this work team's ARN when creating your flow definition. To learn how to set up a private workforce and create a work team, see Create a Private Workforce (Amazon SageMaker Console). If you have already set up a private workforce, see Create a Work Team Using the SageMaker Console to learn how to add a work team to that workforce.

If you are using Amazon A2I with one of the built-in task types, you can create worker instructions using a default worker task template provided by Augmented AI while creating a human review workflow in the console. To see samples of the default templates provided by Augmented AI, see the built-in task types in Use Cases and Examples Using Amazon A2I.

To create flow definition (console)

2. In the navigation pane, under the Augmented AI section, choose Human review workflows and then choose Create human review workflow.
3. In Overview, do the following:
   a. For Name, enter a unique workflow name. The name must be lowercase, unique within the AWS Region in your account, and can have up to 63 characters. Valid characters include: a-z, 0-9, and - (hyphen).
   b. For S3 location for output, enter the S3 bucket where you want to store the human review results. The bucket must be located in the same AWS Region as the workflow.
   c. For IAM role, choose the role that has the required permissions. If you choose a built-in task type and want to preview your worker template in the console, provide a role with the type of policy described in Enable Worker Task Template Previews attached.
4. For Task type, choose the task type that you want the human worker to perform.
5. If you chose the Amazon Rekognition or Amazon Textract task type, specify the conditions that invoke human review.

- For Amazon Rekognition image moderation tasks, choose an inference confidence score threshold interval that initiates human review.

- For Amazon Textract tasks, you can initiate a human review when specific form keys are missing or when form key detection confidence is low. You can also initiate a human review if, after evaluating all of the form keys in the text, confidence is lower than your required threshold for any form key. Two variables specify your confidence thresholds: **Identification confidence** and **Qualification confidence**. To learn more about these variables, see [Use Amazon Augmented AI with Amazon Textract](#).

- For both task types, you can randomly send a percentage of data objects (images or forms) and their labels to humans for review.

6. Configure and specify your worker task template:

a. If you are using the Amazon Rekognition or Amazon Textract task type:

   - In the **Create template** section:
     - To create instructions for your workers using the Amazon A2I default template for Amazon Rekognition and Amazon Textract task types, choose **Build from a default template**.
     - If you choose **Build from a default template**, create your instructions under **Worker task design**:
       - Provide a **Template name** that is unique in the AWS Region you are in.
       - In the **Instructions** section, provide detailed instructions on how to complete your task. To help workers achieve greater accuracy, provide good and bad examples.
       - (Optional) In **Additional instructions**, provide your workers with additional information and instructions.

       For information on creating effective instructions, see [Creating Good Worker Instructions](#).

     - To select a custom template that you've created, choose it from the **Template** menu and provide a **Task description** to briefly describe the task for your workers. To learn how to create a custom template, see [Create a Worker Task Template](#).
b. If you are using the custom task type:

- In the **Worker task template** section, select your template from the list. All of the templates that you have created in the SageMaker console appear in this list. To learn how to create a template for custom task types, see [Create and Manage Worker Task Templates](#).

7. (Optional) Preview your worker template:

For Amazon Rekognition and Amazon Textract task types, you have the option to choose **See a sample worker task** to preview your worker task UI.

If you are creating a flow definition for a custom task type, you can preview your worker task UI using the `RenderUiTemplate` operation. For more information, see [Preview a Worker Task Template](#).

8. For **Workers**, choose a workforce type.

9. Choose **Create**.

**Next Steps**

After you've created a human review workflow, it appears in the console under **Human review workflows**. To see your flow definition's Amazon Resource Name (ARN) and configuration details, choose the workflow by selecting its name.

If you are using a built-in task type, you can use the flow definition ARN to start a human loop using that AWS service's API (for example, the Amazon Textract API). For custom task types, you can use the ARN to start a human loop using the Amazon Augmented AI Runtime API. To learn more about both options, see [Create and Start a Human Loop](#).

**Create a Human Review Workflow (API)**

To create a flow definition using the SageMaker API, you use the `CreateFlowDefinition` operation. After you complete the [Prerequisites to Using Augmented AI](#), use the following procedure to learn how to use this API operation.

For an overview of the `CreateFlowDefinition` operation, and details about each parameter, see [CreateFlowDefinition](#).
To create a flow definition (API)

1. For FlowDefinitionName, enter a unique name. The name must be unique within the AWS Region in your account, and can have up to 63 characters. Valid characters include: a-z, 0-9, and - (hyphen).

2. For RoleArn, enter the ARN of the role that you configured to grant access to your data sources.

3. For HumanLoopConfig, enter information about the workers and what they should see. For information about each parameter in HumanLoopConfig, see HumanLoopConfig.

4. (Optional) If you are using a built-in task type, provide conditions that initiate a human loop in HumanLoopActivationConfig. To learn how to create the input required for the HumanLoopActivationConfig parameter, see JSON Schema for Human Loop Activation Conditions in Amazon Augmented AI. If you do not specify conditions here, when you provide a flow definition to the AWS service associated with a built-in task type (for example, Amazon Textract or Amazon Rekognition), that service sends every task to a human worker for review.

   If you are using a custom task type, HumanLoopActivationConfig is disabled. To learn how to control when tasks are sent to human workers using a custom task type, see Use Amazon Augmented AI with Custom Task Types.

5. (Optional) If you are using a built-in task type, specify the integration source (for example, Amazon Rekognition or Amazon Textract) in the HumanLoopRequestSource parameter.

6. For OutputConfig, indicate where in Amazon Simple Storage Service (Amazon S3) to store the output of the human loop.

7. (Optional) Use Tags to enter key-value pairs to help you categorize and organize a flow definition. Each tag consists of a key and a value, both of which you define.

Amazon Textract – Key-value pair extraction

The following is an example of a request to create an Amazon Textract human review workflow (flow definition) using the AWS SDK for Python (Boto3). You must use 'AWS/Textextract/AnalyzeDocument/Forms/V1' to create a Amazon Textract human loop. Only include PublicWorkforceTaskPrice if you are using the Mechanical Turk workforce.

```python
sagemaker_client = boto3.client('sagemaker', aws_region)
response = sagemaker_client.create_flow_definition(
    FlowDefinitionName='ExampleFlowDefinition',
```
Create a Human Review Workflow
Amazon Rekognition – Image moderation

The following is an example of a request to create an Amazon Rekognition human review workflow (flow definition) using the AWS SDK for Python (Boto3). You must use 'AWS/Rekognition/DetectModerationLabels/Image/V3' to create an Amazon Rekognition flow definition. Only include PublicWorkforceTaskPrice if you are using the Mechanical Turk workforce.

```python
sagemaker_client = boto3.client('sagemaker', aws_region)

response = sagemaker_client.create_flow_definition(
    FlowDefinitionName='ExampleFlowDefinition',
    HumanLoopRequestSource={
        'AwsManagedHumanLoopRequestSource': 'AWS/Rekognition/DetectModerationLabels/Image/V3'
    },
    HumanLoopActivationConfig={
        'HumanLoopActivationConditionsConfig': {
            'HumanLoopActivationConditions': '{...}'
        }
    },
    HumanLoopConfig={
        'TaskTitle': 'Example task title',
        'TaskDescription': 'Example task description.',
        'TaskCount': 123,
        'TaskAvailabilityLifetimeInSeconds': 123,
        'TaskTimeLimitInSeconds': 123,
        'TaskKeywords': ['Keyword1','Keyword2'],
        'PublicWorkforceTaskPrice': {
            'AmountInUsd': {
                'Dollars': 123,
                'Cents': 123,
                'TenthFractionsOfACent': 123
            }
        }
    },
    OutputConfig={
```

Create a Human Review Workflow
Custom Workflow

The following is an example of a request to create a human review workflow (flow definition) for a custom integration. To create this type of human review workflow, omit HumanLoopRequestSource from the flow definition request. You only need to include PublicWorkforceTaskPrice if you are using the Mechanical Turk workforce.

```python
sagemaker_client = boto3.client('sagemaker', **aws_region)
response = sagemaker_client.create_flow_definition(
    FlowDefinitionName='ExampleFlowDefinition',
    HumanLoopActivationConfig={
        'HumanLoopActivationConditionsConfig': {
            'HumanLoopActivationConditions': '{...}'
        }
    },
    HumanLoopConfig={
        'WorkteamArn': 'arn:aws:sagemaker:**aws_region**:aws_account_number:workteam/private-crowd/workteam_name',
        'HumanTaskUiArn': 'arn:aws:sagemaker:**aws_region**:aws_account_number:human-task-ui/template_name',
        'TaskTitle': 'Example task title',
        'TaskDescription': 'Example task description.',
        'TaskCount': 123,
        'TaskAvailabilityLifetimeInSeconds': 123,
        'TaskTimeLimitInSeconds': 123,
        'TaskKeywords': [
            'Keyword1', 'Keyword2'
        ],
        'PublicWorkforceTaskPrice': {
            'AmountInUsd': {
                'S3OutputPath': 's3://bucket/path/',
                'KmsKeyId': '1234abcd-12ab-34cd-56ef-1234567890ab'
            },
            RoleArn='arn:aws:iam::aws_account_number:role/role_name',
            Tags=[
                {'Key': 'KeyName',
                'Value': 'ValueName'}
            ],
        }
    }
)
```
Next Steps

The return value of a successful call of the CreateFlowDefinition API operation is a flow definition Amazon Resource Name (ARN).

If you are using a built-in task type, you can use the flow definition ARN to start a human loop using that AWS service's API (i.e. the Amazon Textract API). For custom task types, you can use the ARN to start a human loop using the Amazon Augmented AI Runtime API. To learn more about both of these options, see Create and Start a Human Loop.

JSON Schema for Human Loop Activation Conditions in Amazon Augmented AI

The HumanLoopActivationConditions is an input parameter of the CreateFlowDefinition API. This parameter is a JSON-formatted string. The JSON models the conditions under which a human loop is created when those conditions are evaluated against the response from an integrating AI service API (such as Rekognition.DetectModerationLabels or Textract.AnalyzeDocument). This response is referred to as an inference. For example, Amazon Rekognition sends an inference of a moderation label with an associated confidence score. In this example, the inference is the model's best estimate of the appropriate label for an image. For Amazon Textract, inference is made on the association between blocks of text (key-value pairs),
such as the association between Name: and Sue in a form as well as content within a block of text, or *word block*, such as 'Name'.

The following is the schema for the JSON. At the top level, the `HumanLoopActivationConditions` has a JSON array, `Conditions`. Each member of this array is an independent condition that, if evaluated to `true`, results in Amazon A2I creating a human loop. Each such independent condition can be a simple condition or a complex condition. A simple condition has the following attributes:

- **ConditionType**: This attribute identifies the type of condition. Each AWS AI service API that integrates with Amazon A2I defines its own set of allowed `ConditionTypes`.
  - **Rekognition DetectModerationLabels** – This API supports the `ModerationLabelConfidenceCheck` and `Sampling` `ConditionType` values.
  - **Textract AnalyzeDocument** – This API supports the `ImportantFormKeyConfidenceCheck`, `MissingImportantFormKey`, and `Sampling` `ConditionType` values.
  - **ConditionParameters** – This is a JSON object that parameterizes the condition. The set of allowed attributes of this object is dependent on the value of the `ConditionType`. Each `ConditionType` defines its own set of `ConditionParameters`.

A member of the `Conditions` array can model a complex condition. This is accomplished by logically connecting simple conditions using the `And` and `Or` logical operators and nesting the underlying simple conditions. Up to two levels of nesting are supported.

```json
{
  "$schema": "http://json-schema.org/draft-07/schema#",
  "definitions": {
    "Condition": {
      "type": "object",
      "properties": {
        "ConditionType": {
          "type": "string"
        },
        "ConditionParameters": {
          "type": "object"
        }
      },
      "required": ["ConditionType"]
    }
  }
}
```
Create a Human Review Workflow
Note

Human loop activation conditions aren't available for human review workflows that are integrated with custom task types. The HumanLoopActivationConditions parameter is disabled for custom task types.

Topics

• Use Human Loop Activation Conditions JSON Schema with Amazon Textract
• Use Human Loop Activation Conditions JSON Schema with Amazon Rekognition

Use Human Loop Activation Conditions JSON Schema with Amazon Textract

When used with Amazon A2I, the AnalyzeDocument operation supports the following inputs in the ConditionType parameter:

• ImportantFormKeyConfidenceCheck – Use this condition to create a human loop when inference confidence is within a specified range for document form keys and word blocks. A form key is any word in a document that is associated with an input. The input is called a value. Together, form keys and values are referred to as key-value pairs. A word block refers to the words that Amazon Textract recognizes inside of a detected block of text. To learn more about Amazon Textract document blocks, see Documents and Block Objects in the Amazon Textract Developer Guide.

• MissingImportantFormKey – Use this condition to create a human loop when Amazon Textract did not identify the key or its associated aliases within the document.

• Sampling – Use this condition to specify a percentage of forms to send to humans for review, regardless of inference confidence scores. Use this condition to do the following:
  • Audit your ML model by randomly sampling all forms analyzed by your model and sending a specified percentage to humans for review.
• Using the `ImportantFormKeyConfidenceCheck` condition, randomly sample a percentage of the inferences that met the conditions specified in `ImportantFormKeyConfidenceCheck` to start a human loop and send only the specified percentage to humans for review.

**Note**

If you send the same request to `AnalyzeDocument` multiple times, the result of `Sampling` does not change for the inference of that input. For example, if you make an `AnalyzeDocument` request once, and `Sampling` doesn't initiate a human loop, subsequent requests to `AnalyzeDocument` with the same configuration do not initiate a human loop.

**ImportantFormKeyConfidenceCheck Inputs and Results**

The `ImportantFormKeyConfidenceCheck` `ConditionType` supports the following `ConditionParameters`:

• `ImportantFormKey` – A string representing a key in a key-value pair detected by Amazon Textract that needs to be reviewed by human workers. If the value of this parameter is the special catch-all value (*), then all keys are considered to be matched to the condition. You can use this to model the case where any key-value pair satisfying certain confidence thresholds needs human review.

• `ImportantFormKeyAliases` – An array that represents alternate spellings or logical equivalents for the important form key.

• `KeyValueBlockConfidenceEquals`

• `KeyValueBlockConfidenceLessThan`

• `KeyValueBlockConfidenceLessThanEquals`

• `KeyValueBlockConfidenceGreaterThan`

• `KeyValueBlockConfidenceGreaterThanEquals`

• `WordBlockConfidenceEquals`

• `WordBlockConfidenceLessThan`

• `WordBlockConfidenceLessThanEquals`

• `WordBlockConfidenceGreaterThan`
• WordBlockConfidenceGreaterThanEquals

When you use the ImportantFormKeyConfidenceCheck ConditionType, Amazon A2I sends the key-value block and word block inferences of the key-value blocks and associated aliases that you specified in ImportantFormKey and ImportantFormKeyAliases for human review.

When creating a flow definition, if you use the default worker task template that is provided in the Human review workflows section of the Amazon SageMaker console, key-value and block inferences sent for human review by this activation condition are included in the worker UI. If you use a custom worker task template, you need to include the `{{ task.input.selectedAiServiceResponse.blocks }}` element to include initial-value input data (inferences) from Amazon Textract. For an example of a custom template that uses this input element, see Custom Template Example for Amazon Textract.

**MissingImportantFormKey Inputs and Results**

The MissingImportantFormKey ConditionType supports the following ConditionParameters:

- **ImportantFormKey** – A string representing a key in a key-value pair detected by Amazon Textract that needs to be reviewed by human workers.

- **ImportantFormKeyAliases** – An array that represents alternate spellings or logical equivalents for the important form key.

When you use the MissingImportantFormKey ConditionType, if the key in ImportantFormKey or aliases in ImportantFormKeyAliases are not included in the Amazon Textract inference, that form is sent to human for review and no predicted key-value pairs are included. For example, if Amazon Textract only identified Address and Phone in a form, but was missing the ImportantFormKey Name (in the MissingImportantFormKey condition type) that form would be sent to humans for review without any of the form keys detected (Address and Phone).

If you use the default worker task template that is provided in the SageMaker console, a task is created asking workers to identify the key in ImportantFormKey and associated value. If you use a custom worker task template, you need to include the `<task.input.humanLoopContext>` custom HTML element to configure this task.
Sampling Inputs and Results

The Sampling ConditionType supports the RandomSamplingPercentage ConditionParameters. The input for RandomSamplingPercentage must be real number between 0.01 and 100. This number represents the percentage of data that qualifies for a human review and is sent to humans for review. If you use the Sampling condition without any other conditions, this number represents the percentage of all resulting inferences made by the AnalyzeDocument operation from a single request that is sent to humans for review.

If you specify the Sampling condition without any other condition type, all key-value and block inferences are sent to workers for review.

When creating a flow definition, if you use the default worker task template that is provided in the Human review workflows section of the SageMaker console, all key-value and block inferences sent for human review by this activation condition are included in the worker UI. If you use a custom worker task template, you need to include the 

```json
{{ task.input.selectedAiServiceResponse.blocks }}
```

element to include initial-value input data (inferences) from Amazon Textract. For an example of a custom template that uses this input element, see Custom Template Example for Amazon Textract.

Examples

While only one condition needs to evaluate to true to initiate a human loop, Amazon A2I evaluates all conditions for each object analyzed by Amazon Textract. The human reviewers are asked to review the important form keys for all the conditions that evaluated to true.

Example 1: Detect important form keys with confidence scores in a specified range that initiate a human loop

The following example shows a HumanLoopActivationConditions JSON that initiates a human loop if any one of the following three conditions is met:

- The Amazon Textract AnalyzeDocument API returns a key-value pair whose key is one of Employee Name, Name, or EmployeeName, with the confidence of the key-value block being less than 60 and the confidences of each of the word blocks making up the key and value being less than 85.
- The Amazon Textract AnalyzeDocument API returns a key-value pair whose key is one of Pay Date, PayDate, DateOfPay, or pay-date, with the confidence of the key-value block being less than 65 and the confidences of each of the word blocks making up the key and value being less than 85.
The Amazon Textract AnalyzeDocument API returns a key-value pair whose key is one of Gross Pay, GrossPay, or GrossAmount, with the confidence of the key-value block being less than 60 and the confidences of each of the word blocks making up the key and value being less than 85.

```json
{
   "Conditions": [
       {
           "ConditionType": "ImportantFormKeyConfidenceCheck",
           "ConditionParameters": {
               "ImportantFormKey": "Employee Name",
               "ImportantFormKeyAliases": ["Name", "EmployeeName"],
               "KeyValueBlockConfidenceLessThan": 60,
               "WordBlockConfidenceLessThan": 85
           }
       },
       {
           "ConditionType": "ImportantFormKeyConfidenceCheck",
           "ConditionParameters": {
               "ImportantFormKey": "Pay Date",
               "ImportantFormKeyAliases": ["PayDate", "DateOfPay", "pay-date"],
               "KeyValueBlockConfidenceLessThan": 65,
               "WordBlockConfidenceLessThan": 85
           }
       },
       {
           "ConditionType": "ImportantFormKeyConfidenceCheck",
           "ConditionParameters": {
               "ImportantFormKey": "Gross Pay",
               "ImportantFormKeyAliases": ["GrossPay", "GrossAmount"],
               "KeyValueBlockConfidenceLessThan": 60,
               "WordBlockConfidenceLessThan": 85
           }
       }
   ]
}```
Example 2: Use ImportantFormKeyConfidenceCheck

In the following example, if Amazon Textract detects a key-value pair whose confidence for the key-value block is less than 60 and is less than 90 for any underlying word blocks, it creates a human loop. The human reviewers are asked to review all the form key-value pairs that matched the confidence value comparisons.

```json
{
  "Conditions": [
    {
      "ConditionType": "ImportantFormKeyConfidenceCheck",
      "ConditionParameters": {
        "ImportantFormKey": "*",
        "KeyValueBlockConfidenceLessThan": 60,
        "WordBlockConfidenceLessThan": 90
      }
    }
  ]
}
```

Example 3: Use Sampling

In the following example, 5% of inferences resulting from an Amazon Textract AnalyzeDocument request are sent to human workers for review. All detected key-value pairs returned by Amazon Textract are sent to workers for review.

```json
{
  "Conditions": [
    {
      "ConditionType": "Sampling",
      "ConditionParameters": {
        "RandomSamplingPercentage": 5
      }
    }
  ]
}
```

Example 4: Use MissingImportantFormKey

Create a Human Review Workflow
In the following example, if Mailing Address or its alias, Mailing Address:, is missing from keys detected by Amazon Textract, a human review is initiated. When using the default worker task template, the worker UI asks workers to identify the key Mailing Address or Mailing Address: and its associated value.

```json
{
   "ConditionType": "MissingImportantFormKey",
   "ConditionParameters": {
      "ImportantFormKey": "Mailing Address",
      "ImportantFormKeyAliases": ["Mailing Address:"
   }
}
```

**Example 5: Use Sampling and ImportantFormKeyConfidenceCheck with the And operator**

In this example, 5% of key-value pairs detected by Amazon Textract whose key is one of Pay Date, PayDate, DateOfPay, or pay-date, with the confidence of the key-value block less than 65 and the confidences of each of the word blocks making up the key and value less than 85, are sent to workers for review.

```json
{
   "Conditions": [
      {
         "And": [
            {
               "ConditionType": "Sampling",
               "ConditionParameters": {
                  "RandomSamplingPercentage": 5
               }
            },
            {
               "ConditionType": "ImportantFormKeyConfidenceCheck",
               "ConditionParameters": {
                  "ImportantFormKey": "Pay Date",
                  "ImportantFormKeyAliases": ["PayDate",
                                               "DateOfPay",
                                               "pay-date"
                  ],
                  "KeyValueBlockConfidenceLessThan": 65,
                  "WordBlockConfidenceLessThan": 85
               }
            }
      }
   ]
}
```
Example 6: Use Sampling and ImportantFormKeyConfidenceCheck with the And operator

Use this example to configure your human review workflow to always send low confidence inferences of a specified key-value pair for human review and sample high confidence inference of a key-value pair at a specified rate.

In the following example, a human review is initiated in one of the following ways:

- Key-value pairs detected whose key is one of Pay Date, PayDate, DateOfPay, or pay-date, with key-value and word block confidences less than 60, are sent for human review. Only the Pay Date form key (and its aliases) and associated values are sent to workers to review.
- 5% of key-value pairs detected whose key is one of Pay Date, PayDate, DateOfPay, or pay-date, with key-value and word block confidences greater than 90, are sent for human review. Only the Pay Date form key (and its aliases) and associated values are sent to workers to review.

```json
{
    "Conditions": [
    {
        "Or": [
        {
            "ConditionType": "ImportantFormKeyConfidenceCheck",
            "ConditionParameters": {
                "ImportantFormKey": "Pay Date",
                "ImportantFormKeyAliases": [
                "PayDate",
                "DateOfPay",
                "pay-date"
                ],
                "KeyValueBlockConfidenceLessThan": 60,
                "WordBlockConfidenceLessThan": 60
            }
        },
    {
        "And": [
```
Example 7: Use Sampling and ImportantFormKeyConfidenceCheck with the Or operator

In the following example, the Amazon Textract AnalyzeDocument operation returns a key-value pair whose key is one of Pay Date, PayDate, DateOfPay, or pay-date, with the confidence of the key-value block less than 65 and the confidences of each of the word blocks making up the key and value less than 85. Additionally, 5% of all other forms initiate a human loop. For each form randomly chosen, all key-value pairs detected for that form are sent to humans for review.

```json
{
   "Conditions": [
      {
         "Or": [
            {
               "ConditionType": "Sampling",
               "ConditionParameters": {
                  "RandomSamplingPercentage": 5
               }
            },
            {
               "ConditionType": "ImportantFormKeyConfidenceCheck",
               "ConditionParameters": {
                  "ImportantFormKey": "Pay Date",
                  "ImportantFormKeyAliases": [
                     "PayDate",
                     "DateOfPay",
                     "pay-date"
                  ],
                  "KeyValueBlockConfidenceLessThan": 90
               }
            }
         ]
      }
   ]
}
```
Use Human Loop Activation Conditions JSON Schema with Amazon Rekognition

When used with Amazon A2I, the Amazon Rekognition DetectModerationLabels operation supports the following inputs in the ConditionType parameters:

- **ModerationLabelConfidenceCheck** – Use this condition type to create a human loop when inference confidence is low for one or more specified labels.
- **Sampling** – Use this condition to specify a percentage of all inferences to send to humans for review. Use this condition to do the following:
  - Audit your ML model by randomly sampling all of your model's inferences and sending a specified percentage to humans for review.
  - Using the **ModerationLabelConfidenceCheck** condition, randomly sample a percentage of the inferences that met the conditions specified in **ModerationLabelConfidenceCheck** to start a human loop and send only the specified percentage to humans for review.

**Note**

If you send the same request to DetectModerationLabels multiple times, the result of Sampling does not change for the inference of that input. For example, if you make a
DetectModerationLabels request once, and Sampling does not initiate a human loop, subsequent requests to DetectModerationLabels with the same configuration don't initiate a human loop.

When creating a flow definition, if you use the default worker task template that is provided in the Human review workflows section of the Amazon SageMaker console, inferences sent for human review by these activation conditions are included in the worker UI when a worker opens your task. If you use a custom worker task template, you need to include the <task.input.selectedAiServiceResponse.blocks> custom HTML element to access these inferences. For an example of a custom template that uses this HTML element, see Custom Template Example for Amazon Rekognition.

**ModerationLabelConfidenceCheck Inputs**

For the ModerationLabelConfidenceCheck ConditionType, the following ConditionParameters are supported:

- **ModerationLabelName** – The exact (case-sensitive) name of a ModerationLabel detected by the Amazon Rekognition DetectModerationLabels operation. You can specify the special catch-all value (*) to denote any moderation label.
- **ConfidenceEquals**
- **ConfidenceLessThan**
- **ConfidenceLessThanEquals**
- **ConfidenceGreaterThan**
- **ConfidenceGreaterThanEquals**

When you use the ModerationLabelConfidenceCheck ConditionType, Amazon A2I sends label inferences for the labels that you specified in ModerationLabelName for human review.

**Sampling Inputs**

The Sampling ConditionType supports the RandomSamplingPercentage ConditionParameters. The input for the RandomSamplingPercentage parameter should be a real number between 0.01 and 100. This number represents the percentage of inferences that qualifies for a human review that are sent to humans for review. If you use the Sampling condition...
without any other conditions, this number represents the percentage of all inferences that result from a single DetectModerationLabel request that are sent to humans for review.

**Examples**

**Example 1: Use ModerationLabelConfidenceCheck with the And operator**

The following example of a HumanLoopActivationConditions condition initiates a human loop when one or more of the following conditions are met:

- Amazon Rekognition detects the Graphic Male Nudity moderation label with a confidence between 90 and 99.
- Amazon Rekognition detects the Graphic Female Nudity moderation label with a confidence between 80 and 99.

Note the use of the Or and And logical operators to model this logic.

Although only one of the two conditions under the Or operator needs to evaluate to true for a human loop to be created, Amazon Augmented AI evaluates all conditions. Human reviewers are asked to review the moderation labels for all the conditions that evaluated to true.

```json
{
   "Conditions": [
      {
         "Or": [
            {
               "And": [
                  {
                     "ConditionType": "ModerationLabelConfidenceCheck",
                     "ConditionParameters": {
                        "ModerationLabelName": "Graphic Male Nudity",
                        "ConfidenceLessThanEquals": 99
                     }
                  },
                  {
                     "ConditionType": "ModerationLabelConfidenceCheck",
                     "ConditionParameters": {
                        "ModerationLabelName": "Graphic Male Nudity",
                        "ConfidenceGreaterThanEquals": 90
                     }
                  }
               ]
            }
      },
      {
         "ConditionType": "ModerationLabelConfidenceCheck",
         "ConditionParameters": {
            "ModerationLabelName": "Graphic Male Nudity",
            "ConfidenceLessThanEquals": 99
         }
      }
   ]
}
```
"And": [{
  "ConditionType": "ModerationLabelConfidenceCheck",
  "ConditionParameters": {
    "ModerationLabelName": "Graphic Female Nudity",
    "ConfidenceLessThanEquals": 99
  }
}, {
  "ConditionType": "ModerationLabelConfidenceCheck",
  "ConditionParameters": {
    "ModerationLabelName": "Graphic Female Nudity",
    "ConfidenceGreaterThanEquals": 80
  }
}]
}

Example 2: Use ModerationLabelConfidenceCheck with the catch-all value (*)

In the following example, if any moderation label with a confidence greater than or equal to 75 is detected, a human loop is initiated. Human reviewers are asked to review all moderation labels with confidence scores greater than or equal to 75.

{
  "Conditions": [
    {
      "ConditionType": "ModerationLabelConfidenceCheck",
      "ConditionParameters": {
        "ModerationLabelName": "*",
        "ConfidenceGreaterThanEquals": 75
      }
    }
  ]
}

Example 3: Use Sampling

In the following example, 5% of Amazon Rekognition inferences from a DetectModerationLabels request are sent to human workers. When using the default worker
task template provided in the SageMaker console, all moderation labels returned by Amazon Rekognition are sent to workers for review.

```json
{
    "Conditions": [
        {
            "ConditionType": "Sampling",
            "ConditionParameters": {
                "RandomSamplingPercentage": 5
            }
        }
    ]
}
```

**Example 4: Use Sampling and ModerationLabelConfidenceCheck with the And operator**

In this example, 5% of Amazon Rekognition inferences of the Graphic Male Nudity moderation label with a confidence greater than 50 are sent workers for review. When using the default worker task template provided in the SageMaker console, only the inferences of the Graphic Male Nudity label are sent to workers for review.

```json
{
    "Conditions": [
        {
            "And": [
                {
                    "ConditionType": "Sampling",
                    "ConditionParameters": {
                        "RandomSamplingPercentage": 5
                    }
                },
                {
                    "ConditionType": "ModerationLabelConfidenceCheck",
                    "ConditionParameters": {
                        "ModerationLabelName": "Graphic Male Nudity",
                        "ConfidenceGreaterThan": 50
                    }
                }
            ]
        }
    ]
}
```
Example 5: Use Sampling and ModerationLabelConfidenceCheck with the And operator

Use this example to configure your human review workflow to always send low-confidence inferences of a specified label for human review and sample high-confidence inferences of a label at a specified rate.

In the following example, a human review is initiated in one of the following ways:

- Inferences for the Graphic Male Nudity moderation label the with confidence scores less than 60 are always sent for human review. Only the Graphic Male Nudity label is sent to workers to review.
- 5% of all inferences for the Graphic Male Nudity moderation label the with confidence scores greater than 90 are sent for human review. Only the Graphic Male Nudity label is sent to workers to review.

```json
{
  "Conditions": [
    {
      "Or": [
        {
          "ConditionType": "ModerationLabelConfidenceCheck",
          "ConditionParameters": {
            "ModerationLabelName": "Graphic Male Nudity",
            "ConfidenceLessThan": 60
          }
        },
        {
          "And": [
            {
              "ConditionType": "Sampling",
              "ConditionParameters": {
                "RandomSamplingPercentage": 5
              }
            },
            {
              "ConditionType": "ModerationLabelConfidenceCheck",
              "ConditionParameters": {
                "ModerationLabelName": "Graphic Male Nudity",
                "ConfidenceGreaterThan": 90
              }
            }
          ]
        }
      ]
    }
  ]
}
```
Example 6: Use Sampling and ModerationLabelConfidenceCheck with the Or operator

In the following example, a human loop is created if the Amazon Rekognition inference response contains the 'Graphic Male Nudity' label with inference confidence greater than 50. Additionally, 5% of all other inferences initiate a human loop.

```
{
  "Conditions": [
    {
      "Or": [
        {
          "ConditionType": "Sampling",
          "ConditionParameters": {
            "RandomSamplingPercentage": 5
          }
        },
        {
          "ConditionType": "ModerationLabelConfidenceCheck",
          "ConditionParameters": {
            "ModerationLabelName": "Graphic Male Nudity",
            "ConfidenceGreaterThan": 50
          }
        }
      ]
    }
  ]
}
```

Delete a Human Review Workflow

When you delete a human review workflow or you delete your AWS account while a human loop is in process, your human review workflow status changes to Deleting. Amazon A2I automatically stops and deletes all associated human loops if workers have not started tasks created by those human loops. If human workers are already working on a task, that task continues to be available.
until it is completed or expires. As long as workers are still working on a task, your human review workflow's status is **Deleting**. If these tasks are completed, the results are stored in the Amazon S3 bucket specified in your flow definition.

Deleting a flow definition does not remove any worker answers from your S3 bucket. If the tasks are completed, but you deleted your AWS account, the results are stored in the Augmented AI service bucket for 30 days and then permanently deleted.

After all human loops have been deleted, the human review workflow is permanently deleted. When a human review workflow has been deleted, you can reuse its name to create a new human review workflow.

You might want to delete a human review workflow for any of the following reasons:

- You have sent data to a set of human reviewers and you want to delete all non-started human loops because you do not want those workers to work on those tasks any longer.
- The worker task template used to generate your worker UI does not render correctly or is not functioning as expected.

After you delete a human review workflow, the following changes occur:

- The human review workflow no longer appears on the **Human review workflows** page in the Augmented AI area of the Amazon SageMaker console.
- When you use the human review workflow name as input to the API operations **DescribeFlowDefinition** or **DeleteFlowDefinition**, Augmented AI returns a **ResourceNotFound** error.
- When you use **ListFlowDefinitions**, deleted human review workflows aren't included in the results.
- When you use the human review workflow ARN as input to the Augmented AI Runtime API operation **ListHumanLoops**, Augmented AI returns a **ResourceNotFoundException**.

### Delete a Flow Definition Using the Console or the SageMaker API

You can delete a human review workflow on the **Human review workflows** page in the Augmented AI area of the SageMaker console or by using the SageMaker API.

Flow definitions can only be deleted if their status is **Active**.
Delete a human review workflow (console)

1. Navigate to the Augmented AI console at https://console.aws.amazon.com/a2i/.
2. In the navigation pane, under the Augmented AI section, choose Human review workflows.
3. Select the hyperlinked name of the human review workflow that you want to delete.
4. On the Summary page of your human review workflow, choose Delete.
5. In the dialog box asking you to confirm that you want to delete your human review workflow, choose Delete.

You're automatically redirected to the Human review workflows page. While your human review workflow is being deleted, the status Deleting appears in the status column for that workflow. After it's deleted, it doesn't appear in the list of workflows on this page.

Delete a human review workflow (API)

You can delete a human review workflow (flow definition) using the SageMaker DeleteFlowDefinition API operation. This API operation is supported through the AWS CLI and a variety of language specific SDKs. The following table shows example requests using SDK for Python (Boto3) and the AWS CLI to delete the human review workflow, example-flow-definition.

AWS SDK for Python (Boto3)

The following request example uses the SDK for Python (Boto3) to delete the human review workflow. For more information, see delete_flow_definition in the AWS SDK for Python (Boto) API Reference.

```python
import boto3
sagemaker_client = boto3.client('sagemaker')
response = sagemaker_client.delete_flow_definition(FlowDefinitionName='example-flow-definition')
```

AWS CLI

The following request example uses the AWS CLI to delete the human review workflow. For more information, see delete-flow-definition in the AWS CLI Command Reference.
If the action is successful, Augmented AI sends back an HTTP 200 response with an empty HTTP body.

**Create and Start a Human Loop**

A *human loop* starts your human review workflow and sends data review tasks to human workers. When you use one of the Amazon A2I built-in task types, the corresponding AWS service creates and starts a human loop on your behalf when the conditions specified in your flow definition are met. If no conditions are specified in your flow definition, a human loop is created for each object. When using Amazon A2I for a custom task, a human loop starts when your application calls `StartHumanLoop`.

Use the following instructions to configure a human loop with Amazon Rekognition or Amazon Textract built-in task types and custom task types.

**Prerequisites**

To create and start a human loop, you must attach the `AmazonAugmentedAIFullAccess` policy to the AWS Identity and Access Management (IAM) user or role that configures or starts the human loop. This is the identity that you use to configure the human loop using `HumanLoopConfig` for built-in task types. For custom task types, this is the identity that you use to call `StartHumanLoop`.

Additionally, when using a built-in task type, your user or role must have permission to invoke API operations of the AWS service associated with your task type. For example, if you are using Amazon Rekognition with Augmented AI, you must attach permissions required to call `DetectModerationLabels`. For examples of identity-based policies you can use to grant these permissions, see [Amazon Rekognition Identity-Based Policy Examples](http://example.com) and [Amazon Textract Identity-Based Policy Examples](http://example.com). You can also use the more general policy `AmazonAugmentedAIIntegratedAPIAccess` to grant these permissions. For more information, see [Create a User With Permissions to Invoke Amazon A2I, Amazon Textract, and Amazon Rekognition API Operations](http://example.com).

To create and start a human loop, you need a flow definition ARN. To learn how to create a flow definition (or human review workflow), see [Create a Human Review Workflow](http://example.com).
Important

Amazon A2I requires all S3 buckets that contain human loop input image data to have a CORS policy attached. To learn more about this change, see CORS Permission Requirement.

Create and Start a Human Loop for a Built-in Task Type

To start a human loop using a built-in task type, use the corresponding service's API to provide your input data and to configure the human loop. For Amazon Textract, you use the AnalyzeDocument API operation. For Amazon Rekognition, you use the DetectModerationLabels API operation. You can use the AWS CLI or a language-specific SDK to create requests using these API operations.

Important

When you create a human loop using a built-in task type, you can use DataAttributes to specify a set of ContentClassifiers related to the input provided to the StartHumanLoop operation. Use content classifiers to declare that your content is free of personally identifiable information or adult content. To use Amazon Mechanical Turk, ensure your data is free of personally identifiable information, including protected health information under HIPAA. Include the FreeOfPersonallyIdentifiableInformation content classifier. If you do not use this content classifier, SageMaker does not send your task to Mechanical Turk. If your data is free of adult content, also include the 'FreeOfAdultContent' classifier. If you do not use these content classifiers, SageMaker may restrict the Mechanical Turk workers that can view your task.

After you start your ML job using your built-in task type's AWS service API, Amazon A2I monitors the inference results of that service. For example, when running a job with Amazon Rekognition, Amazon A2I checks the inference confidence score for each image and compares it to the confidence thresholds specified in your flow definition. If the conditions to start a human review task are satisfied, or if you didn't specify conditions in your flow definition, a human review task is sent to workers.
Create an Amazon Textract Human Loop

Amazon A2I integrates with Amazon Textract so that you can configure and start a human loop using the Amazon Textract API. To send a document file to Amazon Textract for document analysis, you use the Amazon Textract AnalyzeDocument API operation. To add a human loop to this document analysis job, you must configure the parameter HumanLoopConfig.

When you configure your human loop, the flow definition you specify in FlowDefinitionArn of HumanLoopConfig must be located in the same AWS Region as the bucket identified in Bucket of the Document parameter.

The following table shows examples of how to use this operation with the AWS CLI and AWS SDK for Python (Boto3).

AWS SDK for Python (Boto3)

The following request example uses the SDK for Python (Boto3). For more information, see analyze_document in the AWS SDK for Python (Boto) API Reference.

```python
import boto3
textract = boto3.client('textract', aws_region)
response = textract.analyze_document(
    Document={'S3Object': {'Bucket': bucket_name, 'Name': document_name}},
    FeatureTypes=['TABLES', 'FORMS'],
    HumanLoopConfig={
        'FlowDefinitionArn':
            'arn:aws:sagemaker:aws_region:aws_account_number:flow-definition/flow_def_name',
        'HumanLoopName': 'human_loop_name',
        'DataAttributes': {'ContentClassifiers':
            ['FreeOfPersonallyIdentifiableInformation','FreeOfAdultContent']}
    }
)
```

AWS CLI

The following request example uses the AWS CLI. For more information, see analyze-document in the AWS CLI Command Reference.

```
$ aws textract analyze-document \
```

Create and Start a Human Loop
After you run `AnalyzeDocument` with a human loop configured, Amazon A2I monitors the results from `AnalyzeDocument` and checks it against the flow definition's activation conditions. If the Amazon Textract inference confidence score for one or more key-value pairs meets the conditions for review, Amazon A2I starts a human review loop and includes the `HumanLoopActivationOutput` object in the `AnalyzeDocument` response.

**Create an Amazon Rekognition Human Loop**

Amazon A2I integrates with Amazon Rekognition so that you can configure and start a human loop using the Amazon Rekognition API. To send images to Amazon Rekognition for content moderation, you use the Amazon Rekognition `DetectModerationLabels` API operation. To configure a human loop, set the `HumanLoopConfig` parameter when you configure `DetectModerationLabels`.

When you configure your human loop, the flow definition you specify in `FlowDefinitionArn` of `HumanLoopConfig` must be located in the same AWS Region as the S3 bucket identified in `Bucket` of the `Image` parameter.

The following table shows examples of how to use this operation with the AWS CLI and AWS SDK for Python (Boto3).
AWS SDK for Python (Boto3)

The following request example uses the SDK for Python (Boto3). For more information, see `detect_moderation_labels` in the *AWS SDK for Python (Boto) API Reference*.

```python
import boto3

rekognition = boto3.client("rekognition", aws_region)

response = rekognition.detect_moderation_labels( 
    Image={'S3Object': {'Bucket': bucket_name, 'Name': image_name}}, 
    HumanLoopConfig={ 
        'HumanLoopName': 'human_loop_name', 
        'FlowDefinitionArn': 
            "arn:aws:sagemaker:aws_region:aws_account_number:flow-definition/flow_def_name" 
        } 
    'DataAttributes': [{'ContentClassifiers': ['FreeOfPersonallyIdentifiableInformation','FreeOfAdultContent']}
)
```

AWS CLI

The following request example uses the AWS CLI. For more information, see `detect-moderation-labels` in the *AWS CLI Command Reference*.

```bash
$ aws rekognition detect-moderation-labels \
    --image "S3Object={Bucket='bucket_name',Name='image_name'}" \
    --human-loop-config \
    HumanLoopName="human_loop_name",FlowDefinitionArn="arn:aws:sagemaker:aws_region:aws_account_number:flow-definition/flow_def_name",DataAttributes='{ContentClassifiers=["FreeOfPersonallyIdentifiableInformation","FreeOfAdultContent"]}'
```

```bash
$ aws rekognition detect-moderation-labels \
    --image "S3Object={Bucket='bucket_name',Name='image_name'}" \
    --human-loop-config \
    "{"HumanLoopName": "human_loop_name", "FlowDefinitionArn": "arn:aws:sagemaker:aws_region:aws_account_number:flow-definition/flow_def_name", "DataAttributes": ["ContentClassifiers": ["FreeOfPersonallyIdentifiableInformation", "FreeOfAdultContent"]}]"
```
After you run DetectModerationLabels with a human loop configured, Amazon A2I monitors the results from DetectModerationLabels and checks it against the flow definition's activation conditions. If the Amazon Rekognition inference confidence score for an image meets the conditions for review, Amazon A2I starts a human review loop and includes the response element HumanLoopActivationOutput in the DetectModerationLabels response.

**Create and Start a Human Loop for a Custom Task Type**

To configure a human loop for a custom human review task, use the StartHumanLoop operation within your application. This section provides an example of a human loop request using the AWS SDK for Python (Boto3) and the AWS Command Line Interface (AWS CLI). For documentation on other language-specific SDKs that support StartHumanLoop, use the See Also section of StartHumanLoop in the Amazon Augmented AI Runtime API documentation. Refer to **Use Cases and Examples Using Amazon A2I** to see examples that demonstrate how to use Amazon A2I with a custom task type.

**Prerequisites**

To complete this procedure, you need:

- Input data formatted as a string representation of a JSON-formatted file
- The Amazon Resource Name (ARN) of your flow definition

**To configure the human loop**

1. For DataAttributes, specify a set of ContentClassifiers related to the input provided to the StartHumanLoop operation. Use content classifiers to declare that your content is free of personally identifiable information or adult content.

   To use Amazon Mechanical Turk, ensure your data is free of personally identifiable information, including protected health information under HIPAA, and include the FreeOfPersonallyIdentifiableInformation content classifier. If you do not use this content classifier, SageMaker does not send your task to Mechanical Turk. If your data is free of adult content, also include the 'FreeOfAdultContent' classifier. If you do not use these content classifiers, SageMaker may restrict the Mechanical Turk workers that can view your task.

2. For FlowDefinitionArn, enter the Amazon Resource Name (ARN) of your flow definition.
3. For HumanLoopInput, enter your input data as a string representation of a JSON-formatted file. Structure your input data and custom worker task template so that your input data is properly displayed to human workers when you start your human loop. See [Preview a Worker Task Template](#) to learn how to preview your custom worker task template.

4. For HumanLoopName, enter a name for the human loop. The name must be unique within the Region in your account and can have up to 63 characters. Valid characters are a-z, 0-9, and -(hyphen).

**To start a human loop**

- To start a human loop, submit a request similar to the following examples using your preferred language-specific SDK.

**AWS SDK for Python (Boto3)**

The following request example uses the SDK for Python (Boto3). For more information, see [Boto 3 Augmented AI Runtime](#) in the *AWS SDK for Python (Boto) API Reference*.

```python
import boto3

a2i_runtime_client = boto3.client('sagemaker-a2i-runtime')

response = a2i_runtime_client.start_human_loop(
    HumanLoopName='human_loop_name',
    HumanLoopInput={'InputContent': '{"prompt": "What is the answer?"}'},
    DataAttributes={'ContentClassifiers': ['FreeOfPersonallyIdentifiableInformation','FreeOfAdultContent']}
)
```

**AWS CLI**

The following request example uses the AWS CLI. For more information, see [start-human-loop](#) in the *AWS CLI Command Reference*. 

Create and Start a Human Loop
When you successfully start a human loop by invoking StartHumanLoop directly, the response includes a HumanLoopARN and a HumanLoopActivationResults object which is set to NULL. You can use this the human loop name to monitor and manage your human loop.

**Next Steps:**

After starting a human loop, you can manage and monitor it with the Amazon Augmented AI Runtime API and Amazon CloudWatch Events. To learn more, see [Monitor and Manage Your Human Loop](#).

**Delete a Human Loop**

When you delete a human loop, the status changes to Deleting. When the human loop is deleted, the associated human review task is no longer available to workers. You might want to delete a human loop in one of the following circumstances:

- The worker task template used to generate your worker user interface does not render correctly or is not functioning as expected.
- A single data object was accidentally sent to workers multiple times.
- You no longer need a data object reviewed by a human.

If the status of a human loop is InProgress, you must stop the human loop before deleting it. When you stop a human loop, the status changes to Stopping while it is being stopped. When the status changes to Stopped, you can delete the human loop.

If human workers are already working on a task when you stop the associated human loop, that task continues to be available until it is completed or expires. As long as workers are still working on a task, your human loop's status is Stopping. If these tasks are completed, the results are stored in the Amazon S3 bucket URI specified in your human review workflow. If the worker leaves
the task without submitting work, it is stopped and the worker can't return to the task. If no worker has started working on the task, it is stopped immediately.

If you delete the AWS account used to create the human loop, it is stopped and deleted automatically.

**Human Loop Data Retention and Deletion**

When a human worker completes a human review task, the results are stored in the Amazon S3 output bucket you specified in the human review workflow used to create the human loop. Deleting or stopping a human loop does not remove any worker answers from your S3 bucket.

Additionally, Amazon A2I temporarily stores human loop input and output data internally for the following reasons:

- If you configure your human loops so that a single data object is sent to multiple workers for review, Amazon A2I does not write output data to your S3 bucket until all workers have completed the review task. Amazon A2I stores partial answers—answers from individual workers—internally so that it can write full results to your S3 bucket.
- If you report a low-quality human review result, Amazon A2I can investigate and respond to your issue.
- If you lose access to or delete the output S3 bucket specified in the human review workflow used to create a human loop, and the task has already been sent to one or more workers, Amazon A2I needs a place to temporarily store human review results.

Amazon A2I deletes this data internally 30 days after a human loop’s status changes to one of the following: Deleted, Stopped, or Completed. In other words, data is deleted 30 days after the human loop has been completed, stopped, or deleted. Additionally, this data is deleted after 30 days if you close the AWS account used to create associated human loops.

**Stop and Delete a Flow Definition Using the Console or the Amazon A2I API**

You can stop and delete a human loop in the Augmented AI console or by using the SageMaker API. When the human loop has been deleted, the status changes to Deleted.

**Delete a human loop (console)**

1. Navigate to the Augmented AI console at [https://console.aws.amazon.com/a2i/](https://console.aws.amazon.com/a2i/).
2. In the navigation pane, under the Augmented AI section, choose Human review workflows.
3. Choose the hyperlinked name of the human review workflow you used to create the human loop you want to delete.

4. In the **Human loops** section at the bottom of the page, select the human loop you want to stop and delete.

5. If the human loop status is Completed, Stopped, or Failed, select **Delete**.
   
   If the human loop **Status** is InProgress, select **Stop**. When the status changes to **Stopped**, select **Delete**.

---

**Delete a human loop (API)**

1. Check the status of your human loop using the Augmented AI Runtime API operation **DescribeHumanLoop**. See examples using this operation in the following table.

   **AWS SDK for Python (Boto3)**
   
   The following example uses the SDK for Python (Boto3) to describe the human loop named **example-human-loop**. For more information, see **describe_human_loop** in the **AWS SDK for Python (Boto) API Reference**.

   ```python
   import boto3
   a2i_runtime_client = boto3.client('sagemaker-a2i-runtime')
   response = a2i_runtime_client.describe_human_loop(HumanLoopName='example-human-loop')
   human_loop_status = response['HumanLoopStatus']
   print(f'example-human-loop status is: {human_loop_status}')
   ```

   **AWS CLI**
   
   The following example uses the AWS CLI to describe the human loop named **example-human-loop**. For more information, see **describe-human-loop** in the **AWS CLI Command Reference**.

   ```shell
   $ aws sagemaker-a2i-runtime describe-human-loop --human-loop-name 'example-human-loop'
   ```

2. If the flow definition status is Completed, Stopped, or Failed, delete the flow definition using the Augmented AI Runtime API operation **DeleteHumanLoop**.
The following example uses the SDK for Python (Boto3) to delete the human loop named `example-human-loop`. For more information, see `delete_human_loop` in the AWS SDK for Python (Boto) API Reference.

```python
import boto3

a2i_runtime_client = boto3.client('sagemaker-a2i-runtime')
response = a2i_runtime_client.delete_human_loop(HumanLoopName='example-human-loop')
```

If the human loop status is InProgress, stop the human loop using `StopHumanLoop` and then use `DeleteHumanLoop` to delete it.

The following example uses the AWS CLI to delete the human loop named `example-human-loop`. For more information, see `delete-human-loop` in the AWS CLI Command Reference.

```
$ aws sagemaker-a2i-runtime delete-human-loop --human-loop-name 'example-human-loop'
```

The following example uses the SDK for Python (Boto3) to describe the human loop named `example-human-loop`. For more information, see `stop_human_loop` in the AWS SDK for Python (Boto) API Reference.

```python
import boto3

a2i_runtime_client = boto3.client('sagemaker-a2i-runtime')
response = a2i_runtime_client.stop_human_loop(HumanLoopName='example-human-loop')
```
AWS CLI

The following example uses the AWS CLI to describe the human loop named *example-human-loop*. For more information, see [stop-human-loop](https://aws.amazon.com/documentation/sagemaker/) in the [AWS CLI Command Reference](https://docs.aws.amazon.com/cli/latest/reference/sagemaker-a2i-runtime/stop-human-loop).

```
$ aws sagemaker-a2i-runtime stop-human-loop --human-loop-name 'example-human-loop'
```

Create and Manage Worker Task Templates

You can create a task user interface for your workers by creating a *worker task template*. A worker task template is an HTML file that is used to display your input data and instructions to help workers complete your task.

For Amazon Rekognition or Amazon Textract task types, you can customize a pre-made worker task template using a graphical user interface (GUI) and avoid interacting with HTML code. For this option, use the instructions in [Create a Human Review Workflow (Console)](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-console-create-human-loop.html) to create a human review workflow and customize your worker task template in the Amazon SageMaker console. Once you create a template using these instructions, it appears on the worker task templates page of the [Augmented AI console](https://console.aws.amazon.com/sagemaker/a2i-console).

If you are creating a human review workflow for a custom task type, you must create a *custom worker task template* using HTML code. For more information, see [Create Custom Worker Task Templates](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-console-create-custom-workflow.html).

If you create your template using HTML, you must use this template to generate an Amazon A2I *human task UI Amazon Resource Name (ARN)* in the Amazon A2I console. This ARN has the following format: `arn:aws:sagemaker:<aws-region>:<aws-account-number>:human-task-ui/<template-name>`. This ARN is associated with a worker task template resource that you can use in one or more human review workflows (flow definitions).

Generate a human task UI ARN using a worker task template by following the instructions found in [Create a Worker Task Template](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-console-create-worker-task-template.html) or by using the [CreateHumanTaskUi](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateHumanTaskUi.html) API operation.

Topics

- [Create and Delete Worker Task Templates](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-console-create-delete-workflow.html)
- [Create Custom Worker Task Templates](https://docs.aws.amazon.com/sagemaker/latest/dg/a2i-console-create-custom-workflow.html)
Creating Good Worker Instructions

Create and Delete Worker Task Templates

You can use a worker template to customize the interface and instructions that your workers see when working on your tasks. Use the instructions on this page to create a worker task template in the Augmented AI area of the Amazon SageMaker console. A starter template is provided for Amazon Textract and Amazon Rekognition tasks. To learn how to customize your template using HTML crowd elements, see Create Custom Worker Task Templates.

When you create a worker template in the worker task templates page of the Augmented AI area of the SageMaker console, a worker task template ARN is generated. Use this ARN as the input to HumanTaskUiArn when you create a flow definition using the API operation CreateFlowDefinition. You can choose this template when creating a human review workflow on the human review workflows page of the console.

If you are creating a worker task template resource for an Amazon Textract or Amazon Rekognition task type, you can preview the worker UI that is generated from your template on the worker task templates console page. You must attach the policy described in Enable Worker Task Template Previews to the IAM role that you use to preview the template.

Create a Worker Task Template

You can create a worker task template using the SageMaker console and using the SageMaker API operation CreateHumanTaskUi.

Create a worker task template (console)

1. Open the Amazon A2I console at https://console.aws.amazon.com/a2i/.
2. Under Amazon Augmented AI in the left navigation pane, choose Worker task templates.
3. Choose Create template.
4. In Template name, enter a unique name.
5. (Optional) Enter an IAM role that grants Amazon A2I the permissions necessary to call services on your behalf.
6. In Template type, choose a template type from the dropdown list. If you are creating a template for a Textract-form extraction or Rekognition-image moderation task, choose the appropriate option.
7. Enter your custom template elements as follows:

- If you selected the Amazon Textract or Amazon Rekognition task template, the Template editor autopopulates with a default template that you can customize.
- If you are using a custom template, enter your predefined template in the editor.

8. (Optional) To complete this step, you must provide an IAM role ARN with permission to read Amazon S3 objects that get rendered on your user interface in Step 5.

You can only preview your template if you are creating templates for Amazon Textract or Amazon Rekognition.

Choose See preview to preview the interface and instructions that workers see. This is an interactive preview. After you complete the sample task and choose Submit, you see the resulting output from the task that you just performed.

If you are creating a worker task template for a custom task type, you can preview your worker task UI using RenderUiTemplate. For more information, see Preview a Worker Task Template.

9. When you're satisfied with your template, choose Create.

After you've created your template, you can select that template when you create a human review workflow in the console. Your template also appears in the Amazon Augmented AI section of the SageMaker console under Worker task templates. Choose your template to view its ARN. Use this ARN when using the CreateFlowDefinition API operation.

Create a worker task template using a worker task template (API)

To generate a worker task template using the SageMaker API operation CreateHumanTaskUi, specify a name for your UI in HumanTaskUiName and input your HTML template in Content under UiTemplate. Find documentation on language-specific SDKs that support this API operation in the See Also section of the CreateHumanTaskUi.

Delete a Worker Task Template

Once you have created a worker task template, you can delete it using the SageMaker console or the SageMaker API operation DeleteHumanTaskUi.

When you delete a worker task template, you are not able to use human review workflows (flow definitions) created using that template to start human loops. Any human loops that have already
been created using the worker task template that you delete continue to be processed until completion and are not impacted.

**Delete a worker task template (console)**

1. Open the Amazon A2I console at [https://console.aws.amazon.com/a2i/](https://console.aws.amazon.com/a2i/).
2. Under Amazon Augmented AI in the left navigation pane, choose **Worker task templates**.
3. Select the template that you want to delete.
4. Select **Delete**.
5. A modal appears to confirm your choice. Select **Delete**.

**Delete a worker task template (API)**

To delete a worker task template using the SageMaker API operation **DeleteHumanTaskUi**, specify a name of your UI in **HumanTaskUiName**.

**Create Custom Worker Task Templates**

*Crowd HTML Elements* are web components that provide a number of task widgets and design elements that you can tailor to the question you want to ask. You can use these crowd elements to create a custom worker template and integrate it with an Amazon Augmented AI (Amazon A2I) human review workflow to customize the worker console and instructions.

For a list of all HTML crowd elements available to Amazon A2I users, see [Crowd HTML Elements Reference](https://console.aws.amazon.com/a2i/). For examples of templates, see the [AWS GitHub repository](https://github.com/aws-samples/a2i-template-examples), which contains over 60 sample custom task templates.

**Develop Templates Locally**

When in the console to test how your template processes incoming data, you can test the look and feel of your template's HTML and custom elements in your browser by adding the following code to the top of your HTML file.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
```

This loads the necessary code to render the custom HTML elements. Use this code if you want to develop your template's look and feel in your preferred editor instead of in the console.
This code won't parse your variables. You might want to replace them with sample content while developing locally.

**Use External Assets**

Amazon Augmented AI custom templates enable you to embed external scripts and style sheets. For example, the following header embeds a text/css style sheet name stylesheet located at https://www.example.com/my-enhancement-styles.css into the custom template.

**Example**

```html
<script src="https://www.example.com/my-enhancement-script.js"></script>
<link rel="stylesheet" type="text/css" href="https://www.example.com/my-enhancement-styles.css">
```

If you encounter errors, ensure that your originating server is sending the correct MIME type and encoding headers with the assets.

For example, the MIME and encoding type for remote scripts is application/javascript;CHARSET=UTF-8.

The MIME and encoding type for remote stylesheets is text/css;CHARSET=UTF-8.

**Track Your Variables**

When building a custom template, you must add variables to it to represent the pieces of data that might change from task to task, or worker to worker. If you're starting with one of the sample templates, you need to make sure you're aware of the variables it already uses.

For example, for a custom template that integrates an Augmented AI human review loop with a Amazon Textract text review task, {{ task.input.selectedAiServiceResponse.blocks }} is used for initial-value input data. For Amazon Augmented AI (Amazon A2I) integration with Amazon Rekognition, {{ task.input.selectedAiServiceResponse.moderationLabels }} is used.

For a custom task type, you need to determine the input parameter for your task type. Use {{ task.input.customInputValuesForStartHumanLoop }} where you specify customInputValuesForStartHumanLoop.

**Custom Template Example for Amazon Textract**

All custom templates begin and end with the `<crowd-form> </crowd-form>` elements. Like standard HTML `<form>` elements, all of your form code should go between these elements.
For an Amazon Textract document analysis task, use the `<crowd-textract-analyze-document>` element. It uses the following attributes:

- `src` – Specifies the URL of the image file to be annotated.
- `initialValue` – Sets initial values for attributes found in the worker UI.
- `blockTypes (required)` – Determines the kind of analysis that the workers can do. Only `KEY_VALUE_SET` is currently supported.
- `keys (required)` – Specifies new keys and the associated text value that the worker can add.
- `no-key-edit (required)` – Prevents the workers from editing the keys of annotations passed through `initialValue`.
- `no-geometry-edit` – Prevents workers from editing the polygons of annotations passed through `initialValue`.

For children of the `<crowd-textract-analyze-document>` element, you must have two Regions. You can use arbitrary HTML and CSS elements in these Regions.

- `<full-instructions>` – Instructions that are available from the View full instructions link in the tool. You can leave this blank, but we recommend that you provide complete instructions to get better results.
- `<short-instructions>` – A brief description of the task that appears in the tool's sidebar. You can leave this blank, but we recommend that you provide complete instructions to get better results.

An Amazon Textract template would look similar to the following.

**Example**

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
{% capture s3_uri %}http://s3.amazonaws.com/
{{ task.input.aiServiceRequest.document.s3Object.bucket }}/\n{{ task.input.aiServiceRequest.document.s3Object.name }}{% endcapture %}

<crowd-form>
  <crowd-textract-analyze-document
    src="{{ s3_uri | grant_read_access }}"
    initial-value="{{ task.input.selectedAiServiceResponse.blocks }}"
  />
</crowd-form>
```
Review the key-value pairs listed on the right and correct them if they don't match the following document.

Choose a key-value block to highlight the corresponding key-value pair in the document. If it is a valid key-value pair, review the content for the value. If the content is incorrect, correct it. The text of the value is incorrect, correct it.

A wrong value is identified, correct it.

If it is not a valid key-value relationship, choose No.

If you can’t find the key in the document, choose Key not found.

If the content of a field is empty, choose Value is blank.

Examples
Key and value are often displayed next to or below to each other.

Key and value displayed in one line.
Custom Template Example for Amazon Rekognition

All custom templates begin and end with the `<crowd-form> </crowd-form>` elements. Like standard HTML `<form>` elements, all of your form code should go between these elements. For an Amazon Rekognition custom task template, use the `<crowd-rekognition-detect-moderation-labels>` element. This element supports the following attributes:

- **categories** – An array of strings or an array of objects where each object has a name field.
  - If the categories come in as objects, the following applies:
    - The displayed categories are the value of the name field.
    - The returned answer contains the *full* objects of any selected categories.
  - If the categories come in as strings, the following applies:
    - The returned answer is an array of all the strings that were selected.
- **exclusion-category** – By setting this attribute, you create a button underneath the categories in the UI. When a user selects the button, all categories are deselected and disabled. If the worker selects the button again, you re-enable users to choose categories. If the worker submits the task by selecting **Submit** after you select the button, that task returns an empty array.

For children of the `<crowd-rekognition-detect-moderation-labels>` element, you must have two Regions.
• `<full-instructions>` – Instructions that are available from the View full instructions link in the tool. You can leave this blank, but we recommend that you provide complete instructions to get better results.

• `<short-instructions>` – Brief description of the task that appears in the tool's sidebar. You can leave this blank, but we recommend that you provide complete instructions to get better results.

A template using these elements would look similar to the following.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
{% capture s3_uri %}http://s3.amazonaws.com/
{{ task.input.aiServiceRequest.image.s3Object.bucket }}/%
{{ task.input.aiServiceRequest.image.s3Object.name }}{% endcapture %}

<crowd-form>
  <crowd-rekognition-detect-moderation-labels
    categories='[
      {% for label in task.input.selectedAiServiceResponse.moderationLabels %}
        {
          name: "{{ label.name }}",
          parentName: "{{ label.parentName }}",
        },
      {% endfor %}
    ]'
    src="{{ s3_uri | grant_read_access }}"
    header="Review the image and choose all applicable categories.">
    <short-instructions header="Instructions">
      <style>
        .instructions {
          white-space: pre-wrap;
        }
      </style>
      <p class='instructions'>Review the image and choose all applicable categories. If no categories apply, choose None.

      <b>Nudity</b>
      Visuals depicting nude male or female person or persons

      <b>Graphic Male Nudity</b>
      Visuals depicting full frontal male nudity, often close ups
  </short-instructions>
</crowd-form>
```
<b>Graphic Female Nudity</b>
Visuals depicting full frontal female nudity, often close ups

<b>Sexual Activity</b>
Visuals depicting various types of explicit sexual activities and pornography

<b>Illustrated Nudity or Sexual Activity</b>
Visuals depicting animated or drawn sexual activity, nudity, or pornography

<b>Adult Toys</b>
Visuals depicting adult toys, often in a marketing context

<b>Female Swimwear or Underwear</b>
Visuals depicting female person wearing only swimwear or underwear

<b>Male Swimwear Or Underwear</b>
Visuals depicting male person wearing only swimwear or underwear

<b>Partial Nudity</b>
Visuals depicting covered up nudity, for example using hands or pose

<b>Revealing Clothes</b>
Visuals depicting revealing clothes and poses, such as deep cut dresses

<b>Graphic Violence or Gore</b>
Visuals depicting prominent blood or bloody injuries

<b>Physical Violence</b>
Visuals depicting violent physical assault, such as kicking or punching

<b>Weapon Violence</b>
Visuals depicting violence using weapons like firearms or blades, such as shooting

<b>Weapons</b>
Visuals depicting weapons like firearms and blades

<b>Self Injury</b>
Visuals depicting self-inflicted cutting on the body, typically in distinctive patterns using sharp objects

<b>Emaciated Bodies</b>
Visuals depicting extremely malnourished human bodies

<b>Corpses</b>
Add Automation with Liquid

The custom template system uses Liquid for automation. Liquid is an open-source inline markup language. For more information and documentation, see the Liquid homepage.

In Liquid, the text between single curly braces and percent symbols is an instruction or tag that performs an operation like control flow or iteration. Text between double curly braces is a variable or object that outputs its value. The following list includes two types of liquid tags that you may find useful to automate template input data processing. If you select one of the following tag-types, you are redirected to the Liquid documentation.

- **Control flow**: Includes programming logic operators like if/else, unless, and case/when.
- **Iteration**: Enables you to run blocks of code repeatedly using statements like for loops.

For example, the following code example demonstrates how you can use the Liquid for tag to create a for loop. This example loops through the moderationLabels returned from Amazon Rekognition and displays the moderationLabels attributes name and parentName for workers to review:

```liquid
{% for label in task.input.selectedAiServiceResponse.moderationLabels %}
    {
        name: &quot;{{ label.name }}&quot;,
        parentName: &quot;{{ label.parentName }}&quot;
    },
{% endfor %}
```
Use Variable Filters

In addition to the standard Liquid filters and actions, Amazon Augmented AI (Amazon A2I) offers additional filters. You apply filters by placing a pipe (|) character after the variable name, and then specifying a filter name. To chain filters, use the following format.

Example

```liquid
{{ <content> | <filter> | <filter> }}
```

Autoescape and Explicit Escape

By default, inputs are HTML-escaped to prevent confusion between your variable text and HTML. You can explicitly add the escape filter to make it more obvious to someone reading the source of your template that escaping is being done.

escape_once

escape_once ensures that if you've already escaped your code, it doesn't get re-escaped again. For example, it ensures that `&` doesn't become `&amp;`.

skip_autoescape

skip_autoescape is useful when your content is meant to be used as HTML. For example, you might have a few paragraphs of text and some images in the full instructions for a bounding box.

Note

Use skip_autoescape sparingly. As a best practice for templates, avoid passing in functional code or markup with skip_autoescape unless you are absolutely sure that you have strict control over what's being passed. If you're passing user input, you could be opening your workers up to a cross-site scripting attack.

to_json

to_json encodes data that you provide to JavaScript Object Notation (JSON). If you provide an object, it serializes it.
grant_read_access

grant_read_access takes an Amazon Simple Storage Service (Amazon S3) URI and encodes it into an HTTPS URL with a short-lived access token for that resource. This makes it possible to display photo, audio, or video objects stored in S3 buckets that are not otherwise publicly accessible to workers.

Example Example of the to_json and grant_read_access filters

Input

```liquid
auto-escape: {{ "Have you read 'James & the Giant Peach'?" }}
explicit escape: {{ "Have you read 'James & the Giant Peach'?" | escape }}
explicit escape_once: {{ "Have you read 'James & the Giant Peach'?" | escape_once }}
skip_autoescape: {{ "Have you read 'James & the Giant Peach'?" | skip_autoescape }}
to_json: {{ jsObject | to_json }}
grant_read_access: {{ "s3://examplebucket/myphoto.png" | grant_read_access }}
```

Example

Output

```liquid
auto-escape: Have you read 'James & the Giant Peach'??
explicit escape: Have you read 'James & the Giant Peach'??
explicit escape_once: Have you read 'James & the Giant Peach'??
skip_autoescape: Have you read 'James & the Giant Peach'??
to_json: { "point_number": 8, "coords": [ 59, 76 ] }
grant_read_access: https://s3.amazonaws.com/examplebucket/myphoto.png?access token and other params>
```

Example Example of an automated classification template.

To automate this simple text classification sample, include the Liquid tag
`{{ task.input.source }}`. This example uses the `crowd-classifier` element.

```html
<script src="https://assets.crowd.aws/crowd-html-elements.js"></script>
<crowd-form>
    <crowd-classifier
        name="tweetFeeling"
        categories="['positive', 'negative', 'neutral', 'cannot determine']"
        header="Which term best describes this tweet?"
    >
</crowd-form>
```
Preview a Worker Task Template

To preview a custom worker task template, use the SageMaker RenderUiTemplate operation. You can use the RenderUiTemplate operation with the AWS CLI or your preferred AWS SDK. For documentation on the supported language specific SDKs for this API operation, see the See Also section of the RenderUiTemplate.

Prerequisites

To preview your worker task template, the AWS Identity and Access Management (IAM) role Amazon Resource Name (ARN), or RoleArn, that you use must have permission to access to the S3 objects that are used by the template. To learn how to configure your role or user see Enable Worker Task Template Previews.

To preview your worker task template using the RenderUiTemplate operation:

1. Provide a RoleArn of the role with required policies attached to preview your custom template.

2. In the Input parameter of Task, provide a JSON object that contains values for the variables defined in the template. These are the variables that are substituted for the task.input.source variable. For example, if you define a task.input.text variable in your template, you can supply the variable in the JSON object as text: sample text.

3. In the Content parameter of UiTemplate, insert your template.
Once you've configured `RenderUiTemplate`, use your preferred SDK or the AWS CLI to submit a request to render your template. If your request was successful, the response includes `RenderedContent`, a Liquid template that renders the HTML for the worker UI.

⚠️ **Important**

To preview your template, you need an IAM role with permissions to read Amazon S3 objects that get rendered on your user interface. For a sample policy that you can attach to your IAM role to grant these permissions, see [Enable Worker Task Template Previews](#).

---

### Creating Good Worker Instructions

Creating good instructions for your human review jobs improves your worker's accuracy in completing their task. You can modify the default instructions that are provided in the console when creating a human review workflow, or you can use the console to create a custom worker template and include your instructions in this template. The instructions are shown to the worker on the UI page where they complete their labeling task.

**Create Good Worker Instructions**

There are three kinds of instructions in the Amazon Augmented AI console:

- **Task Description** – The description should provide a succinct explanation of the task.
- **Instructions** – These instructions are shown on the same webpage where workers complete a task. These instructions should provide an easy reference to show the worker the correct way to complete the task.
- **Additional Instructions** – These instructions are shown in a dialog box that appears when a worker chooses [View full instructions](#). We recommend that you provide detailed instructions for completing the task, and include several examples showing edge cases and other difficult situations for labeling objects.

**Add Example Images to Your Instructions**

Images provide useful examples for your workers. To add a publicly accessible image to your instructions, do the following:

1. Place the cursor where the image should go in the instructions editor.
2. Choose the image icon in the editor toolbar.

3. Enter the URL of your image.

If your instruction image is in an S3 bucket that isn't publicly accessible, do the following:

- For the image URL, enter: {{ 'https://s3.amazonaws.com/your-bucket-name/image-file-name' | grant_read_access }}.

This renders the image URL with a short-lived, one-time access code that's appended so the worker's browser can display it. A broken image icon is displayed in the instructions editor, but previewing the tool displays the image in the rendered preview. See grant_read_access for more information about the grand_read_access element.

**Monitor and Manage Your Human Loop**

Once you've started a human review loop, you can check the results of tasks sent to the loop and manage it using the Amazon Augmented AI Runtime API. Additionally, Amazon A2I integrates with Amazon EventBridge (also known as Amazon CloudWatch Events) to alert you when a human review loop status changes to Completed, Failed, or Stopped. This event delivery is guaranteed at least once, which means all events created when human loops finish are successfully delivered to EventBridge.

Use the procedures below to learn how to use the Amazon A2I Runtime API to monitor and manage your human loops. See Use Amazon CloudWatch Events in Amazon Augmented AI to learn how Amazon A2I integrates with Amazon EventBridge.

**To check your output data:**

1. Check the results of your human loop by calling the DescribeHumanLoop operation. The result of this API operation contains information about the reason for and outcome of the loop activation.

2. Check the output data from your human loop in Amazon Simple Storage Service (Amazon S3). In the path to the data, YYYY/MM/DD/hh/mm/ss represents the human loop creation date with year (YYYY), month (MM), and day (DD), and the creation time with hour (hh), minute (mm), and second (ss).
You can integrate this structure with AWS Glue or Amazon Athena to partition and analyze your output data. For more information, see [Managing Partitions for ETL Output in AWS Glue](#).

To learn more about Amazon A2I output data format, see [Amazon A2I Output Data](#).

To stop and delete your human loop:

1. Once a human loop has been started, you can stop your human loop by calling the `StopHumanLoop` operation using the `HumanLoopName`. If a human loop was successfully stopped, the server sends back an HTTP 200 response.

2. To delete a human loop for which the status equals Failed, Completed, or Stopped, use the `DeleteHumanLoop` operation.

To list human loops:

1. You can list all active human loops by calling the `ListHumanLoops` operation. You can filter human loops by the creation date of the loop using the `CreationTimeAfter` and `CreateTimeBefore` parameters.

2. If successful, `ListHumanLoops` returns `HumanLoopSummaries` and `NextToken` objects in the response element. `HumanLoopSummaries` contains information about a single human loop. For example, it lists a loop's status and, if applicable, its failure reason.

   Use the string returned in `NextToken` as an input in a subsequent call to `ListHumanLoops` to see the next page of human loops.

### Amazon A2I Output Data

When your machine learning workflow sends Amazon A2I a data object, a human loop is created and human reviewers receive a task to review that data object. The output data from each human review task is stored in the Amazon Simple Storage Service (Amazon S3) output bucket you specify in your human review workflow. In the path to the data, `YYYY/MM/DD/hh/mm/ss` represents the human loop creation date with year (YYYY), month (MM), and day (DD), and the creation time with hour (hh), minute (mm), and second (ss).
The content of your output data depends on the type of task type (built-in or custom) and the type of workforce you use. Your output data always includes the response from the human worker. Additionally, output data may include metadata about the human loop, the human reviewer (worker), and the data object.

Use the following sections to learn more about Amazon A2I output data format for different task types and workforces.

### Output Data From Built-In Task Types

Amazon A2I built-in task types include Amazon Textract and Amazon Rekognition. In addition to human responses, the output data from one of these tasks includes details about the reason the human loop was created and information about the integrated service used to create the human loop. Use the following table to learn more about the output data schema for all built-in task types. The value for each of these parameters depends on the service you use with Amazon A2I. Refer to the second table in this section for more information about these service-specific values.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value Type</th>
<th>Example Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>awsManagedHumanLoopRequestSource</td>
<td>String</td>
<td>AWS/Rekognition/DetectModerationLabels/Image/V3 or AWS/Textract/AnalyzeDocument/Forms/V1</td>
<td>The API operation and associated AWS services that requested that Amazon A2I create the a human loop. This is the API operation you use to configure your Amazon A2I human loop.</td>
</tr>
<tr>
<td>flowDefinitionArn</td>
<td>String</td>
<td>arn:aws:sagemaker:us-west-2:1111222223</td>
<td>The Amazon Resource Number (ARN) of the human review workflow</td>
</tr>
<tr>
<td>Parameter</td>
<td>Value Type</td>
<td>Example Values</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------</td>
<td>--------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>333:flow-definition/flow-definition-name</td>
<td>String</td>
<td>(flow definition) used to create the human loop.</td>
<td></td>
</tr>
</tbody>
</table>
| humanAnswers      | List of JSON objects | `{
  "answerContent": {
    "AWS/Rekognition/DetectionLabels/Image/V3": {
      "moderationLabels": [...]
    }
  },
| or                |                  | {                                                            | A list of JSON objects that contain worker responses in answerContent.
|                   |                  |   "answerContent": {
|                   |                  |     "AWS/Textract/AnalyzeDocument/Forms/V1": {
|                   |                  |       "blocks": [...]}
|                   |                  | },
|                   |                  | }                                                                 | This object also contains submission details and, if a private workforce was used, worker metadata. To learn more, see Track Worker Activity.
<p>|                   |                  |                                                                     | For human loop output data produced from Amazon Rekognition DetectionLabel review tasks, this parameter only contains positive responses. For example, if workers select No content, this response is not included. |
| humanLoopName     | String           | 'human-loop-name'                                                     | The name of the human loop.                                                                                                                 |</p>
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value Type</th>
<th>Example Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>inputContent</td>
<td>JSON object</td>
<td>{</td>
<td>The input content the AWS service sent to Amazon A2I when it requested a human loop be created.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;aiServiceRequest&quot;: {...},</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;aiServiceResponse&quot;: {...},</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;humanTaskActivationResults&quot;: {...},</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;selectedAiServiceResponse&quot;: {...}</td>
<td></td>
</tr>
<tr>
<td>aiServiceRequest</td>
<td>JSON object</td>
<td>{</td>
<td>The original request sent to the AWS service integrated with Amazon A2I. For example, if you use Amazon Rekognition with Amazon A2I, this includes the request made through the API operation DetectModerationLabels . For Amazon Textract integrations, this includes the request made through AnalyzeDocument .</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;document&quot;: {...},</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;featureTypes&quot;: [...],</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;humanLoopConfig&quot;: {...}</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>or</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;image&quot;: {...},</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;humanLoopConfig&quot;: {...}</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Value Type</td>
<td>Example Values</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------</td>
<td>-------------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>aiServiceResponse</td>
<td>JSON object</td>
<td>{</td>
<td>The full response from the AWS service. This is the data that is used to determine if a human review is required. This object may contain metadata about the data object that is not shared with human reviewers.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;moderationLabels&quot;:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>[...],</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;moderationModelVersion&quot;: &quot;3.0&quot;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>or</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>{</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;blocks&quot;:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>[...],</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&quot;documentMetadata&quot;: {}</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Value Type</td>
<td>Example Values</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------------</td>
<td>------------------</td>
<td>-------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><code>selectedAiServiceResponse</code></td>
<td>JSON object</td>
<td><code>{   &quot;moderationLabels&quot;: [...],   &quot;moderationModelVersion&quot;: &quot;3.0&quot; }</code></td>
<td>The subset of the aiServiceResponse that matches the activation conditions in ActivationConditions. All data objects listed in aiServiceResponse are listed in selectedAiServiceResponse when inferences are randomly sampled, or all inferences initiated activation conditions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>or <code>{   &quot;blocks&quot;: [...],   &quot;documentMetadata&quot;: {} }</code></td>
<td></td>
</tr>
</tbody>
</table>

**Output Data**
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value Type</th>
<th>Example Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>humanTask ActivationConditionResults</td>
<td>JSON object</td>
<td><code>{  &quot;Conditions&quot;: [ ...] }</code></td>
<td>A JSON object in inputContent that contains the reason a human loop was created. This includes a list of the activation conditions (Conditions) included in your human review workflow (flow definition), and the evaluation result for each condition—this result is either true or false. To learn more about activation conditions, see <a href="https://aws.amazon.com">JSON Schema for Human Loop Activation Conditions in Amazon Augmented AI</a>.</td>
</tr>
</tbody>
</table>

Select a tab on the following table to learn about the task type–specific parameters and see an example output-data code block for each of the built-in task types.

## Amazon Textract Task Type Output Data

When you use the Amazon Textract built-in integration, you see 'AWS/Textract/AnalyzeDocument/Forms/V1' as the value for awsManagedHumanLoopRequestSource in your output data.

The `answerContent` parameter contains a Block object that includes human responses for all blocks sent to Amazon A2I.
The aiServiceResponse parameter also includes a Block object with Amazon Textract's response to the original request sent using to AnalyzeDocument.

To learn more about the parameters you see in the block object, refer to Block in the Amazon Textract Developer Guide.

The following is an example of the output data from an Amazon A2I human review of Amazon Textract document analysis inferences.

```json
{
    "awsManagedHumanLoopRequestSource": "AWS/Text\text{ract}/AnalyzeDocument/Forms/V1",
    "humanAnswers": [
        {
            "answerContent": {
                "AWS/Text\text{ract}/AnalyzeDocument/Forms/V1": {
                    "blocks": [...]
                }
            },
            "submissionTime": "2020-09-28T19:17:59.880Z",
            "workerId": "111122223333",
            "workerMetadata": {
                "identityData": {
                    "identityProviderType": "Cognito",
                    "sub": "c6aa8e7b-9944-42e9-a6b9-111122223333"
                }
            }
        }
    ],
    "humanLoopName": "human-loop-name",
    "inputContent": {
        "aiServiceRequest": {
            "document": {
                "s3Object": {
                    "bucket": "DOC-EXAMPLE-BUCKET1",
                    "name": "document-demo.jpg"
                }
            },
            "featureTypes": [
                "TABLES",
```
"FORMS",
"humanLoopConfig": {
  "dataAttributes": {
    "contentClassifiers": [
      "FreeOfPersonallyIdentifiableInformation"
    ],
  }
},
"humanTaskActivationConditionResults": {
  "Conditions": [
    {
      "EvaluationResult": true,
      "Or": [
        {
          "ConditionParameters": {
            "ImportantFormKey": "Mail address",
            "ImportantFormKeyAliases": [
              "Mail Address:",
              "Mail address:",
              "Mailing Add:",
              "Mailing Addresses"
            ],
            "KeyValueBlockConfidenceLessThan": 100,
            "WordBlockConfidenceLessThan": 100
          }
        },
        {
          "ConditionParameters": {
            "ImportantFormKey": "Mail address",
            "ImportantFormKeyAliases": [
              "Mail Address:",
              "Mail address:",
              "Mailing Add:",
              "Mailing Addresses"
            ],
            "KeyValueBlockConfidenceLessThan": 100,
            "WordBlockConfidenceLessThan": 100
          }
        }
      ]
    }
  ]
}
}
Amazon Rekognition Task Type Output Data

When you use the Amazon Textract built-in integration, you see the string 'AWS/Rekognition/DetectModerationLabels/Image/V3' as the value for awsManagedHumanLoopRequestSource in your output data.

The answerContent parameter contains a moderationLabels object that contains human responses for all moderation labels sent to Amazon A2I.

The aiServiceResponse parameter also includes a moderationLabels object with Amazon Rekognition's response to the original request sent to DetectModerationLabels.

To learn more about the parameters you see in the block object, refer to ModerationLabel in the Amazon Rekognition Developer Guide.

The following is an example of the output data from an Amazon A2I human review of Amazon Rekognition image moderation inferences.

```json
{
    "awsManagedHumanLoopRequestSource": "AWS/Rekognition/DetectModerationLabels/Image/V3",
    "humanAnswers": [
        {
```
"answerContent": {
  "AWS/Rekognition/DetectModerationLabels/Image/V3": {
    "moderationLabels": [...] 
  
  },
  "submissionTime": "2020-09-28T19:22:35.508Z",
  "workerId": "ef7294f850a3d9d1",
  "workerMetadata": {
    "identityData": {
      "identityProviderType": "Cognito",
      "sub": "c6aa8eb7-9944-42e9-a6b9-111122223333"
    }
  }
},
"humanLoopName": "humnan-loop-name",
"inputContent": {
  "aiServiceRequest": {
    "humanLoopConfig": {
      "humanLoopName": "humnan-loop-name"
    },
    "image": {
      "s3Object": {
        "bucket": "DOC-EXAMPLE-BUCKET1",
        "name": "example-image.jpg"
      }
    }
  }
},
"aiServiceResponse": {
  "moderationLabels": [...],
  "moderationModelVersion": "3.0"
},
"humanTaskActivationConditionResults": {
  "Conditions": [
    {
      "EvaluationResult": true,
      "Or": [
        {
          "ConditionParameters": {
            "ConfidenceLessThan": 98,
          }
        }
      
    }
  ]
}
Output Data From Custom Task Types

When you add Amazon A2I to a custom human review workflow, you see the following parameters in the output data returned from human review tasks.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>flowDefinitionArn</td>
<td>String</td>
<td>The Amazon Resource Number (ARN) of the human review workflow (flow</td>
</tr>
<tr>
<td>Parameter</td>
<td>Value Type</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>--------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>humanAnswers</td>
<td>List of JSON objects</td>
<td>A list of JSON objects that contain worker responses in answerContent. The value in this parameter is determined by the output received from your worker task template. If you are using a private workforce, worker metadata is included. To learn more, see Track Worker Activity.</td>
</tr>
<tr>
<td>humanLoopName</td>
<td>String</td>
<td>The name of the human loop.</td>
</tr>
<tr>
<td>inputContent</td>
<td>JSON Object</td>
<td>The input content sent to Amazon A2I in the request to StartHumanLoop.</td>
</tr>
</tbody>
</table>

The following is an example of output data from a custom integration with Amazon A2I and Amazon Transcribe. In this example, the inputContent consists of:

- A path to an .mp4 file in Amazon S3 and the video title
- The transcription returned from Amazon Transcribe (parsed from Amazon Transcribe output data)
- A start and end time used by the worker task template to clip the .mp4 file and show workers a relevant portion of the video

```json
{
    "humanAnswers": [
        {
```
Track Worker Activity

Amazon A2I provides information that you can use to track individual workers in task output data. To identify the worker that worked on the human review task, use the following from the output data in Amazon S3:

- The `acceptanceTime` is the time that the worker accepted the task. The format of this date and time stamp is `YYYY-MM-DDTHH:MM:SS.mmmZ` for the year (YYYY), month (MM), day (DD), hour (HH), minute (MM), second (SS), and millisecond (mmm). The date and time are separated by a T.

- The `submissionTime` is the time that the worker submitted their annotations using the Submit button. The format of this date and time stamp is `YYYY-MM-DDTHH:MM:SS.mmmZ` for the year (YYYY), month (MM), day (DD), hour (HH), minute (MM), second (SS), and millisecond (mmm). The date and time are separated by a T.
- timeSpentIn Seconds reports the total time, in seconds, that a worker actively worked on that task. This metric does not include time when a worker paused or took a break.

- The workerId is unique to each worker.

- If you use a private workforce, in workerMetadata, you see the following.
  - The identityProviderType is the service used to manage the private workforce.
  - The issuer is the Amazon Cognito user pool or OpenID Connect (OIDC) Identity Provider (IdP) issuer associated with the work team assigned to this human review task.
  - A unique sub identifier refers to the worker. If you create a workforce using Amazon Cognito, you can retrieve details about this worker (such as the name or user name) associated with this ID using Amazon Cognito. To learn how, see Managing and Searching for User Accounts in Amazon Cognito Developer Guide.

The following is an example of the output you may see if you use Amazon Cognito to create a private workforce. This is identified in the identityProviderType.

```json
"submissionTime": "2020-12-28T18:59:58.321Z",
"acceptanceTime": "2020-12-28T18:59:15.191Z",
"timeSpentInSeconds": 40.543,
"workerId": "a12b3cdefg4h5i67",
"workerMetadata": {
    "identityData": {
        "identityProviderType": "Cognito",
        "sub": "aaaaaaaa-bbbb-cccc-dddd-eeeeeeeeee"
    }
}
```

The following is an example of the output you may see if you use your own OIDC IdP to create a private workforce:

```json
"workerMetadata": {
    "identityData": {
        "identityProviderType": "Oidc",
        "issuer": "https://example-oidc-ipd.com/adfs",
        "sub": "aaaaaaaa-bbbb-cccc-dddd-eeeeeeeeee"
    }
}
```
To learn more about using private workforces, see Use a Private Workforce.

Permissions and Security in Amazon Augmented AI

When using Amazon Augmented AI (Amazon A2I) to create a human review workflow for your ML/AI application, you create and configure resources in Amazon SageMaker such as a human workforce and worker task templates. To configure and start a human loop, you either integrate Amazon A2I with other AWS services such as Amazon Textract or Amazon Rekognition, or use the Amazon Augmented AI Runtime API. To create a human review workflow and start a human loop, you must attach certain policies to your AWS Identity and Access Management (IAM) role or user. Specifically:

- When you start a human loop using image input data on or after January 12th, 2020, you must add a CORS header policy to the Amazon S3 bucket that contains your input data. See CORS Permission Requirement to learn more.

- When you create a flow definition, you need to provide a role that grants Amazon A2I permission to access Amazon S3 both for reading objects that are rendered in a human task UI and for writing the results of the human review.

  This role must also have a trust policy attached to give SageMaker permission to assume the role. This allows Amazon A2I to perform actions in accordance with permissions that you attach to the role.

  See Add Permissions to the IAM Role Used to Create a Flow Definition for example policies that you can modify and attach to the role you use to create a flow definition. These are the policies that are attached to the IAM role that is created in the Human review workflows section of the Amazon A2I area of the SageMaker console.

- To create and start human loops, you either use an API operation from a built-in task type (such as DetectModerationLabel or AnalyzeDocument) or the Amazon A2I Runtime API operation StartHumanLoop in a custom ML application. You need to attach the AmazonAugmentedAIFullAccess managed policy to the user that invokes these API operations to grant permission to these services to use Amazon A2I operations. To learn how, see Create a User That Can Invoke Amazon A2I API Operations.

  This policy does not grant permission to invoke the API operations of the AWS service associated with built-in task types. For example, AmazonAugmentedAIFullAccess does not grant permission to call the Amazon Rekognition DetectModerationLabel API operation or Amazon Textract AnalyzeDocument API operation. You can use the more general policy,
AmazonAugmentedAIIntegratedAPIAccess, to grant these permissions. For more information, see Create a User With Permissions to Invoke Amazon A2I, Amazon Textract, and Amazon Rekognition API Operations. This is a good option when you want to grant a user broad permissions to use Amazon A2I and integrated AWS services' API operations.

If you want to configure more granular permissions, see Amazon Rekognition Identity-Based Policy Examples and Amazon Textract Identity-Based Policy Examples for identity-based policies you can use to grant permission to use these individual services.

- To preview your custom worker task UI template, you need an IAM role with permissions to read Amazon S3 objects that get rendered on your user interface. See a policy example in Enable Worker Task Template Previews.

Topics

- CORS Permission Requirement
- Add Permissions to the IAM Role Used to Create a Flow Definition
- Create a User That Can Invoke Amazon A2I API Operations
- Create a User With Permissions to Invoke Amazon A2I, Amazon Textract, and Amazon Rekognition API Operations
- Enable Worker Task Template Previews
- Using Amazon A2I with AWS KMS Encrypted Buckets
- Additional Permissions and Security Resources

CORS Permission Requirement

Earlier in 2020, widely used browsers like Chrome and Firefox changed their default behavior for rotating images based on image metadata, referred to as EXIF data. Previously, images would always display in browsers exactly how they are stored on disk, which is typically unrotated. After the change, images now rotate according to a piece of image metadata called orientation value. This has important implications for the entire machine learning (ML) community. For example, if the EXIF orientation is not considered, applications that are used to annotate images may display images in unexpected orientations and result in incorrect labels.

Starting with Chrome 89, AWS can no longer automatically prevent the rotation of images because the web standards group W3C has decided that the ability to control rotation of images violates the web’s Same-Origin Policy. Therefore, to ensure human workers annotate your input images in
a predictable orientation when you submit requests to create a human loop, you must add a CORS header policy to the S3 buckets that contain your input images.

⚠️ **Important**
If you do not add a CORS configuration to the S3 buckets that contains your input data, human review tasks for those input data objects fail.

You can add a CORS policy to an S3 bucket that contains input data in the Amazon S3 console. To set the required CORS headers on the S3 bucket that contains your input images in the S3 console, follow the directions detailed in [How do I add cross-domain resource sharing with CORS?](#). Use the following CORS configuration code for the buckets that host your images. If you use the Amazon S3 console to add the policy to your bucket, you must use the JSON format.

**JSON**

```json
[{
    "AllowedHeaders": [],
    "AllowedMethods": ["GET"],
    "AllowedOrigins": ["*"],
    "ExposeHeaders": []
}]
```

**XML**

```xml
<CORSConfiguration>
    <CORSRule>
        <AllowedOrigin>*</AllowedOrigin>
        <AllowedMethod>GET</AllowedMethod>
    </CORSRule>
</CORSConfiguration>
```

**Add Permissions to the IAM Role Used to Create a Flow Definition**

To create a flow definition, attach the policies in this section to the role that you use when creating a human review workflow in the SageMaker console, or when using the CreateFlowDefinition API operation.
• If you are using the console to create a human review workflow, enter the role Amazon Resource Name (ARN) in the IAM role field when creating a human review workflow in the console.

• When creating a flow definition using the API, attach these policies to the role that is passed to the RoleArn parameter of the CreateFlowDefinition operation.

When you create a human review workflow (flow definition), Amazon A2I invokes Amazon S3 to complete your task. To grant Amazon A2I permission to retrieve and store your files in your Amazon S3 bucket, create the following policy and attach it to your role. For example, if the images, documents, and other files that you are sending for human review are stored in an S3 bucket named my_input_bucket, and if you want the human reviews to be stored in a bucket named my_output_bucket, create the following policy.

```
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": ["s3:GetObject"],
         "Resource": ["arn:aws:s3:::my_input_bucket/*"]
      },
      {
         "Effect": "Allow",
         "Action": ["s3:PutObject"],
         "Resource": ["arn:aws:s3:::my_output_bucket/*"]
      }
   ]
}
```

In addition, the IAM role must have the following trust policy to give SageMaker permission to assume the role. To learn more about IAM trust policies, see Resource-Based Policies section of Policies and Permissions in the AWS Identity and Access Management documentation.
For more information about creating and managing IAM roles and policies, see the following topics in the *AWS Identity and Access Management User Guide*:

- To create an IAM role, see [Creating a Role to Delegate Permissions to an IAM User](#).
- To learn how to create IAM policies, see [Creating IAM Policies](#).
- To learn how to attach an IAM policy to a role, see [Adding and Removing IAM Identity Permissions](#).

## Create a User That Can Invoke Amazon A2I API Operations

To use Amazon A2I to create and start human loops for Amazon Rekognition, Amazon Textract, or the Amazon A2I runtime API, you must use a user that has permissions to invoke Amazon A2I operations. To do this, use the IAM console to attach the `AmazonAugmentedAIFullAccess` managed policy to a new or existing user.

This policy grants permission to a user to invoke API operations from the SageMaker API for flow definition creation and management and the Amazon Augmented AI Runtime API for human loop creation and management. To learn more about these API operations, see [Use APIs in Amazon Augmented AI](#).

`AmazonAugmentedAIFullAccess` does not grant permissions to use Amazon Rekognition or Amazon Textract API operations.
Note

You can also attach the AmazonAugmentedAIFullAccess policy to an IAM role that is used to create and start a human loop.

To provide access, add permissions to your users, groups, or roles:

- Users and groups in AWS IAM Identity Center:
  
  Create a permission set. Follow the instructions in Create a permission set in the AWS IAM Identity Center User Guide.

- Users managed in IAM through an identity provider:
  
  Create a role for identity federation. Follow the instructions in Creating a role for a third-party identity provider (federation) in the IAM User Guide.

- IAM users:
  
  - Create a role that your user can assume. Follow the instructions in Creating a role for an IAM user in the IAM User Guide.
  
  - (Not recommended) Attach a policy directly to a user or add a user to a user group. Follow the instructions in Adding permissions to a user (console) in the IAM User Guide.

For more information, see Adding and Removing IAM Identity Permissions in the AWS Identity and Access Management User Guide.

Create a User With Permissions to Invoke Amazon A2I, Amazon Textract, and Amazon Rekognition API Operations

To create a user that has permission to invoke the API operations used by the built-in task types (that is, DetectModerationLables for Amazon Rekognition and AnalyzeDocument for Amazon Textract) and permission to use all Amazon A2I API operations, attach the IAM managed policy, AmazonAugmentedAIIIntegratedAPIAccess. You may want to use this policy when you want to grant broad permissions to a user using Amazon A2I with more than one task type. To learn more about these API operations, see Use APIs in Amazon Augmented AI.
To provide access, add permissions to your users, groups, or roles:

- Users and groups in AWS IAM Identity Center:

  Create a permission set. Follow the instructions in Create a permission set in the AWS IAM Identity Center User Guide.

- Users managed in IAM through an identity provider:

  Create a role for identity federation. Follow the instructions in Creating a role for a third-party identity provider (federation) in the IAM User Guide.

- IAM users:

  - Create a role that your user can assume. Follow the instructions in Creating a role for an IAM user in the IAM User Guide.
  
  - (Not recommended) Attach a policy directly to a user or add a user to a user group. Follow the instructions in Adding permissions to a user (console) in the IAM User Guide.

For more information, see Adding and Removing IAM Identity Permissions in the AWS Identity and Access Management User Guide.

### Enable Worker Task Template Previews

To customize the interface and instructions that your workers see when working on your tasks, you create a worker task template. You can create the template using the CreateHumanTaskUi operation or the SageMaker console.

To preview your template, you need an IAM role with the following permissions to read Amazon S3 objects that get rendered on your user interface.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
```
For Amazon Rekognition and Amazon Textract task types, you can preview your template using the Amazon Augmented AI section of the SageMaker console. For custom task types, you preview your template by invoking the `RenderUiTemplate` operation. To preview your template, follow the instructions for your task type:

- Amazon Rekognition and Amazon Textract task types – In the SageMaker console, use the role's Amazon Resource Name (ARN) in the procedure documented in [Create a Worker Task Template](#).
- Custom task types – In the `RenderUiTemplate` operation, use the role's ARN in the `RoleArn` parameter.

### Using Amazon A2I with AWS KMS Encrypted Buckets

If you specify an AWS Key Management Service (AWS KMS) customer managed key to encrypt output data in `OutputConfig` of `CreateFlowDefinition`, you must add an IAM policy similar to the following to that key. This policy gives the IAM execution role that you use to create your human loops permission to use this key to perform all of the actions listed in "Action". To learn more about these actions, see [AWS KMS permissions](#) in the AWS Key Management Service Developer Guide.

To use this policy, replace the IAM service-role ARN in "Principal" with the ARN of the execution role you use to create the human review workflow (flow definition). When you create a labeling job using `CreateFlowDefinition`, this is the ARN you specify for `RoleArn`. Note that you cannot provide a `KmsKeyId` when you create a flow definition in the console.

```json
{
    "Sid": "AllowUseOfKmsKey",
    "Effect": "Allow",
    "Principal": {
        "AWS": "arn:aws:iam::111122223333:role/service-role/example-role"
    }
}
```
Additional Permissions and Security Resources

- the section called “Control Access to SageMaker Resources by Using Tags”.
- the section called “SageMaker Identity-Based Policies”
- the section called “Control Creation of SageMaker Resources with Condition Keys”
- the section called “Amazon SageMaker API Permissions Reference”
- Configure security in Amazon SageMaker

Use Amazon CloudWatch Events in Amazon Augmented AI

Amazon Augmented AI uses Amazon CloudWatch Events to alert you when a human review loop status changes to Completed, Failed, or Stopped. This event delivery is guaranteed at least once, which means all events created when human loops finish are successfully delivered to CloudWatch Events (Amazon EventBridge). When a review loop changes to one of these states, Augmented AI sends an event to CloudWatch Events similar to the following.

```json
{
    "version":"0",
    "id":"12345678-1111-2222-3333-12345EXAMPLE",
    "detail-type":"SageMaker A2I HumanLoop Status Change",
    "source":"aws.sagemaker",
    "account":"111111111111",
    "time":"2019-11-14T17:49:25Z",
    "region":"us-east-1",
    "resources": ["arn:aws:sagemaker:us-east-1:111111111111:human-loop/humanloop-nov-14-1"],
    "detail":{
        "creationTime":"2019-11-14T17:36.740Z",
        "failureCode":null,
```
The details in the JSON output include the following:

creationTime

   The timestamp when Augmented AI created the human loop.

failureCode

   A failure code denoting a specific type of failure.

failureReason

   The reason why a human loop has failed. The failure reason is only returned when the human review loop status is failed.

flowDefinitionArn

   The Amazon Resource Name (ARN) of the flow definition, or human review workflow.

humanLoopArn

   The Amazon Resource Name (ARN) of the human loop.

humanLoopName

   The name of the human loop.

humanLoopOutput

   An object containing information about the output of the human loop.

outputS3Uri

   The location of the Amazon S3 object where Augmented AI stores your human loop output.
humanLoopStatus

The status of the human loop.

Send Events from Your Human Loop to CloudWatch Events

To configure a CloudWatch Events rule to get status updates, or events, for your Amazon A2I human loops, use the AWS Command Line Interface (AWS CLI) put-rule command. When using the put-rule command, specify the following to receive human loop statuses:

- "source": ["aws.sagemaker"]
- "detail-type": ["SageMaker A2I HumanLoop Status Change"]

To configure a CloudWatch Events rule to watch for all status changes, use the following command and replace the placeholder text. For example, replace "A2IHumanLoopStatusChanges" with a unique CloudWatch Events rule name and "arn:aws:iam::111122223333:role/MyRoleForThisRule" with the Amazon Resource Number (ARN) of an IAM role with an events.amazonaws.com trust policy attached. Replace region with the AWS Region in which you want to create the rule.

```
aws events put-rule --name "A2IHumanLoopStatusChanges"
    --event-pattern "{"source": ["aws.sagemaker"], "detail-type": ["SageMaker A2I HumanLoop Status Change"]}"
    --role-arn "arn:aws:iam::111122223333:role/MyRoleForThisRule"
    --region "region"
```

To learn more about the put-rule request, see Event Patterns in CloudWatch Events in the Amazon CloudWatch Events User Guide.

Set Up a Target to Process Events

To process events, you need to set up a target. For example, if you want to receive an email when a human loop status changes, use a procedure in Setting Up Amazon SNS Notifications in the Amazon CloudWatch User Guide to set up an Amazon SNS topic and subscribe your email to it. Once you have created a topic, you can use it to create a target.

To add a target to your CloudWatch Events rule

1. Open the CloudWatch console: https://console.aws.amazon.com/cloudwatch/home
2. In the navigation pane, choose Rules.
3. Choose the rule to which you want to add a target.
4. Choose Actions, and then choose Edit.
5. Under Targets, choose Add Target and choose the AWS service you want to act when a human loop status change event is detected.
6. Configure your target. For instructions, see the topic for configuring a target in the AWS documentation for that service.
7. Choose Configure details.
8. For Name, enter a name and, optionally, provide details about the purpose of the rule in Description.
9. Make sure that the check box next to State is selected so that your rule is listed as Enabled.
10. Choose Update rule.

Use Human Review Output

After you receive human review results, you can analyze the results and compare them to machine learning predictions. The JSON that is stored in the Amazon S3 bucket contains both the machine learning predictions and the human review results.

More Information

Automating Amazon SageMaker with Amazon EventBridge

Use APIs in Amazon Augmented AI

You can create a human review workflow or a worker task template programmatically. The APIs you use depend on whether you are creating a Amazon Rekognition, Amazon Textract, or custom task type. This topic provides links to API reference documentation for each task type and programming task.

The following APIs can be used with Augmented AI:

Amazon Augmented AI

Use the Augmented AI API to start, stop, and delete human review loops. You can also list all human review loops and return information about human review loops in your account.
Learn more about human review loop APIs in the [Amazon Augmented AI Runtime API Reference](https://docs.aws.amazon.com/a2i/latest/dg/).

### Amazon Rekognition

Use the `HumanLoopConfig` parameter of the [DetectModerationLabels](https://docs.aws.amazon.com/rekognition/latest/dg/API_DetectModerationLabels.html) API to initiate a human review workflow using Amazon Rekognition.

### Amazon SageMaker

Use the Amazon SageMaker API to create a `FlowDefinition`, also known as a *human review workflow*. You can also create a `HumanTaskUi` or *worker task template*.

For more information, see the [CreateFlowDefinition](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateFlowDefinition.html) or the [CreateHumanTaskUi](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateHumanTaskUi.html) API documentation.

### Amazon Textract

Use the `HumanLoopConfig` parameter of the [AnalyzeDocument](https://docs.aws.amazon.com/textract/latest/dg/API_AnalyzeDocument.html) API to initiate a human review workflow using Amazon Textract.

### Programmatic Tutorials

The following tutorials provide example code and step-by-step instructions for creating human review workflows and worker task templates programmatically.

- [Tutorial: Get Started Using the Amazon A2I API](https://docs.aws.amazon.com/a2i/latest/dg/)
- [Create a Human Review Workflow (API)](https://docs.aws.amazon.com/a2i/latest/dg/)
- [Create and Start a Human Loop](https://docs.aws.amazon.com/a2i/latest/dg/)
- [Using Amazon Augmented AI with Amazon Rekognition](https://docs.aws.amazon.com/rekognition/latest/dg/) in the *Amazon Rekognition Developer Guide*
- [Using Amazon Augmented AI with Amazon Textract AnalyzeDocument](https://docs.aws.amazon.com/textract/latest/dg/) in the *Amazon Textract Developer Guide*
Prepare data

Data preparation in machine learning refers to the process of collecting, preprocessing, and organizing raw data to make it suitable for analysis and modeling. This step ensures that the data is in a format that machine learning algorithms can effectively learn from. Data preparation tasks may include handling missing values, removing outliers, scaling features, encoding categorical variables, assessing potential biases and taking steps to mitigate them, splitting data into training and testing sets, labeling, and other necessary transformations to optimize the quality and usability of the data for subsequent machine learning tasks.

Amazon SageMaker provides several built-in features for performing data preparation tasks such as cleaning, transforming, and labeling datasets before model training.

- For **low code data preparation**, you can use Amazon SageMaker Data Wrangler to create data flows that define your ML data pre-processing and feature engineering workflows using little to no coding. Import data from sources such as Amazon S3, Amazon Redshift, or Snowflake to engineer features. You can use built-in visualizations and analyses to get insights from your data. After preparing your data, you can export the finished output to Amazon S3, Amazon SageMaker Feature Store, or SageMaker Pipelines. Data Wrangler exists within Amazon SageMaker Canvas and Amazon SageMaker Studio Classic. We recommend using it within SageMaker Canvas for the latest features. For more information about Data Wrangler within SageMaker Canvas, see the section called “Prepare data”. For information about Data Wrangler within Studio Classic, see the section called “Prepare Data with Data Wrangler”.

- For **data preparation at scale** using open-source frameworks such as Apache Spark, Apache Hive, or Presto, Amazon SageMaker Studio Classic provides a built-in integration with Amazon EMR. You can use SageMaker Studio Classic to connect or provision Amazon EMR clusters from your notebooks for petabyte-scale data processing, interactive analytics, and machine learning. For more information about using Amazon EMR from SageMaker Studio Classic, see Prepare data using Amazon EMR.

  Alternatively, you can use the Apache Spark-based serverless engine from AWS Glue interactive sessions to aggregate, transform, and prepare data from multiple sources in SageMaker Studio Classic. For more information about using AWS Glue interactive sessions within SageMaker Studio Classic, see Prepare data using AWS Glue Interactive Sessions.

- For **feature discovery and storage**, the Amazon SageMaker Feature Store has capabilities to search, discover, and retrieve features for model training and provides a centralized repository to store feature data in a standardized format. Storing curated features in the Feature Store allows
reuse of existing features for new ML projects. The Feature Store manages the full lifecycle of features including tracking lineage, calculating statistics, and maintaining audit trails. For more information on feature data storage for your ML pipelines, refer to the Create, store, and share features section in this guide.

- **For bias detection**, you can use Amazon SageMaker Clarify to analyze your data and detect potential biases across multiple facets. For example, you can use SageMaker Clarify to detect if your training data contains imbalanced representations or labeling biases between groups such as gender, race, or age. SageMaker Clarify can help you identify these biases before training a model to avoid propagating biases into the model's predictions. For information about using SageMaker Clarify to uncover biases, refer to the the section called “Detect Pre-training Data Bias” section in this guide.

- **For data labeling**, you can use SageMaker Ground Truth to manage the data labeling workflows of your training datasets. For information about how to use Ground Truth for your labeling tasks, refer to the Label data with a human-in-the-loop section in this guide.

After performing exploratory data analysis and creating your data transformations steps, you can productionize your transformation code using SageMaker Processing jobs and automate your preparation workflow using SageMaker Model Building Pipelines.

For information about the SageMaker Processing API, see Amazon SageMaker processing jobs.

For information about automating your transformation steps, see SageMaker Model Building Pipelines.

**Topics**

- Prepare ML Data with Amazon SageMaker Data Wrangler
- Prepare Data at Scale with Studio Classic using Amazon EMR or AWS Glue

**Prepare ML Data with Amazon SageMaker Data Wrangler**

⚠️ **Important**

Amazon SageMaker Data Wrangler has been integrated into Amazon SageMaker Canvas. Within the new Data Wrangler experience in SageMaker Canvas, you can use a natural language interface to explore and transform your data in addition to the visual interface. For more information about Data Wrangler in SageMaker Canvas, see Prepare data.
Amazon SageMaker Data Wrangler (Data Wrangler) is a feature of Amazon SageMaker Studio Classic that provides an end-to-end solution to import, prepare, transform, featurize, and analyze data. You can integrate a Data Wrangler data preparation flow into your machine learning (ML) workflows to simplify and streamline data pre-processing and feature engineering using little to no coding. You can also add your own Python scripts and transformations to customize workflows.

Data Wrangler provides the following core functionalities to help you analyze and prepare data for machine learning applications.

- **Import** – Connect to and import data from Amazon Simple Storage Service (Amazon S3), Amazon Athena (Athena), Amazon Redshift, Snowflake, and Databricks.

- **Data Flow** – Create a data flow to define a series of ML data prep steps. You can use a flow to combine datasets from different data sources, identify the number and types of transformations you want to apply to datasets, and define a data prep workflow that can be integrated into an ML pipeline.

- **Transform** – Clean and transform your dataset using standard transforms like string, vector, and numeric data formatting tools. Featurize your data using transforms like text and date/time embedding and categorical encoding.

- **Generate Data Insights** – Automatically verify data quality and detect abnormalities in your data with Data Wrangler Data Insights and Quality Report.

- **Analyze** – Analyze features in your dataset at any point in your flow. Data Wrangler includes built-in data visualization tools like scatter plots and histograms, as well as data analysis tools like target leakage analysis and quick modeling to understand feature correlation.

- **Export** – Export your data preparation workflow to a different location. The following are example locations:
  - Amazon Simple Storage Service (Amazon S3) bucket
  - Amazon SageMaker Model Building Pipelines – Use SageMaker Pipelines to automate model deployment. You can export the data that you've transformed directly to the pipelines.
  - Amazon SageMaker Feature Store – Store the features and their data in a centralized store.
  - Python script – Store the data and their transformations in a Python script for your custom workflows.

To start using Data Wrangler, see [Get Started with Data Wrangler](#).
### Important

Data Wrangler no longer supports Jupyter Lab Version 1 (JL1). To access the latest features and updates, update to Jupyter Lab Version 3. For more information about upgrading, see View and update the JupyterLab version of an application from the console.

### Important

The information and procedures in this guide use the latest version of Amazon SageMaker Studio Classic. For information about updating Studio Classic to the latest version, see Amazon SageMaker Studio Classic UI Overview.

You must use Studio Classic version 1.3.0 or later. Use the following procedure to open Amazon SageMaker Studio Classic and see which version you're running.

To open Studio Classic and check its version, see the following procedure.

1. Use the steps in Prerequisites to access Data Wrangler through Amazon SageMaker Studio Classic.
2. Next to the user you want to use to launch Studio Classic, select Launch app.
3. Choose Studio.
4. After Studio Classic loads, select File, then New, and then Terminal.
5. Once you have launched Studio Classic, select **File**, then **New**, and then **Terminal**.

6. Enter `cat /opt/conda/share/jupyter/lab/staging/yarn.lock | grep -A 1 "@amzn/sagemaker-ui-data-prep-plugin@"` to print the version of your Studio Classic instance. You must have Studio Classic version 1.3.0 to use Snowflake.

You can update Amazon SageMaker Studio Classic from within the AWS Management Console. For more information about updating Studio Classic, see [Amazon SageMaker Studio Classic UI Overview](#).

**Topics**

- [Get Started with Data Wrangler](#)
Get Started with Data Wrangler

Amazon SageMaker Data Wrangler is a feature in Amazon SageMaker Studio Classic. Use this section to learn how to access and get started using Data Wrangler. Do the following:

1. Complete each step in Prerequisites.
2. Follow the procedure in Access Data Wrangler to start using Data Wrangler.

Prerequisites

To use Data Wrangler, you must complete the following prerequisites.

1. To use Data Wrangler, you need access to an Amazon Elastic Compute Cloud (Amazon EC2) instance. For more information about the Amazon EC2 instances that you can use, see Instances. To learn how to view your quotas and, if necessary, request a quota increase, see AWS service quotas.
2. Configure the required permissions described in Security and Permissions.

3. If your organization is using a firewall that blocks internet traffic, you must have access to the following URLs:

   • https://ui.prod-1.data-wrangler.sagemaker.aws/
   • https://ui.prod-2.data-wrangler.sagemaker.aws/
   • https://ui.prod-3.data-wrangler.sagemaker.aws/
   • https://ui.prod-4.data-wrangler.sagemaker.aws/

To use Data Wrangler, you need an active Studio Classic instance. To learn how to launch a new instance, see Amazon SageMaker domain overview. When your Studio Classic instance is Ready, use the instructions in Access Data Wrangler.

Access Data Wrangler

The following procedure assumes you have completed the Prerequisites.

To access Data Wrangler in Studio Classic, do the following.

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.

2. Choose Studio.

3. Choose Launch app.

4. From the dropdown list, select Studio.

5. Choose the Home icon.

6. Choose Data.

7. Choose Data Wrangler.

8. You can also create a Data Wrangler flow by doing the following.

   a. In the top navigation bar, select File.

   b. Select New.

   c. Select Data Wrangler Flow.
9. (Optional) Rename the new directory and the .flow file.

10. When you create a new .flow file in Studio Classic, you might see a carousel that introduces you to Data Wrangler.

    **This may take a few minutes.**

This messaging persists as long as the KernelGateway app on your User Details page is Pending. To see the status of this app, in the SageMaker console on the Amazon SageMaker Studio Classic page, select the name of the user you are using to access Studio Classic. On the User Details page, you see a KernelGateway app under Apps. Wait until this app status is Ready to start using Data Wrangler. This can take around 5 minutes the first time you launch Data Wrangler.
11. To get started, choose a data source and use it to import a dataset. See Import to learn more.

When you import a dataset, it appears in your data flow. To learn more, see Create and Use a Data Wrangler Flow.

12. After you import a dataset, Data Wrangler automatically infers the type of data in each column. Choose + next to the Data types step and select Edit data types.

⚠️ Important
After you add transforms to the Data types step, you cannot bulk-update column types using Update types.

13. Use the data flow to add transforms and analyses. To learn more see Transform Data and Analyze and Visualize.

14. To export a complete data flow, choose Export and choose an export option. To learn more, see Export.

15. Finally, choose the Components and registries icon, and select Data Wrangler from the dropdown list to see all the .flow files that you've created. You can use this menu to find and move between data flows.

After you have launched Data Wrangler, you can use the following section to walk through how you might use Data Wrangler to create an ML data prep flow.
Update Data Wrangler

We recommend that you periodically update the Data Wrangler Studio Classic app to access the latest features and updates. The Data Wrangler app name starts with `sagemaker-data-wrang`. To learn how to update a Studio Classic app, see Shut down and Update Studio Classic Apps.

Demo: Data Wrangler Titanic Dataset Walkthrough

The following sections provide a walkthrough to help you get started using Data Wrangler. This walkthrough assumes that you have already followed the steps in Access Data Wrangler and have a new data flow file open that you intend to use for the demo. You may want to rename this .flow file to something similar to `titanic-demo.flow`.

This walkthrough uses the Titanic dataset. It's a modified version of the Titanic dataset that you can import into your Data Wrangler flow more easily. This data set contains the survival status, age, gender, and class (which serves as a proxy for economic status) of passengers aboard the maiden voyage of the RMS Titanic in 1912.

In this tutorial, you perform the following steps.

1. Do one of the following:
   - Open your Data Wrangler flow and choose Use Sample Dataset.
   - Upload the Titanic dataset to Amazon Simple Storage Service (Amazon S3), and then import this dataset into Data Wrangler.
2. Analyze this dataset using Data Wrangler analyses.
3. Define a data flow using Data Wrangler data transforms.
4. Export your flow to a Jupyter Notebook that you can use to create a Data Wrangler job.
5. Process your data, and kick off a SageMaker training job to train a XGBoost Binary Classifier.

Upload Dataset to S3 and Import

To get started, you can use one of the following methods to import the Titanic dataset into Data Wrangler:

- Importing the dataset directly from the Data Wrangler flow
- Uploading the dataset to Amazon S3 and then importing it into Data Wrangler

To import the dataset directly into Data Wrangler, open the flow and choose Use Sample Dataset.
Uploading the dataset to Amazon S3 and importing it into Data Wrangler is closer to the experience you have importing your own data. The following information tells you how to upload your dataset and import it.

Before you start importing the data into Data Wrangler, download the Titanic dataset and upload it to an Amazon S3 (Amazon S3) bucket in the AWS Region in which you want to complete this demo.

If you are a new user of Amazon S3, you can do this using drag and drop in the Amazon S3 console. To learn how, see Uploading Files and Folders by Using Drag and Drop in the Amazon Simple Storage Service User Guide.

⚠️ Important

Upload your dataset to an S3 bucket in the same AWS Region you want to use to complete this demo.

When your dataset has been successfully uploaded to Amazon S3, you can import it into Data Wrangler.

**Import the Titanic dataset to Data Wrangler**

1. Choose the Import data button in your Data flow tab or choose the Import tab.
2. Select Amazon S3.
3. Use the Import a dataset from S3 table to find the bucket to which you added the Titanic dataset. Choose the Titanic dataset CSV file to open the Details pane.
4. Under Details, the File type should be CSV. Check First row is header to specify that the first row of the dataset is a header. You can also name the dataset something more friendly, such as Titanic-train.
5. Choose the Import button.

When your dataset is imported into Data Wrangler, it appears in your Data Flow tab. You can double click on a node to enter the node detail view, which allows you to add transformations or analysis. You can use the plus icon for a quick access to the navigation. In the next section, you use this data flow to add analysis and transform steps.
Data Flow

In the data flow section, the only steps in the data flow are your recently imported dataset and a Data type step. After applying transformations, you can come back to this tab and see what the data flow looks like. Now, add some basic transformations under the Prepare and Analyze tabs.

Prepare and Visualize

Data Wrangler has built-in transformations and visualizations that you can use to analyze, clean, and transform your data.

The Data tab of the node detail view lists all built-in transformations in the right panel, which also contains an area in which you can add custom transformations. The following use case showcases how to use these transformations.

To get information that might help you with data exploration and feature engineering, create a data quality and insights report. The information from the report can help you clean and process your data. It gives you information such as the number of missing values and the number of outliers. If you have issues with your data, such as target leakage or imbalance, the insights report can bring those issues to your attention. For more information about creating a report, see Get Insights On Data and Data Quality.

Data Exploration

First, create a table summary of the data using an analysis. Do the following:

1. Choose the + next to the Data type step in your data flow and select Add analysis.
2. In the Analysis area, select Table summary from the dropdown list.
3. Give the table summary a Name.
4. Select Preview to preview the table that will be created.
5. Choose Save to save it to your data flow. It appears under All Analyses.

Using the statistics you see, you can make observations similar to the following about this dataset:

- Fare average (mean) is around $33, while the max is over $500. This column likely has outliers.
- This dataset uses ? to indicate missing values. A number of columns have missing values: cabin, embarked, and home.dest
- The age category is missing over 250 values.
Next, clean your data using the insights gained from these stats.

**Drop Unused Columns**

Using the analysis from the previous section, clean up the dataset to prepare it for training. To add a new transform to your data flow, choose + next to the Data type step in your data flow and choose Add transform.

First, drop columns that you don't want to use for training. You can use pandas data analysis library to do this, or you can use one of the built-in transforms.

Use the following procedure to drop the unused columns.

To drop the unused columns.

1. Open the Data Wrangler flow.
2. There are two nodes in your Data Wrangler flow. Choose the + to the right of the Data types node.
3. Choose Add transform.
4. In the All steps column, choose Add step.
5. In the Standard transform list, choose Manage Columns. The standard transformations are ready-made, built-in transformations. Make sure that Drop column is selected.
6. Under Columns to drop, check the following column names:
   - cabin
   - ticket
   - name
   - sibsp
   - parch
   - home.dest
   - boat
   - body
7. Choose Preview.
8. Verify that the columns have been dropped, then choose Add.

To do this using pandas, follow these steps.
1. In the All steps column, choose Add step.

2. In the Custom transform list, choose Custom transform.

3. Provide a name for your transformation, and choose Python (Pandas) from the dropdown list.

4. Enter the following Python script in the code box.

```python
cols = ['name', 'ticket', 'cabin', 'sibsp', 'parch', 'home.dest', 'boat', 'body']
df = df.drop(cols, axis=1)
```

5. Choose Preview to preview the change, and then choose Add to add the transformation.

### Clean up Missing Values

Now, clean up missing values. You can do this with the Handling missing values transform group.

A number of columns have missing values. Of the remaining columns, age and fare contain missing values. Inspect this using a Custom Transform.

Using the Python (Pandas) option, use the following to quickly review the number of entries in each column:

```python
df.info()
```
To drop rows with missing values in the \textit{age} category, do the following:

1. Choose \textbf{Handle missing}.
2. Choose \textbf{Drop missing} for the \textbf{Transformer}.
3. Choose \textit{age} for the \textbf{Input column}.
4. Choose \textbf{Preview} to see the new data frame, and then choose \textbf{Add} to add the transform to your flow.
5. Repeat the same process for \textit{fare}.

You can use \texttt{df.info()} in the \textbf{Custom transform} section to confirm that all rows now have 1,045 values.

\textbf{Custom Pandas: Encode}

Try flat encoding using Pandas. Encoding categorical data is the process of creating a numerical representation for categories. For example, if your categories are Dog and Cat, you may encode this information into two vectors: $[1, 0]$ to represent Dog, and $[0, 1]$ to represent Cat.

1. In the \textbf{Custom Transform} section, choose \textbf{Python (Pandas)} from the dropdown list.
2. Enter the following in the code box.

```python
import pandas as pd

dummies = []
cols = ['pclass', 'sex', 'embarked']
for col in cols:
    dummies.append(pd.get_dummies(df[col]))

encoded = pd.concat(dummies, axis=1)
df = pd.concat((df, encoded), axis=1)
```

3. Choose Preview to preview the change. The encoded version of each column is added to the dataset.

4. Choose Add to add the transformation.

**Custom SQL: SELECT Columns**

Now, select the columns you want to keep using SQL. For this demo, select the columns listed in the following SELECT statement. Because survived is your target column for training, put that column first.

1. In the Custom Transform section, select SQL (PySpark SQL) from the dropdown list.
2. Enter the following in the code box.

```sql
SELECT survived, age, fare, 1, 2, 3, female, male, C, Q, S FROM df;
```

3. Choose Preview to preview the change. The columns listed in your SELECT statement are the only remaining columns.

4. Choose Add to add the transformation.

**Export to a Data Wrangler Notebook**

When you've finished creating a data flow, you have a number of export options. The following section explains how to export to a Data Wrangler job notebook. A Data Wrangler job is used to process your data using the steps defined in your data flow. To learn more about all export options, see Export.
Export to Data Wrangler Job Notebook

When you export your data flow using a Data Wrangler job, the process automatically creates a Jupyter Notebook. This notebook automatically opens in your Studio Classic instance and is configured to run a SageMaker processing job to run your Data Wrangler data flow, which is referred to as a Data Wrangler job.

1. Save your data flow. Select File and then select Save Data Wrangler Flow.
2. Back to the Data Flow tab, select the last step in your data flow (SQL), then choose the + to open the navigation.
3. Choose Export, and Amazon S3 (via Jupyter Notebook). This opens a Jupyter Notebook.
4. Choose any Python 3 (Data Science) kernel for the Kernel.
5. When the kernel starts, run the cells in the notebook book until Kick off SageMaker Training Job (Optional).
6. Optionally, you can run the cells in Kick off SageMaker Training Job (Optional) if you want to create a SageMaker training job to train an XGBoost classifier. You can find the cost to run a SageMaker training job in Amazon SageMaker Pricing.

Alternatively, you can add the code blocks found in Training XGBoost Classifier to the notebook and run them to use the XGBoost open source library to train an XGBoost classifier.
7. Uncomment and run the cell under Cleanup and run it to revert the SageMaker Python SDK to its original version.

You can monitor your Data Wrangler job status in the SageMaker console in the Processing tab. Additionally, you can monitor your Data Wrangler job using Amazon CloudWatch. For additional information, see Monitor Amazon SageMaker Processing Jobs with CloudWatch Logs and Metrics.
If you kicked off a training job, you can monitor its status using the SageMaker console under **Training jobs** in the **Training section**.

**Training XGBoost Classifier**

You can train an XGBoost Binary Classifier using either a Jupyter notebook or a Amazon SageMaker Autopilot. You can use Autopilot to automatically train and tune models on the data that you've transformed directly from your Data Wrangler flow. For information about Autopilot, see [Automatically Train Models on Your Data Flow](#).

In the same notebook that kicked off the Data Wrangler job, you can pull the data and train an XGBoost Binary Classifier using the prepared data with minimal data preparation.

1. First, upgrade necessary modules using `pip` and remove the `_SUCCESS` file (this last file is problematic when using `awswrangler`).

   ```bash
   ! pip install --upgrade awscli awswrangler boto sklearn
   ! aws s3 rm {output_path} --recursive --exclude "*" --include "*_SUCCESS*"
   ```

2. Read the data from Amazon S3. You can use `awswrangler` to recursively read all the CSV files in the S3 prefix. The data is then split into features and labels. The label is the first column of the dataframe.

   ```python
   import awswrangler as wr
   
   df = wr.s3.read_csv(path=output_path, dataset=True)
   X, y = df.iloc[:,:-1], df.iloc[:,-1]
   ```

   - Finally, create DMatrices (the XGBoost primitive structure for data) and do cross-validation using the XGBoost binary classification.

     ```python
     import xgboost as xgb
     
     dmatrix = xgb.DMatrix(data=X, label=y)
     
     params = {"objective":"binary:logistic","learning_rate": 0.1, 'max_depth': 5, 'alpha': 10}
     
     xgb.cv(
         dtrain=dmatrix,
         params=params,
     ```
nfold=3,
num_boost_round=50,
early_stopping_rounds=10,
metrics="rmse",
    as_pandas=True,
    seed=123)

**Shut down Data Wrangler**

When you are finished using Data Wrangler, we recommend that you shut down the instance it runs on to avoid incurring additional charges. To learn how to shut down the Data Wrangler app and associated instance, see [Shut Down Data Wrangler](#).

**Import**

You can use Amazon SageMaker Data Wrangler to import data from the following *data sources*: Amazon Simple Storage Service (Amazon S3), Amazon Athena, Amazon Redshift, and Snowflake. The dataset that you import can include up to 1000 columns.

**Topics**

- [Import data from Amazon S3](#)
- [Import data from Athena](#)
- [Import data from Amazon Redshift](#)
- [Import data from Amazon EMR](#)
- [Import data from Databricks (JDBC)](#)
- [Import data from Salesforce Data Cloud](#)
- [Import data from Snowflake](#)
- [Import Data From Software as a Service (SaaS) Platforms](#)
- [Imported Data Storage](#)

Some data sources allow you to add multiple *data connections*:

- You can connect to multiple Amazon Redshift clusters. Each cluster becomes a data source.
- You can query any Athena database in your account to import data from that database.
When you import a dataset from a data source, it appears in your data flow. Data Wrangler automatically infers the data type of each column in your dataset. To modify these types, select the **Data types** step and select **Edit data types**.

When you import data from Athena or Amazon Redshift, the imported data is automatically stored in the default SageMaker S3 bucket for the AWS Region in which you are using Studio Classic. Additionally, Athena stores data you preview in Data Wrangler in this bucket. To learn more, see [Imported Data Storage](#).

⚠️ **Important**

The default Amazon S3 bucket may not have the least permissive security settings, such as bucket policy and server-side encryption (SSE). We strongly recommend that you [Add a Bucket Policy To Restrict Access to Datasets Imported to Data Wrangler](#).

⚠️ **Important**

In addition, if you use the managed policy for SageMaker, we strongly recommend that you scope it down to the most restrictive policy that allows you to perform your use case. For more information, see [Grant an IAM Role Permission to Use Data Wrangler](#).

All data sources except for Amazon Simple Storage Service (Amazon S3) require you to specify a SQL query to import your data. For each query, you must specify the following:

- **Data catalog**
- **Database**
- **Table**

You can specify the name of the database or the data catalog in either the drop down menus or within the query. The following are example queries:

- `select * from example-data-catalog-name.example-database-name.example-table-name` – The query doesn't use anything specified in the dropdown menus of the user-interface (UI) to run. It queries `example-table-name` within `example-database-name` within `example-data-catalog-name`. 

---
• select * from `example-database-name.example-table-name` – The query uses the data catalog that you've specified in the Data catalog dropdown menu to run. It queries `example-table-name` within `example-database-name` within the data catalog that you've specified.

• select * from `example-table-name` – The query requires you to select fields for both the Data catalog and Database name dropdown menus. It queries `example-table-name` within the data catalog within the database and data catalog that you've specified.

The link between Data Wrangler and the data source is a connection. You use the connection to import data from your data source.

There are the following types of connections:

• Direct
• Cataloged

Data Wrangler always has access to the most recent data in a direct connection. If the data in the data source has been updated, you can use the connection to import the data. For example, if someone adds a file to one of your Amazon S3 buckets, you can import the file.

A cataloged connection is the result of a data transfer. The data in the cataloged connection doesn't necessarily have the most recent data. For example, you might set up a data transfer between Salesforce and Amazon S3. If there's an update to the Salesforce data, you must transfer the data again. You can automate the process of transferring data. For more information about data transfers, see Import Data From Software as a Service (SaaS) Platforms.

**Import data from Amazon S3**

You can use Amazon Simple Storage Service (Amazon S3) to store and retrieve any amount of data, at any time, from anywhere on the web. You can accomplish these tasks using the AWS Management Console, which is a simple and intuitive web interface, and the Amazon S3 API. If you've stored your dataset locally, we recommend that you add it to an S3 bucket for import into Data Wrangler. To learn how, see Uploading an object to a bucket in the Amazon Simple Storage Service User Guide.

Data Wrangler uses S3 Select to allow you to preview your Amazon S3 files in Data Wrangler. You incur standard charges for each file preview. To learn more about pricing, see the Requests & data retrievals tab on Amazon S3 pricing.
Important
If you plan to export a data flow and launch a Data Wrangler job, ingest data into a SageMaker feature store, or create a SageMaker pipeline, be aware that these integrations require Amazon S3 input data to be located in the same AWS region.

Important
If you're importing a CSV file, make sure it meets the following requirements:

- A record in your dataset can't be longer than one line.
- A backslash, \, is the only valid escape character.
- Your dataset must use one of the following delimiters:
  - Comma – ,
  - Colon – :
  - Semicolon – ;
  - Pipe – |
  - Tab – [TAB]

To save space, you can import compressed CSV files.

Data Wrangler gives you the ability to either import the entire dataset or sample a portion of it. For Amazon S3, it provides the following sampling options:

- None – Import the entire dataset.
- First K – Sample the first K rows of the dataset, where K is an integer that you specify.
- Randomized – Takes a random sample of a size that you specify.
- Stratified – Takes a stratified random sample. A stratified sample preserves the ratio of values in a column.

After you've imported your data, you can also use the sampling transformer to take one or more samples from your entire dataset. For more information about the sampling transformer, see Sampling.
You can use one of the following resource identifiers to import your data:

- An Amazon S3 URI that uses an Amazon S3 bucket or Amazon S3 access point
- An Amazon S3 access point alias
- An Amazon Resource Name (ARN) that uses an Amazon S3 access point or Amazon S3 bucket

Amazon S3 access points are named network endpoints that are attached to the buckets. Each access point has distinct permissions and network controls that you can configure. For more information about access points, see Managing data access with Amazon S3 access points.

Important

If you're using an Amazon Resource Name (ARN) to import your data, it must be for a resource located in the same AWS Region that you're using to access Amazon SageMaker Studio Classic.

You can import either a single file or multiple files as a dataset. You can use the multifile import operation when you have a dataset that is partitioned into separate files. It takes all of the files from an Amazon S3 directory and imports them as a single dataset. For information on the types of files that you can import and how to import them, see the following sections.

Single File Import

You can import single files in the following formats:

- Comma Separated Values (CSV)
- Parquet
- Javascript Object Notation (JSON)
- Optimized Row Columnar (ORC)
- Image – Data Wrangler uses OpenCV to import images. For more information about supported image formats, see Image file reading and writing.

For files formatted in JSON, Data Wrangler supports both JSON lines (.jsonl) and JSON documents (.json). When you preview your data, it automatically shows the JSON in tabular format. For nested JSON documents that are larger than 5 MB, Data Wrangler shows the schema for the structure and the arrays as values in the dataset. Use the Flatten structured and...
Explode array operators to display the nested values in tabular format. For more information, see Unnest JSON Data and Explode Array.

When you choose a dataset, you can rename it, specify the file type, and identify the first row as a header.

You can import a dataset that you've partitioned into multiple files in an Amazon S3 bucket in a single import step.

To import a dataset into Data Wrangler from a single file that you've stored in Amazon S3:

1. If you are not currently on the Import tab, choose Import.
2. Under Available, choose Amazon S3.
3. From the Import tabular, image, or time-series data from S3, do one of the following:
   • Choose an Amazon S3 bucket from the tabular view and navigate to the file that you're importing.
   • For S3 source, specify an Amazon S3 bucket or an Amazon S3 URI and select Go. The Amazon S3 URIs can be in one of the following formats:
     - s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file
     - example-access-point-aqfqprnsten7aefdfbarligizwgyfouse1a-s3alias/datasets/example-file
4. Choose the dataset to open the Import settings pane.
5. If your CSV file has a header, select the checkbox next to Add header to table.
6. Use the Preview table to preview your dataset. This table shows up to 100 rows.
7. In the Details pane, verify or change the Name and File Type for your dataset. If you add a Name that contains spaces, these spaces are replaced with underscores when your dataset is imported.
8. Specify the sampling configuration that you'd like to use.
9. Choose Import.

Multifile Import

The following are the requirements for importing multiple files:
The files must be in the same folder of your Amazon S3 bucket.

The files must either share the same header or have no header.

Each file must be in one of the following formats:

- CSV
- Parquet
- Optimized Row Columnar (ORC)
- Image – Data Wrangler uses OpenCV to import images. For more information about supported image formats, see [Image file reading and writing](#).

Use the following procedure to import multiple files.

**To import a dataset into Data Wrangler from multiple files that you've stored in an Amazon S3 directory**

1. If you are not currently on the **Import** tab, choose **Import**.
2. Under **Available**, choose **Amazon S3**.
3. From the **Import tabular, image, or time-series data from S3**,
   do one of the following:
   - Choose an Amazon S3 bucket from the tabular view and navigate to the folder containing the files that you're importing.
   - For **S3 source**, specify the Amazon S3 bucket or an Amazon S3 URI with your files and select **Go**. The following are valid URIs:
     - `s3://DOC-EXAMPLE-BUCKET/example-prefix/example-prefix`
     - `example-access-point-aqfqprnstan7aefdfbarligizwgyfouse1a-s3alias/example-prefix/`
     - `s3://arn:aws:s3:AWS-Region:111122223333:accesspoint/example-prefix`
4. Select the folder containing the files that you want to import. Each file must be in one of the supported formats. Your files must be the same data type.
5. If your folder contains CSV files with headers, select the checkbox next to **First row is header**.
6. If your files are nested within other folders, select the checkbox next to Include nested directories.

7. (Optional) Choose Add filename column add a column to the dataset that shows the filename for each observation.

8. (Optional) By default, Data Wrangler doesn't show you a preview of a folder. You can activate previewing by choosing the blue Preview off button. A preview shows the first 10 rows of the first 10 files in the folder.

9. In the Details pane, verify or change the Name and File Type for your dataset. If you add a Name that contains spaces, these spaces are replaced with underscores when your dataset is imported.

10. Specify the sampling configuration that you'd like to use.

11. Choose Import dataset.

You can also use parameters to import a subset of files that match a pattern. Parameters help you more selectively pick the files that you're importing. To start using parameters, edit the data source and apply them to the path that you're using to import the data. For more information, see Reusing Data Flows for Different Datasets.

**Import data from Athena**

Use Amazon Athena to import your data from Amazon Simple Storage Service (Amazon S3) into Data Wrangler. In Athena, you write standard SQL queries to select the data that you're importing from Amazon S3. For more information, see What is Amazon Athena?

You can use the AWS Management Console to set up Amazon Athena. You must create at least one database in Athena before you start running queries. For more information about getting started with Athena, see Getting started.

Athena is directly integrated with Data Wrangler. You can write Athena queries without having to leave the Data Wrangler UI.

In addition to writing simple Athena queries in Data Wrangler, you can also use:

- Athena workgroups for query result management. For more information about workgroups, see Managing query results.
- Lifecycle configurations for setting data retention periods. For more information about data retention, see Setting data retention periods.
Query Athena within Data Wrangler

Note
Data Wrangler does not support federated queries.

If you use AWS Lake Formation with Athena, make sure your Lake Formation IAM permissions do not override IAM permissions for the database sagemaker_data_wrangler.

Data Wrangler gives you the ability to either import the entire dataset or sample a portion of it. For Athena, it provides the following sampling options:

- None – Import the entire dataset.
- First K – Sample the first K rows of the dataset, where K is an integer that you specify.
- Randomized – Takes a random sample of a size that you specify.
- Stratified – Takes a stratified random sample. A stratified sample preserves the ratio of values in a column.

The following procedure shows how to import a dataset from Athena into Data Wrangler.

**To import a dataset into Data Wrangler from Athena**

1. Sign into [Amazon SageMaker Console](https://aws.amazon.com/sagemaker/).
2. Choose **Studio**.
3. Choose **Launch app**.
4. From the dropdown list, select **Studio**.
5. Choose the Home icon.
6. Choose **Data**.
7. Choose **Data Wrangler**.
8. Choose **Import data**.
9. Under **Available**, choose **Amazon Athena**.
10. For **Data Catalog**, choose a data catalog.
11. Use the **Database** dropdown list to select the database that you want to query. When you select a database, you can preview all tables in your database using the **Tables** listed under **Details**.
12. (Optional) Choose **Advanced configuration**.

   a. Choose a **Workgroup**.

   b. If your workgroup hasn't enforced the Amazon S3 output location or if you don't use a workgroup, specify a value for **Amazon S3 location of query results**.

   c. (Optional) For **Data retention period**, select the checkbox to set a data retention period and specify the number of days to store the data before it's deleted.

   d. (Optional) By default, Data Wrangler saves the connection. You can choose to deselect the checkbox and not save the connection.

13. For **Sampling**, choose a sampling method. Choose **None** to turn off sampling.

14. Enter your query in the query editor and use the **Run** button to run the query. After a successful query, you can preview your result under the editor.

   **Note**

   Salesforce data uses the `timestamptz` type. If you're querying the timestamp column that you've imported to Athena from Salesforce, cast the data in the column to the `timestamp` type. The following query casts the timestamp column to the correct type.

   ```sql
   # cast column timestamptz_col as timestamp type, and name it as timestamp_col
   select cast(timestamptz_col as timestamp) as timestamp_col from table
   ```

15. To import the results of your query, select **Import**.

After you complete the preceding procedure, the dataset that you've queried and imported appears in the Data Wrangler flow.

By default, Data Wrangler saves the connection settings as a new connection. When you import your data, the query that you've already specified appears as a new connection. The saved connections store information about the Athena workgroups and Amazon S3 buckets that you're using. When you're connecting to the data source again, you can choose the saved connection.
Managing query results

Data Wrangler supports using Athena workgroups to manage the query results within an AWS account. You can specify an Amazon S3 output location for each workgroup. You can also specify whether the output of the query can go to different Amazon S3 locations. For more information, see Using Workgroups to Control Query Access and Costs.

Your workgroup might be configured to enforce the Amazon S3 query output location. You can't change the output location of the query results for those workgroups.

If you don't use a workgroup or specify an output location for your queries, Data Wrangler uses the default Amazon S3 bucket in the same AWS Region in which your Studio Classic instance is located to store Athena query results. It creates temporary tables in this database to move the query output to this Amazon S3 bucket. It deletes these tables after data has been imported; however the database, sagemaker_data_wrangler, persists. To learn more, see Imported Data Storage.

To use Athena workgroups, set up the IAM policy that gives access to workgroups. If you're using a SageMaker-Execution-Role, we recommend adding the policy to the role. For more information about IAM policies for workgroups, see IAM policies for accessing workgroups. For example workgroup policies, see Workgroup example policies.

Setting data retention periods

Data Wrangler automatically sets a data retention period for the query results. The results are deleted after the length of the retention period. For example, the default retention period is five days. The results of the query are deleted after five days. This configuration is designed to help you clean up data that you're no longer using. Cleaning up your data prevents unauthorized users from gaining access. It also helps control the costs of storing your data on Amazon S3.

If you don't set a retention period, the Amazon S3 lifecycle configuration determines the duration that the objects are stored. The data retention policy that you've specified for the lifecycle configuration removes any query results that are older than the lifecycle configuration that you've specified. For more information, see Setting lifecycle configuration on a bucket.

Data Wrangler uses Amazon S3 lifecycle configurations to manage data retention and expiration. You must give your Amazon SageMaker Studio Classic IAM execution role permissions to manage bucket lifecycle configurations. Use the following procedure to give permissions.

To give permissions to manage the lifecycle configuration do the following.
1. Sign in to the AWS Management Console and open the IAM console at https://console.aws.amazon.com/iam/.

2. Choose **Roles**.

3. In the search bar, specify the Amazon SageMaker execution role that Amazon SageMaker Studio Classic is using.

4. Choose the role.

5. Choose **Add permissions**.

6. Choose **Create inline policy**.

7. For **Service**, specify **S3** and choose it.

8. Under the **Read** section, choose **GetLifecycleConfiguration**.

9. Under the **Write** section, choose **PutLifecycleConfiguration**.

10. For **Resources**, choose **Specific**.

11. For **Actions**, select the arrow icon next to **Permissions management**.

12. Choose **PutResourcePolicy**.

13. For **Resources**, choose **Specific**.

14. Choose the checkbox next to **Any in this account**.

15. Choose **Review policy**.

16. For **Name**, specify a name.

17. Choose **Create policy**.

**Import data from Amazon Redshift**

Amazon Redshift is a fully managed, petabyte-scale data warehouse service in the cloud. The first step to create a data warehouse is to launch a set of nodes, called an Amazon Redshift cluster. After you provision your cluster, you can upload your dataset and then perform data analysis queries.

You can connect to and query one or more Amazon Redshift clusters in Data Wrangler. To use this import option, you must create at least one cluster in Amazon Redshift. To learn how, see [Getting started with Amazon Redshift](#).

You can output the results of your Amazon Redshift query in one of the following locations:

- The default Amazon S3 bucket
• An Amazon S3 output location that you specify

You can either import the entire dataset or sample a portion of it. For Amazon Redshift, it provides the following sampling options:

• None – Import the entire dataset.
• First K – Sample the first K rows of the dataset, where K is an integer that you specify.
• Randomized – Takes a random sample of a size that you specify.
• Stratified – Takes a stratified random sample. A stratified sample preserves the ratio of values in a column.

The default Amazon S3 bucket is in the same AWS Region in which your Studio Classic instance is located to store Amazon Redshift query results. For more information, see Imported Data Storage.

For either the default Amazon S3 bucket or the bucket that you specify, you have the following encryption options:

• The default AWS service-side encryption with an Amazon S3 managed key (SSE-S3)
• An AWS Key Management Service (AWS KMS) key that you specify

An AWS KMS key is an encryption key that you create and manage. For more information on KMS keys, see AWS Key Management Service.

You can specify an AWS KMS key using either the key ARN or the ARN of your AWS account.

If you use the IAM managed policy, AmazonSageMakerFullAccess, to grant a role permission to use Data Wrangler in Studio Classic, your Database User name must have the prefix sagemaker_access.

Use the following procedures to learn how to add a new cluster.

**Note**

Data Wrangler uses the Amazon Redshift Data API with temporary credentials. To learn more about this API, refer to Using the Amazon Redshift Data API in the Amazon Redshift Management Guide.
To connect to a Amazon Redshift cluster

1. Sign into Amazon SageMaker Console.
2. Choose Studio.
3. Choose Launch app.
4. From the dropdown list, select Studio.
5. Choose the Home icon.
6. Choose Data.
7. Choose Data Wrangler.
8. Choose Import data.
10. Choose Amazon Redshift.
11. Choose Temporary credentials (IAM) for Type.
12. Enter a Connection Name. This is a name used by Data Wrangler to identify this connection.
13. Enter the Cluster Identifier to specify to which cluster you want to connect. Note: Enter only the cluster identifier and not the full endpoint of the Amazon Redshift cluster.
14. Enter the Database Name of the database to which you want to connect.
15. Enter a Database User to identify the user you want to use to connect to the database.
16. For UNLOAD IAM Role, enter the IAM role ARN of the role that the Amazon Redshift cluster should assume to move and write data to Amazon S3. For more information about this role, see Authorizing Amazon Redshift to access other AWS services on your behalf in the Amazon Redshift Management Guide.
17. Choose Connect.
18. (Optional) For Amazon S3 output location, specify the S3 URI to store the query results.
19. (Optional) For KMS key ID, specify the ARN of the AWS KMS key or alias. The following image shows you where you can find either key in the AWS Management Console.
The following image shows all the fields from the preceding procedure.
After your connection is successfully established, it appears as a data source under Data Import. Select this data source to query your database and import data.

**To query and import data from Amazon Redshift**

1. Select the connection that you want to query from **Data Sources**.
2. Select a Schema. To learn more about Amazon Redshift Schemas, see Schemas in the Amazon Redshift Database Developer Guide.

3. (Optional) Under Advanced configuration, specify the Sampling method that you'd like to use.

4. Enter your query in the query editor and choose Run to run the query. After a successful query, you can preview your result under the editor.

5. Select Import dataset to import the dataset that has been queried.

6. Enter a Dataset name. If you add a Dataset name that contains spaces, these spaces are replaced with underscores when your dataset is imported.

7. Choose Add.

To edit a dataset, do the following.

1. Navigate to your Data Wrangler flow.
2. Choose the + next to Source - Sampled.
3. Change the data that you're importing.
4. Choose Apply

**Import data from Amazon EMR**

You can use Amazon EMR as a data source for your Amazon SageMaker Data Wrangler flow. Amazon EMR is a managed cluster platform that you can use process and analyze large amounts of data. For more information about Amazon EMR, see What is Amazon EMR? To import a dataset from EMR, you connect to it and query it.

⚠️ **Important**

You must meet the following prerequisites to connect to an Amazon EMR cluster:

**Prerequisites**

- **Network configurations**
  - You have an Amazon VPC in the Region that you're using to launch Amazon SageMaker Studio Classic and Amazon EMR.
• Both Amazon EMR and Amazon SageMaker Studio Classic must be launched in private subnets. They can be in the same subnet or in different ones.

• Amazon SageMaker Studio Classic must be in VPC-only mode.

For more information about creating a VPC, see Create a VPC.

For more information about creating a VPC, see Connect SageMaker Studio Classic Notebooks in a VPC to External Resources.

• The Amazon EMR clusters that you're running must be in the same Amazon VPC.

• The Amazon EMR clusters and the Amazon VPC must be in the same AWS account.

• Your Amazon EMR clusters are running Hive or Presto.
  • Hive clusters must allow inbound traffic from Studio Classic security groups on port 10000.
  • Presto clusters must allow inbound traffic from Studio Classic security groups on port 8889.

  ⚠️ Note
  The port number is different for Amazon EMR clusters using IAM roles.
  Navigate to the end of the prerequisites section for more information.

• SageMaker Studio Classic

• Amazon SageMaker Studio Classic must run Jupyter Lab Version 3. For information about updating the Jupyter Lab Version, see View and update the JupyterLab version of an application from the console.

• Amazon SageMaker Studio Classic has an IAM role that controls user access. The default IAM role that you're using to run Amazon SageMaker Studio Classic doesn't have policies that can give you access to Amazon EMR clusters. You must attach the policy granting permissions to the IAM role. For more information, see Configure the discoverability of Amazon EMR clusters (for administrators).

• The IAM role must also have the following policy attached secretsmanager:PutResourcePolicy.

• If you're using a Studio Classic domain that you've already created, make sure that its AppNetworkAccessType is in VPC-only mode. For information about updating a domain to use VPC-only mode, see Shut down and Update SageMaker Studio Classic.
• Amazon EMR clusters
  • You must have Hive or Presto installed on your cluster.
  • The Amazon EMR release must be version 5.5.0 or later.

**Note**
Amazon EMR supports auto termination. Auto termination stops idle clusters from running and prevents you from incurring costs. The following are the releases that support auto termination:
  • For 6.x releases, version 6.1.0 or later.
  • For 5.x releases, version 5.30.0 or later.

• Amazon EMR clusters using IAM runtime roles
  • Use the following pages to set up IAM runtime roles for the Amazon EMR cluster. You must enable in-transit encryption when you're using runtime roles:
    • Prerequisites for launching an Amazon EMR cluster with a runtime role
    • Launch an Amazon EMR cluster with role-based access control
  • You must Lake Formation as a governance tool for the data within your databases. You must also use external data filtering for access control.
    • For more information about Lake Formation, see What is AWS Lake Formation?
    • For more information about integrating Lake Formation into Amazon EMR, see Integrating third-party services with Lake Formation.
  • The version of your cluster must be 6.9.0 or later.
  • Access to AWS Secrets Manager. For more information about Secrets Manager see What is AWS Secrets Manager?
  • Hive clusters must allow inbound traffic from Studio Classic security groups on port 10000.

An Amazon VPC is a virtual network that is logically isolated from other networks on the AWS cloud. Amazon SageMaker Studio Classic and your Amazon EMR cluster only exist within the Amazon VPC.

Use the following procedure to launch Amazon SageMaker Studio Classic in an Amazon VPC.
To launch Studio Classic within a VPC, do the following.

2. Choose **Launch SageMaker Studio Classic**.
3. Choose **Standard setup**.
4. For **Default execution role**, choose the IAM role to set up Studio Classic.
5. Choose the VPC where you've launched the Amazon EMR clusters.
6. For **Subnet**, choose a private subnet.
7. For **Security group(s)**, specify the security groups that you're using to control between your VPC.
8. Choose **VPC Only**.
9. (Optional) AWS uses a default encryption key. You can specify an AWS Key Management Service key to encrypt your data.
10. Choose **Next**.
11. Under **Studio settings**, choose the configurations that are best suited to you.
12. Choose **Next** to skip the SageMaker Canvas settings.
13. Choose **Next** to skip the RStudio settings.

If you don't have an Amazon EMR cluster ready, you can use the following procedure to create one. For more information about Amazon EMR, see [What is Amazon EMR?](https://docs.aws.amazon.com/elasticmapreduce/latest/EMR-release-note/emr-release-note.html)

To create a cluster, do the following.

1. Navigate to the AWS Management Console.
2. In the search bar, specify **Amazon EMR**.
3. Choose **Create cluster**.
4. For **Cluster name**, specify the name of your cluster.
5. For **Release**, select the release version of the cluster.

### Note
Amazon EMR supports auto termination for the following releases:

- For 6.x releases, releases 6.1.0 or later
• For 5.x releases, releases 5.30.0 or later

Auto termination stops idle clusters from running and prevents you from incurring costs.

6. (Optional) For Applications, choose Presto.

7. Choose the application that you're running on the cluster.

8. Under Networking, for Hardware configuration, specify the hardware configuration settings.

**Important**
For Networking, choose the VPC that is running Amazon SageMaker Studio Classic and choose a private subnet.

9. Under Security and access, specify the security settings.

10. Choose Create.

For a tutorial about creating an Amazon EMR cluster, see Getting started with Amazon EMR. For information about best practices for configuring a cluster, see Considerations and best practices.

**Note**
For security best practices, Data Wrangler can only connect to VPCs on private subnets. You can’t connect to the master node unless you use AWS Systems Manager for your Amazon EMR instances. For more information, see Securing access to EMR clusters using AWS Systems Manager.

You can currently use the following methods to access an Amazon EMR cluster:

• No authentication
• Lightweight Directory Access Protocol (LDAP)
• IAM (Runtime role)

Not using authentication or using LDAP can require you to create multiple clusters and Amazon EC2 instance profiles. If you're an administrator, you might need to provide groups of users with
different levels of access to the data. These methods can result in administrative overhead that makes it more difficult to manage your users.

We recommend using an IAM runtime role that gives multiple users the ability to connect to the same Amazon EMR cluster. A runtime role is an IAM role that you can assign to a user who is connecting to an Amazon EMR cluster. You can configure the runtime IAM role to have permissions that are specific to each group of users.

Use the following sections to create a Presto or Hive Amazon EMR cluster with LDAP activated.

Presto

⚠️ Important
To use AWS Glue as a metastore for Presto tables, select **Use** for **Presto table metadata** to store the results of your Amazon EMR queries in a AWS Glue data catalog when you're launching an EMR cluster. Storing the query results in a AWS Glue data catalog can save you from incurring charges.

To query large datasets on Amazon EMR clusters, you must add the following properties to the Presto configuration file on your Amazon EMR clusters:

```json
["classification":"presto-config","properties":{
  "http-server.max-request-header-size":"5MB",
  "http-server.max-response-header-size":"5MB"
}]
```

You can also modify the configuration settings when you launch the Amazon EMR cluster.

The configuration file for your Amazon EMR cluster is located under the following path: `/etc/presto/conf/config.properties`.

Use the following procedure to create a Presto cluster with LDAP activated.

To create a cluster, do the following.

1. Navigate to the AWS Management Console.
2. In the search bar, specify **Amazon EMR**.
3. Choose **Create cluster**.

4. For **Cluster name**, specify the name of your cluster.

5. For **Release**, select the release version of the cluster.

   ![Note]

   **Note**

   Amazon EMR supports auto termination for the following releases:

   - For 6.x releases, releases 6.1.0 or later
   - For 5.x releases, releases 5.30.0 or later

   Auto termination stops idle clusters from running and prevent you from incurring costs.

6. Choose the application that you're running on the cluster.

7. Under **Networking**, for **Hardware configuration**, specify the hardware configuration settings.

   ![Important]

   **Important**

   For **Networking**, choose the VPC that is running Amazon SageMaker Studio Classic and choose a private subnet.

8. Under **Security and access**, specify the security settings.

9. Choose **Create**.

Hive

![Important]

**Important**

To use AWS Glue as a metastore for Hive tables, select **Use** for **Hive table metadata** to store the results of your Amazon EMR queries in a AWS Glue data catalog when you're launching an EMR cluster. Storing the query results in a AWS Glue data catalog can save you from incurring charges.

To be able to query large datasets on Amazon EMR clusters, add the following properties to Hive configuration file on your Amazon EMR clusters:
You can also modify the configuration settings when you launch the Amazon EMR cluster. The configuration file for your Amazon EMR cluster is located under the following path: /etc/hive/conf/hive-site.xml. You can specify the following property and restart the cluster:

```xml
<property>
  <name>hive.resultset.use.unique.column.names</name>
  <value>false</value>
</property>
```

Use the following procedure to create a Hive cluster with LDAP activated.

To create a Hive cluster with LDAP activated, do the following.

1. Navigate to the AWS Management Console.
2. In the search bar, specify Amazon EMR.
3. Choose Create cluster.
4. Choose Go to advanced options.
5. For Release, select an Amazon EMR release version.
6. The Hive configuration option is selected by default. Make sure the Hive option has a checkbox next to it.
7. (Optional) You can also select Presto as a configuration option to activate both Hive and Presto on your cluster.
8. (Optional) Select Use for Hive table metadata to store the results of your Amazon EMR queries in a AWS Glue data catalog. Storing the query results in a AWS Glue catalog can save you from incurring charges. For more information, see Using the AWS Glue Data Catalog as the metastore for Hive.
9. **Under Enter configuration**, specify the following JSON:

```json
[
  {
    "classification": "hive-site",
    "properties": {
      "hive.server2.authentication.ldap.baseDN": "dc=example,dc=org",
      "hive.server2.authentication": "LDAP",
      "hive.server2.authentication.ldap.url": "ldap://ldap-server-dns-name:389"
    }
  }
]
```

10. **Note**

As a security best practice, we recommend enabling SSL for HiveServer by adding a few properties in the preceding hive-site JSON. For more information, see [Enable SSL on HiveServer2](#).

Use the following sections to use LDAP authentication for Amazon EMR clusters that you've already created.

**LDAP for Presto**

Using LDAP on a cluster running Presto requires access to the Presto coordinator through HTTPS. Do the following to provide access:

- Activate access on port 636
- Enable SSL for the Presto coordinator
Use the following template to configure Presto:

- Classification: presto-config
  ConfigurationProperties:
  - http-server.authentication.type: 'PASSWORD'
  - http-server.https.enabled: 'true'
  - http-server.https.port: '8889'
  - http-server.http.port: '8899'
  - node-scheduler.include-coordinator: 'true'
  - http-server.https.keystore.path: '/path/to/keystore/path/for/presto'
  - http-server.https.keystore.key: 'keystore-key-password'
  - discovery.uri: 'http://master-node-dns-name:8899'

- Classification: presto-password-authenticator
  ConfigurationProperties:
  - password-authenticator.name: 'ldap'
  - ldap.url: !Sub 'ldaps://ldap-server-dns-name:636'
  - ldap.user-bind-pattern: "uid=${USER},dc=example,dc=org"
  - internal-communication.authentication.ldap.user: "ldap-user-name"
  - internal-communication.authentication.ldap.password: "ldap-password"

For information about setting up LDAP in Presto, see the following resources:

- [LDAP Authentication](#)
- [Using LDAP Authentication for Presto on Amazon EMR](#)

**Note**

As a security best practice, we recommend enabling SSL for Presto. For more information, see [Secure Internal Communication](#).

**LDAP for Hive**

To use LDAP for Hive for a cluster that you've created, use the following procedure [Reconfigure an instance group in the console](#).

You're specifying the name of the cluster to which you're connecting.
Use the following procedure to import data from a cluster.

To import data from a cluster, do the following.

1. Open a Data Wrangler flow.
2. Choose **Create Connection**.
3. Choose **Amazon EMR**.
4. Do one of the following.
   - (Optional) For **Secrets ARN**, specify the Amazon Resource Number (ARN) of the database within the cluster. Secrets provide additional security. For more information about secrets, see [What is AWS Secrets Manager?](#) For information about creating a secret for your cluster, see [Creating a AWS Secrets Manager secret for your cluster](#).
   
     | Important |
     |-----------|
     | You must specify a secret if you’re using an IAM runtime role for authentication. |
   
   - From the dropdown table, choose a cluster.
5. Choose **Next**.
6. For **Select an endpoint for example-cluster-name cluster**, choose a query engine.
7. (Optional) Select **Save connection**.
8. Choose **Next, select login** and choose one of the following:
   - No authentication
- LDAP
- IAM

9. For **Login** into `example-cluster-name` **cluster**, specify the **Username** and **Password** for the cluster.

10. Choose **Connect**.

11. In the query editor specify a SQL query.

12. Choose **Run**.

13. Choose **Import**.

**Creating a AWS Secrets Manager secret for your cluster**

If you're using an IAM runtime role to access your Amazon EMR cluster, you must store the credentials that you're using to access the Amazon EMR as a Secrets Manager secret. You store all the credentials that you use to access the cluster within the secret.

You must store the following information in the secret:

- JDBC endpoint – `jdbc:hive2://`
- DNS name – The DNS name of your Amazon EMR cluster. It's either the endpoint for the primary node or the hostname.
- Port – 8446

You can also store the following additional information within the secret:

- IAM role – The IAM role that you're using to access the cluster. Data Wrangler uses your SageMaker execution role by default.
- Truststore path – By default, Data Wrangler creates a truststore path for you. You can also use your own truststore path. For more information about truststore paths, see [In-transit encryption in HiveServer2](#).
- Truststore password – By default, Data Wrangler creates a truststore password for you. You can also use your own truststore path. For more information about truststore paths, see [In-transit encryption in HiveServer2](#).

Use the following procedure to store the credentials within a Secrets Manager secret.
To store your credentials as a secret, do the following.

1. Navigate to the AWS Management Console.
2. In the search bar, specify Secrets Manager.
3. Choose **AWS Secrets Manager**.
4. Choose **Store a new secret**.
5. For **Secret type**, choose **Other type of secret**.
6. Under **Key/value** pairs, select **Plaintext**.
7. For clusters running Hive, you can use the following template for IAM authentication.

   ```json
   {
     "jdbcURL": "",
     "iam_auth": {
       "endpoint": "jdbc:hive2://", #required
       "dns": "ip-xx-xxx-xxx-xxx.ec2.internal", #required
       "port": "10000", #required
       "cluster_id": "j-yyyyyyyyy", #required
       "iam_role": "arn:aws:iam::xxxxxx:role/xxxxxxxxxxxxx", #optional
       "truststore_path": "/etc/alternatives/jre/lib/security/cacerts",
       #optional
       "truststore_password": "changeit" #optional
     }
   }
   ```

**Note**

After you import your data, you apply transformations to them. You then export the data that you've transformed to a specific location. If you're using a Jupyter notebook to export your transformed data to Amazon S3, you must use the truststore path specified in the preceding example.

A Secrets Manager secret stores the JDBC URL of the Amazon EMR cluster as a secret. Using a secret is more secure than directly entering in your credentials.

Use the following procedure to store the JDBC URL as a secret.

To store the JDBC URL as a secret, do the following.
1. Navigate to the AWS Management Console.
2. In the search bar, specify Secrets Manager.
3. Choose **AWS Secrets Manager**.
4. Choose **Store a new secret**.
5. For **Secret type**, choose **Other type of secret**.
6. For **Key/value pairs**, specify `jdbcURL` as the key and a valid JDBC URL as the value.

The format of a valid JDBC URL depends on whether you use authentication and whether you use Hive or Presto as the query engine. The following list shows the valid JDBC URL formats for the different possible configurations.

- **Hive, LDAP authentication** – `jdbc:hive2://emr-cluster-master-public-dns-name:10000/;AuthMech=3;UID=david;PWD=welcome123`;
- For **Hive with SSL enabled**, the JDBC URL format depends on whether you use a Java Keystore File for the TLS configuration. The Java Keystore File helps verify the identity of the master node of the Amazon EMR cluster. To use a Java Keystore File, generate it on an EMR cluster and upload it to Data Wrangler. To generate a file, use the following command on the Amazon EMR cluster, `keytool -genkey -alias hive -keyalg RSA -keysize 1024 -keystore hive.jks`. For information about running commands on an Amazon EMR cluster, see [Securing access to EMR clusters using AWS Systems Manager](https://docs.aws.amazon.com/sagemaker/latest/dg/secure-emr-ssl.html). To upload a file, choose the upward arrow on the left-hand navigation of the Data Wrangler UI.

The following are the valid JDBC URL formats for Hive with SSL enabled:

- **Without a Java Keystore File** – `jdbc:hive2://emr-cluster-master-public-dns:10000/;AuthMech=3;UID=**user-name**;PWD=**password**;SSL=1;AllowSelfSignedCerts=1`;
- **With a Java Keystore File** – `jdbc:hive2://emr-cluster-master-public-dns:10000/;AuthMech=3;UID=**user-name**;PWD=**password**;SSL=1;SSLKeyStore=/home/sagemaker-user/data/Java-keystore-file-name;SSLKeyStorePw=Java-keystore-file-passsword`;
- For **Presto with LDAP authentication and SSL enabled**, the JDBC URL format depends on whether you use a Java Keystore File for the TLS configuration. The Java Keystore File helps verify the identity of the master node of the Amazon EMR cluster. To use a Java Keystore...
File, generate it on an EMR cluster and upload it to Data Wrangler. To upload a file, choose the upward arrow on the left-hand navigation of the Data Wrangler UI. For information about creating a Java Keystore File for Presto, see [Java Keystore File for TLS](#). For information about running commands on an Amazon EMR cluster, see [Securing access to EMR clusters using AWS Systems Manager](#).

- Without a Java Keystore File – `jdbc:presto://emr-cluster-master-public-dns:8889;SSL=1;AuthenticationType=LDAP Authentication;UID=user-name;PWD=password;AllowSelfSignedServerCert=1;AllowHostNameCNMismatch=1;`

- With a Java Keystore File – `jdbc:presto://emr-cluster-master-public-dns:8889;SSL=1;AuthenticationType=LDAP Authentication;SSLTrustStorePath=/home/sagemaker-user/data/Java-keystore-file-name;SSLTrustStorePwd=Java-keystore-file-password;UID=user-name;PWD=password;`

Throughout the process of importing data from an Amazon EMR cluster, you might run into issues. For information about troubleshooting them, see [Troubleshooting issues with Amazon EMR](#).

### Import data from Databricks (JDBC)

You can use Databricks as a data source for your Amazon SageMaker Data Wrangler flow. To import a dataset from Databricks, use the JDBC (Java Database Connectivity) import functionality to access to your Databricks database. After you access the database, specify a SQL query to get the data and import it.

We assume that you have a running Databricks cluster and that you've configured your JDBC driver to it. For more information, see the following Databricks documentation pages:

- [JDBC driver](#)
- [JDBC configuration and connection parameters](#)
- [Authentication parameters](#)

Data Wrangler stores your JDBC URL in AWS Secrets Manager. You must give your Amazon SageMaker Studio Classic IAM execution role permissions to use Secrets Manager. Use the following procedure to give permissions.

To give permissions to Secrets Manager, do the following.
1. Sign in to the AWS Management Console and open the IAM console at https://console.aws.amazon.com/iam/.

2. Choose Roles.

3. In the search bar, specify the Amazon SageMaker execution role that Amazon SageMaker Studio Classic is using.

4. Choose the role.

5. Choose Add permissions.

6. Choose Create inline policy.

7. For Service, specify Secrets Manager and choose it.

8. For Actions, select the arrow icon next to Permissions management.


10. For Resources, choose Specific.

11. Choose the checkbox next to Any in this account.


13. For Name, specify a name.

14. Choose Create policy.

You can use partitions to import your data more quickly. Partitions give Data Wrangler the ability to process the data in parallel. By default, Data Wrangler uses 2 partitions. For most use cases, 2 partitions give you near-optimal data processing speeds.

If you choose to specify more than 2 partitions, you can also specify a column to partition the data. The type of the values in the column must be numeric or date.

We recommend using partitions only if you understand the structure of the data and how it's processed.

You can either import the entire dataset or sample a portion of it. For a Databricks database, it provides the following sampling options:

- None – Import the entire dataset.
- First K – Sample the first K rows of the dataset, where K is an integer that you specify.
- Randomized – Takes a random sample of a size that you specify.
- Stratified – Takes a stratified random sample. A stratified sample preserves the ratio of values in a column.
Use the following procedure to import your data from a Databricks database.

To import data from Databricks, do the following.

1. Sign into Amazon SageMaker Console.
2. Choose Studio.
3. Choose Launch app.
4. From the dropdown list, select Studio.
5. From the Import data tab of your Data Wrangler flow, choose Databricks.
6. Specify the following fields:
   - **Dataset name** – A name that you want to use for the dataset in your Data Wrangler flow.
   - **Driver** – `com.simba.spark.jdbc.Driver`.
   - **JDBC URL** – The URL of the Databricks database. The URL formatting can vary between Databricks instances. For information about finding the URL and the specifying the parameters within it, see [JDBC configuration and connection parameters](#). The following is an example of how a URL can be formatted: 
     
     ```
     jdbc:spark://aws-sagemaker-datawrangler.cloud.databricks.com:443/
     default;transportMode=http;httpPath=sql/protocolv1/
     o/3122619508517275/0909-200301-cut318;AuthMech=3;UID=token;PWD=personal-access-token.
     ```
   
   **Note**
   You can specify a secret ARN that contains the JDBC URL instead of specifying the JDBC URL itself. The secret must contain a key-value pair with the following format: `jdbcURL: JDBC-URL`. For more information, see [What is Secrets Manager?](#).

7. Specify a SQL SELECT statement.

   **Note**
   Data Wrangler doesn't support Common Table Expressions (CTE) or temporary tables within a query.

8. For Sampling, choose a sampling method.
10. (Optional) For the **PREVIEW**, choose the gear to open the **Partition settings**.
   The gear for the additional settings is located to the far right of the **PREVIEW** title.

   - Specify the number of partitions. You can partition by column if you specify the number of partitions:
     - **Enter number of partitions** – Specify a value greater than 2.
     - (Optional) **Partition by column** – Specify the following fields. You can only partition by a column if you've specified a value for **Enter number of partitions**.
       - **Select column** – Select the column that you’re using for the data partition. The data type of the column must be numeric or date.
       - **Upper bound** – From the values in the column that you've specified, the upper bound is the value that you're using in the partition. The value that you specify doesn’t change the data that you're importing. It only affects the speed of the import. For the best performance, specify an upper bound that's close to the column's maximum.
       - **Lower bound** – From the values in the column that you've specified, the lower bound is the value that you're using in the partition. The value that you specify doesn't change the data that you're importing. It only affects the speed of the import. For the best performance, specify a lower bound that's close to the column's minimum.

11. Choose **Import**.

**Import data from Salesforce Data Cloud**

You can use Salesforce Data Cloud as a data source in Amazon SageMaker Data Wrangler to prepare the data in your Salesforce Data Cloud for machine learning.

With Salesforce Data Cloud as a data source in Data Wrangler, you can quickly connect to your Salesforce data without writing a single line of code. You can join your Salesforce data with data from any other data source in Data Wrangler.

After you connect to the data cloud, you can do the following:

- Visualize your data with built-in visualizations
- Understand data and identify potential errors and extreme values
- Transform data with more than 300 built-in transformations
- Export the data that you've transformed
## Topics

- Administrator setup
- Data Scientist Guide

### Administrator setup

⚠️ **Important**

Before you get started, make sure that your users are running Amazon SageMaker Studio Classic version 1.3.0 or later. For information about checking the version of Studio Classic and updating it, see [Prepare ML Data with Amazon SageMaker Data Wrangler](#).

When you're setting up access to Salesforce Data Cloud, you must complete the following tasks:

- Getting your Salesforce Domain URL. Salesforce also refers to the Domain URL as your org's URL.
- Getting OAuth credentials from Salesforce.
- Getting the authorization URL and token URL for your Salesforce Domain.
- Creating a AWS Secrets Manager secret with the OAuth configuration.
- Creating a lifecycle configuration that Data Wrangler uses to read the credentials from the secret.
- Giving Data Wrangler permissions to read the secret.

After you perform the preceding tasks, your users can log into the Salesforce Data Cloud using OAuth.

⚠️ **Note**

Your users might run into issues after you've set everything up. For information about troubleshooting, see [Troubleshooting with Salesforce](#).

Use the following procedure to get the Domain URL.

1. Navigate to the Salesforce login page.
After you get the Salesforce Domain URL, you can use the following procedure to get the login credentials from Salesforce and allow Data Wrangler to access your Salesforce data.

To get the login credentials from Salesforce and provide access to Data Wrangler, do the following.

1. Navigate to your Salesforce Domain URL and log into your account.
2. Choose the gear icon.
3. In the search bar that appears, specify App Manager.
5. Specify the following fields:
   - Connected App Name – You can specify any name, but we recommend choosing a name that includes Data Wrangler. For example, you can specify Salesforce Data Cloud Data Wrangler Integration.
   - API name – Use the default value.
   - Contact Email – Specify your email address.
   - Under API heading (Enable OAuth Settings), select the checkbox to activate OAuth settings.
   - For Callback URL specify the Amazon SageMaker Studio Classic URL. To get the URL for Studio Classic, access it from the AWS Management Console and copy the URL.
6. Under Selected OAuth Scopes, move the following from the Available OAuth Scopes to Selected OAuth Scopes:
   - Manage user data via APIs (api)
   - Perform requests at any time (refresh_token, offline_access)
   - Perform ANSI SQL queries on Salesforce Data Cloud data (cdp_query_api)
   - Manage Salesforce Customer Data Platform profile data (cdp_profile_api)
7. Choose Save. After you save your changes, Salesforce opens a new page.
8. Choose Continue
10. Choose **Manage Consumer Details**. Salesforce redirects you to a new page where you might have to pass two-factor authentication.

11. **Important**

   Copy the Consumer Key and Consumer Secret to a text editor. You need this information to connect the data cloud to Data Wrangler.

12. Navigate back to **Manage Connected Apps**.

13. Navigate to **Connected App Name** and the name of your application.

14. Choose **Manage**.

   a. Select **Edit Policies**.
   
   b. Change **IP Relaxation** to **Relax IP restrictions**.

   c. Choose **Save**.

After you provide access to your Salesforce Data Cloud, you need to provide permissions for your users. Use the following procedure to provide them with permissions.

To provide your users with permissions, do the following.

1. Navigate to the setup home page.

2. On the left-hand navigation, search for **Users** and choose the **Users** menu item.

3. Choose the hyperlink with your user name.

4. Navigate to **Permission Set Assignments**.

5. Choose **Edit Assignments**.

6. Add the following permissions:

   - **Customer Data Platform Admin**
   - **Customer Data Platform Data Aware Specialist**

7. Choose **Save**.

After you get the information for your Salesforce Domain, you must get the authorization URL and the token URL for the AWS Secrets Manager secret that you're creating.

Use the following procedure to get the authorization URL and the token URL.
To get the authorization URL and token URL

1. Navigate to your Salesforce Domain URL.

2. Use one of the following methods to get the URLs. If you are on a Linux distribution with curl and jq installed, we recommend using the method that only works on Linux.

   - (Linux only) Specify the following command in your terminal.

     ```
     curl salesforce-domain-URL/.well-known/openid-configuration | \
     jq '. | { authorization_url: .authorization_endpoint,
        token_url: .token_endpoint }' | \
     jq '. += { identity_provider: "SALESFORCE", client_id: "example-client-id",
        client_secret: "example-client-secret" }'
     ```

   - b. Copy the authorization_endpoint and token_endpoint to a text editor.
   - c. Create the following JSON object:

     ```
     {
       "identity_provider": "SALESFORCE",
       "authorization_url": "example-authorization-endpoint",
       "token_url": "example-token-endpoint",
       "client_id": "example-consumer-key",
       "client_secret": "example-consumer-secret"
     }
     ```

After you create the OAuth configuration object, you can create a AWS Secrets Manager secret that stores it. Use the following procedure to create the secret.

To create a secret, do the following.

1. Navigate to the AWS Secrets Manager console.
2. Choose Store a secret.
3. Select Other type of secret.
4. Under Key/value pairs select Plaintext.
5. Replace the empty JSON with the following configuration settings.

```json
{
    "identity_provider": "SALESFORCE",
    "authorization_url": "example-authorization-endpoint",
    "token_url": "example-token-endpoint",
    "client_id": "example-consumer-key",
    "client_secret": "example-consumer-secret"
}
```

6. Choose Next.

7. For Secret Name, specify the name of the secret.

8. Under Tags, choose Add.

   - For the Key, specify sagemaker:partner. For Value, we recommend specifying a value that might be useful for your use case. However, you can specify anything.

   **Important**
   You must create the key. You can't import your data from Salesforce if you don't create it.

9. Choose Next.

10. Choose Store.

11. Choose the secret you've created.

12. Make a note of the following fields:

   - The Amazon Resource Number (ARN) of the secret
   - The name of the secret

After you've created the secret, you must add permissions for Data Wrangler to read the secret. Use the following procedure to add permissions.

To add read permissions for Data Wrangler, do the following.

1. Navigate to the Amazon SageMaker console.

2. Choose domains.
3. Choose the domain that you're using to access Data Wrangler.

4. Choose your **User Profile**.

5. Under **Details**, find the **Execution role**. Its ARN is in the following format:
   `arn:aws:iam::111122223333:role/example-role`. Make a note of the SageMaker execution role. Within the ARN, it's everything after `role/`.

6. Navigate to the **IAM console**.

7. In the **Search IAM** search bar, specify the name of the SageMaker execution role.

8. Choose the role.

9. Choose **Add permissions**.

10. Choose **Create inline policy**.

11. Choose the JSON tab.

12. Specify the following policy within the editor.

   ```json
   {
   "Version": "2012-10-17",
   "Statement": [
   {
   "Effect": "Allow",
   "Action": [
   "secretsmanager:GetSecretValue",
   "secretsmanager:PutSecretValue"
   ],
   "Resource": "arn:aws:secretsmanager:*:*:secret:*",
   "Condition": {
   "ForAnyValue:StringLike": {
   "aws:ResourceTag/sagemaker:partner": "*
   }
   }
   },
   {
   "Effect": "Allow",
   "Action": [
   "secretsmanager:UpdateSecret"
   ],
   "Resource": "arn:aws:secretsmanager:*:*:secret:AmazonSageMaker-*"
   }
   ]
   }
   ```
13. Choose **Review Policy**.
14. For **Name**, specify a name.
15. Choose **Create policy**.

After you've given Data Wrangler permissions to read the secret, you must add a Lifecycle Configuration that uses your Secrets Manager secret to your Amazon SageMaker Studio Classic user profile.

Use the following procedure to create a lifecycle configuration and add it to the Studio Classic profile.

To create a lifecycle configuration and add it to the Studio Classic profile, do the following.

1. Navigate to the [Amazon SageMaker console](https://aws.amazon.com/sagemaker).
2. Choose **domains**.
3. Choose the domain that you're using to access Data Wrangler.
4. Choose your **User Profile**.
5. If you see the following applications, delete them:
   - KernelGateway
   - JupyterKernel

   **Note**
   Deleting the applications updates Studio Classic. It can take a while for the updates to happen.

6. While you're waiting for updates to happen, choose **Lifecycle configurations**.
7. Make sure the page you're on says **Studio Classic Lifecycle configurations**.
8. Choose **Create configuration**.
9. Make sure **Jupyter server app** has been selected.
10. Choose **Next**.
11. For **Name**, specify a name for the configuration.
12. For **Scripts**, specify the following script:

```bash
#!/bin/bash
set -eux

cat > ~/.sfgenie_identity_provider_oauth_config <<EOL
{
   "secret_arn": "secrets-arn-containing-salesforce-credentials"
}
EOL
```

13. Choose **Submit**.

14. On the left hand navigation, choose **domains**.

15. Choose your domain.

16. Choose **Environment**.

17. Under **Lifecycle configurations for personal Studio Classic apps**, choose **Attach**.

18. Select **Existing configuration**.

19. Under **Studio Classic Lifecycle configurations** select the lifecycle configuration that you've created.

20. Choose **Attach to domain**.

21. Select the checkbox next to the lifecycle configuration that you've attached.

22. Select **Set as default**.

You might run into issues when you set up your lifecycle configuration. For information about debugging them, see [Debug lifecycle configurations](#).

**Data Scientist Guide**

Use the following to connect Salesforce Data Cloud and access your data in Data Wrangler.
Important

Your administrator needs to use the information in the preceding sections to set up Salesforce Data Cloud. If you're running into issues, contact them for troubleshooting help.

To open Studio Classic and check its version, see the following procedure.

1. Use the steps in Prerequisites to access Data Wrangler through Amazon SageMaker Studio Classic.
2. Next to the user you want to use to launch Studio Classic, select Launch app.
3. Choose Studio.

To create a dataset in Data Wrangler with data from the Salesforce Data Cloud

1. Sign into Amazon SageMaker Console.
2. Choose Studio.
3. Choose Launch app.
4. From the dropdown list, select Studio.
5. Choose the Home icon.
6. Choose Data.
7. Choose Data Wrangler.
8. Choose Import data.
10. For Connection name, specify a name for your connection to the Salesforce Data Cloud.
11. For Org URL, specify the organization URL in your Salesforce account. You can get the URL from your administrator.
12. Choose Connect.
13. Specify your credentials to log into Salesforce.

You can begin creating a dataset using data from Salesforce Data Cloud after you've connected to it.
After you select a table, you can write queries and run them. The output of your query shows under **Query results**.

After you have settled on the output of your query, you can then import the output of your query into a Data Wrangler flow to perform data transformations.

After you’ve created a dataset, navigate to the **Data flow** screen to start transforming your data.

**Import data from Snowflake**

You can use Snowflake as a data source in SageMaker Data Wrangler to prepare data in Snowflake for machine learning.

With Snowflake as a data source in Data Wrangler, you can quickly connect to Snowflake without writing a single line of code. You can join your data in Snowflake with data from any other data source in Data Wrangler.

Once connected, you can interactively query data stored in Snowflake, transform data with more than 300 preconfigured data transformations, understand data and identify potential errors and extreme values with a set of robust preconfigured visualization templates, quickly identify inconsistencies in your data preparation workflow, and diagnose issues before models are deployed into production. Finally, you can export your data preparation workflow to Amazon S3 for use with other SageMaker features such as Amazon SageMaker Autopilot, Amazon SageMaker Feature Store and Amazon SageMaker Model Building Pipelines.

You can encrypt the output of your queries using an AWS Key Management Service key that you’ve created. For more information about AWS KMS, see [AWS Key Management Service](https://aws.amazon.com/kms/).

**Topics**

- [Administrator Guide](#)
- [Data Scientist Guide](#)

**Administrator Guide**

⚠️ **Important**

To learn more about granular access control and best practices, see [Security Access Control](#).
This section is for Snowflake administrators who are setting up access to Snowflake from within SageMaker Data Wrangler.

⚠️ Important

You are responsible for managing and monitoring the access control within Snowflake. Data Wrangler does not add a layer of access control with respect to Snowflake. Access control includes the following:

- The data that a user accesses
- (Optional) The storage integration that provides Snowflake the ability to write query results to an Amazon S3 bucket
- The queries that a user can run

(Optional) Configure Snowflake Data Import Permissions

By default, Data Wrangler queries the data in Snowflake without creating a copy of it in an Amazon S3 location. Use the following information if you're configuring a storage integration with Snowflake. Your users can use a storage integration to store their query results in an Amazon S3 location.

Your users might have different levels of access of sensitive data. For optimal data security, provide each user with their own storage integration. Each storage integration should have its own data governance policy.

This feature is currently not available in the opt-in Regions.

Snowflake requires the following permissions on an S3 bucket and directory to be able to access files in the directory:

- s3:GetObject
- s3:GetObjectVersion
- s3:ListBucket
- s3:ListObjects
- s3:GetBucketLocation

Create an IAM policy
You must create an IAM policy to configure access permissions for Snowflake to load and unload data from an Amazon S3 bucket.

The following is the JSON policy document that you use to create the policy:

```json
# Example policy for S3 write access
# This needs to be updated
{
"Version": "2012-10-17",
"Statement": [
{
"Effect": "Allow",
"Action": [
"s3:PutObject",
"s3:GetObject",
"s3:GetObjectVersion",
"s3:DeleteObject",
"s3:DeleteObjectVersion"
],
"Resource": "arn:aws:s3:::bucket/prefix/*"
},
{
"Effect": "Allow",
"Action": ["s3:ListBucket"
],
"Resource": "arn:aws:s3:::bucket/",
"Condition": {
"StringLike": {
"s3:prefix": ["prefix/*"]
}
}
]
}
```

For information and procedures about creating policies with policy documents, see Creating IAM policies.

For documentation that provides an overview of using IAM permissions with Snowflake, see the following resources:

- What is IAM?
• **Create the IAM Role in AWS**

• **Create a Cloud Storage Integration in Snowflake**

• **Retrieve the AWS IAM User for your Snowflake Account**

• **Grant the IAM User Permissions to Access Bucket.**

To grant the data scientist's Snowflake role usage permission to the storage integration, you must run
\[
\text{GRANT USAGE ON INTEGRATION integration\_name TO snowflake\_role;}.
\]

- **integration\_name** is the name of your storage integration.
- **snowflake\_role** is the name of the default [Snowflake role](#) given to the data scientist user.

**Setting up Snowflake OAuth Access**

Instead of having your users directly enter their credentials into Data Wrangler, you can have them use an identity provider to access Snowflake. The following are links to the Snowflake documentation for the identity providers that Data Wrangler supports.

• **Azure AD**

• **Okta**

• **Ping Federate**

Use the documentation from the preceding links to set up access to your identity provider. The information and procedures in this section help you understand how to properly use the documentation to access Snowflake within Data Wrangler.

Your identity provider needs to recognize Data Wrangler as an application. Use the following procedure to register Data Wrangler as an application within the identity provider:

1. Select the configuration that starts the process of registering Data Wrangler as an application.
2. Provide the users within the identity provider access to Data Wrangler.
3. Turn on OAuth client authentication by storing the client credentials as an AWS Secrets Manager secret.
4. Specify a redirect URL using the following format: https://domain-ID.studio.AWS Region.sagemaker.aws/jupyter/default/lab
Important

You're specifying the Amazon SageMaker domain ID and AWS Region that you're using to run Data Wrangler.

Important

You must register a URL for each Amazon SageMaker domain and AWS Region where you're running Data Wrangler. Users from a domain and AWS Region that don't have redirect URLs set up for them won't be able to authenticate with the identity provider to access the Snowflake connection.

5. Make sure that the authorization code and refresh token grant types are allowed for the Data Wrangler application.

Within your identity provider, you must set up a server that sends OAuth tokens to Data Wrangler at the user level. The server sends the tokens with Snowflake as the audience.

Snowflake uses the concept of roles that are distinct role the IAM roles used in AWS. You must configure the identity provider to use any role to use the default role associated with the Snowflake account. For example, if a user has systems administrator as the default role in their Snowflake profile, the connection from Data Wrangler to Snowflake uses systems administrator as the role.

Use the following procedure to set up the server.

To set up the server, do the following. You're working within Snowflake for all steps except the last one.

1. Start setting up the server or API.
2. Configure the authorization server to use the authorization code and refresh token grant types.
3. Specify the lifetime of the access token.
4. Set the refresh token idle timeout. The idle timeout is the time that the refresh token expires if it's not used.
If you're scheduling jobs in Data Wrangler, we recommend making the idle timeout time greater than the frequency of the processing job. Otherwise, some processing jobs might fail because the refresh token expired before they could run. When the refresh token expires, the user must re-authenticate by accessing the connection that they've made to Snowflake through Data Wrangler.

5. Specify `session:role-any` as the new scope.

For Azure AD, copy the unique identifier for the scope. Data Wrangler requires you to provide it with the identifier.

6. **Important**
   Within the External OAuth Security Integration for Snowflake, enable `external_oauth_any_role_mode`.

**Important**
Data Wrangler doesn't support rotating refresh tokens. Using rotating refresh tokens might result in access failures or users needing to log in frequently.

**Important**
If the refresh token expires, your users must reauthenticate by accessing the connection that they've made to Snowflake through Data Wrangler.

After you've set up the OAuth provider, you provide Data Wrangler with the information it needs to connect to the provider. You can use the documentation from your identity provider to get values for the following fields:
- Token URL – The URL of the token that the identity provider sends to Data Wrangler.
- Client ID – The ID of the identity provider.
- Client secret – The secret that only the authorization server or API recognizes.
- (Azure AD only) The OAuth scope credentials that you've copied.

You store the fields and values in an AWS Secrets Manager secret and add it to the Amazon SageMaker Studio Classic lifecycle configuration that you're using for Data Wrangler. A Lifecycle Configuration is a shell script. Use it to make the Amazon Resource Name (ARN) of the secret accessible to Data Wrangler. For information about creating secrets see Move hardcoded secrets to AWS Secrets Manager. For information about using lifecycle configurations in Studio Classic, see Use lifecycle configurations with Amazon SageMaker Studio Classic.

**Important**

Before you create a Secrets Manager secret, make sure that the SageMaker execution role that you're using for Amazon SageMaker Studio Classic has permissions to create and update secrets in Secrets Manager. For more information about adding permissions, see Example: Permission to create secrets.

For Okta and Ping Federate, the following is the format of the secret:

```json
{
    "token_url": "https://identityprovider.com/oauth2/example-portion-of-URL-path/v2/token",
    "client_id": "example-client-id",
    "client_secret": "example-client-secret",
    "identity_provider": "OKTA" | "PING_FEDERATE",
    "authorization_url": "https://identityprovider.com/oauth2/example-portion-of-URL-path/v2/authorize"
}
```

For Azure AD, the following is the format of the secret:

```json
{
    "token_url": "https://identityprovider.com/oauth2/example-portion-of-URL-path/v2/token",
    "client_id": "example-client-id",
    "client_secret": "example-client-secret",
    "identity_provider": "AZURE_AD",
    "authorization_url": "https://identityprovider.com/oauth2/example-portion-of-URL-path/v2/authorize"
}
```
You must have a lifecycle configuration that uses the Secrets Manager secret that you've created. You can either create the lifecycle configuration or modify one that has already been created. The configuration must use the following script.

```
#!/bin/bash

set -eux

## Script Body

cat > ~/.snowflake_identity_provider_oauth_config <<EOL
{
    "secret_arn": "example-secret-arn"
}
EOL
```

For information about setting up lifecycle configurations, see [Create and associate a lifecycle configuration](#). When you're going through the process of setting up, do the following:

- Set the application type of the configuration to Jupyter Server.
- Attach the configuration to the Amazon SageMaker domain that has your users.
- Have the configuration run by default. It must run every time a user logs into Studio Classic. Otherwise, the credentials saved in the configuration won't be available to your users when they're using Data Wrangler.
- The lifecycle configuration creates a file with the name, `snowflake_identity_provider_oauth_config` in the user's home folder. The file contains
the Secrets Manager secret. Make sure that it's in the user's home folder every time the Jupyter Server's instance is initialized.

Private Connectivity between Data Wrangler and Snowflake via AWS PrivateLink

This section explains how to use AWS PrivateLink to establish a private connection between Data Wrangler and Snowflake. The steps are explained in the following sections.

Create a VPC

If you do not have a VPC set up, then follow the Create a new VPC instructions to create one.

Once you have a chosen VPC you would like to use for establishing a private connection, provide the following credentials to your Snowflake Administrator to enable AWS PrivateLink:

- VPC ID
- AWS Account ID
- Your corresponding account URL you use to access Snowflake

⚠️ Important

As described in Snowflake's documentation, enabling your Snowflake account can take up to two business days.

Set up Snowflake AWS PrivateLink Integration

After AWS PrivateLink is activated, retrieve the AWS PrivateLink configuration for your Region by running the following command in a Snowflake worksheet. Log into your Snowflake console and enter the following under Worksheets: select SYSTEM$GET_PRIVATELINK_CONFIG();

1. Retrieve the values for the following: privatelink-account-name, privatelink_ocsp-url, privatelink-account-url, and privatelink_ocsp-url from the resulting JSON object. Examples of each value are shown in the following snippet. Store these values for later use.

```plaintext
privatelink-account-name: xxxxxxxx.region.privatelink
privatelink-vpce-id: com.amazonaws.vpce.region.vpce-svc-xxxxxxxxxxxxxxxxxxxxx
```
2. Switch to your AWS Console and navigate to the VPC menu.

3. From the left side panel, choose the **Endpoints** link to navigate to the **VPC Endpoints** setup.

   Once there, choose **Create Endpoint**.

4. Select the radio button for **Find service by name**, as shown in the following screenshot.

   ![Create Endpoint Screen](image)

   **Create Endpoint**

   A VPC endpoint enables you to securely connect your VPC to another service.
   
   There are three types of VPC endpoints - Interface endpoints, Gateway Load Balancer endpoints, and gateway endpoints. Interface endpoints and Gateway Load Balancer endpoints are powered by AWS PrivateLink, and use an elastic network interface (ENI) as an entry point for traffic destined to the service.
   
   Interface endpoints are typically accessed using the public or private DNS name associated with the service, while gateway endpoints and Gateway Load Balancer endpoints serve as a target for a route in your route table for traffic destined for the service.

   **Service category**
   
   - AWS services
     - Find service by name
     - Your AWS Marketplace services

   **Service Name**
   
   Enter private service name and verify.

   ![Service Name Input](image)

5. In the **Service Name** field, paste in the value for `privatelink-vpce-id` that you retrieved in the preceding step and choose **Verify**.

   If the connection is successful, a green alert saying **Service name found** appears on your screen and the **VPC** and **Subnet** options automatically expand, as shown in the following screenshot.

   Depending on your targeted Region, your resulting screen may show another AWS Region name.
6. Select the same VPC ID that you sent to Snowflake from the VPC dropdown list.

7. If you have not yet created a subnet, then perform the following set of instructions on creating a subnet.

8. Select Subnets from the VPC dropdown list. Then select Create subnet and follow the prompts to create a subnet in your VPC. Ensure you select the VPC ID you sent Snowflake.


10. Provide a name for the new security group (such as datawrangler-doc-snowflake-privatelink-connection) and a description. Be sure to select the VPC ID you have used in previous steps.

11. Add two rules to allow traffic from within your VPC to this VPC endpoint.

   Navigate to your VPC under Your VPCs in a separate tab, and retrieve your CIDR block for your VPC. Then choose Add Rule in the Inbound Rules section. Select HTTPS for the type, leave the Source as Custom in the form, and paste in the value retrieved from the preceding describe-vpcs call (such as 10.0.0.0/16).

12. Choose Create Security Group. Retrieve the Security Group ID from the newly created security group (such as sg-xxxxxxxxxxxxxxxxxxx).
13. In the VPC Endpoint configuration screen, remove the default security group. Paste in the security group ID in the search field and select the checkbox.


15. If the endpoint creation is successful, you see a page that has a link to your VPC endpoint configuration, specified by the VPC ID. Select the link to view the configuration in full.

Retrieve the topmost record in the DNS names list. This can be differentiated from other DNS names because it only includes the Region name (such as us-west-2), and no Availability Zone letter notation (such as us-west-2a). Store this information for later use.

Configure DNS for Snowflake Endpoints in your VPC

This section explains how to configure DNS for Snowflake endpoints in your VPC. This allows your VPC to resolve requests to the Snowflake AWS PrivateLink endpoint.

1. Navigate to the Route 53 menu within your AWS console.
2. Select the Hosted Zones option (if necessary, expand the left-hand menu to find this option).
3. Choose Create Hosted Zone.
a. In the **Domain name** field, reference the value that was stored for `privatellink-account-url` in the preceding steps. In this field, your Snowflake account ID is removed from the DNS name and only uses the value starting with the Region identifier. A **Resource Record Set** is also created later for the subdomain, such as, `region.privatellink.snowflakecomputing.com`.

b. Select the radio button for **Private Hosted Zone** in the **Type** section. Your Region code may not be `us-west-2`. Reference the DNS name returned to you by Snowflake.

c. In the **VPCs to associate with the hosted zone** section, select the Region in which your VPC is located and the VPC ID used in previous steps.
d. Choose **Create hosted zone**.

4. Next, create two records, one for `privatelink-account-url` and one for `privatelink_ocsp-url`.

   - In the **Hosted Zone** menu, choose **Create Record Set**.
     
     a. Under **Record name**, enter your Snowflake Account ID only (the first 8 characters in `privatelink-account-url`).

     b. Under **Record type**, select **CNAME**.

     c. Under **Value**, enter the DNS name for the regional VPC endpoint you retrieved in the last step of the **Set up the Snowflake AWS PrivateLink Integration** section.
d. Choose Create records.
e. Repeat the preceding steps for the OCSP record we notated as privatelink-ocsp-url, starting with ocsp through the 8-character Snowflake ID for the record name (such as ocsp.xxxxxxxx).

Configure Route 53 Resolver Inbound Endpoint for your VPC

This section explains how to configure Route 53 resolvers inbound endpoints for your VPC.

1. Navigate to the Route 53 menu within your AWS console.
   - In the left hand panel in the Security section, select the Security Groups option.
   - Provide a name for your security group (such as datawranger-doc-route53-resolver-sg) and a description.
   - Select the VPC ID used in previous steps.
   - Create rules that allow for DNS over UDP and TCP from within the VPC CIDR block.
Choose **Create Security Group**. Note the **Security Group ID** because adds a rule to allow traffic to the VPC endpoint security group.

3. Navigate to the **Route 53 menu** within your AWS console.
   - In the **Resolver** section, select the **Inbound Endpoint** option.

4. Choose **Create Inbound Endpoint**.
   - Provide an endpoint name.
   - From the **VPC in the Region** dropdown list, select the VPC ID you have used in all previous steps.
   - In the **Security group for this endpoint** dropdown list, select the security group ID from Step 2 in this section.

   ![General settings for inbound endpoint](image)

   - In the **IP Address** section, select an Availability Zones, select a subnet, and leave the radio selector for **Use an IP address that is selected automatically** selected for each IP address.
5. Select the **Inbound endpoint** after it has been created.

6. Once the inbound endpoint is created, note the two IP addresses for the resolvers.
This section explains how to create VPC endpoints for the following: Amazon SageMaker Studio Classic, SageMaker Notebooks, the SageMaker API, SageMaker Runtime Runtime, and Amazon SageMaker Feature Store Runtime.

Create a security group that is applied to all endpoints.

1. Navigate to the **EC2 menu** in the AWS Console.
2. In the **Network & Security** section, select the **Security groups** option.
3. Choose **Create security group**.
4. Provide a security group name and description (such as `datawrangler-doc-sagemaker-vpce-sg`). A rule is added later to allow traffic over HTTPS from SageMaker to this group.

Creating the endpoints

1. Navigate to the **VPC menu** in the AWS console.
2. Select the **Endpoints** option.
3. Choose **Create Endpoint**.
4. Search for the service by entering its name in the **Search** field.
5. From the **VPC** dropdown list, select the VPC in which your Snowflake AWS PrivateLink connection exists.
6. In the **Subnets** section, select the subnets which have access to the Snowflake PrivateLink connection.
7. Leave the **Enable DNS Name** checkbox selected.
8. In the **Security Groups** section, select the security group you created in the preceding section.
9. Choose **Create Endpoint**.
Configure Studio Classic and Data Wrangler

This section explains how to configure Studio Classic and Data Wrangler.

1. Configure the security group.
   a. Navigate to the Amazon EC2 menu in the AWS Console.
   b. Select the **Security Groups** option in the **Network & Security** section.
   c. Choose **Create Security Group**.
   d. Provide a name and description for your security group (such as datawrangler-doc-sagemaker-studio).
   e. Create the following inbound rules.
      * The HTTPS connection to the security group you provisioned for the Snowflake PrivateLink connection you created in the **Set up the Snowflake PrivateLink Integration** step.
      * The HTTP connection to the security group you provisioned for the Snowflake PrivateLink connection you created in the **Set up the Snowflake PrivateLink Integration step**.
      * The UDP and TCP for DNS (port 53) to Route 53 Resolver Inbound Endpoint security group you create in step 2 of **Configure Route 53 Resolver Inbound Endpoint for your VPC**.
   f. Choose **Create Security Group** button in the lower right hand corner.

2. Configure Studio Classic.
   • Navigate to the SageMaker menu in the AWS console.
   • From the left hand console, Select the **SageMaker Studio Classic** option.
   • If you do not have any domains configured, the **Get Started** menu is present.
   • Select the **Standard Setup** option from the **Get Started** menu.
   • Under **Authentication method**, select **AWS Identity and Access Management (IAM)**.
   • From the **Permissions** menu, you can create a new role or use a pre-existing role, depending on your use case.
      * If you choose **Create a new role**, you are presented the option to provide an S3 bucket name, and a policy is generated for you.
      * If you already have a role created with permissions for the S3 buckets to which you require access, select the role from the dropdown list. This role should have the **AmazonSageMakerFullAccess** policy attached to it.
   • Select the **Network and Storage** dropdown list to configure the VPC, security, and subnets SageMaker uses.
• Under **VPC**, select the VPC in which your Snowflake PrivateLink connection exists.

• Under **Subnet(s)**, select the subnets which have access to the Snowflake PrivateLink connection.

• Under **Network Access for Studio Classic**, select **VPC Only**.

• Under **Security Group(s)**, select the security group you created in step 1.

• Choose **Submit**.

3. Edit the SageMaker security group.

• Create the following inbound rules:

  • Port 2049 to the inbound and outbound NFS Security Groups created automatically by SageMaker in step 2 (the security group names contain the Studio Classic domain ID).

  • Access to all TCP ports to itself (required for SageMaker for VPC Only).

4. Edit the VPC Endpoint Security Groups:

• Navigate to the Amazon EC2 menu in the AWS console.

• Locate the security group you created in a preceding step.

• Add an inbound rule allowing for HTTPS traffic from the security group created in step 1.

5. Create a user profile.

• From the **SageMaker Studio Classic Control Panel**, choose **Add User**.

• Provide a user name.

• For the **Execution Role**, choose to create a new role or to use a pre-existing role.

  • If you choose **Create a new role**, you are presented the option to provide an Amazon S3 bucket name, and a policy is generated for you.

  • If you already have a role created with permissions to the Amazon S3 buckets to which you require access, select the role from the dropdown list. This role should have the AmazonSageMakerFullAccess policy attached to it.

• Choose **Submit**.

6. Create a data flow (follow the data scientist guide outlined in a preceding section).

• When adding a Snowflake connection, enter the value of `privatelink-account-name` (from the **Set up Snowflake PrivateLink Integration** step) into the **Snowflake account name (alphanumeric)** field, instead of the plain Snowflake account name. Everything else is left unchanged.
Provide information to the data scientist

Provide the data scientist with the information that they need to access Snowflake from Amazon SageMaker Data Wrangler.

⚠️ Important

Your users need to run Amazon SageMaker Studio Classic version 1.3.0 or later. For information about checking the version of Studio Classic and updating it, see Prepare ML Data with Amazon SageMaker Data Wrangler.

1. To allow your data scientist to access Snowflake from SageMaker Data Wrangler, provide them with one of the following:

   - For Basic Authentication, a Snowflake account name, user name, and password.
   - For OAuth, a user name and password in the identity provider.
   - For ARN, the Secrets Manager secret Amazon Resource Name (ARN).
   - A secret created with AWS Secrets Manager and the ARN of the secret. Use the following procedure below to create the secret for Snowflake if you choose this option.

⚠️ Important

If your data scientists use the Snowflake Credentials (User name and Password) option to connect to Snowflake, you can use Secrets Manager to store the credentials in a secret. Secrets Manager rotates secrets as part of a best practice security plan. The secret created in Secrets Manager is only accessible with the Studio Classic role configured when you set up a Studio Classic user profile. This requires you to add this permission, secretsmanager:PutResourcePolicy, to the policy that is attached to your Studio Classic role.

We strongly recommend that you scope the role policy to use different roles for different groups of Studio Classic users. You can add additional resource-based permissions for the Secrets Manager secrets. See Manage Secret Policy for condition keys you can use.

For information about creating a secret, see Create a secret. You're charged for the secrets that you create.
2. (Optional) Provide the data scientist with the name of the storage integration that you created using the following procedure Create a Cloud Storage Integration in Snowflake. This is the name of the new integration and is called integration_name in the CREATE INTEGRATION SQL command you ran, which is shown in the following snippet:

```
CREATE STORAGE INTEGRATION integration_name
TYPE = EXTERNAL_STAGE
STORAGE_PROVIDER = S3
ENABLED = TRUE
STORAGE_AWS_ROLE_ARN = 'iam_role'
[ STORAGE_AWS_OBJECT_ACL = 'bucket-owner-full-control' ]
STORAGE_ALLOWED_LOCATIONS = ('s3://bucket/path/','s3://bucket/path/')
[ STORAGE_BLOCKED_LOCATIONS = ('s3://bucket/path/','s3://bucket/path/') ]
```

---

### Data Scientist Guide

Use the following to connect Snowflake and access your data in Data Wrangler.

⚠️ Important

Your administrator needs to use the information in the preceding sections to set up Snowflake. If you're running into issues, contact them for troubleshooting help.

---

You can connect to Snowflake in one of the following ways:

- Specifying your Snowflake credentials (account name, user name, and password) in Data Wrangler.
- Providing an Amazon Resource Name (ARN) of a secret containing the credentials.
- Using an open standard for access delegation (OAuth) provider that connects to Snowflake. Your administrator can give you access to one of the following OAuth providers:
  - Azure AD
  - Okta
  - Ping Federate
Talk to your administrator about the method that you need to use to connect to Snowflake.

The following sections have information about how you can connect to Snowflake using the preceding methods.

Specifying your Snowflake Credentials

To import a dataset into Data Wrangler from Snowflake using your credentials

1. Sign into Amazon SageMaker Console.
2. Choose Studio.
3. Choose Launch app.
4. From the dropdown list, select Studio.
5. Choose the Home icon.
6. Choose Data.
7. Choose Data Wrangler.
8. Choose Import data.
10. For Connection name, specify a name that uniquely identifies the connection.
11. For Authentication method, choose Basic Username-Password.
12. For Snowflake account name (alphanumeric), specify the full name of the Snowflake account.
13. For Username, specify the username that you use to access the Snowflake account.
14. For Password, specify the password associated with the username.
15. (Optional) For Advanced settings, specify the following:
   - Role – A role within Snowflake. Some roles have access to different datasets. If you don't specify a role, Data Wrangler uses the default role in your Snowflake account.
   - Storage integration – When you specify and run a query, Data Wrangler creates a temporary copy of the query results in memory. To store a permanent copy of the query results, specify the Amazon S3 location for the storage integration. Your administrator provided you with the S3 URI.
   - KMS key ID – A KMS key that you've created. You can specify its ARN to encrypt the output of the Snowflake query. Otherwise, Data Wrangler uses the default encryption.
16. Choose Connect.
Providing an Amazon Resource Name (ARN)

**To import a dataset into Data Wrangler from Snowflake using an ARN**

1. Sign into [Amazon SageMaker Console](https://aws.amazon.com/sagemaker/).
2. Choose **Studio**.
3. Choose **Launch app**.
4. From the dropdown list, select **Studio**.
5. Choose the Home icon.
6. Choose **Data**.
7. Choose **Data Wrangler**.
8. Choose **Import data**.
9. Under **Available**, choose **Snowflake**.
10. For **Connection name**, specify a name that uniquely identifies the connection.
11. For **Authentication method**, choose **ARN**.
12. **Secrets Manager ARN** – The ARN of the AWS Secrets Manager secret used to store the credentials used to connect to Snowflake.
13. (Optional) For **Advanced settings**, specify the following:
   - **Role** – A role within Snowflake. Some roles have access to different datasets. If you don’t specify a role, Data Wrangler uses the default role in your Snowflake account.
   - **Storage integration** – When you specify and run a query, Data Wrangler creates a temporary copy of the query results in memory. To store a permanent copy of the query results, specify the Amazon S3 location for the storage integration. Your administrator provided you with the S3 URI.
   - **KMS key ID** – A KMS key that you’ve created. You can specify its ARN to encrypt the output of the Snowflake query. Otherwise, Data Wrangler uses the default encryption.
14. Choose **Connect**.
Using an OAuth Connection

⚠️ Important

Your administrator customized your Studio Classic environment to provide the functionality you're using to use an OAuth connection. You might need to restart the Jupyter server application to use the functionality.

Use the following procedure to update the Jupyter server application.

1. Within Studio Classic, choose File
2. Choose Shut down.
3. Choose Shut down server.
4. Close the tab or window that you're using to access Studio Classic.
5. From the Amazon SageMaker console, open Studio Classic.

To import a dataset into Data Wrangler from Snowflake using your credentials

1. Sign into Amazon SageMaker Console.
2. Choose Studio.
3. Choose Launch app.
4. From the dropdown list, select Studio.
5. Choose the Home icon.
6. Choose Data.
7. Choose Data Wrangler.
8. Choose Import data.
10. For Connection name, specify a name that uniquely identifies the connection.
11. For Authentication method, choose OAuth.
12. (Optional) For Advanced settings, specify the following:
   - Role – A role within Snowflake. Some roles have access to different datasets. If you don't specify a role, Data Wrangler uses the default role in your Snowflake account.
   - Storage integration – When you specify and run a query, Data Wrangler creates a temporary copy of the query results in memory. To store a permanent copy of the query
results, specify the Amazon S3 location for the storage integration. Your administrator provided you with the S3 URI.

- **KMS key ID** – A KMS key that you've created. You can specify its ARN to encrypt the output of the Snowflake query. Otherwise, Data Wrangler uses the default encryption.

13. Choose **Connect**.

You can begin the process of importing your data from Snowflake after you've connected to it.

Within Data Wrangler, you can view your data warehouses, databases, and schemas, along with the eye icon with which you can preview your table. After you select the **Preview Table** icon, the schema preview of that table is generated. You must select a warehouse before you can preview a table.

⚠️ **Important**

If you're importing a dataset with columns of type `TIMESTAMP_TZ` or `TIMESTAMP_LTZ`, add `::string` to the column names of your query. For more information, see [How To: Unload TIMESTAMP_TZ and TIMESTAMP_LTZ data to a Parquet file](#).

After you select a data warehouse, database and schema, you can now write queries and run them. The output of your query shows under **Query results**.

After you have settled on the output of your query, you can then import the output of your query into a Data Wrangler flow to perform data transformations.

After you've imported your data, navigate to your Data Wrangler flow and start adding transformations to it. For a list of available transforms, see [Transform Data](#).

### Import Data From Software as a Service (SaaS) Platforms

You can use Data Wrangler to import data from more than forty software as a service (SaaS) platforms. To import your data from your SaaS platform, you or your administrator must use Amazon AppFlow to transfer the data from the platform to Amazon S3 or Amazon Redshift. For more information about Amazon AppFlow, see [What is Amazon AppFlow?](#) If you don't need to use Amazon Redshift, we recommend transferring the data to Amazon S3 for a simpler process.

Data Wrangler supports transferring data from the following SaaS platforms:
• Amplitude
• Asana
• Braintree
• CircleCI
• DocuSign Monitor
• Delighted
• Domo
• Datadog
• Dynatrace
• Facebook Ads
• Facebook Page Insights
• Google Ads
• Google Analytics 4
• Google Calendar
• Google Search Console
• GitHub
• GitLab
• Infor Nexus
• Instagram Ads
• Intercom
• JDBC (Sync)
• Jira Cloud
• LinkedIn Ads
• Mailchimp
• Marketo
• Microsoft Dynamics 365
• Microsoft Teams
• Mixpanel
• Okta
• Oracle HCM
The preceding list has links to more information about setting up your data source. You or your administrator can refer to the preceding links after you've read the following information.

When you navigate to the **Import** tab of your Data Wrangler flow, you see data sources under the following sections:

- **Available**
- **Set up data sources**
You can connect to data sources under **Available** without needing additional configuration. You can choose the data source and import your data.

Data sources under **Set up data sources**, require you or your administrator to use Amazon AppFlow to transfer the data from the SaaS platform to Amazon S3 or Amazon Redshift. For information about performing a transfer, see [Using Amazon AppFlow to transfer your data](#).

After you perform the data transfer, the SaaS platform appears as a data source under **Available**. You can choose it and import the data that you've transferred into Data Wrangler. The data that you've transferred appears as tables that you can query.

**Using Amazon AppFlow to transfer your data**

Amazon AppFlow is a platform that you can use to transfer data from your SaaS platform to Amazon S3 or Amazon Redshift without having to write any code. To perform a data transfer, you use the AWS Management Console.

⚠️ **Important**

You must make sure you've set up the permissions to perform a data transfer. For more information, see [Amazon AppFlow Permissions](#).

After you've added permissions, you can transfer the data. Within Amazon AppFlow, you create a **flow** to transfer the data. A flow is a series of configurations. You can use it to specify whether you're running the data transfer on a schedule or whether you're partitioning the data into separate files. After you've configured the flow, you run it to transfer the data.

For information about creating a flow, see [Creating flows in Amazon AppFlow](#). For information about running a flow, see [Activate an Amazon AppFlow flow](#).

After the data has been transferred, use the following procedure to access the data in Data Wrangler.

⚠️ **Important**

Before you try to access your data, make sure your IAM role has the following policy:
By default, the IAM role that you use to access Data Wrangler is the SageMakerExecutionRole. For more information about adding policies, see Adding IAM identity permissions (console).

To connect to a data source, do the following.

1. Sign into Amazon SageMaker Console.
2. Choose Studio.
3. Choose Launch app.
4. From the dropdown list, select Studio.
5. Choose the Home icon.
6. Choose Data.
7. Choose Data Wrangler.
8. Choose Import data.
9. Under Available, choose the data source.
10. For the Name field, specify the name of the connection.
11. (Optional) Choose Advanced configuration.
   a. Choose a Workgroup.
   b. If your workgroup hasn't enforced the Amazon S3 output location or if you don't use a workgroup, specify a value for Amazon S3 location of query results.
c. (Optional) For Data retention period, select the checkbox to set a data retention period
and specify the number of days to store the data before it's deleted.

d. (Optional) By default, Data Wrangler saves the connection. You can choose to deselect the
checkbox and not save the connection.

12. Choose Connect.

13. Specify a query.

[Note]
To help you specify a query, you can choose a table on the left-hand navigation panel.
Data Wrangler shows the table name and a preview of the table. Choose the icon next
to the table name to copy the name. You can use the table name in the query.


15. Choose Import query.

16. For Dataset name, specify the name of the dataset.

17. Choose Add.

When you navigate to the Import data screen, you can see the connection that you've created. You
can use the connection to import more data.

Imported Data Storage

[Important]
We strongly recommend that you follow the best practices around protecting your Amazon
S3 bucket by following Security best practices.

When you query data from Amazon Athena or Amazon Redshift, the queried dataset is
automatically stored in Amazon S3. Data is stored in the default SageMaker S3 bucket for the AWS
Region in which you are using Studio Classic.

Default S3 buckets have the following naming convention: sagemaker-region-account
number. For example, if your account number is 111122223333 and you are using Studio Classic in
us-east-1, your imported datasets are stored in sagemaker-us-east-1-111122223333.
Data Wrangler flows depend on this Amazon S3 dataset location, so you should not modify this dataset in Amazon S3 while you are using a dependent flow. If you do modify this S3 location, and you want to continue using your data flow, you must remove all objects in `trained_parameters` in your .flow file. To do this, download the .flow file from Studio Classic and for each instance of `trained_parameters`, delete all entries. When you are done, `trained_parameters` should be an empty JSON object:

```
"trained_parameters": {}
```

When you export and use your data flow to process your data, the .flow file you export refers to this dataset in Amazon S3. Use the following sections to learn more.

**Amazon Redshift Import Storage**

Data Wrangler stores the datasets that result from your query in a Parquet file in your default SageMaker S3 bucket.

This file is stored under the following prefix (directory): `redshift/uuid/data/`, where `uuid` is a unique identifier that gets created for each query.

For example, if your default bucket is `sagemaker-us-east-1-111122223333`, a single dataset queried from Amazon Redshift is located in `s3://sagemaker-us-east-1-111122223333/redshift/uuid/data/`.

**Amazon Athena Import Storage**

When you query an Athena database and import a dataset, Data Wrangler stores the dataset, as well as a subset of that dataset, or *preview files*, in Amazon S3.

The dataset you import by selecting **Import dataset** is stored in Parquet format in Amazon S3.

Preview files are written in CSV format when you select **Run** on the Athena import screen, and contain up to 100 rows from your queried dataset.

The dataset you query is located under the prefix (directory): `athena/uuid/data/`, where `uuid` is a unique identifier that gets created for each query.

For example, if your default bucket is `sagemaker-us-east-1-111122223333`, a single dataset queried from Athena is located in `s3://sagemaker-us-east-1-111122223333/athena/uuid/data/example_dataset.parquet`. 
The subset of the dataset that is stored to preview dataframes in Data Wrangler is stored under the prefix: athena/.

**Create and Use a Data Wrangler Flow**

Use an Amazon SageMaker Data Wrangler flow, or a *data flow*, to create and modify a data preparation pipeline. The data flow connects the datasets, transformations, and analyses, or *steps*, you create and can be used to define your pipeline.

**Instances**

When you create a Data Wrangler flow in Amazon SageMaker Studio Classic, Data Wrangler uses an Amazon EC2 instance to run the analyses and transformations in your flow. By default, Data Wrangler uses the m5.4xlarge instance. m5 instances are general purpose instances that provide a balance between compute and memory. You can use m5 instances for a variety of compute workloads.

Data Wrangler also gives you the option of using r5 instances. r5 instances are designed to deliver fast performance that processes large datasets in memory.

We recommend that you choose an instance that is best optimized around your workloads. For example, the r5.8xlarge might have a higher price than the m5.4xlarge, but the r5.8xlarge might be better optimized for your workloads. With better optimized instances, you can run your data flows in less time at lower cost.

The following table shows the instances that you can use to run your Data Wrangler flow.

<table>
<thead>
<tr>
<th>Standard Instances</th>
<th>vCPU</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.m5.4xlarge</td>
<td>16</td>
<td>64 GiB</td>
</tr>
<tr>
<td>ml.m5.8xlarge</td>
<td>32</td>
<td>128 GiB</td>
</tr>
<tr>
<td>ml.m5.16xlarge</td>
<td>64</td>
<td>256 GiB</td>
</tr>
<tr>
<td>ml.m5.24xlarge</td>
<td>96</td>
<td>384 GiB</td>
</tr>
<tr>
<td>r5.4xlarge</td>
<td>16</td>
<td>128 GiB</td>
</tr>
<tr>
<td>r5.8xlarge</td>
<td>32</td>
<td>256 GiB</td>
</tr>
</tbody>
</table>
### Standard Instances

<table>
<thead>
<tr>
<th>Instance Type</th>
<th>vCPU</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>r5.24xlarge</td>
<td>96</td>
<td>768 GiB</td>
</tr>
</tbody>
</table>

For more information about r5 instances, see [Amazon EC2 R5 Instances](#). For more information about m5 instances, see [Amazon EC2 M5 Instances](#).

Each Data Wrangler flow has an Amazon EC2 instance associated with it. You might have multiple flows that are associated with a single instance.

For each flow file, you can seamlessly switch the instance type. If you switch the instance type, the instance that you used to run the flow continues to run.

To switch the instance type of your flow, do the following.

1. Choose the home icon,

2. Navigate to the instance that you're using and choose it.

3. Choose the instance type that you want to use.
4. Choose **Save**.

You are charged for all running instances. To avoid incurring additional charges, shut down the instances that you aren't using manually. To shut down an instance that is running, use the following procedure.

To shut down a running instance.

1. Choose the instance icon. The following image shows you where to select the **RUNNING INSTANCES** icon.

![Instance selection](image.png)

2. Choose **Shut down** next to the instance that you want to shut down.

If you shut down an instance used to run a flow, you temporarily can't access the flow. If you get an error while attempting to open the flow running an instance you previously shut down, wait for 5 minutes and try opening it again.

When you export your data flow to a location such as Amazon Simple Storage Service or Amazon SageMaker Feature Store, Data Wrangler runs an Amazon SageMaker processing job. You can use one of the following instances for the processing job. For more information on exporting your data, see **Export**.

<table>
<thead>
<tr>
<th>Standard Instances</th>
<th>vCPU</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.m5.4xlarge</td>
<td>16</td>
<td>64 GiB</td>
</tr>
<tr>
<td>ml.m5.12xlarge</td>
<td>48</td>
<td>192 GiB</td>
</tr>
<tr>
<td>ml.m5.24xlarge</td>
<td>96</td>
<td>384 GiB</td>
</tr>
</tbody>
</table>
For more information about the cost per hour for using the available instance types, see SageMaker Pricing.

The Data Flow UI

When you import a dataset, the original dataset appears on the data flow and is named Source. If you turned on sampling when you imported your data, this dataset is named Source - sampled. Data Wrangler automatically infers the types of each column in your dataset and creates a new dataframe named Data types. You can select this frame to update the inferred data types. You see results similar to those shown in the following image after you upload a single dataset:

![Data Flow UI](image)

Each time you add a transform step, you create a new dataframe. When multiple transform steps (other than Join or Concatenate) are added to the same dataset, they are stacked.

Join and Concatenate create standalone steps that contain the new joined or concatenated dataset.

The following diagram shows a data flow with a join between two datasets, as well as two stacks of steps. The first stack (Steps (2)) adds two transforms to the type inferred in the Data types dataset. The downstream stack, or the stack to the right, adds transforms to the dataset resulting from a join named demo-join.
The small, gray box in the bottom right corner of the data flow provides an overview of number of stacks and steps in the flow and the layout of the flow. The lighter box inside the gray box indicates the steps that are within the UI view. You can use this box to see sections of your data flow that fall outside of the UI view. Use the fit screen icon to fit all steps and datasets into your UI view.

The bottom left navigation bar includes icons that you can use to zoom in and out of your data flow and resize the data flow to fit the screen. Use the lock icon to lock and unlock the location of each step on the screen.
Add a Step to Your Data Flow

Select + next to any dataset or previously added step and then select one of the following options:

- **Edit data types** (For a Data types step only): If you have not added any transforms to a Data types step, you can select **Edit data types** to update the data types Data Wrangler inferred when importing your dataset.

- **Add transform**: Adds a new transform step. See [Transform Data](#) to learn more about the data transformations you can add.

- **Add analysis**: Adds an analysis. You can use this option to analyze your data at any point in the data flow. When you add one or more analyses to a step, an analysis icon appears on that step. See [Analyze and Visualize](#) to learn more about the analyses you can add.

- **Join**: Joins two datasets and adds the resulting dataset to the data flow. To learn more, see [Join Datasets](#).

- **Concatenate**: Concatenates two datasets and adds the resulting dataset to the data flow. To learn more, see [Concatenate Datasets](#).

Delete a Step from Your Data Flow

To delete a step, select the step and select **Delete**. If the node is a node that has a single input, you delete only the step that you select. Deleting a step that has a single input doesn't delete the steps that follow it. If you're deleting a step for a source, join, or concatenate node, all the steps that follow it are also deleted.

To delete a step from a stack of steps, select the stack and then select the step you want to delete.

You can use one of the following procedures to delete a step without deleting the downstream steps.

**Delete a step in the Data Wrangler flow**

You can delete an individual step for nodes in your data flow that have a single input. You can't delete individual steps for source, join, and concatenate nodes.

Use the following procedure to delete a step in the Data Wrangler flow.
1. Choose the group of steps that has the step that you're deleting.
2. Choose the icon next to the step.
3. Choose **Delete step**.

Delete a step in the table view

Use the following procedure to delete a step in the table view.

You can delete an individual step for nodes in your data flow that have a single input. You can't delete individual steps for source, join, and concatenate nodes.

1. Choose the step and open the table view for the step.
2. Move your cursor over the step so the ellipsis icon appears.
3. Choose the icon next to the step.
4. Choose **Delete**.
Edit a Step in Your Data Wrangler Flow

You can edit each step that you've added in your Data Wrangler flow. By editing steps, you can change the transformations or the data types of the columns. You can edit the steps to make changes with which you can perform better analyses.

There are many ways that you can edit a step. Some examples include changing the imputation method or changing the threshold for considering a value to be an outlier.

Use the following procedure to edit a step.

To edit a step, do the following.

1. Choose a step in the Data Wrangler flow to open the table view.
2. Choose a step in the data flow.
3. Edit the step.

The following image shows an example of editing a step.

Note
You can use the shared spaces within your Amazon SageMaker domain to work collaboratively on your Data Wrangler flows. Within a shared space, you and your
collaborators can edit a flow file in real-time. However, neither you nor your collaborators can see the changes in real-time. When anyone makes a change to the Data Wrangler flow, they must save it immediately. When someone saves a file, a collaborator won’t be able to see it unless they close the file and reopen it. Any changes that aren’t saved by one person are overwritten by the person who saved their changes.

Get Insights On Data and Data Quality

Use the Data Quality and Insights Report to perform an analysis of the data that you’ve imported into Data Wrangler. We recommend that you create the report after you import your dataset. You can use the report to help you clean and process your data. It gives you information such as the number of missing values and the number of outliers. If you have issues with your data, such as target leakage or imbalance, the insights report can bring those issues to your attention.

Use the following procedure to create a Data Quality and Insights report. It assumes that you've already imported a dataset into your Data Wrangler flow.

To create a Data Quality and Insights report

1. Choose a + next to a node in your Data Wrangler flow.
2. Select Get data insights.
3. For Analysis name, specify a name for the insights report.
4. (Optional) For Target column, specify the target column.
5. For Problem type, specify Regression or Classification.
6. For Data size, specify one of the following:
   - 50 K – Uses the first 50000 rows of the dataset that you've imported to create the report.
   - Entire dataset – Uses the entire dataset that you've imported to create the report.

Note

Creating a Data Quality and Insights report on the entire dataset uses an Amazon SageMaker processing job. A SageMaker processing job provisions the additional compute resources required to get insights for all of your data. For more information about SageMaker processing jobs, see Process data.
7. Choose **Create**.

The following topics show the sections of the report:

**Topics**

- **Summary**
- **Target column**
- **Quick model**
- **Feature summary**
- **Samples**
- **Definitions**

You can either download the report or view it online. To download the report, choose the download button at the top right corner of the screen. The following image shows the button.

![Download button](image)

**Summary**

The insights report has a brief summary of the data that includes general information such as missing values, invalid values, feature types, outlier counts, and more. It can also include high severity warnings that point to probable issues with the data. We recommend that you investigate the warnings.

The following is an example of a report summary.
Target column

When you create the data quality and insights report, Data Wrangler gives you the option to select a target column. A target column is a column that you're trying to predict. When you choose a target column, Data Wrangler automatically creates a target column analysis. It also ranks the features in the order of their predictive power. When you select a target column, you must specify whether you're trying to solve a regression or a classification problem.

For classification, Data Wrangler shows a table and a histogram of the most common classes. A class is a category. It also presents observations, or rows, with a missing or invalid target value.

The following image shows an example target column analysis for a classification problem.
For regression, Data Wrangler shows a histogram of all the values in the target column. It also presents observations, or rows, with a missing, invalid, or outlier target value.

The following image shows an example target column analysis for a regression problem.
Quick model

The **Quick model** provides an estimate of the expected predicted quality of a model that you train on your data.

Data Wrangler splits your data into training and validation folds. It uses 80% of the samples for training and 20% of the values for validation. For classification, the sample is stratified split. For a stratified split, each data partition has the same ratio of labels. For classification problems, it's important to have the same ratio of labels between the training and classification folds. Data Wrangler trains the XGBoost model with the default hyperparameters. It applies early stopping on the validation data and performs minimal feature preprocessing.

For classification models, Data Wrangler returns both a model summary and a confusion matrix.

The following is an example of a classification model summary. To learn more about the information that it returns, see [Definitions](#).
The following is an example of a confusion matrix that the quick model returns.

<table>
<thead>
<tr>
<th>Class</th>
<th>Precision</th>
<th>Recall</th>
<th>F1-Score</th>
<th>Support</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.9698795180722891</td>
<td>0.9938271604938271</td>
<td>0.9817073170731707</td>
<td>162.0</td>
</tr>
<tr>
<td>1</td>
<td>0.9895833333333334</td>
<td>0.95</td>
<td>0.9693877551020408</td>
<td>100.0</td>
</tr>
</tbody>
</table>

A confusion matrix gives you the following information:

- The number of times the predicted label matches the true label.
- The number of times the predicted label doesn't match the true label.
The true label represents an actual observation in your data. For example, if you're using a model to detect fraudulent transactions, the true label represents a transaction that is actually fraudulent or non-fraudulent. The predicted label represents the label that your model assigns to the data.

You can use the confusion matrix to see how well the model predicts the presence or the absence of a condition. If you're predicting fraudulent transactions, you can use the confusion matrix to get a sense of both the sensitivity and the specificity of the model. The sensitivity refers to the model's ability to detect fraudulent transactions. The specificity refers to the model's ability to avoid detecting non-fraudulent transactions as fraudulent.

The following is an example of the quick model outputs for a regression problem.

**Feature summary**

When you specify a target column, Data Wrangler orders the features by their prediction power. Prediction power is measured on the data after it was split into 80% training and 20% validation folds. Data Wrangler fits a model for each feature separately on the training fold. It applies minimal feature preprocessing and measures prediction performance on the validation data.

It normalizes the scores to the range [0,1]. Higher prediction scores indicate columns that are more useful for predicting the target on their own. Lower scores point to columns that aren’t predictive of the target column.

It’s uncommon for a column that isn’t predictive on its own to be predictive when it's used in tandem with other columns. You can confidently use the prediction scores to determine whether a feature in your dataset is predictive.
A low score usually indicates the feature is redundant. A score of 1 implies perfect predictive abilities, which often indicates target leakage. Target leakage usually happens when the dataset contains a column that isn’t available at the prediction time. For example, it could be a duplicate of the target column.

The following are examples of the table and the histogram that show the prediction value of each feature.
Data Wrangler provides information about whether your samples are anomalous or if there are duplicates in your dataset.

Data Wrangler detects anomalous samples using the isolation forest algorithm. The isolation forest associates an anomaly score with each sample (row) of the dataset. Low anomaly scores indicate anomalous samples. High scores are associated with non-anomalous samples. Samples with a negative anomaly score are usually considered anomalous and samples with positive anomaly score are considered non-anomalous.

When you look at a sample that might be anomalous, we recommend that you pay attention to unusual values. For example, you might have anomalous values that result from errors in gathering and processing the data. The following is an example of the most anomalous samples according to the Data Wrangler’s implementation of the isolation forest algorithm. We recommend using domain knowledge and business logic when you examine the anomalous samples.

Data Wrangler detects duplicate rows and calculates the ratio of duplicate rows in your data. Some data sources could include valid duplicates. Other data sources could have duplicates that point to problems in data collection. Duplicate samples that result from faulty data collection could interfere with machine learning processes that rely on splitting the data into independent training and validation folds.
The following are elements of the insights report that can be impacted by duplicated samples:

- Quick model
- Prediction power estimation
- Automatic hyperparameter tuning

You can remove duplicate samples from the dataset using the **Drop duplicates** transform under **Manage rows**. Data Wrangler shows you the most frequently duplicated rows.

**Definitions**

The following are definitions for the technical terms that are used in the data insights report.

**Feature types**

The following are the definitions for each of the feature types:

- **Numeric** – Numeric values can be either floats or integers, such as age or income. The machine learning models assume that numeric values are ordered and a distance is defined over them. For example, 3 is closer to 4 than to 10 and $3 < 4 < 10$.

- **Categorical** – The column entries belong to a set of unique values, which is usually much smaller than the number of entries in the column. For example, a column of length 100 could contain the unique values Dog, Cat, and Mouse. The values could be numeric, text, or a combination of both. Horse, House, 8, Love, and $3.1$ would all be valid values and could be found in the same categorical column. The machine learning model does not assume order or distance on the values of categorical features, as opposed to numeric features, even when all the values are numbers.

- **Binary** – Binary features are a special categorical feature type in which the cardinality of the set of unique values is 2.

- **Text** – A text column contains many non-numeric unique values. In extreme cases, all the elements of the column are unique. In an extreme case, no two entries are the same.

- **Datetime** – A datetime column contains information about the date or time. It can have information about both the date and time.

**Feature statistics**

The following are definitions for each of the feature statistics:
• **Prediction power** – Prediction power measures how useful the column is in predicting the target.

• **Outliers** (in numeric columns) – Data Wrangler detects outliers using two statistics that are robust to outliers: median and robust standard deviation (RSTD). RSTD is derived by clipping the feature values to the range [5 percentile, 95 percentile] and calculating the standard deviation of the clipped vector. All values larger than median + 5 * RSTD or smaller than median - 5 * RSTD are considered to be outliers.

• **Skew** (in numeric columns) – Skew measures the symmetry of the distribution and is defined as the third moment of the distribution divided by the third power of the standard deviation. The skewness of the normal distribution or any other symmetric distribution is zero. Positive values imply that the right tail of the distribution is longer than the left tail. Negative values imply that the left tail of the distribution is longer than the right tail. As a rule of thumb, a distribution is considered skewed when the absolute value of the skew is larger than 3.

• **Kurtosis** (in numeric columns) – Pearson's kurtosis measures the heaviness of the tail of the distribution. It's defined as the fourth moment of the distribution divided by the square of the second moment. The kurtosis of the normal distribution is 3. Kurtosis values lower than 3 imply that the distribution is concentrated around the mean and the tails are lighter than the tails of the normal distribution. Kurtosis values higher than 3 imply heavier tails or outliers.

• **Missing values** – Null-like objects, empty strings and strings composed of only white spaces are considered missing.

• **Valid values for numeric features or regression target** – All values that you can cast to finite floats are valid. Missing values are not valid.

• **Valid values for categorical, binary, or text features, or for classification target** – All values that are not missing are valid.

• **Datetime features** – All values that you can cast to a datetime object are valid. Missing values are not valid.

• **Invalid values** – Values that are either missing or you can't properly cast. For example, in a numeric column, you can't cast the string "six" or a null value.

Quick model metrics for regression

The following are the definitions for the quick model metrics:

• **R2 or coefficient of determination** – R2 is the proportion of the variation in the target that is predicted by the model. R2 is in the range of [-infty, 1]. 1 is the score of the model that
predicts the target perfectly and 0 is the score of the trivial model that always predicts the target mean.

- **MSE** or mean squared error – MSE is in the range \([0, \infty]\). 0 is the score of the model that predicts the target perfectly.
- **MAE** or mean absolute error – MAE is in the range \([0, \infty]\) where 0 is the score of the model that predicts the target perfectly.
- **RMSE** or root mean square error – RMSE is in the range \([0, \infty]\) where 0 is the score of the model that predicts the target perfectly.
- **Max error** – The maximum absolute value of the error over the dataset. Max error is in the range \([0, \infty]\). 0 is the score of the model that predicts the target perfectly.
- **Median absolute error** – Median absolute error is in the range \([0, \infty]\). 0 is the score of the model that predicts the target perfectly.

Quick model metrics for classification

The following are the definitions for the quick model metrics:

- **Accuracy** – Accuracy is the ratio of samples that are predicted accurately. Accuracy is in the range \([0, 1]\). 0 is the score of the model that predicts all samples incorrectly and 1 is the score of the perfect model.
- **Balanced accuracy** – Balanced accuracy is the ratio of samples that are predicted accurately when the class weights are adjusted to balance the data. All classes are given the same importance, regardless of their frequency. Balanced accuracy is in the range \([0, 1]\). 0 is the score of the model that predicts all samples wrong. 1 is the score of the perfect model.
- **AUC (binary classification)** – This is the area under the receiver operating characteristic curve. AUC is in the range \([0, 1]\) where a random model returns a score of 0.5 and the perfect model returns a score of 1.
- **AUC (OVR)** – For multiclass classification, this is the area under the receiver operating characteristic curve calculated separately for each label using one versus rest. Data Wrangler reports the average of the areas. AUC is in the range \([0, 1]\) where a random model returns a score of 0.5 and the perfect model returns a score of 1.
- **Precision** – Precision is defined for a specific class. Precision is the fraction of true positives out of all the instances that the model classified as that class. Precision is in the range \([0, 1]\). 1 is the score of the model that has no false positives for the class. For binary classification, Data Wrangler reports the precision of the positive class.
• **Recall** – Recall is defined for a specific class. Recall is the fraction of the relevant class instances that are successfully retrieved. Recall is in the range [0, 1]. 1 is the score of the model that classifies all the instances of the class correctly. For binary classification, Data Wrangler reports the recall of the positive class.

• **F1** – F1 is defined for a specific class. It's the harmonic mean of the precision and recall. F1 is in the range [0, 1]. 1 is the score of the perfect model. For binary classification, Data Wrangler reports the F1 for classes with positive values.

**Textual patterns**

*Patterns* describe the textual format of a string using an easy to read format. The following are examples of textual patterns:

- "{digits:4-7}" describes a sequence of digits that have a length between 4 and 7.
- "{alnum:5}" describes an alpha-numeric string with a length of exactly 5.

Data Wrangler infers the patterns by looking at samples of non-empty strings from your data. It can describe many of the commonly used patterns. The confidence expressed as a percentage indicates how much of the data is estimated to match the pattern. Using the textual pattern, you can see which rows in your data you need to correct or drop.

The following describes the patterns that Data Wrangler can recognize:

<table>
<thead>
<tr>
<th>Pattern</th>
<th>Textual Format</th>
</tr>
</thead>
<tbody>
<tr>
<td>{alnum}</td>
<td>Alphanumeric strings</td>
</tr>
<tr>
<td>{any}</td>
<td>Any string of word characters</td>
</tr>
<tr>
<td>{digits}</td>
<td>A sequence of digits</td>
</tr>
<tr>
<td>{lower}</td>
<td>A lowercase word</td>
</tr>
<tr>
<td>{mixed}</td>
<td>A mixed-case word</td>
</tr>
<tr>
<td>{name}</td>
<td>A word beginning with a capital letter</td>
</tr>
<tr>
<td>{upper}</td>
<td>An uppercase word</td>
</tr>
</tbody>
</table>
A word character is either an underscore or a character that might appear in a word in any language. For example, the strings 'Hello_word' and 'écoute' both consist of word characters. 'H' and 'é' are both examples of word characters.

**Automatically Train Models on Your Data Flow**

You can use Amazon SageMaker Autopilot to automatically train, tune, and deploy models on the data that you've transformed in your data flow. Amazon SageMaker Autopilot can go through several algorithms and use the one that works best with your data. For more information about Amazon SageMaker Autopilot, see [SageMaker Autopilot](#).

When you train and tune a model, Data Wrangler exports your data to an Amazon S3 location where Amazon SageMaker Autopilot can access it.

You can prepare and deploy a model by choosing a node in your Data Wrangler flow and choosing **Export and Train** in the data preview. You can use this method to view your dataset before you choose to train a model on it.

You can also train and deploy a model directly from your data flow.

The following procedure prepares and deploys a model from the data flow. For Data Wrangler flows with multi-row transforms, you can't use the transforms from the Data Wrangler flow when you're deploying the model. You can use the following procedure to process the data before you use it to perform inference.

To train and deploy a model directly from your data flow, do the following.

1. Choose the + next to the node containing the training data.
2. Choose **Train model**.
3. (Optional) Specify a AWS KMS key or ID. For more information about creating and controlling cryptographic keys to protect your data, see [AWS Key Management Service](#).
4. Choose **Export and train**.
5. After Amazon SageMaker Autopilot trains the model on the data that Data Wrangler exported, specify a name for **Experiment name**.

6. Under **Input data**, choose **Preview** to verify that Data Wrangler properly exported your data to Amazon SageMaker Autopilot.

7. For **Target**, choose the target column.

8. (Optional) For **S3 location** under **Output data**, specify an Amazon S3 location other than the default location.

9. Choose **Next: Training method**.

10. Choose a training method. For more information, see **Training modes**.

11. (Optional) For **Auto deploy endpoint**, specify a name for the endpoint.

12. For **Deployment option**, choose a deployment method. You can choose to deploy with or without the transformations that you've made to your data.

    ⚠ **Important**
    
    You can't deploy an Amazon SageMaker Autopilot model with the transformations that you've made in your Data Wrangler flow. For more information about those transformations, see **Export to an Inference Endpoint**.

13. Choose **Next: Review and create**.

14. Choose **Create experiment**.

For more information about model training and deployment, see **Create a regression or classification job for tabular data using the AutoML API**. Autopilot shows you analyses about the best model's performance. For more information about model performance, see **View an Autopilot Model Performance Report**.

**Transform Data**

Amazon SageMaker Data Wrangler provides numerous ML data transforms to streamline cleaning, transforming, and featurizing your data. When you add a transform, it adds a step to the data flow. Each transform you add modifies your dataset and produces a new dataframe. All subsequent transforms apply to the resulting dataframe.

Data Wrangler includes built-in transforms, which you can use to transform columns without any code. You can also add custom transformations using PySpark, Python (User-Defined Function),
pandas, and PySpark SQL. Some transforms operate in place, while others create a new output column in your dataset.

You can apply transforms to multiple columns at once. For example, you can delete multiple columns in a single step.

You can apply the **Process numeric** and **Handle missing** transforms only to a single column.

Use this page to learn more about these built-in and custom transforms.

### Transform UI

Most of the built-in transforms are located in the **Prepare** tab of the Data Wrangler UI. You can access the join and concatenate transforms through the data flow view. Use the following table to preview these two views.

#### Transform

You can add a transform to any step in your data flow. Use the following procedure to add a transform to your data flow.

To add a step to your data flow, do the following.

1. Choose the + next to the step in the data flow.
2. Choose **Add transform**.
3. Choose **Add step**.
4. Choose a transform.

5. (Optional) You can search for the transform that you want to use. Data Wrangler highlights the query in the results.
Join View

To join two datasets, select the first dataset in your data flow and choose Join. When you choose Join, you see results similar to those shown in the following image. Your left and right datasets are displayed in the left panel. The main panel displays your data flow, with the newly joined dataset added.
When you choose **Configure** to configure your join, you see results similar to those shown in the following image. Your join configuration is displayed in the left panel. You can use this panel to choose the joined dataset name, join type, and columns to join. The main panel displays three tables. The top two tables display the left and right datasets on the left and right respectively. Under this table, you can preview the joined dataset.
See [Join Datasets](#) to learn more.

**Concatenate View**

To concatenate two datasets, you select the first dataset in your data flow and choose [Concatenate](#). When you select Concatenate, you see results similar to those shown in the following image. Your left and right datasets are displayed in the left panel. The main panel displays your data flow, with the newly concatenated dataset added.
When you choose **Configure** to configure your concatenation, you see results similar to those shown in the following image. Your concatenate configuration displays in the left panel. You can use this panel to choose the concatenated dataset's name, and choose to remove duplicates after concatenation and add columns to indicate the source data frame. The main panel displays three tables. The top two tables display the left and right datasets on the left and right respectively. Under this table, you can preview the concatenated dataset.
Join Datasets

You join dataframes directly in your data flow. When you join two datasets, the resulting joined dataset appears in your flow. The following join types are supported by Data Wrangler.

- **Left Outer** – Include all rows from the left table. If the value for the column joined on a left table row does not match any right table row values, that row contains null values for all right table columns in the joined table.

- **Left Anti** – Include rows from the left table that do not contain values in the right table for the joined column.

- **Left semi** – Include a single row from the left table for all identical rows that satisfy the criteria in the join statement. This excludes duplicate rows from the left table that match the criteria of the join.

See [Concatenate Datasets](#) to learn more.
- **Right Outer** – Include all rows from the right table. If the value for the joined column in a right table row does not match any left table row values, that row contains null values for all left table columns in the joined table.

- **Inner** – Include rows from left and right tables that contain matching values in the joined column.

- **Full Outer** – Include all rows from the left and right tables. If the row value for the joined column in either table does not match, separate rows are created in the joined table. If a row doesn’t contain a value for a column in the joined table, null is inserted for that column.

- **Cartesian Cross** – Include rows which combine each row from the first table with each row from the second table. This is a [Cartesian product](#) of rows from tables in the join. The result of this product is the size of the left table times the size of the right table. Therefore, we recommend caution in using this join between very large datasets.

Use the following procedure to join two dataframes.

1. Select + next to the left dataframe that you want to join. The first dataframe you select is always the left table in your join.
2. Choose **Join**.
3. Select the right dataframe. The second dataframe you select is always the right table in your join.
4. Choose **Configure** to configure your join.
5. Give your joined dataset a name using the **Name** field.
6. Select a **Join type**.
7. Select a column from the left and right tables to join.
8. Choose **Apply** to preview the joined dataset on the right.
9. To add the joined table to your data flow, choose **Add**.

### Concatenate Datasets

**Concatenate two datasets:**

1. Choose + next to the left dataframe that you want to concatenate. The first dataframe you select is always the left table in your concatenate.
2. Choose **Concatenate**.
3. Select the right dataframe. The second dataframe you select is always the right table in your concatenate.

4. Choose **Configure** to configure your concatenate.

5. Give your concatenated dataset a name using the **Name** field.

6. (Optional) Select the checkbox next to **Remove duplicates after concatenation** to remove duplicate columns.

7. (Optional) Select the checkbox next to **Add column to indicate source dataframe** if, for each column in the new dataset, you want to add an indicator of the column's source.

8. Choose **Apply** to preview the new dataset.

9. Choose **Add** to add the new dataset to your data flow.

### Balance Data

You can balance the data for datasets with an underrepresented category. Balancing a dataset can help you create better models for binary classification.

**Note**

You can't balance datasets containing column vectors.

You can use the **Balance data** operation to balance your data using one of the following operators:

- **Random oversampling** – Randomly duplicates samples in the minority category. For example, if you're trying to detect fraud, you might only have cases of fraud in 10% of your data. For an equal proportion of fraudulent and non-fraudulent cases, this operator randomly duplicates fraud cases within the dataset 8 times.

- **Random undersampling** – Roughly equivalent to random oversampling. Randomly removes samples from the overrepresented category to get the proportion of samples that you desire.

- **Synthetic Minority Oversampling Technique (SMOTE)** – Uses samples from the underrepresented category to interpolate new synthetic minority samples. For more information about SMOTE, see the following description.

You can use all transforms for datasets containing both numeric and non-numeric features. SMOTE interpolates values by using neighboring samples. Data Wrangler uses the R-squared distance...
to determine the neighborhood to interpolate the additional samples. Data Wrangler only uses numeric features to calculate the distances between samples in the underrepresented group.

For two real samples in the underrepresented group, Data Wrangler interpolates the numeric features by using a weighted average. It randomly assigns weights to those samples in the range of [0, 1]. For numeric features, Data Wrangler interpolates samples using a weighted average of the samples. For samples A and B, Data Wrangler could randomly assign a weight of 0.7 to A and 0.3 to B. The interpolated sample has a value of 0.7A + 0.3B.

Data Wrangler interpolates non-numeric features by copying from either of the interpolated real samples. It copies the samples with a probability that it randomly assigns to each sample. For samples A and B, it can assign probabilities 0.8 to A and 0.2 to B. For the probabilities it assigned, it copies A 80% of the time.

**Custom Transforms**

The **Custom Transforms** group allows you to use Python (User-Defined Function), PySpark, pandas, or PySpark (SQL) to define custom transformations. For all three options, you use the variable `df` to access the dataframe to which you want to apply the transform. To apply your custom code to your dataframe, assign the dataframe with the transformations that you've made to the `df` variable. If you're not using Python (User-Defined Function), you don't need to include a return statement. Choose **Preview** to preview the result of the custom transform. Choose **Add** to add the custom transform to your list of **Previous steps**.

You can import the popular libraries with an `import` statement in the custom transform code block, such as the following:

- NumPy version 1.19.0
- scikit-learn version 0.23.2
- SciPy version 1.5.4
- pandas version 1.0.3
- PySpark version 3.0.0

⚠️ **Important**

**Custom transform** doesn't support columns with spaces or special characters in the name. We recommend that you specify column names that only have alphanumeric characters and underscores. You can use the **Rename column** transform in the **Manage columns**
transform group to remove spaces from a column's name. You can also add a **Python (Pandas) Custom transform** similar to the following to remove spaces from multiple columns in a single step. This example changes columns named A column and B column to A_column and B_column respectively.

```python
df.rename(columns={"A column": "A_column", "B column": "B_column"})
```

If you include print statements in the code block, the result appears when you select **Preview**. You can resize the custom code transformer panel. Resizing the panel provides more space to write code. The following image shows the resizing of the panel.

The following sections provide additional context and examples for writing custom transform code.

**Python (User-Defined Function)**

The Python function gives you the ability to write custom transformations without needing to know Apache Spark or pandas. Data Wrangler is optimized to run your custom code quickly. You get similar performance using custom Python code and an Apache Spark plugin.

To use the Python (User-Defined Function) code block, you specify the following:
• **Input column** – The input column where you’re applying the transform.

• **Mode** – The scripting mode, either pandas or Python.

• **Return type** – The data type of the value that you’re returning.

Using the pandas mode gives better performance. The Python mode makes it easier for you to write transformations by using pure Python functions.

The following video shows an example of how to use custom code to create a transformation. It uses the Titanic dataset to create a column with the person’s salutation.

**PySpark**

The following example extracts date and time from a timestamp.

```python
from pyspark.sql.functions import from_unixtime, to_date, date_format
df = df.withColumn('DATE_TIME', from_unixtime('TIMESTAMP'))
df = df.withColumn('EVENT_DATE', to_date('DATE_TIME')).withColumn('EVENT_TIME', date_format('DATE_TIME', 'HH:mm:ss'))
```

**pandas**

```python```
The following example provides an overview of the dataframe to which you are adding transforms.

```python
df.info()
```

**PySpark (SQL)**

The following example creates a new dataframe with four columns: `name`, `fare`, `pclass`, `survived`.

```sql
SELECT name, fare, pclass, survived FROM df
```

If you don't know how to use PySpark, you can use custom code snippets to help you get started.

Data Wrangler has a searchable collection of code snippets. You can use to code snippets to perform tasks such as dropping columns, grouping by columns, or modelling.

To use a code snippet, choose **Search example snippets** and specify a query in the search bar. The text you specify in the query doesn’t have to match the name of the code snippet exactly.

The following example shows a **Drop duplicate rows** code snippet that can delete rows with similar data in your dataset. You can find the code snippet by searching for one of the following:

- Duplicates
- Identical
- Remove

The following snippet has comments to help you understand the changes that you need to make. For most snippets, you must specify the column names of your dataset in the code.

```python
# Specify the subset of columns
# all rows having identical values in these columns will be dropped

subset = ["col1", "col2", "col3"]

df = df.dropDuplicates(subset)

# to drop the full-duplicate rows run
# df = df.dropDuplicates()
```
To use a snippet, copy and paste its content into the **Custom transform** field. You can copy and paste multiple code snippets into the custom transform field.

**Custom Formula**

Use **Custom formula** to define a new column using a Spark SQL expression to query data in the current dataframe. The query must use the conventions of Spark SQL expressions.

⚠️ **Important**

**Custom formula** doesn't support columns with spaces or special characters in the name. We recommend that you specify column names that only have alphanumeric characters and underscores. You can use the **Rename column** transform in the **Manage columns** transform group to remove spaces from a column's name. You can also add a **Python (Pandas) Custom transform** similar to the following to remove spaces from multiple columns in a single step. This example changes columns named *A column* and *B column* to *A_column* and *B_column* respectively.

```python
df.rename(columns={"A column": "A_column", "B column": "B_column"})
```

You can use this transform to perform operations on columns, referencing the columns by name. For example, assuming the current dataframe contains columns named *col_a* and *col_b*, you can use the following operation to produce an **Output column** that is the product of these two columns with the following code:

```python
col_a * col_b
```

Other common operations include the following, assuming a dataframe contains *col_a* and *col_b* columns:

- Concatenate two columns: `concat(col_a, col_b)`
- Add two columns: `col_a + col_b`
- Subtract two columns: `col_a - col_b`
- Divide two columns: `col_a / col_b`
- Take the absolute value of a column: `abs(col_a)`
For more information, see the Spark documentation on selecting data.

**Reduce Dimensionality within a Dataset**

Reduce the dimensionality in your data by using Principal Component Analysis (PCA). The dimensionality of your dataset corresponds to the number of features. When you use dimensionality reduction in Data Wrangler, you get a new set of features called components. Each component accounts for some variability in the data.

The first component accounts for the largest amount of variation in the data. The second component accounts for the second largest amount of variation in the data, and so on.

You can use dimensionality reduction to reduce the size of the data sets that you use to train models. Instead of using the features in your dataset, you can use the principal components instead.

To perform PCA, Data Wrangler creates axes for your data. An axis is an affine combination of columns in your dataset. The first principal component is the value on the axis that has the largest amount of variance. The second principal component is the value on the axis that has the second largest amount of variance. The nth principal component is the value on the axis that has the nth largest amount of variance.

You can configure the number of principal components that Data Wrangler returns. You can either specify the number of principal components directly or you can specify the variance threshold percentage. Each principal component explains an amount of variance in the data. For example, you might have a principal component with a value of 0.5. The component would explain 50% of the variation in the data. When you specify a variance threshold percentage, Data Wrangler returns the smallest number of components that meet the percentage that you specify.

The following are example principal components with the amount of variance that they explain in the data.

- Component 1 – 0.5
- Component 2 – 0.45
- Component 3 – 0.05

If you specify a variance threshold percentage of 94 or 95, Data Wrangler returns Component 1 and Component 2. If you specify a variance threshold percentage of 96, Data Wrangler returns all three principal components.
You can use the following procedure to run PCA on your dataset.

To run PCA on your dataset, do the following.

1. Open your Data Wrangler data flow.
2. Choose the +, and select Add transform.
3. Choose Add step.
4. Choose Dimensionality Reduction.
5. For Input Columns, choose the features that you’re reducing into the principal components.
6. (Optional) For Number of principal components, choose the number of principal components that Data Wrangler returns in your dataset. If specify a value for the field, you can't specify a value for Variance threshold percentage.
7. (Optional) For Variance threshold percentage, specify the percentage of variation in the data that you want explained by the principal components. Data Wrangler uses the default value of 95 if you don't specify a value for the variance threshold. You can't specify a variance threshold percentage if you've specified a value for Number of principal components.
8. (Optional) Deselect Center to not use the mean of the columns as the center of the data. By default, Data Wrangler centers the data with the mean before scaling.
9. (Optional) Deselect Scale to not scale the data with the unit standard deviation.
10. (Optional) Choose Columns to output the components to separate columns. Choose Vector to output the components as a single vector.
11. (Optional) For Output column, specify a name for an output column. If you're outputting the components to separate columns, the name that you specify is a prefix. If you're outputting the components to a vector, the name that you specify is the name of the vector column.
12. (Optional) Select Keep input columns. We don't recommend selecting this option if you plan on only using the principal components to train your model.

**Encode Categorical**

Categorical data is usually composed of a finite number of categories, where each category is represented with a string. For example, if you have a table of customer data, a column that indicates the country a person lives in is categorical. The categories would be Afghanistan, Albania,
Algeria, and so on. Categorical data can be **nominal** or **ordinal**. Ordinal categories have an inherent order, and nominal categories do not. The highest degree obtained (High school, Bachelors, Masters, and so on) is an example of ordinal categories.

Encoding categorical data is the process of creating a numerical representation for categories. For example, if your categories are Dog and Cat, you may encode this information into two vectors, [1, 0] to represent Dog, and [0, 1] to represent Cat.

When you encode ordinal categories, you may need to translate the natural order of categories into your encoding. For example, you can represent the highest degree obtained with the following map: {"High school": 1, "Bachelors": 2, "Masters":3}.

Use categorical encoding to encode categorical data that is in string format into arrays of integers.

The Data Wrangler categorical encoders create encodings for all categories that exist in a column at the time the step is defined. If new categories have been added to a column when you start a Data Wrangler job to process your dataset at time t, and this column was the input for a Data Wrangler categorical encoding transform at time t-1, these new categories are considered **missing** in the Data Wrangler job. The option you select for **Invalid handling strategy** is applied to these missing values. Examples of when this can occur are:

- When you use a .flow file to create a Data Wrangler job to process a dataset that was updated after the creation of the data flow. For example, you may use a data flow to regularly process sales data each month. If that sales data is updated weekly, new categories may be introduced into columns for which an encode categorical step is defined.
- When you select **Sampling** when you import your dataset, some categories may be left out of the sample.

In these situations, these new categories are considered missing values in the Data Wrangler job.

You can choose from and configure an **ordinal** and a **one-hot encode**. Use the following sections to learn more about these options.

Both transforms create a new column named **Output column name**. You specify the output format of this column with **Output style**:

- Select **Vector** to produce a single column with a sparse vector.
- Select **Columns** to create a column for every category with an indicator variable for whether the text in the original column contains a value that is equal to that category.
Ordinal Encode

Select **Ordinal encode** to encode categories into an integer between 0 and the total number of categories in the **Input column** you select.

**Invalid handing strategy**: Select a method to handle invalid or missing values.

- Choose **Skip** if you want to omit the rows with missing values.
- Choose **Keep** to retain missing values as the last category.
- Choose **Error** if you want Data Wrangler to throw an error if missing values are encountered in the **Input column**.
- Choose **Replace with NaN** to replace missing with NaN. This option is recommended if your ML algorithm can handle missing values. Otherwise, the first three options in this list may produce better results.

One-Hot Encode

Select **One-hot encode** for **Transform** to use one-hot encoding. Configure this transform using the following:

- **Drop last category**: If True, the last category does not have a corresponding index in the one-hot encoding. When missing values are possible, a missing category is always the last one and setting this to True means that a missing value results in an all zero vector.
- **Invalid handing strategy**: Select a method to handle invalid or missing values.
  - Choose **Skip** if you want to omit the rows with missing values.
  - Choose **Keep** to retain missing values as the last category.
  - Choose **Error** if you want Data Wrangler to throw an error if missing values are encountered in the **Input column**.
- **Is input ordinal encoded**: Select this option if the input vector contains ordinal encoded data. This option requires that input data contain non-negative integers. If True, input \( i \) is encoded as a vector with a non-zero in the \( i \)th location.

Similarity encode

Use similarity encoding when you have the following:

- A large number of categorical variables
Noisy data

The similarity encoder creates embeddings for columns with categorical data. An embedding is a mapping of discrete objects, such as words, to vectors of real numbers. It encodes similar strings to vectors containing similar values. For example, it creates very similar encodings for "California" and "Calfornia".

Data Wrangler converts each category in your dataset into a set of tokens using a 3-gram tokenizer. It converts the tokens into an embedding using min-hash encoding.

The following example shows how the similarity encoder creates vectors from strings.
The similarity encodings that Data Wrangler creates:

- Have low dimensionality
- Are scalable to a large number of categories
- Are robust and resistant to noise

For the preceding reasons, similarity encoding is more versatile than one-hot encoding.

To add the similarity encoding transform to your dataset, use the following procedure.

To use similarity encoding, do the following.

1. Sign in to the Amazon SageMaker Console.
2. Choose Open Studio Classic.
3. Choose Launch app.
4. Choose Studio.
5. Specify your data flow.
6. Choose a step with a transformation.
7. Choose Add step.
8. Choose Encode categorical.
9. Specify the following:
• **Transform – Similarity encode**

• **Input column** – The column containing the categorical data that you’re encoding.

• **Target dimension** – (Optional) The dimension of the categorical embedding vector. The default value is 30. We recommend using a larger target dimension if you have a large dataset with many categories.

• **Output style** – Choose Vector for a single vector with all of the encoded values. Choose Column to have the encoded values in separate columns.

• **Output column** – (Optional) The name of the output column for a vector encoded output. For a column-encoded output, this is the prefix of the column names followed by listed number.

**Featurize Text**

Use the **Featurize Text** transform group to inspect string-typed columns and use text embedding to featurize these columns.

This feature group contains two features, **Character statistics** and Vectorize. Use the following sections to learn more about these transforms. For both options, the **Input column** must contain text data (string type).

**Character Statistics**

Use **Character statistics** to generate statistics for each row in a column containing text data.

This transform computes the following ratios and counts for each row, and creates a new column to report the result. The new column is named using the input column name as a prefix and a suffix that is specific to the ratio or count.

• **Number of words**: The total number of words in that row. The suffix for this output column is -stats_word_count.

• **Number of characters**: The total number of characters in that row. The suffix for this output column is -stats_char_count.

• **Ratio of upper**: The number of uppercase characters, from A to Z, divided by all characters in the column. The suffix for this output column is -stats_capital_ratio.

• **Ratio of lower**: The number of lowercase characters, from a to z, divided by all characters in the column. The suffix for this output column is -stats_lower_ratio.
• **Ratio of digits**: The ratio of digits in a single row over the sum of digits in the input column. The suffix for this output column is `-stats_digit_ratio`.

• **Special characters ratio**: The ratio of non-alphanumeric (characters like #$&%:@) characters to over the sum of all characters in the input column. The suffix for this output column is `-stats_special_ratio`.

**Vectorize**

Text embedding involves mapping words or phrases from a vocabulary to vectors of real numbers. Use the Data Wrangler text embedding transform to tokenize and vectorize text data into term frequency–inverse document frequency (TF-IDF) vectors.

When TF-IDF is calculated for a column of text data, each word in each sentence is converted to a real number that represents its semantic importance. Higher numbers are associated with less frequent words, which tend to be more meaningful.

When you define a Vectorize transform step, Data Wrangler uses the data in your dataset to define the count vectorizer and TF-IDF methods. Running a Data Wrangler job uses these same methods.

You configure this transform using the following:

• **Output column name**: This transform creates a new column with the text embedding. Use this field to specify a name for this output column.

• **Tokenizer**: A tokenizer converts the sentence into a list of words, or tokens.

  Choose **Standard** to use a tokenizer that splits by white space and converts each word to lowercase. For example, "Good dog" is tokenized to ["good", "dog"].

  Choose **Custom** to use a customized tokenizer. If you choose **Custom**, you can use the following fields to configure the tokenizer:

  • **Minimum token length**: The minimum length, in characters, for a token to be valid. Defaults to 1. For example, if you specify 3 for minimum token length, words like a, at, in are dropped from the tokenized sentence.

  • **Should regex split on gaps**: If selected, regex splits on gaps. Otherwise, it matches tokens. Defaults to `True`.

  • **Regex pattern**: Regex pattern that defines the tokenization process. Defaults to `' \s+'`.

  • **To lowercase**: If chosen, Data Wrangler converts all characters to lowercase before tokenization. Defaults to `True`. 
To learn more, see the Spark documentation on [Tokenizer](#).

- **Vectorizer**: The vectorizer converts the list of tokens into a sparse numeric vector. Each token corresponds to an index in the vector and a non-zero indicates the existence of the token in the input sentence. You can choose from two vectorizer options, Count and Hashing.

- **Count vectorize** allows customizations that filter infrequent or too common tokens. **Count vectorize parameters** include the following:
  - **Minimum term frequency**: In each row, terms (tokens) with smaller frequency are filtered. If you specify an integer, this is an absolute threshold (inclusive). If you specify a fraction between 0 (inclusive) and 1, the threshold is relative to the total term count. Defaults to 1.
  - **Minimum document frequency**: Minimum number of rows in which a term (token) must appear to be included. If you specify an integer, this is an absolute threshold (inclusive). If you specify a fraction between 0 (inclusive) and 1, the threshold is relative to the total term count. Defaults to 1.
  - **Maximum document frequency**: Maximum number of documents (rows) in which a term (token) can appear to be included. If you specify an integer, this is an absolute threshold (inclusive). If you specify a fraction between 0 (inclusive) and 1, the threshold is relative to the total term count. Defaults to 0.999.
  - **Maximum vocabulary size**: Maximum size of the vocabulary. The vocabulary is made up of all terms (tokens) in all rows of the column. Defaults to 262144.
  - **Binary outputs**: If selected, the vector outputs do not include the number of appearances of a term in a document, but rather are a binary indicator of its appearance. Defaults to False.

To learn more about this option, see the Spark documentation on [CountVectorizer](#).

- **Hashing** is computationally faster. **Hash vectorize parameters** includes the following:
  - **Number of features during hashing**: A hash vectorizer maps tokens to a vector index according to their hash value. This feature determines the number of possible hash values. Large values result in fewer collisions between hash values but a higher dimension output vector.

To learn more about this option, see the Spark documentation on [FeatureHasher](#).

- **Apply IDF** applies an IDF transformation, which multiplies the term frequency with the standard inverse document frequency used for TF-IDF embedding. **IDF parameters** include the following:
  - **Minimum document frequency**: Minimum number of documents (rows) in which a term (token) must appear to be included. If `count_vectorize` is the chosen vectorizer, we
recommend that you keep the default value and only modify the `min_doc_freq` field in `Count vectorize parameters`. Defaults to 5.

- **Output format**: The output format of each row.
  - Select **Vector** to produce a single column with a sparse vector.
  - Select **Flattened** to create a column for every category with an indicator variable for whether the text in the original column contains a value that is equal to that category. You can only choose flattened when **Vectorizer** is set as **Count vectorizer**.

**Transform Time Series**

In Data Wrangler, you can transform time series data. The values in a time series dataset are indexed to specific time. For example, a dataset that shows the number of customers in a store for each hour in a day is a time series dataset. The following table shows an example of a time series dataset.

**Hourly number of customers in a store**

<table>
<thead>
<tr>
<th>Number of customers</th>
<th>Time (hour)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>09:00</td>
</tr>
<tr>
<td>10</td>
<td>10:00</td>
</tr>
<tr>
<td>14</td>
<td>11:00</td>
</tr>
<tr>
<td>25</td>
<td>12:00</td>
</tr>
<tr>
<td>20</td>
<td>13:00</td>
</tr>
<tr>
<td>18</td>
<td>14:00</td>
</tr>
</tbody>
</table>

For the preceding table, the **Number of Customers** column contains the time series data. The time series data is indexed on the hourly data in the **Time (hour)** column.

You might need to perform a series of transformations on your data to get it in a format that you can use for your analysis. Use the **Time series** transform group to transform your time series data. For more information about the transformations that you can perform, see the following sections.

**Topics**
Group by a Time Series

You can use the group by operation to group time series data for specific values in a column.

For example, you have the following table that tracks the average daily electricity usage in a household.

**Average daily household electricity usage**

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Daily timestamp</th>
<th>Electricity usage (kWh)</th>
<th>Number of household occupants</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>1/1/2020</td>
<td>30</td>
<td>2</td>
</tr>
<tr>
<td>household_0</td>
<td>1/2/2020</td>
<td>40</td>
<td>2</td>
</tr>
<tr>
<td>household_0</td>
<td>1/4/2020</td>
<td>35</td>
<td>3</td>
</tr>
<tr>
<td>household_1</td>
<td>1/2/2020</td>
<td>45</td>
<td>3</td>
</tr>
<tr>
<td>household_1</td>
<td>1/3/2020</td>
<td>55</td>
<td>4</td>
</tr>
</tbody>
</table>

If you choose to group by ID, you get the following table.
Electricity usage grouped by household ID

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
<th>Number of household occupants series</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35]</td>
<td>[2, 2, 3]</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, 55]</td>
<td>[3, 4]</td>
</tr>
</tbody>
</table>

Each entry in the time series sequence is ordered by the corresponding timestamp. The first element of the sequence corresponds to the first timestamp of the series. For household_0, 30 is the first value of the Electricity Usage Series. The value of 30 corresponds to the first timestamp of 1/1/2020.

You can include the starting timestamp and ending timestamp. The following table shows how that information appears.

Electricity usage grouped by household ID

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
<th>Number of household occupants series</th>
<th>Start_time</th>
<th>End_time</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35]</td>
<td>[2, 2, 3]</td>
<td>1/1/2020</td>
<td>1/4/2020</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, 55]</td>
<td>[3, 4]</td>
<td>1/2/2020</td>
<td>1/3/2020</td>
</tr>
</tbody>
</table>

You can use the following procedure to group by a time series column.

1. Open your Data Wrangler data flow.
2. If you haven't imported your dataset, import it under the Import data tab.
3. In your data flow, under Data types, choose the +, and select Add transform.
5. Choose Time Series.
7. Specify a column in Group by this column.
8. For Apply to columns, specify a value.
9. Choose **Preview** to generate a preview of the transform.

10. Choose **Add** to add the transform to the Data Wrangler data flow.

### Resample Time Series Data

Time series data usually has observations that aren't taken at regular intervals. For example, a dataset could have some observations that are recorded hourly and other observations that are recorded every two hours.

Many analyses, such as forecasting algorithms, require the observations to be taken at regular intervals. Resampling gives you the ability to establish regular intervals for the observations in your dataset.

You can either upsample or downsample a time series. Downsampling increases the interval between observations in the dataset. For example, if you downsample observations that are taken either every hour or every two hours, each observation in your dataset is taken every two hours. The hourly observations are aggregated into a single value using an aggregation method such as the mean or median.

Upsampling reduces the interval between observations in the dataset. For example, if you upsample observations that are taken every two hours into hourly observations, you can use an interpolation method to infer hourly observations from the ones that have been taken every two hours. For information on interpolation methods, see `pandas.DataFrame.interpolate`.

You can resample both numeric and non-numeric data.

Use the **Resample** operation to resample your time series data. If you have multiple time series in your dataset, Data Wrangler standardizes the time interval for each time series.

The following table shows an example of downsampling time series data by using the mean as the aggregation method. The data is downsampled from every two hours to every hour.

### Hourly temperature readings over a day before downsampling

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Temperature (Celsius)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:00</td>
<td>30</td>
</tr>
<tr>
<td>1:00</td>
<td>32</td>
</tr>
<tr>
<td>Timestamp</td>
<td>Temperature (Celsius)</td>
</tr>
<tr>
<td>-----------</td>
<td>----------------------</td>
</tr>
<tr>
<td>2:00</td>
<td>35</td>
</tr>
<tr>
<td>3:00</td>
<td>32</td>
</tr>
<tr>
<td>4:00</td>
<td>30</td>
</tr>
</tbody>
</table>

Temperature readings downsampled to every two hours

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Temperature (Celsius)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:00</td>
<td>30</td>
</tr>
<tr>
<td>2:00</td>
<td>33.5</td>
</tr>
<tr>
<td>4:00</td>
<td>35</td>
</tr>
</tbody>
</table>

You can use the following procedure to resample time series data.

1. Open your Data Wrangler data flow.
2. If you haven’t imported your dataset, import it under the **Import data** tab.
3. In your data flow, under **Data types**, choose the +, and select **Add transform**.
4. Choose **Add step**.
5. Choose **Resample**.
6. For **Timestamp**, choose the timestamp column.
7. For **Frequency unit**, specify the frequency that you're resampling.
8. (Optional) Specify a value for **Frequency quantity**.
9. Configure the transform by specifying the remaining fields.
10. Choose **Preview** to generate a preview of the transform.
11. Choose **Add** to add the transform to the Data Wrangler data flow.

**Handle Missing Time Series Data**

If you have missing values in your dataset, you can do one of the following:
For datasets that have multiple time series, drop the time series that have missing values that are greater than a threshold that you specify.

Impute the missing values in a time series by using other values in the time series.

Imputing a missing value involves replacing the data by either specifying a value or by using an inferential method. The following are the methods that you can use for imputation:

- **Constant value** – Replace all the missing data in your dataset with a value that you specify.
- **Most common value** – Replace all the missing data with the value that has the highest frequency in the dataset.
- **Forward fill** – Use a forward fill to replace the missing values with the non-missing value that precedes the missing values. For the sequence: [2, 4, 7, NaN, NaN, NaN, 8], all of the missing values are replaced with 7. The sequence that results from using a forward fill is [2, 4, 7, 7, 7, 7, 8].
- **Backward fill** – Use a backward fill to replace the missing values with the non-missing value that follows the missing values. For the sequence: [2, 4, 7, NaN, NaN, NaN, 8], all of the missing values are replaced with 8. The sequence that results from using a backward fill is [2, 4, 7, 8, 8, 8, 8].
- **Interpolate** – Uses an interpolation function to impute the missing values. For more information on the functions that you can use for interpolation, see `pandas.DataFrame.interpolate`.

Some of the imputation methods might not be able to impute all the missing value in your dataset. For example, a **Forward fill** can’t impute a missing value that appears at the beginning of the time series. You can impute the values by using either a forward fill or a backward fill.

You can either impute missing values within a cell or within a column.

The following example shows how values are imputed within a cell.

**Electricity usage with missing values**

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35, NaN, NaN]</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, NaN, 55]</td>
</tr>
</tbody>
</table>
Electricity usage with values imputed using a forward fill

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage series (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>[30, 40, 35, 35, 35]</td>
</tr>
<tr>
<td>household_1</td>
<td>[45, 45, 55]</td>
</tr>
</tbody>
</table>

The following example shows how values are imputed within a column.

**Average daily household electricity usage with missing values**

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>30</td>
</tr>
<tr>
<td>household_0</td>
<td>40</td>
</tr>
<tr>
<td>household_0</td>
<td>NaN</td>
</tr>
<tr>
<td>household_1</td>
<td>NaN</td>
</tr>
<tr>
<td>household_1</td>
<td>NaN</td>
</tr>
</tbody>
</table>

**Average daily household electricity usage with values imputed using a forward fill**

<table>
<thead>
<tr>
<th>Household ID</th>
<th>Electricity usage (kWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td>household_0</td>
<td>30</td>
</tr>
<tr>
<td>household_0</td>
<td>40</td>
</tr>
<tr>
<td>household_0</td>
<td>40</td>
</tr>
<tr>
<td>household_1</td>
<td>40</td>
</tr>
<tr>
<td>household_1</td>
<td>40</td>
</tr>
</tbody>
</table>

You can use the following procedure to handle missing values.
1. Open your Data Wrangler data flow.
2. If you haven't imported your dataset, import it under the Import data tab.
3. In your data flow, under Data types, choose the +, and select Add transform.
5. Choose Handle missing.
6. For Time series input type, choose whether you want to handle missing values inside of a cell or along a column.
7. For Impute missing values for this column, specify the column that has the missing values.
8. For Method for imputing values, select a method.
9. Configure the transform by specifying the remaining fields.
10. Choose Preview to generate a preview of the transform.
11. If you have missing values, you can specify a method for imputing them under Method for imputing values.
12. Choose Add to add the transform to the Data Wrangler data flow.

Validate the Timestamp of Your Time Series Data

You might have time stamp data that is invalid. You can use the Validate time stamp function to determine whether the timestamps in your dataset are valid. Your timestamp can be invalid for one or more of the following reasons:

- Your timestamp column has missing values.
- The values in your timestamp column are not formatted correctly.

If you have invalid timestamps in your dataset, you can't perform your analysis successfully. You can use Data Wrangler to identify invalid timestamps and understand where you need to clean your data.

The time series validation works in one of the two ways:

You can configure Data Wrangler to do one of the following if it encounters missing values in your dataset:

- Drop the rows that have the missing or invalid values.
• Identify the rows that have the missing or invalid values.

• Throw an error if it finds any missing or invalid values in your dataset.

You can validate the timestamps on columns that either have the timestamp type or the string type. If the column has the string type, Data Wrangler converts the type of the column to timestamp and performs the validation.

You can use the following procedure to validate the timestamps in your dataset.

1. Open your Data Wrangler data flow.
2. If you haven’t imported your dataset, import it under the Import data tab.
3. In your data flow, under Data types, choose the +, and select Add transform.
5. Choose Validate timestamps.
6. For Timestamp Column, choose the timestamp column.
7. For Policy, choose whether you want to handle missing timestamps.
8. (Optional) For Output column, specify a name for the output column.
9. If the date time column is formatted for the string type, choose Cast to datetime.
10. Choose Preview to generate a preview of the transform.
11. Choose Add to add the transform to the Data Wrangler data flow.

**Standardizing the Length of the Time Series**

If you have time series data stored as arrays, you can standardize each time series to the same length. Standardizing the length of the time series array might make it easier for you to perform your analysis on the data.

You can standardize your time series for data transformations that require the length of your data to be fixed.

Many ML algorithms require you to flatten your time series data before you use them. Flattening time series data is separating each value of the time series into its own column in a dataset. The number of columns in a dataset can't change, so the lengths of the time series need to be standardized between you flatten each array into a set of features.
Each time series is set to the length that you specify as a quantile or percentile of the time series set. For example, you can have three sequences that have the following lengths:

- 3
- 4
- 5

You can set the length of all of the sequences as the length of the sequence that has the 50th percentile length.

Time series arrays that are shorter than the length you've specified have missing values added. The following is an example format of standardizing the time series to a longer length: [2, 4, 5, NaN, NaN, NaN].

You can use different approaches to handle the missing values. For information on those approaches, see [Handle Missing Time Series Data](#).

The time series arrays that are longer than the length that you specify are truncated.

You can use the following procedure to standardize the length of the time series.

1. Open your Data Wrangler data flow.
2. If you haven’t imported your dataset, import it under the Import data tab.
3. In your data flow, under Data types, choose the +, and select Add transform.
5. Choose Standardize length.
6. For Standardize the time series length for the column, choose a column.
7. (Optional) For Output column, specify a name for the output column. If you don’t specify a name, the transform is done in place.
8. If the datetime column is formatted for the string type, choose Cast to datetime.
9. Choose Cutoff quantile and specify a quantile to set the length of the sequence.
10. Choose Flatten the output to output the values of the time series into separate columns.
11. Choose Preview to generate a preview of the transform.
12. Choose Add to add the transform to the Data Wrangler data flow.
Extract Features from Your Time Series Data

If you're running a classification or a regression algorithm on your time series data, we recommend extracting features from the time series before running the algorithm. Extracting features might improve the performance of your algorithm.

Use the following options to choose how you want to extract features from your data:

- **Use Minimal subset** to specify extracting 8 features that you know are useful in downstream analyses. You can use a minimal subset when you need to perform computations quickly. You can also use it when your ML algorithm has a high risk of overfitting and you want to provide it with fewer features.

- **Use Efficient subset** to specify extracting the most features possible without extracting features that are computationally intensive in your analyses.

- **Use All features** to specify extracting all features from the time series.

- **Use Manual subset** to choose a list of features that you think explain the variation in your data well.

Use the following the procedure to extract features from your time series data.

1. Open your Data Wrangler data flow.
2. If you haven't imported your dataset, import it under the **Import data** tab.
3. In your data flow, under **Data types**, choose the +, and select **Add transform**.
4. Choose **Add step**.
5. Choose **Extract features**.
6. For **Extract features for this column**, choose a column.
7. (Optional) Select **Flatten** to output the features into separate columns.
8. For **Strategy**, choose a strategy to extract the features.
9. Choose **Preview** to generate a preview of the transform.
10. Choose **Add** to add the transform to the Data Wrangler data flow.

Use Lagged Features from Your Time Series Data

For many use cases, the best way to predict the future behavior of your time series is to use its most recent behavior.
The most common uses of lagged features are the following:

- Collecting a handful of past values. For example, for time, $t + 1$, you collect $t$, $t - 1$, $t - 2$, and $t - 3$.
- Collecting values that correspond to seasonal behavior in the data. For example, to predict the occupancy in a restaurant at 1:00 PM, you might want to use the features from 1:00 PM on the previous day. Using the features from 12:00 PM or 11:00 AM on the same day might not be as predictive as using the features from previous days.

1. Open your Data Wrangler data flow.
2. If you haven’t imported your dataset, import it under the Import data tab.
3. In your data flow, under Data types, choose the +, and select Add transform.
5. Choose Lag features.
6. For Generate lag features for this column, choose a column.
7. For Timestamp Column, choose the column containing the timestamps.
8. For Lag, specify the duration of the lag.
9. (Optional) Configure the output using one of the following options:
   - Include the entire lag window
   - Flatten the output
   - Drop rows without history
10. Choose Preview to generate a preview of the transform.
11. Choose Add to add the transform to the Data Wrangler data flow.

**Create a Datetime Range In Your Time Series**

You might have time series data that don’t have timestamps. If you know that the observations were taken at regular intervals, you can generate timestamps for the time series in a separate column. To generate timestamps, you specify the value for the start timestamp and the frequency of the timestamps.

For example, you might have the following time series data for the number of customers at a restaurant.
Time series data on the number of customers at a restaurant

<table>
<thead>
<tr>
<th>Number of customers</th>
<th>Timestamp</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>1:00 PM</td>
</tr>
<tr>
<td>14</td>
<td>2:00 PM</td>
</tr>
<tr>
<td>24</td>
<td>3:00 PM</td>
</tr>
<tr>
<td>40</td>
<td>4:00 PM</td>
</tr>
<tr>
<td>30</td>
<td>5:00 PM</td>
</tr>
<tr>
<td>20</td>
<td>6:00 PM</td>
</tr>
</tbody>
</table>

If you know that the restaurant opened at 5:00 PM and that the observations are taken hourly, you can add a timestamp column that corresponds to the time series data. You can see the timestamp column in the following table.

Time series data on the number of customers at a restaurant

<table>
<thead>
<tr>
<th>Number of customers</th>
<th>Timestamp</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>1:00 PM</td>
</tr>
<tr>
<td>14</td>
<td>2:00 PM</td>
</tr>
<tr>
<td>24</td>
<td>3:00 PM</td>
</tr>
<tr>
<td>40</td>
<td>4:00 PM</td>
</tr>
<tr>
<td>30</td>
<td>5:00 PM</td>
</tr>
<tr>
<td>20</td>
<td>6:00 PM</td>
</tr>
</tbody>
</table>

Use the following procedure to add a datetime range to your data.

1. Open your Data Wrangler data flow.
2. If you haven't imported your dataset, import it under the **Import data** tab.
3. In your data flow, under **Data types**, choose the +, and select **Add transform**.
4. Choose **Add step**.
5. Choose **Datetime range**.
6. For **Frequency type**, choose the unit used to measure the frequency of the timestamps.
7. For **Starting timestamp**, specify the start timestamp.
8. For **Output column**, specify a name for the output column.
9. (Optional) Configure the output using the remaining fields.
10. Choose **Preview** to generate a preview of the transform.
11. Choose **Add** to add the transform to the Data Wrangler data flow.

**Use a Rolling Window In Your Time Series**

You can extract features over a time period. For example, for time, \( t \), and a time window length of 3, and for the row that indicates the \( t \)th timestamp, we append the features that are extracted from the time series at times \( t - 3 \), \( t - 2 \), and \( t - 1 \). For information on extracting features, see [Extract Features from Your Time Series Data](#).

You can use the following procedure to extract features over a time period.

1. Open your Data Wrangler data flow.
2. If you haven't imported your dataset, import it under the **Import data** tab.
3. In your data flow, under **Data types**, choose the +, and select **Add transform**.
4. Choose **Add step**.
5. Choose **Rolling window features**.
6. For **Generate rolling window features for this column**, choose a column.
7. For **Timestamp Column**, choose the column containing the timestamps.
8. (Optional) For **Output Column**, specify the name of the output column.
9. For **Window size**, specify the window size.
10. For **Strategy**, choose the extraction strategy.
11. Choose **Preview** to generate a preview of the transform.
12. Choose **Add** to add the transform to the Data Wrangler data flow.
Featurize Datetime

Use Featurize date/time to create a vector embedding representing a datetime field. To use this transform, your datetime data must be in one of the following formats:

- Strings describing datetime: For example, "January 1st, 2020, 12:44pm".
- A Unix timestamp: A Unix timestamp describes the number of seconds, milliseconds, microseconds, or nanoseconds from 1/1/1970.

You can choose to Infer datetime format and provide a Datetime format. If you provide a datetime format, you must use the codes described in the Python documentation. The options you select for these two configurations have implications for the speed of the operation and the final results.

- The most manual and computationally fastest option is to specify a Datetime format and select No for Infer datetime format.
- To reduce manual labor, you can choose Infer datetime format and not specify a datetime format. It is also a computationally fast operation; however, the first datetime format encountered in the input column is assumed to be the format for the entire column. If there are other formats in the column, these values are NaN in the final output. Inferring the datetime format can give you unparsed strings.
- If you don't specify a format and select No for Infer datetime format, you get the most robust results. All the valid datetime strings are parsed. However, this operation can be an order of magnitude slower than the first two options in this list.

When you use this transform, you specify an Input column which contains datetime data in one of the formats listed above. The transform creates an output column named Output column name. The format of the output column depends on your configuration using the following:

- Vector: Outputs a single column as a vector.
- Columns: Creates a new column for every feature. For example, if the output contains a year, month, and day, three separate columns are created for year, month, and day.

Additionally, you must choose an Embedding mode. For linear models and deep networks, we recommend choosing cyclic. For tree-based algorithms, we recommend choosing ordinal.
Format String

The Format string transforms contain standard string formatting operations. For example, you can use these operations to remove special characters, normalize string lengths, and update string casing.

This feature group contains the following transforms. All transforms return copies of the strings in the Input column and add the result to a new, output column.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left pad</td>
<td>Left-pad the string with a given Fill character to the given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Right pad</td>
<td>Right-pad the string with a given Fill character to the given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Center (pad on either side)</td>
<td>Center-pad the string (add padding on both sides of the string) with a given Fill character to the given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Prepend zeros</td>
<td>Left-fill a numeric string with zeros, up to a given width. If the string is longer than width, the return value is shortened to width characters.</td>
</tr>
<tr>
<td>Strip left and right</td>
<td>Returns a copy of the string with the leading and trailing characters removed.</td>
</tr>
<tr>
<td>Strip characters from left</td>
<td>Returns a copy of the string with leading characters removed.</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Strip characters from right</td>
<td>Returns a copy of the string with trailing characters removed.</td>
</tr>
<tr>
<td>Lower case</td>
<td>Convert all letters in text to lowercase.</td>
</tr>
<tr>
<td>Upper case</td>
<td>Convert all letters in text to uppercase.</td>
</tr>
<tr>
<td>Capitalize</td>
<td>Capitalize the first letter in each sentence.</td>
</tr>
<tr>
<td>Swap case</td>
<td>Converts all uppercase characters to lowercase and all lowercase characters to uppercase characters of the given string, and returns it.</td>
</tr>
<tr>
<td>Add prefix or suffix</td>
<td>Adds a prefix and a suffix the string column. You must specify at least one of Prefix and Suffix.</td>
</tr>
<tr>
<td>Remove symbols</td>
<td>Removes given symbols from a string. All listed characters are removed. Defaults to white space.</td>
</tr>
</tbody>
</table>

### Handle Outliers

Machine learning models are sensitive to the distribution and range of your feature values. Outliers, or rare values, can negatively impact model accuracy and lead to longer training times. Use this feature group to detect and update outliers in your dataset.

When you define a **Handle outliers** transform step, the statistics used to detect outliers are generated on the data available in Data Wrangler when defining this step. These same statistics are used when running a Data Wrangler job.

Use the following sections to learn more about the transforms this group contains. You specify an **Output name** and each of these transforms produces an output column with the resulting data.

#### Robust standard deviation numeric outliers

This transform detects and fixes outliers in numeric features using statistics that are robust to outliers.
You must define an **Upper quantile** and a **Lower quantile** for the statistics used to calculate outliers. You must also specify the number of **Standard deviations** from which a value must vary from the mean to be considered an outlier. For example, if you specify 3 for **Standard deviations**, a value must fall more than 3 standard deviations from the mean to be considered an outlier.

The **Fix method** is the method used to handle outliers when they are detected. You can choose from the following:

- **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.
- **Remove**: Use this option to remove rows with outliers from the dataframe.
- **Invalidate**: Use this option to replace outliers with invalid values.

**Standard Deviation Numeric Outliers**

This transform detects and fixes outliers in numeric features using the mean and standard deviation.

You specify the number of **Standard deviations** a value must vary from the mean to be considered an outlier. For example, if you specify 3 for **Standard deviations**, a value must fall more than 3 standard deviations from the mean to be considered an outlier.

The **Fix method** is the method used to handle outliers when they are detected. You can choose from the following:

- **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.
- **Remove**: Use this option to remove rows with outliers from the dataframe.
- **Invalidate**: Use this option to replace outliers with invalid values.

**Quantile Numeric Outliers**

Use this transform to detect and fix outliers in numeric features using quantiles. You can define an **Upper quantile** and a **Lower quantile**. All values that fall above the upper quantile or below the lower quantile are considered outliers.

The **Fix method** is the method used to handle outliers when they are detected. You can choose from the following:

- **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.
• **Remove**: Use this option to remove rows with outliers from the dataframe.
• **Invalidate**: Use this option to replace outliers with invalid values.

**Min-Max Numeric Outliers**

This transform detects and fixes outliers in numeric features using upper and lower thresholds. Use this method if you know threshold values that demark outliers.

You specify a **Upper threshold** and a **Lower threshold**, and if values fall above or below those thresholds respectively, they are considered outliers.

The **Fix method** is the method used to handle outliers when they are detected. You can choose from the following:

• **Clip**: Use this option to clip the outliers to the corresponding outlier detection bound.
• **Remove**: Use this option to remove rows with outliers from the dataframe.
• **Invalidate**: Use this option to replace outliers with invalid values.

**Replace Rare**

When you use the **Replace rare** transform, you specify a threshold and Data Wrangler finds all values that meet that threshold and replaces them with a string that you specify. For example, you may want to use this transform to categorize all outliers in a column into an "Others" category.

• **Replacement string**: The string with which to replace outliers.
• **Absolute threshold**: A category is rare if the number of instances is less than or equal to this absolute threshold.
• **Fraction threshold**: A category is rare if the number of instances is less than or equal to this fraction threshold multiplied by the number of rows.
• **Max common categories**: Maximum not-rare categories that remain after the operation. If the threshold does not filter enough categories, those with the top number of appearances are classified as not rare. If set to 0 (default), there is no hard limit to the number of categories.

**Handle Missing Values**

Missing values are a common occurrence in machine learning datasets. In some situations, it is appropriate to impute missing data with a calculated value, such as an average or categorically
common value. You can process missing values using the **Handle missing values** transform group. This group contains the following transforms.

### Fill Missing

Use the **Fill missing** transform to replace missing values with a **Fill value** you define.

### Impute Missing

Use the **Impute missing** transform to create a new column that contains imputed values where missing values were found in input categorical and numerical data. The configuration depends on your data type.

For numeric data, choose an imputing strategy, the strategy used to determine the new value to impute. You can choose to impute the mean or the median over the values that are present in your dataset. Data Wrangler uses the value that it computes to impute the missing values.

For categorical data, Data Wrangler imputes missing values using the most frequent value in the column. To impute a custom string, use the **Fill missing** transform instead.

### Add Indicator for Missing

Use the **Add indicator for missing** transform to create a new indicator column, which contains a Boolean "false" if a row contains a value, and "true" if a row contains a missing value.

### Drop Missing

Use the **Drop missing** option to drop rows that contain missing values from the **Input column**.

### Manage Columns

You can use the following transforms to quickly update and manage columns in your dataset:

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drop Column</td>
<td>Delete a column.</td>
</tr>
<tr>
<td>Duplicate Column</td>
<td>Duplicate a column.</td>
</tr>
<tr>
<td>Rename Column</td>
<td>Rename a column.</td>
</tr>
<tr>
<td>Move Column</td>
<td>Move a column's location in the dataset. Choose to move your column to the start or...</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>--------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>end of the dataset, before or after a reference column, or to a specific index.</td>
</tr>
</tbody>
</table>

**Manage Rows**

Use this transform group to quickly perform sort and shuffle operations on rows. This group contains the following:

- **Sort**: Sort the entire dataframe by a given column. Select the check box next to **Ascending order** for this option; otherwise, deselect the check box and descending order is used for the sort.
- **Shuffle**: Randomly shuffle all rows in the dataset.

**Manage Vectors**

Use this transform group to combine or flatten vector columns. This group contains the following transforms.

- **Assemble**: Use this transform to combine Spark vectors and numeric data into a single column. For example, you can combine three columns: two containing numeric data and one containing vectors. Add all the columns you want to combine in **Input columns** and specify a **Output column name** for the combined data.
- **Flatten**: Use this transform to flatten a single column containing vector data. The input column must contain PySpark vectors or array-like objects. You can control the number of columns created by specifying a **Method to detect number of outputs**. For example, if you select **Length of first vector**, the number of elements in the first valid vector or array found in the column determines the number of output columns that are created. All other input vectors with too many items are truncated. Inputs with too few items are filled with NaNs.

  You also specify an **Output prefix**, which is used as the prefix for each output column.

**Process Numeric**

Use the **Process Numeric** feature group to process numeric data. Each scalar in this group is defined using the Spark library. The following scalars are supported:
- **Standard Scaler**: Standardize the input column by subtracting the mean from each value and scaling to unit variance. To learn more, see the Spark documentation for `StandardScaler`.

- **Robust Scaler**: Scale the input column using statistics that are robust to outliers. To learn more, see the Spark documentation for `RobustScaler`.

- **Min Max Scaler**: Transform the input column by scaling each feature to a given range. To learn more, see the Spark documentation for `MinMaxScaler`.

- **Max Absolute Scaler**: Scale the input column by dividing each value by the maximum absolute value. To learn more, see the Spark documentation for `MaxAbsScaler`.

**Sampling**

After you've imported your data, you can use the **Sampling** transformer to take one or more samples of it. When you use the sampling transformer, Data Wrangler samples your original dataset.

You can choose one of the following sample methods:

- **Limit**: Samples the dataset starting from the first row up to the limit that you specify.
- **Randomized**: Takes a random sample of a size that you specify.
- **Stratified**: Takes a stratified random sample.

You can stratify a randomized sample to make sure that it represents the original distribution of the dataset.

You might be performing data preparation for multiple use cases. For each use case, you can take a different sample and apply a different set of transformations.

The following procedure describes the process of creating a random sample.

To take a random sample from your data.

1. Choose the + to the right of the dataset that you've imported. The name of your dataset is located below the +.
2. Choose Add transform.
3. Choose Sampling.
4. For Sampling method, choose the sampling method.
5. For **Approximate sample size**, choose the approximate number of observations that you want in your sample.

6. (Optional) Specify an integer for **Random seed** to create a reproducible sample.

The following procedure describes the process of creating a stratified sample.

To take a stratified sample from your data.

1. Choose the + to the right of the dataset that you've imported. The name of your dataset is located below the +.
2. Choose **Add transform**.
3. Choose **Sampling**.
4. For **Sampling method**, choose the sampling method.
5. For **Approximate sample size**, choose the approximate number of observations that you want in your sample.
6. For **Stratify column**, specify the name of the column that you want to stratify on.
7. (Optional) Specify an integer for **Random seed** to create a reproducible sample.

**Search and Edit**

Use this section to search for and edit specific patterns within strings. For example, you can find and update strings within sentences or documents, split strings by delimiters, and find occurrences of specific strings.

The following transforms are supported under **Search and edit**. All transforms return copies of the strings in the **Input column** and add the result to a new output column.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Find substring</td>
<td>Returns the index of the first occurrence of the <strong>Substring</strong> for which you searched. You can start and end the search at <strong>Start</strong> and <strong>End</strong> respectively.</td>
</tr>
<tr>
<td>Find substring (from right)</td>
<td>Returns the index of the last occurrence of the <strong>Substring</strong> for which you searched. You</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Matches prefix</td>
<td>Returns a Boolean value if the string contains a given Pattern. A pattern can be a character sequence or regular expression. Optionally, you can make the pattern case sensitive.</td>
</tr>
<tr>
<td>Find all occurrences</td>
<td>Returns an array with all occurrences of a given pattern. A pattern can be a character sequence or regular expression.</td>
</tr>
<tr>
<td>Extract using regex</td>
<td>Returns a string that matches a given Regex pattern.</td>
</tr>
<tr>
<td>Extract between delimiters</td>
<td>Returns a string with all characters found between Left delimiter and Right delimiter.</td>
</tr>
<tr>
<td>Extract from position</td>
<td>Returns a string, starting from Start position in the input string, that contains all characters up to the start position plus Length.</td>
</tr>
<tr>
<td>Find and replace substring</td>
<td>Returns a string with all matches of a given Pattern (regular expression) replaced by Replacement string.</td>
</tr>
<tr>
<td>Replace between delimiters</td>
<td>Returns a string with the substring found between the first appearance of a Left delimiter and the last appearance of a Right delimiter replaced by Replacement string. If no match is found, nothing is replaced.</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>---------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Replace from position</td>
<td>Returns a string with the substring between Start position and Start position plus Length replaced by Replacement string. If Start position plus Length is greater than the length of the replacement string, the output contains ....</td>
</tr>
<tr>
<td>Convert regex to missing</td>
<td>Converts a string to None if invalid and returns the result. Validity is defined with a regular expression in Pattern.</td>
</tr>
<tr>
<td>Split string by delimiter</td>
<td>Returns an array of strings from the input string, split by Delimiter, with up to Max number of splits (optional). The delimiter defaults to white space.</td>
</tr>
</tbody>
</table>

**Split data**

Use the **Split data** transform to split your dataset into two or three datasets. For example, you can split your dataset into a dataset used to train your model and a dataset used to test it. You can determine the proportion of the dataset that goes into each split. For example, if you’re splitting one dataset into two datasets, the training dataset can have 80% of the data while the testing dataset has 20%.

Splitting your data into three datasets gives you the ability to create training, validation, and test datasets. You can see how well the model performs on the test dataset by dropping the target column.

Your use case determines how much of the original dataset each of your datasets get and the method you use to split the data. For example, you might want to use a stratified split to make sure that the distribution of the observations in the target column are the same across datasets. You can use the following split transforms:

- **Randomized split** — Each split is a random, non-overlapping sample of the original dataset. For larger datasets, using a randomized split might be computationally expensive and take longer than an ordered split.
- **Ordered split** – Splits the dataset based on the sequential order of the observations. For example, for an 80/20 train-test split, the first observations that make up 80% of the dataset go to the training dataset. The last 20% of the observations go to the testing dataset. Ordered splits are effective in keeping the existing order of the data between splits.

- **Stratified split** – Splits the dataset to make sure that the number of observations in the input column have proportional representation. For an input column that has the observations 1, 1, 1, 1, 2, 2, 2, 2, 2, 2, 3, 3, 3, 3, 3, 3, 3, 3, an 80/20 split on the column would mean that approximately 80% of the 1s, 80% of the 2s, and 80% of the 3s go to the training set. About 20% of each type of observation go to the testing set.

- **Split by key** – Avoids data with the same key occurring in more than one split. For example, if you have a dataset with the column 'customer_id' and you're using it as a key, no customer id is in more than one split.

After you split the data, you can apply additional transformations to each dataset. For most use cases, they aren't necessary.

Data Wrangler calculates the proportions of the splits for performance. You can choose an error threshold to set the accuracy of the splits. Lower error thresholds more accurately reflect the proportions that you specify for the splits. If you set a higher error threshold, you get better performance, but lower accuracy.

For perfectly split data, set the error threshold to 0. You can specify a threshold between 0 and 1 for better performance. If you specify a value greater than 1, Data Wrangler interprets that value as 1.

If you have 10000 rows in your dataset and you specify an 80/20 split with an error of 0.001, you would get observations approximating one of the following results:

- 8010 observations in the training set and 1990 in the testing set
- 7990 observations in the training set and 2010 in the testing set

The number of observations for the testing set in the preceding example is in the interval between 8010 and 7990.

By default, Data Wrangler uses a random seed to make the splits reproducible. You can specify a different value for the seed to create a different reproducible split.
Randomized split

Use the following procedure to perform a randomized split on your dataset.

To split your dataset randomly, do the following

1. Choose the + next to the node containing the dataset that you're splitting.
2. Choose Add transform.
3. Choose Split data.
4. (Optional) For Splits, specify the names and proportions of each split. The proportions must sum to 1.
5. (Optional) Choose the + to create an additional split.
   - Specify the names and proportions of all the splits. The proportions must sum to 1.
6. (Optional) Specify a value for Error threshold other than the default value.
7. (Optional) Specify a value for Random seed.

Ordered split

Use the following procedure to perform an ordered split on your dataset.

To make an ordered split in your dataset, do the following.

1. Choose the + next to the node containing the dataset that you're splitting.
2. Choose Add transform.
3. For Transform, choose Ordered split.
4. Choose Split data.
5. (Optional) For Splits, specify the names and proportions of each split. The proportions must sum to 1.
6. (Optional) Choose the + to create an additional split.
   - Specify the names and proportions of all the splits. The proportions must sum to 1.
7. (Optional) Specify a value for Error threshold other than the default value.
8. (Optional) For **Input column**, specify a column with numeric values. Uses the values of the columns to infer which records are in each split. The smaller values are in one split with the larger values in the other splits.

9. (Optional) Select **Handle duplicates** to add noise to duplicate values and create a dataset of entirely unique values.

10. (Optional) Specify a value for **Random seed**.

11. Choose **Preview**.

12. Choose **Add**.

---

**Stratified split**

Use the following procedure to perform a stratified split on your dataset.

To make a stratified split in your dataset, do the following.

1. Choose the + next to the node containing the dataset that you're splitting.

2. Choose **Add transform**.

3. Choose **Split data**.

4. For **Transform**, choose **Stratified split**.

5. (Optional) For **Splits**, specify the names and proportions of each split. The proportions must sum to 1.

6. (Optional) Choose the + to create an additional split.

   - Specify the names and proportions of all the splits. The proportions must sum to 1.

7. For **Input column**, specify a column with up to 100 unique values. Data Wrangler can't stratify a column with more than 100 unique values.

8. (Optional) Specify a value for **Error threshold** other than the default value.

9. (Optional) Specify a value for **Random seed** to specify a different seed.

10. Choose **Preview**.

11. Choose **Add**.

---

**Split by column keys**

Use the following procedure to split by the column keys in your dataset.
To split by the column keys in your dataset, do the following.

1. Choose the + next to the node containing the dataset that you're splitting.
2. Choose Add transform.
3. Choose Split data.
4. For Transform, choose Split by key.
5. (Optional) For Splits, specify the names and proportions of each split. The proportions must sum to 1.
6. (Optional) Choose the + to create an additional split.
   - Specify the names and proportions of all the splits. The proportions must sum to 1.
7. For Key columns, specify the columns with values that you don't want to appear in both datasets.
8. (Optional) Specify a value for Error threshold other than the default value.
10. Choose Add.

Parse Value as Type

Use this transform to cast a column to a new type. The supported Data Wrangler data types are:

- Long
- Float
- Boolean
- Date, in the format dd-MM-yyyy, representing day, month, and year respectively.
- String

Validate String

Use the Validate string transforms to create a new column that indicates that a row of text data meets a specified condition. For example, you can use a Validate string transform to verify that a string only contains lowercase characters. The following transforms are supported under Validate string.
The following transforms are included in this transform group. If a transform outputs a Boolean value, True is represented with a 1 and False is represented with a 0.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>String length</td>
<td>Returns True if a string length equals specified length. Otherwise, returns False.</td>
</tr>
<tr>
<td>Starts with</td>
<td>Returns True if a string starts will a specified prefix. Otherwise, returns False.</td>
</tr>
<tr>
<td>Ends with</td>
<td>Returns True if a string length equals specified length. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is alphanumeric</td>
<td>Returns True if a string only contains numbers and letters. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is alpha (letters)</td>
<td>Returns True if a string only contains letters. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is digit</td>
<td>Returns True if a string only contains digits. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is space</td>
<td>Returns True if a string only contains numbers and letters. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is title</td>
<td>Returns True if a string contains any white spaces. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is lowercase</td>
<td>Returns True if a string only contains lower case letters. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is uppercase</td>
<td>Returns True if a string only contains upper case letters. Otherwise, returns False.</td>
</tr>
<tr>
<td>Is numeric</td>
<td>Returns True if a string only contains numbers. Otherwise, returns False.</td>
</tr>
<tr>
<td>Name</td>
<td>Function</td>
</tr>
<tr>
<td>------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Is decimal</td>
<td>Returns True if a string only contains decimal numbers. Otherwise, returns False.</td>
</tr>
</tbody>
</table>

**Unnest JSON Data**

If you have a .csv file, you might have values in your dataset that are JSON strings. Similarly, you might have nested data in columns of either a Parquet file or a JSON document.

Use the **Flatten structured** operator to separate the first level keys into separate columns. A first level key is a key that isn't nested within a value.

For example, you might have a dataset that has a `person` column with demographic information on each person stored as JSON strings. A JSON string might look like the following.

```json
{"seq": 1,"name": {"first": "Nathaniel","last": "Ferguson"},"age": 59,"city": "Posbotno","state": "WV"}
```

The **Flatten structured** operator converts the following first level keys into additional columns in your dataset:

- seq
- name
- age
- city
- state

Data Wrangler puts the values of the keys as values under the columns. The following shows the column names and values of the JSON.

<table>
<thead>
<tr>
<th>seq, name, age, city, state</th>
</tr>
</thead>
<tbody>
<tr>
<td>1, {&quot;first&quot;: &quot;Nathaniel&quot;,&quot;last&quot;: &quot;Ferguson&quot;}, 59, Posbotno, WV</td>
</tr>
</tbody>
</table>
For each value in your dataset containing JSON, the **Flatten structured** operator creates columns for the first-level keys. To create columns for nested keys, call the operator again. For the preceding example, calling the operator creates the columns:

- name_first
- name_last

The following example shows the dataset that results from calling the operation again.

<table>
<thead>
<tr>
<th>seq, name,</th>
<th>age, city, state, name_first, name_last</th>
</tr>
</thead>
<tbody>
<tr>
<td>1, {&quot;first&quot;: &quot;Nathaniel&quot;,&quot;last&quot;: &quot;Ferguson&quot;}, 59, Posbotno, WV, Nathaniel, Ferguson</td>
<td></td>
</tr>
</tbody>
</table>

Choose **Keys to flatten on** to specify the first-level keys that want to extract as separate columns. If you don't specify any keys, Data Wrangler extracts all the keys by default.

**Explode Array**

Use **Explode array** to expand the values of the array into separate output rows. For example, the operation can take each value in the array, [[1, 2, 3], [4, 5, 6], [7, 8, 9]] and create a new column with the following rows:

- [1, 2, 3]
- [4, 5, 6]
- [7, 8, 9]

Data Wrangler names the new column, input_column_name_flatten.

You can call the **Explode array** operation multiple times to get the nested values of the array into separate output columns. The following example shows the result of calling the operation multiple times on a dataset with a nested array.
Putting the values of a nested array into separate columns

<table>
<thead>
<tr>
<th>id</th>
<th>array</th>
<th>id</th>
<th>array_items</th>
<th>id</th>
<th>array_items</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>[ [cat, dog], [bat, frog] ]</td>
<td>1</td>
<td>[cat, dog]</td>
<td>1</td>
<td>cat</td>
</tr>
<tr>
<td>2</td>
<td>[[rose, petunia], [lily, daisy]]</td>
<td>1</td>
<td>[bat, frog]</td>
<td>1</td>
<td>dog</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>[rose, petunia]</td>
<td>1</td>
<td>bat</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>[lily, daisy]</td>
<td>1</td>
<td>frog</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>rose</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>petunia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>lily</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>2</td>
<td>daisy</td>
</tr>
</tbody>
</table>

Transform Image Data

Use Data Wrangler to import and transform the images that you're using for your machine learning (ML) pipelines. After you've prepared your image data, you can export it from your Data Wrangler flow to your ML pipeline.

You can use the information provided here to familiarize yourself with importing and transforming image data in Data Wrangler. Data Wrangler uses OpenCV to import images. For more information about supported image formats, see [Image file reading and writing](#).

After you've familiarized yourself with the concepts of transforming your image data, go through the following tutorial, [Prepare image data with Amazon SageMaker Data Wrangler](#).

The following industries and use cases are examples where applying machine learning to transformed image data can be useful:
• Manufacturing – Identifying defects in items from the assembly line
• Food – Identifying spoiled or rotten food
• Medicine – Identifying lesions in tissues

When you work with image data in Data Wrangler, you go through the following process:

1. Import – Select the images by choosing the directory containing them in your Amazon S3 bucket.
2. Transform – Use the built-in transformations to prepare the images for your machine learning pipeline.
3. Export – Export the images that you’ve transformed to a location that can be accessed from the pipeline.

Use the following procedure to import your image data.

To import your image data

1. Navigate to the Create connection page.
2. Choose Amazon S3.
3. Specify the Amazon S3 file path that contains the image data.
4. For File type, choose Image.
5. (Optional) Choose Import nested directories to import images from multiple Amazon S3 paths.
6. Choose Import.

Data Wrangler uses the open-source imgaug library for its built-in image transformations. You can use the following built-in transformations:

• ResizeImage
• EnhanceImage
• CorruptImage
• SplitImage
• DropCorruptedImages
- DropImageDuplicates
- Brightness
- ColorChannels
- Grayscale
- Rotate

Use the following procedure to transform your images without writing code.

**To transform the image data without writing code**

1. From your Data Wrangler flow, choose the + next to the node representing the images that you've imported.
2. Choose *Add transform*.
3. Choose *Add step*.
4. Choose the transform and configure it.
5. Choose *Preview*.
6. Choose *Add*.

In addition to using the transformations that Data Wrangler provides, you can also use your own custom code snippets. For more information about using custom code snippets, see [Custom Transforms](#). You can import the OpenCV and imgaug libraries within your code snippets and use the transforms associated with them. The following is an example of a code snippet that detects edges within the images.

```python
# A table with your image data is stored in the `df` variable
import cv2
import numpy as np
from pyspark.sql.functions import column
from sagemaker_dataprep.compute.operators.transforms.image.constants import DEFAULT_IMAGE_COLUMN, IMAGE_COLUMN_TYPE
from sagemaker_dataprep.compute.operators.transforms.image.decorators import BasicImageOperationDecorator, PandasUDFOperationDecorator

@BasicImageOperationDecorator
Transform Data
```
When apply transformations in your Data Wrangler flow, Data Wrangler only applies them to a sample of the images in your dataset. To optimize your experience with the application, Data Wrangler doesn't apply the transforms to all of your images.

To apply the transformations to all of your images, export your Data Wrangler flow to an Amazon S3 location. You can use the images that you've exported in your training or inference pipelines. Use a destination node or a Jupyter Notebook to export your data. You can access either method for exporting your data from the Data Wrangler flow. For information about using these methods, see Export to Amazon S3.

Filter data

Use Data Wrangler to filter the data in your columns. When you filter the data in a column, you specify the following fields:

- **Column name** – The name of the column that you're using to filter the data.
- **Condition** – The type of filter that you're applying to values in the column.
- **Value** – The value or category in the column to which you're applying the filter.

You can filter on the following conditions:

- `=` – Returns values that match the value or category that you specify.
- `!=` – Returns values that don't match the value or category that you specify.
• >= – For Long or Float data, filters for values that are greater than or equal to the value that you specify.
• <= – For Long or Float data, filters for values that are less than or equal to the value that you specify.
• > – For Long or Float data, filters for values that are greater than the value that you specify.
• < – For Long or Float data, filters for values that are less than the value that you specify.

For a column that has the categories, male and female, you can filter out all the male values. You could also filter for all the female values. Because there are only male and female values in the column, the filter returns a column that only has female values.

You can also add multiple filters. The filters can be applied across multiple columns or the same column. For example, if you're creating a column that only has values within a certain range, you add two different filters. One filter specifies that the column must have values greater than the value that you provide. The other filter specifies that the column must have values less than the value that you provide.

Use the following procedure to add the filter transform to your data.

To filter your data

1. From your Data Wrangler flow, choose the + next to the node with the data that you're filtering.
2. Choose Add transform.
3. Choose Add step.
4. Choose Filter data.
5. Specify the following fields:
   - Column name – The column that you're filtering.
   - Condition – The condition of the filter.
   - Value – The value or category in the column to which you're applying the filter.
6. (Optional) Choose + following the filter that you've created.
7. Configure the filter.
Map Columns for Amazon Personalize

Data Wrangler integrates with Amazon Personalize, a fully managed machine learning service that generates item recommendations and user segments. You can use the **Map columns for Amazon Personalize** transform to get your data into a format that Amazon Personalize can interpret. For more information about the transforms specific to Amazon Personalize, see [Importing data using Amazon SageMaker Data Wrangler](#). For more information about Amazon Personalize see [What is Amazon Personalize?](#)

Analyze and Visualize

Amazon SageMaker Data Wrangler includes built-in analyses that help you generate visualizations and data analyses in a few clicks. You can also create custom analyses using your own code.

You add an analysis to a dataframe by selecting a step in your data flow, and then choosing **Add analysis**. To access an analysis you've created, select the step that contains the analysis, and select the analysis.

All analyses are generated using 100,000 rows of your dataset.

You can add the following analysis to a dataframe:

- Data visualizations, including histograms and scatter plots.
- A quick summary of your dataset, including number of entries, minimum and maximum values (for numeric data), and most and least frequent categories (for categorical data).
- A quick model of the dataset, which can be used to generate an importance score for each feature.
- A target leakage report, which you can use to determine if one or more features are strongly correlated with your target feature.
- A custom visualization using your own code.

Use the following sections to learn more about these options.

**Histogram**

Use histograms to see the counts of feature values for a specific feature. You can inspect the relationships between features using the **Color by** option. For example, the following histogram
charts the distribution of user ratings of the best-selling books on Amazon from 2009–2019, colored by genre.

You can use the **Facet by** feature to create histograms of one column, for each value in another column. For example, the following diagram shows histograms of user reviews of best-selling books on Amazon if faceted by year.
Scatter Plot

Use the **Scatter Plot** feature to inspect the relationship between features. To create a scatter plot, select a feature to plot on the X axis and the Y axis. Both of these columns must be numeric typed columns.

You can color scatter plots by an additional column. For example, the following example shows a scatter plot comparing the number of reviews against user ratings of top-selling books on Amazon between 2009 and 2019. The scatter plot is colored by book genre.
Additionally, you can facet scatter plots by features. For example, the following image shows an example of the same review versus user rating scatter plot, faceted by year.
Table Summary

Use the **Table Summary** analysis to quickly summarize your data.

For columns with numerical data, including log and float data, a table summary reports the number of entries (count), minimum (min), maximum (max), mean, and standard deviation (stddev) for each column.

For columns with non-numerical data, including columns with string, Boolean, or date/time data, a table summary reports the number of entries (count), least frequent value (min), and most frequent value (max).

Quick Model

Use the **Quick Model** visualization to quickly evaluate your data and produce importance scores for each feature. A feature importance score score indicates how useful a feature is at predicting a target label. The feature importance score is between [0, 1] and a higher number indicates that the feature is more important to the whole dataset. On the top of the quick model chart, there is a model score. A classification problem shows an F1 score. A regression problem has a mean squared error (MSE) score.

When you create a quick model chart, you select a dataset you want evaluated, and a target label against which you want feature importance to be compared. Data Wrangler does the following:

- Infers the data types for the target label and each feature in the dataset selected.
- Determines the problem type. Based on the number of distinct values in the label column, Data Wrangler determines if this is a regression or classification problem type. Data Wrangler sets a categorical threshold to 100. If there are more than 100 distinct values in the label column, Data Wrangler classifies it as a regression problem; otherwise, it is classified as a classification problem.
- Pre-processes features and label data for training. The algorithm used requires encoding features to vector type and encoding labels to double type.
- Trains a random forest algorithm with 70% of data. Spark's RandomForestRegressor is used to train a model for regression problems. The RandomForestClassifier is used to train a model for classification problems.
- Evaluates a random forest model with the remaining 30% of data. Data Wrangler evaluates classification models using an F1 score and evaluates regression models using an MSE score.
- Calculates feature importance for each feature using the Gini importance method.
The following image shows the user interface for the quick model feature.

Target Leakage

Target leakage occurs when there is data in a machine learning training dataset that is strongly correlated with the target label, but is not available in real-world data. For example, you may have a column in your dataset that serves as a proxy for the column you want to predict with your model.

When you use the **Target Leakage** analysis, you specify the following:

- **Target**: This is the feature about which you want your ML model to be able to make predictions.
- **Problem type**: This is the ML problem type on which you are working. Problem type can either be **classification** or **regression**.
- **(Optional) Max features**: This is the maximum number of features to present in the visualization, which shows features ranked by their risk of being target leakage.

For classification, the target leakage analysis uses the area under the receiver operating characteristic, or AUC - ROC curve for each column, up to **Max features**. For regression, it uses a coefficient of determination, or R2 metric.
The AUC - ROC curve provides a predictive metric, computed individually for each column using cross-validation, on a sample of up to around 1000 rows. A score of 1 indicates perfect predictive abilities, which often indicates target leakage. A score of 0.5 or lower indicates that the information on the column could not provide, on its own, any useful information towards predicting the target. Although it can happen that a column is uninformative on its own but is useful in predicting the target when used in tandem with other features, a low score could indicate the feature is redundant.

For example, the following image shows a target leakage report for a diabetes classification problem, that is, predicting if a person has diabetes or not. An AUC - ROC curve is used to calculate the predictive ability of five features, and all are determined to be safe from target leakage.

Multicollinearity

Multicollinearity is a circumstance where two or more predictor variables are related to each other. The predictor variables are the features in your dataset that you're using to predict a target variable. When you have multicollinearity, the predictor variables are not only predictive of the target variable, but also predictive of each other.
You can use the **Variance Inflation Factor (VIF)**, **Principal Component Analysis (PCA)**, or **Lasso feature selection** as measures for the multicollinearity in your data. For more information, see the following.

**Variance Inflation Factor (VIF)**

The Variance Inflation Factor (VIF) is a measure of collinearity among variable pairs. Data Wrangler returns a VIF score as a measure of how closely the variables are related to each other. A VIF score is a positive number that is greater than or equal to 1.

A score of 1 means that the variable is uncorrelated with the other variables. Scores greater than 1 indicate higher correlation.

Theoretically, you can have a VIF score with a value of infinity. Data Wrangler clips high scores to 50. If you have a VIF score greater than 50, Data Wrangler sets the score to 50.

You can use the following guidelines to interpret your VIF scores:

- A VIF score less than or equal to 5 indicates that the variables are moderately correlated with the other variables.
- A VIF score greater than or equal to 5 indicates that the variables are highly correlated with the other variables.

** Principle Component Analysis (PCA)**

Principal Component Analysis (PCA) measures the variance of the data along different directions in the feature space. The feature space consists of all the predictor variables that you use to predict the target variable in your dataset.

For example, if you're trying to predict who survived on the *RMS Titanic* after it hit an iceberg, your feature space can include the passengers' age, gender, and the fare that they paid.

From the feature space, PCA generates an ordered list of variances. These variances are also known as singular values. The values in the list of variances are greater than or equal to 0. We can use them to determine how much multicollinearity there is in our data.

When the numbers are roughly uniform, the data has very few instances of multicollinearity. When there is a lot of variability among the values, we have many instances of multicollinearity. Before it performs PCA, Data Wrangler normalizes each feature to have a mean of 0 and a standard deviation of 1.
Note

PCA in this circumstance can also be referred to as Singular Value Decomposition (SVD).

Lasso feature selection

Lasso feature selection uses the L1 regularization technique to only include the most predictive features in your dataset.

For both classification and regression, the regularization technique generates a coefficient for each feature. The absolute value of the coefficient provides an importance score for the feature. A higher importance score indicates that it is more predictive of the target variable. A common feature selection method is to use all the features that have a non-zero lasso coefficient.

Detect Anomalies In Time Series Data

You can use the anomaly detection visualization to see outliers in your time series data. To understand what determines an anomaly, you need to understand that we decompose the time series into a predicted term and an error term. We treat the seasonality and trend of the time series as the predicted term. We treat the residuals as the error term.

For the error term, you specify a threshold as the number of standard of deviations the residual can be away from the mean for it to be considered an anomaly. For example, you can specify a threshold as being 3 standard deviations. Any residual greater than 3 standard deviations away from the mean is an anomaly.

You can use the following procedure to perform an **Anomaly detection** analysis.

1. Open your Data Wrangler data flow.
2. In your data flow, under **Data types**, choose the +, and select **Add analysis**.
3. For **Analysis type**, choose **Time Series**.
4. For **Visualization**, choose **Anomaly detection**.
5. For **Anomaly threshold**, choose the threshold that a value is considered an anomaly.
6. Choose **Preview** to generate a preview of the analysis.
7. Choose **Add** to add the transform to the Data Wrangler data flow.
Seasonal Trend Decomposition In Time Series Data

You can determine whether there's seasonality in your time series data by using the Seasonal Trend Decomposition visualization. We use the STL (Seasonal Trend decomposition using LOESS) method to perform the decomposition. We decompose the time series into its seasonal, trend, and residual components. The trend reflects the long term progression of the series. The seasonal component is a signal that recurs in a time period. After removing the trend and the seasonal components from the time series, you have the residual.

You can use the following procedure to perform a Seasonal-Trend decomposition analysis.

1. Open your Data Wrangler data flow.
2. In your data flow, under Data types, choose the +, and select Add analysis.
3. For Analysis type, choose Time Series.
4. For Visualization, choose Seasonal-Trend decomposition.
5. For Anomaly threshold, choose the threshold that a value is considered an anomaly.
6. Choose Preview to generate a preview of the analysis.
7. Choose Add to add the transform to the Data Wrangler data flow.

Bias Report

You can use the bias report in Data Wrangler to uncover potential biases in your data. To generate a bias report, you must specify the target column, or Label, that you want to predict and a Facet, or the column that you want to inspect for biases.

Label: The feature about which you want a model to make predictions. For example, if you are predicting customer conversion, you may select a column containing data on whether or not a customer has placed an order. You must also specify whether this feature is a label or a threshold. If you specify a label, you must specify what a positive outcome looks like in your data. In the customer conversion example, a positive outcome may be a 1 in the orders column, representing the positive outcome of a customer placing an order within the last three months. If you specify a threshold, you must specify a lower bound defining a positive outcome. For example, if your customer orders columns contains the number of orders placed in the last year, you may want to specify 1.

Facet: The column that you want to inspect for biases. For example, if you are trying to predict customer conversion, your facet may be the age of the customer. You may choose this facet
because you believe that your data is biased toward a certain age group. You must identify whether the facet is measured as a value or threshold. For example, if you wanted to inspect one or more specific ages, you select **Value** and specify those ages. If you want to look at an age group, you select **Threshold** and specify the threshold of ages you want to inspect.

After you select your feature and label, you select the types of bias metrics you want to calculate.

To learn more, see [*Generate reports for bias in pre-training data*](#).

### Create Custom Visualizations

You can add an analysis to your Data Wrangler flow to create a custom visualization. Your dataset, with all the transformations you've applied, is available as a [Pandas DataFrame](https://pandas.pydata.org/pandas-docs/stable/). Data Wrangler uses the `df` variable to store the dataframe. You access the dataframe by calling the variable.

You must provide the output variable, `chart`, to store an [Altair](https://altair-viz.github.io/) output chart. For example, you can use the following code block to create a custom histogram using the Titanic dataset.

```python
import altair as alt
df = df.iloc[:30]
df = df.rename(columns={'Age': 'value'})
df = df.assign(count=df.groupby('value').value.transform('count'))
df = df[['value', 'count']]  
base = alt.Chart(df)
bar = base.mark_bar().encode(x=alt.X('value', bin=True, axis=None), y=alt.Y('count'))
rule = base.mark_rule(color='red').encode(
    x='mean(value):Q',
    size=alt.value(5))
chart = bar + rule
```

To create a custom visualization:

1. Next to the node containing the transformation that you'd like to visualize, choose the +.
2. Choose **Add analysis**.
3. For **Analysis type**, choose **Custom Visualization**.
4. For **Analysis name**, specify a name.
5. Enter your code in the code box.
6. Choose **Preview** to preview your visualization.
7. Choose **Save** to add your visualization.
If you don’t know how to use the Altair visualization package in Python, you can use custom code snippets to help you get started.

Data Wrangler has a searchable collection of visualization snippets. To use a visualization snippet, choose **Search example snippets** and specify a query in the search bar.

The following example uses the **Binned scatterplot** code snippet. It plots a histogram for 2 dimensions.

The snippets have comments to help you understand the changes that you need to make to the code. You usually need to specify the column names of your dataset in the code.

```python
import altair as alt

# Specify the number of top rows for plotting
```
rows_number = 1000
df = df.head(rows_number)
# You can also choose bottom rows or randomly sampled rows
# df = df.tail(rows_number)
# df = df.sample(rows_number)

chart = (
    alt.Chart(df)
    .mark_circle()
    .encode(
        # Specify the column names for binning and number of bins for X and Y axis
        x=alt.X("col1:Q", bin=alt.Bin(maxbins=20)),
        y=alt.Y("col2:Q", bin=alt.Bin(maxbins=20)),
        size="count()",
    )
)

# :Q specifies that label column has quantitative type.
# For more details on Altair typing refer to
# https://altair-viz.github.io/user_guide/encoding.html#encoding-data-types

---

**Reusing Data Flows for Different Datasets**

For Amazon Simple Storage Service (Amazon S3) data sources, you can create and use parameters. A parameter is a variable that you've saved in your Data Wrangler flow. Its value can be any portion of the data source's Amazon S3 path. Use parameters to quickly change the data that you're importing into a Data Wrangler flow or exporting to a processing job. You can also use parameters to select and import a specific subset of your data.

After you created a Data Wrangler flow, you might have trained a model on the data that you've transformed. For datasets that have the same schema, you can use parameters to apply the same transformations on a different dataset and train a different model. You can use the new datasets to perform inference with your model or you could be using them to retrain your model.

In general, parameters have the following attributes:

- **Name** – The name you specify for the parameter
- **Type** – The type of value that the parameter represents
- **Default value** – The value of the parameter when you don't specify a new value
Note

Datetime parameters have a time range attribute that they use as the default value.

Data Wrangler uses curly braces, {{}}, to indicate that a parameter is being used in the Amazon S3 path. For example, you can have a URL such as s3://DOC-EXAMPLE-BUCKET1/{{example_parameter_name}}/example-dataset.csv.

You create a parameter when you're editing the Amazon S3 data source that you've imported. You can set any portion of the file path to a parameter value. You can set the parameter value to either a value or a pattern. The following are the available parameter value types in the Data Wrangler flow:

- Number
- String
- Pattern
- Datetime

Note

You can't create a pattern parameter or a datetime parameter for the name of the bucket in the Amazon S3 path.

You must set a number as the default value of a number parameter. You can change the value of the parameter to a different number when you're editing a parameter or when you're launching a processing job. For example, in the S3 path, s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file-1.csv, you can create a number parameter named number_parameter in the place of 1. Your S3 path now appears as s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file-{{number_parameter}}.csv. The path continues to point to the example-file-1.csv dataset until you change the value of the parameter. If you change the value of number_parameter to 2 the path is now s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file-2.csv. You can import example-file-2.csv into Data Wrangler if you've uploaded the file to that Amazon S3 location.
A string parameter stores a string as its default value. For example, in the S3 path, s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file-1.csv, you can create a string parameter named string_parameter in the place of the filename, example-file-1.csv. The path now appears as s3://DOC-EXAMPLE-BUCKET/example-prefix/{{string_parameter}}. It continues to match s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file-1.csv, until you change the value of the parameter.

Instead of specifying the filename as a string parameter, you can create a string parameter using the entire Amazon S3 path. You can specify a dataset from any Amazon S3 location in the string parameter.

A pattern parameter stores a regular expression (Python REGEX) string as its default value. You can use a pattern parameter to import multiple data files at the same time. To import more than one object at a time, specify a parameter value that matches the Amazon S3 objects that you're importing.

You can also create a pattern parameter for the following datasets:

- s3://DOC-EXAMPLE-BUCKET1/example-prefix/example-file-1.csv
- s3://DOC-EXAMPLE-BUCKET1/example-prefix/example-file-2.csv
- s3://DOC-EXAMPLE-BUCKET1/example-prefix/example-file-10.csv
- s3://DOC-EXAMPLE-BUCKET/example-prefix/example-file-0123.csv

For s3://DOC-EXAMPLE-BUCKET1/example-prefix/example-file-1.csv, you can create a pattern parameter in the place of 1, and set the default value of the parameter to \d+. The \d+ REGEX string matches any one or more decimal digits. If you create a pattern parameter named pattern_parameter, your S3 path appears as s3://DOC-EXAMPLE-BUCKET1/example-prefix/example-file-{{pattern_parameter}}.csv.

You can also use pattern parameters to match all CSV objects within your bucket. To match all objects in a bucket, create a pattern parameter with the default value of .* and set the path to s3://DOC-EXAMPLE-BUCKET/{{pattern_parameter}}.csv. The .* character matches any string character in the path.

The s3://DOC-EXAMPLE-BUCKET/{{pattern_parameter}}.csv path can match the following datasets.

- example-file-1.csv
• other-example-file.csv
• example-file-a.csv

A datetime parameter stores the format with the following information:

• A format for parsing strings inside an Amazon S3 path.
• A relative time range to limit the datetime values that match

For example, in the Amazon S3 file path, s3://DOC-EXAMPLE-BUCKET/2020/01/01/example-dataset.csv, 2020/01/01 represents a datetime in the format of year/month/day. You can set the parameter's time range to an interval such as 1 years or 24 hours. An interval of 1 years matches all S3 paths with datetimes that fall between the current time and the time exactly a year before the current time. The current time is the time when you start exporting the transformations that you've made to the data. For more information about exporting data, see Export. If the current date is 2022/01/01 and the time range is 1 years, the S3 path matches datasets such as the following:

• s3://DOC-EXAMPLE-BUCKET/2021/01/01/example-dataset.csv
• s3://DOC-EXAMPLE-BUCKET/2021/06/30/example-dataset.csv
• s3://DOC-EXAMPLE-BUCKET/2021/12/31/example-dataset.csv

The datetime values within a relative time range change as time passes. The S3 paths that fall within the relative time range might also differ.

For the Amazon S3 file path, s3://DOC-EXAMPLE-BUCKET1/20200101/example-dataset.csv, 20220101 is an example of a path that can become a datetime parameter.

To view a table of all parameters that you've created in Data Wrangler flow, choose the `{}` to the right of the text box containing the Amazon S3 path. If you no longer need a parameter that you've created, you can edit or delete. To edit or delete a parameter, choose icons to the right of the parameter.

⚠️ Important

Before you delete a parameter, make sure that you haven't used it anywhere in your Data Wrangler flow. Deleted parameters that are still within the flow cause errors.
You can create parameters for any step of your Data Wrangler flow. You can edit or delete any parameter that you create. If you're applying transformations to data that is no longer relevant to your use case, you can modify the values of parameters. Modifying the values of the parameters changes the data that you're importing.

The following sections provide additional examples and general guidance on using parameters. You can use the sections to understand the parameters that work best for you.

**Note**

The following sections contain procedures that use the Data Wrangler interface to override the parameters and create a processing job. You can also override the parameters by using the following procedures.

To export your Data Wrangler flow and override the value of a parameter, do the following.

1. Choose the + next to the node that you want to export.
2. Choose **Export to**.
3. Choose the location where you're exporting the data.
4. Under **parameter_overrides**, specify different values for the parameters that you've created.
5. Run the Jupyter Notebook.

---

**Applying a Data Wrangler flow to files using patterns**

You can use parameters to apply transformations in your Data Wrangler flow to different files that match a pattern in the Amazon S3 URI path. This helps you specify the files in your S3 bucket that you want to transform with high specificity. For example, you might have a dataset with the path `s3://DOC-EXAMPLE-BUCKET1/example-prefix-0/example-prefix-1/example-prefix-2/example-dataset.csv`. Different datasets named `example-dataset.csv` are stored under many different example prefixes. The prefixes might also be numbered sequentially. You can create patterns for the numbers in the Amazon S3 URI. Pattern parameters use REGEX to select any number of files that match the pattern of the expression. The following are REGEX patterns that might be useful:

- `.*` – Matches zero or more of any character, except newline characters
- `.+` – Matches one or more of any character, excluding newline characters
You can replace each number in the following paths with a single parameter that has a value of \d+.

- s3://DOC-EXAMPLE-BUCKET1/example-prefix-3/example-prefix-4/example-prefix-5/example-dataset.csv
- s3://DOC-EXAMPLE-BUCKET1/example-prefix-8/example-prefix-12/example-prefix-13/example-dataset.csv
- s3://DOC-EXAMPLE-BUCKET1/example-prefix-4/example-prefix-9/example-prefix-137/example-dataset.csv

The following procedure creates a pattern parameter for a dataset with the path s3://DOC-EXAMPLE-BUCKET1/example-prefix-0/example-prefix-1/example-prefix-2/example-dataset.csv.

To create a pattern parameter, do the following.

1. Next to the dataset that you've imported, choose Edit dataset.
2. Highlight the 0 in example-prefix-0.
3. Specify values for the following fields:
   - **Name** – A name for parameter
   - **Type** – Pattern
   - **Value** – \d+ a regular expression that corresponds to one or more digits
4. Choose Create.
5. Replace the 1 and the 2 in S3 URI path with the parameter. The path should have the following format: s3://DOC-EXAMPLE-BUCKET1/example-prefix-{{example_parameter_name}}/example-prefix-{{example_parameter_name}}/example-prefix-{{example_parameter_name}}/example-dataset.csv
The following is a general procedure for creating a pattern parameter.

1. Navigate to your Data Wrangler flow.
2. Next to the dataset that you've imported, choose Edit dataset.
3. Highlight the portion of the URI that you're using as the value of the pattern parameter.
4. Choose Create custom parameter.
5. Specify values for the following fields:
   - Name – A name for parameter
   - Type – Pattern
   - Value – A regular expression containing the pattern that you'd like to store.
6. Choose Create.

Applying a Data Wrangler flow to files using numeric values

You can use parameters to apply transformations in your Data Wrangler flow to different files that have similar paths. For example, you might have a dataset with the path s3://DOC-EXAMPLE-BUCKET1/example-prefix-0/example-prefix-1/example-prefix-2/example-dataset.csv.

You might have the transformations from your Data Wrangler flow that you've applied to datasets under example-prefix-1. You might want to apply the same transformations to example-dataset.csv that falls under example-prefix-10 or example-prefix-20.

You can create a parameter that stores the value 1. If you want to apply the transformations to different datasets, you can create processing jobs that replace the value of the parameter with a different value. The parameter acts as a placeholder for you to change when you want to apply the transformations from your Data Wrangler flow to new data. You can override the value of the parameter when you create a Data Wrangler processing job to apply the transformations in your Data Wrangler flow to different datasets.

Use the following procedure to create numeric parameters for s3://DOC-EXAMPLE-BUCKET1/example-prefix-0/example-prefix-1/example-prefix-2/example-dataset.csv.

To create parameters for the preceding S3 URI path, do the following.

1. Navigate to your Data Wrangler flow.
2. Next to the dataset that you've imported, choose Edit dataset.
3. Highlight the number in an example prefix of example-prefix-number.
4. Choose **Create custom parameter**.
5. For **Name**, specify a name for the parameter.
6. For **Type**, choose **Integer**.
7. For **Value**, specify the number.
8. Create parameters for the remaining numbers by repeating the procedure.

After you've created the parameters, apply the transforms to your dataset and create a destination node for them. For more information about destination nodes, see **Export**.

Use the following procedure to apply the transformations from your Data Wrangler flow to a different time range. It assumes that you've created a destination node for the transformations in your flow.

To change the value of a numeric parameter in a Data Wrangler processing job, do the following.

1. From your Data Wrangler flow, choose **Create job**
2. Select only the destination node that contains the transformations to the dataset containing the datetime parameters.
3. Choose **Configure job**.
4. Choose **Parameters**.
5. Choose the name of a parameter that you've created.
6. Change the value of the parameter.
7. Repeat the procedure for the other parameters.
8. Choose **Run**.

**Applying a Data Wrangler flow to files using strings**

You can use parameters to apply transformations in your Data Wrangler flow to different files that have similar paths. For example, you might have a dataset with the path `s3:///DOC-EXAMPLE-BUCKET1/example-prefix/example-dataset.csv`.

You might have transformations from your Data Wrangler flow that you've applied to datasets under example-prefix. You might want to apply the same transformations to example-dataset.csv under another-example-prefix or example-prefix-20.
You can create a parameter that stores the value `example-prefix`. If you want to apply the transformations to different datasets, you can create processing jobs that replace the value of the parameter with a different value. The parameter acts as a placeholder for you to change when you want to apply the transformations from your Data Wrangler flow to new data. You can override the value of the parameter when you create a Data Wrangler processing job to apply the transformations in your Data Wrangler flow to different datasets.

Use the following procedure to create a string parameter for `s3://DOC-EXAMPLE-BUCKET1/example-prefix/example-dataset.csv`.

To create a parameter for the preceding S3 URI path, do the following.

1. Navigate to your Data Wrangler flow.
2. Next to the dataset that you've imported, choose **Edit dataset**.
3. Highlight the example prefix, `example-prefix`.
4. Choose **Create custom parameter**.
5. For **Name**, specify a name for the parameter.
6. For **Type**, choose **String**.
7. For **Value**, specify the prefix.

After you've created the parameter, apply the transforms to your dataset and create a destination node for them. For more information about destination nodes, see **Export**.

Use the following procedure to apply the transformations from your Data Wrangler flow to a different time range. It assumes that you've created a destination node for the transformations in your flow.

To change the value of a numeric parameter in a Data Wrangler processing job, do the following:

1. From your Data Wrangler flow, choose **Create job**
2. Select only the destination node that contains the transformations to the dataset containing the datetime parameters.
3. Choose **Configure job**.
4. Choose **Parameters**.
5. Choose the name of a parameter that you've created.
6. Change the value of the parameter.
7. Repeat the procedure for the other parameters.


Applying a Data Wrangler flow to different datetime ranges

Use datetime parameters to apply transformations in your Data Wrangler flow to different time ranges. Highlight the portion of the Amazon S3 URI that has a timestamp and create a parameter for it. When you create a parameter, you specify a time range from the current time to a time in the past. For example, you might have an Amazon S3 URI that looks like the following: `s3://DOC-EXAMPLE-BUCKET1/example-prefix/2022/05/15/example-dataset.csv`. You can save 2022/05/15 as a datetime parameter. If you specify a year as the time range, the time range includes the moment that you run the processing job containing the datetime parameter and the time exactly one year ago. If the moment you're running the processing job is September 6th, 2022 or 2022/09/06, the time ranges can include the following:

- `s3://DOC-EXAMPLE-BUCKET1/example-prefix/2022/03/15/example-dataset.csv`
- `s3://DOC-EXAMPLE-BUCKET1/example-prefix/2022/01/08/example-dataset.csv`
- `s3://DOC-EXAMPLE-BUCKET1/example-prefix/2022/07/31/example-dataset.csv`
- `s3://DOC-EXAMPLE-BUCKET1/example-prefix/2021/09/07/example-dataset.csv`

The transformations in the Data Wrangler flow apply to all of the preceding prefixes. Changing the value of the parameter in the processing job doesn't change the value of the parameter in the Data Wrangler flow. To apply the transformations to datasets within a different time range, do the following:

1. Create a destination node containing all the transformations that you'd like to use.
2. Create a Data Wrangler job.
3. Configure the job to use a different time range for the parameter. Changing the value of the parameter in the processing job doesn't change the value of the parameter in the Data Wrangler flow.

For more information about destination nodes and Data Wrangler jobs, see Export.

The following procedure creates a datetime parameter for the Amazon S3 path: `s3://DOC-EXAMPLE-BUCKET1/example-prefix/2022/05/15/example-dataset.csv`. 
To create a datetime parameter for the preceding S3 URI path, do the following.

1. Navigate to your Data Wrangler flow.
2. Next to the dataset that you've imported, choose Edit dataset.
3. Highlight the portion of the URI that you're using as the value of the datetime parameter.
4. Choose Create custom parameter.
5. For Name, specify a name for the parameter.
6. For Type, choose Datetime.

**Note**

By default, Data Wrangler selects Predefined, which provides a dropdown menu for you to select a date format. However, the timestamp format that you're using might not be available. Instead of using Predefined as the default option, you can choose Custom and specify the timestamp format manually.

7. For Date format, open the dropdown menu following Predefined and choose yyyy/MM/dd. The format, yyyy/MM/dd, corresponds to the year/month/day of the timestamp.
8. For Timezone, choose a time zone.

**Note**

The data that you're analyzing might have time stamps taken in a different time zone from your time zone. Make sure that the time zone that you select matches the time zone of the data.

9. For Time range, specify the time range for the parameter.
10. (Optional) Enter a description to describe how you're using the parameter.
11. Choose Create.

After you've created the datetime parameters, apply the transforms to your dataset and create a destination node for them. For more information about destination nodes, see Export.

Use the following procedure to apply the transformations from your Data Wrangler flow to a different time range. It assumes that you've created a destination node for the transformations in your flow.
To change the value of a datetime parameter in a Data Wrangler processing job, do the following:

1. From your Data Wrangler flow, choose Create job
2. Select only the destination node that contains the transformations to the dataset containing the datetime parameters.
3. Choose Configure job.
4. Choose Parameters.
5. Choose the name of the datetime parameter that you've created.
6. For Time range, change the time range for the datasets.
7. Choose Run.

**Export**

In your Data Wrangler flow, you can export some or all of the transformations that you've made to your data processing pipelines.

A *Data Wrangler flow* is the series of data preparation steps that you've performed on your data. In your data preparation, you perform one or more transformations to your data. Each transformation is done using a transform step. The flow has a series of nodes that represent the import of your data and the transformations that you've performed. For an example of nodes, see the following image.
The preceding image shows a Data Wrangler flow with two nodes. The **Source - sampled** node shows the data source from which you've imported your data. The **Data types** node indicates that Data Wrangler has performed a transformation to convert the dataset into a usable format.

Each transformation that you add to the Data Wrangler flow appears as an additional node. For information on the transforms that you can add, see [Transform Data](#). The following image shows a Data Wrangler flow that has a **Rename-column** node to change the name of a column in a dataset.

You can export your data transformations to the following:

- Amazon S3
- SageMaker Pipelines
- Amazon SageMaker Feature Store
- Python Code
Important

We recommend that you use the IAM AmazonSageMakerFullAccess managed policy to grant AWS permission to use Data Wrangler. If you don't use the managed policy, you can use an IAM policy that gives Data Wrangler access to an Amazon S3 bucket. For more information on the policy, see [Security and Permissions](#).

When you export your data flow, you're charged for the AWS resources that you use. You can use cost allocation tags to organize and manage the costs of those resources. You create these tags for your user-profile and Data Wrangler automatically applies them to the resources used to export the data flow. For more information, see [Using Cost Allocation Tags](#).

Export to Amazon S3

Data Wrangler gives you the ability to export your data to a location within an Amazon S3 bucket. You can specify the location using one of the following methods:

- Destination node – Where Data Wrangler stores the data after it has processed it.
- Export to – Exports the data resulting from a transformation to Amazon S3.
- Export data – For small datasets, can quickly export the data that you've transformed.

Use the following sections to learn more about each of these methods.

Destination Node

If you want to output a series of data processing steps that you've performed to Amazon S3, you create a destination node. A destination node tells Data Wrangler where to store the data after you've processed it. After you create a destination node, you create a processing job to output the data. A processing job is an Amazon SageMaker processing job. When you're using a destination node, it runs the computational resources needed to output the data that you've transformed to Amazon S3.

You can use a destination node to export some of the transformations or all of the transformations that you've made in your Data Wrangler flow.

You can use multiple destination nodes to export different transformations or sets of transformations. The following example shows two destination nodes in a single Data Wrangler flow.
You can use the following procedure to create destination nodes and export them to an Amazon S3 bucket.

To export your data flow, you create destination nodes and a Data Wrangler job to export the data. Creating a Data Wrangler job starts a SageMaker processing job to export your flow. You can choose the destination nodes that you want to export after you've created them.

**Note**

You can choose **Create job** in the Data Wrangler flow to view the instructions to use a processing job.

Use the following procedure to create destination nodes.

1. Choose the + next to the nodes that represent the transformations that you want to export.
2. Choose **Add destination**.

3. Choose **Amazon S3**.
4. Specify the following fields.

- **Dataset name** – The name that you specify for the dataset that you're exporting.
- **File type** – The format of the file that you're exporting.
- **Delimiter** (CSV and Parquet files only) – The value used to separate other values.
- **Compression** (CSV and Parquet files only) – The compression method used to reduce the file size. You can use the following compression methods:
  - bzip2
  - deflate
  - gzip
- (Optional) **Amazon S3 location** – The S3 location that you're using to output the files.
- (Optional) **Number of partitions** – The number of datasets that you're writing as the output of the processing job.
- (Optional) **Partition by column** – Writes all data with the same unique value from the column.

- (Optional) **Inference Parameters** – Selecting **Generate inference artifact** applies all of the transformations you've used in the Data Wrangler flow to data coming into your inference pipeline. The model in your pipeline makes predictions on the transformed data.

5. Choose **Add destination**.

Use the following procedure to create a processing job.

Create a job from the **Data flow** page and choose the destination nodes that you want to export.

**Note**

You can choose **Create job** in the Data Wrangler flow to view the instructions for creating a processing job.

1. Choose **Create job**. The following image shows the pane that appears after you select **Create job**.
2. For **Job name**, specify the name of the export job.

3. Choose the destination nodes that you want to export.

4. (Optional) Specify a AWS KMS key ARN. A AWS KMS key is a cryptographic key that you can use to protect your data. For more information about AWS KMS keys, see [AWS Key Management Service](#).

5. (Optional) Under **Trained parameters**, choose **Refit** if you've done the following:
   - Sampled your dataset
   - Applied a transform that uses your data to create a new column in the dataset

For more information about refitting the transformations you've made to an entire dataset, see [Refit Transforms to The Entire Dataset and Export Them](#).
6. Choose **Configure job**. The following image shows the **Configure job** page.

7. (Optional) Configure the Data Wrangler job. You can make the following configurations:
   - Job configuration
   - Spark memory configuration
   - Network configuration
   - Tags
   - Parameters
• Associate Schedules


Export to

As an alternative to using a destination node, you can use the **Export to** option to export your Data Wrangler flow to Amazon S3 using a Jupyter notebook. You can choose any data node in your Data Wrangler flow and export it. Exporting the data node exports the transformation that the node represents and the transformations that precede it.

Use the following procedure to generate a Jupyter notebook and run it to export your Data Wrangler flow to Amazon S3.

1. Choose the + next to the node that you want to export.
2. Choose **Export to**.
3. Choose **Amazon S3 (via Jupyter Notebook)**.
4. Run the Jupyter notebook.

When you run the notebook, it exports your data flow (.flow file) in the same AWS Region as the Data Wrangler flow.
The notebook provides options that you can use to configure the processing job and the data that it outputs.

⚠️ **Important**

We provide you with job configurations to configure the output of your data. For the partitioning and driver memory options, we strongly recommend that you don't specify a configuration unless you already have knowledge about them.

Under **Job Configurations**, you can configure the following:

- **output_content_type** – The content type of the output file. Uses CSV as the default format, but you can specify Parquet.
- **delimiter** – The character used to separate values in the dataset when writing to a CSV file.
- **compression** – If set, compresses the output file. Uses gzip as the default compression format.
- **num_partitions** – The number of partitions or files that Data Wrangler writes as the output.
- **partition_by** – The names of the columns that you use to partition the output.

To change the output file format from CSV to Parquet, change the value from "CSV" to "Parquet". For the rest of the preceding fields, uncomment the lines containing the fields that you want to specify.

Under **(Optional) Configure Spark Cluster Driver Memory** you can configure Spark properties for the job, such as the Spark driver memory, in the config dictionary.

The following shows the config dictionary.

```python
config = json.dumps(
    {
        "Classification": "spark-defaults",
        "Properties": {
            "spark.driver.memory": f"{driver_memory_in_mb}m",
        }
    }
)
```
To apply the configuration to the processing job, uncomment the following lines:

```
# data_sources.append(ProcessingInput(
#     source=config_s3_uri,
#     destination="/opt/ml/processing/input/conf",
#     input_name="spark-config",
#     s3_data_type="S3Prefix",
#     s3_input_mode="File",
#     s3_data_distribution_type="FullyReplicated"
# ))
```

Export data

If you have a transformation on a small dataset that you want to export quickly, you can use the **Export data** method. When you start choose **Export data**, Data Wrangler works synchronously to export the data that you've transformed to Amazon S3. You can't use Data Wrangler until either it finishes exporting your data or you cancel the operation.

For information on using the **Export data** method in your Data Wrangler flow, see the following procedure.

To use the **Export data** method:

1. Choose a node in your Data Wrangler flow by opening (double-clicking on) it.
2. Configure how you want to export the data.

3. Choose Export data.

When you export your data flow to an Amazon S3 bucket, Data Wrangler stores a copy of the flow file in the S3 bucket. It stores the flow file under the `data_wrangler_flows` prefix. If you use the default Amazon S3 bucket to store your flow files, it uses the following naming convention: `sagemaker-region-account number`. For example, if your account number is 111122223333 and you are using Studio Classic in us-east-1, your imported datasets are stored in `sagemaker-us-east-1-111122223333`. In this example, your .flow files created in us-east-1 are stored in `s3://sagemaker-region-account number/data_wrangler_flows/`.

### Export to SageMaker Pipelines

When you want to build and deploy large-scale machine learning (ML) workflows, you can use SageMaker Pipelines to create workflows that manage and deploy SageMaker jobs. With SageMaker Pipelines, you can build workflows that manage your SageMaker data preparation, model training, and model deployment jobs. You can use the first-party algorithms that SageMaker offers by using SageMaker Pipelines. For more information on SageMaker Pipelines, see SageMaker Pipelines.

When you export one or more steps from your data flow to SageMaker Pipelines, Data Wrangler creates a Jupyter notebook that you can use to define, instantiate, run, and manage a pipeline.
Use a Jupyter Notebook to Create a Pipeline

Use the following procedure to create a Jupyter notebook to export your Data Wrangler flow to SageMaker Pipelines.

Use the following procedure to generate a Jupyter notebook and run it to export your Data Wrangler flow to SageMaker Pipelines.

1. Choose the + next to the node that you want to export.
2. Choose Export to.
4. Run the Jupyter notebook.

You can use the Jupyter notebook that Data Wrangler produces to define a pipeline. The pipeline includes the data processing steps that are defined by your Data Wrangler flow.

You can add additional steps to your pipeline by adding steps to the steps list in the following code in the notebook:

```python
pipeline = Pipeline(
```
For more information on defining pipelines, see Define SageMaker Pipeline.

**Export to an Inference Endpoint**

Use your Data Wrangler flow to process data at the time of inference by creating a SageMaker serial inference pipeline from your Data Wrangler flow. An inference pipeline is a series of steps that results in a trained model making predictions on new data. A serial inference pipeline within Data Wrangler transforms the raw data and provides it to the machine learning model for a prediction. You create, run, and manage the inference pipeline from a Jupyter notebook within Studio Classic. For more information about accessing the notebook, see Use a Jupyter Notebook to create an inference endpoint.

Within the notebook, you can either train a machine learning model or specify one that you've already trained. You can either use Amazon SageMaker Autopilot or XGBoost to train the model using the data that you've transformed in your Data Wrangler flow.

The pipeline provides the ability to perform either batch or real-time inference. You can also add the Data Wrangler flow to SageMaker Model Registry. For more information about hosting models, see Host multiple models in one container behind one endpoint.

⚠️ **Important**

You can't export your Data Wrangler flow to an inference endpoint if it has the following transformations:

- Join
- Concatenate
- Group by

If you must use the preceding transforms to prepare your data, use the following procedure.

**To prepare your data for inference with unsupported transforms**

1. Create a Data Wrangler flow.
2. Apply the preceding transforms that aren't supported.
3. Export the data to an Amazon S3 bucket.
4. Create a separate Data Wrangler flow.
5. Import the data that you've exported from the preceding flow.
6. Apply the remaining transforms.
7. Create a serial inference pipeline using the Jupyter notebook that we provide.

For information about exporting your data to an Amazon S3 bucket see Export to Amazon S3. For information about opening the Jupyter notebook used to create the serial inference pipeline, see Use a Jupyter Notebook to create an inference endpoint.

Data Wrangler ignores transforms that remove data at the time of inference. For example, Data Wrangler ignores the Handle Missing Values transform if you use the Drop missing configuration.

If you've refit transforms to your entire dataset, the transforms carry over to your inference pipeline. For example, if you used the median value to impute missing values, the median value from refitting the transform is applied to your inference requests. You can either refit the transforms from your Data Wrangler flow when you're using the Jupyter notebook or when you're exporting your data to an inference pipeline. For information about refitting transforms, see Refit Transforms to The Entire Dataset and Export Them.

The serial inference pipeline supports the following data types for the input and output strings. Each data type has a set of requirements.

**Supported datatypes**

- **text/csv** – the datatype for CSV strings
  - The string can't have a header.
  - Features used for the inference pipeline must be in the same order as features in the training dataset.
  - There must be a comma delimiter between features.
  - Records must be delimited by a newline character.

The following is an example of a validly formatted CSV string that you can provide in an inference request.
The features used in the dataset for the inference pipeline must be in the same order as the features in the training dataset.

The data must have a specific schema. You define schema as a single `instances` object that has a set of `features`. Each `features` object represents an observation.

The following is an example of a validly formatted JSON string that you can provide in an inference request.

```json
{
    "instances": [
        {
            "features": ["abc", 0.0, "Doe, John", 12345]
        },
        {
            "features": ["def", 1.1, "Doe, Jane", 67890]
        }
    ]
}
```

**Use a Jupyter Notebook to create an inference endpoint**

Use the following procedure to export your Data Wrangler flow to create an inference pipeline.

To create an inference pipeline using a Jupyter notebook, do the following.

1. Choose the + next to the node that you want to export.
2. Choose **Export to**.
3. Choose **SageMaker Inference Pipeline (via Jupyter Notebook)**.
4. Run the Jupyter notebook.
When you run the Jupyter notebook, it creates an inference flow artifact. An inference flow artifact is a Data Wrangler flow file with additional metadata used to create the serial inference pipeline. The node that you're exporting encompasses all of the transforms from the preceding nodes.

⚠️ **Important**

Data Wrangler needs the inference flow artifact to run the inference pipeline. You can't use your own flow file as the artifact. You must create it by using the preceding procedure.

**Export to Python Code**

To export all steps in your data flow to a Python file that you can manually integrate into any data processing workflow, use the following procedure.

Use the following procedure to generate a Jupyter notebook and run it to export your Data Wrangler flow to Python Code.

1. Choose the + next to the node that you want to export.
2. Choose **Export to**.
3. Choose **Python Code**.
4. Run the Jupyter notebook.
You might need to configure the Python script to make it run in your pipeline. For example, if you're running a Spark environment, make sure that you are running the script from an environment that has permission to access AWS resources.

**Export to Amazon SageMaker Feature Store**

You can use Data Wrangler to export features you've created to Amazon SageMaker Feature Store. A feature is a column in your dataset. Feature Store is a centralized store for features and their associated metadata. You can use Feature Store to create, share, and manage curated data for machine learning (ML) development. Centralized stores make your data more discoverable and reusable. For more information about Feature Store, see [Amazon SageMaker Feature Store](https://docs.aws.amazon.com/sagemaker/latest/dg/feature-store.html).

A core concept in Feature Store is a feature group. A feature group is a collection of features, their records (observations), and associated metadata. It's similar to a table in a database.

You can use Data Wrangler to do one of the following:

- Update an existing feature group with new records. A record is an observation in the dataset.
- Create a new feature group from a node in your Data Wrangler flow. Data Wrangler adds the observations from your datasets as records in your feature group.
If you're updating an existing feature group, your dataset's schema must match the schema of the feature group. All the records in the feature group are replaced with the observations in your dataset.

You can use either a Jupyter notebook or a destination node to update your feature group with the observations in the dataset.

If your feature groups with the Iceberg table format have a custom offline store encryption key, make sure you grant the IAM that you're using for the Amazon SageMaker Processing job permissions to use it. At a minimum, you must grant it permissions to encrypt the data that you're writing to Amazon S3. To grant the permissions, give the IAM role the ability to use the `GenerateDataKey`. For more information about granting IAM roles permissions to use AWS KMS keys see [https://docs.aws.amazon.com/kms/latest/developerguide/key-policies.html](https://docs.aws.amazon.com/kms/latest/developerguide/key-policies.html)

**Destination Node**

If you want to output a series of data processing steps that you've performed to a feature group, you can create a destination node. When you create and run a destination node, Data Wrangler updates a feature group with your data. You can also create a new feature group from the destination node UI. After you create a destination node, you create a processing job to output the data. A processing job is an Amazon SageMaker processing job. When you're using a destination node, it runs the computational resources needed to output the data that you've transformed to the feature group.

You can use a destination node to export some of the transformations or all of the transformations that you've made in your Data Wrangler flow.

Use the following procedure to create a destination node to update a feature group with the observations from your dataset.

To update a feature group using a destination node, do the following.

1. Choose the + symbol next to the node containing the dataset that you'd like to export.
2. Under **Add destination**, choose **SageMaker Feature Store**.
3. Choose (double-click) the feature group. Data Wrangler checks whether the schema of the feature group matches the schema of the data that you’re using to update the feature group.

4. (Optional) Select **Export to offline store only** for feature groups that have both an online store and an offline store. This option only updates the offline store with observations from your dataset.

5. After Data Wrangler validates the schema of your dataset, choose **Add**.

Use the following procedure to create a new feature group with data from your dataset.

You can store your feature group in one of the following ways:

- **Online** – Low-latency, high-availability cache for a feature group that provides real-time lookup of records. The online store allows quick access to the latest value for a record in a feature group.

- **Offline** – Stores data for your feature group in an Amazon S3 bucket. You can store your data offline when you don't need low-latency (sub-second) reads. You can use an offline store for features used in data exploration, model training, and batch inference.

- **Both online and offline** – Stores your data in both an online store and an offline store.

To create a feature group using a destination node, do the following.
1. Choose the + symbol next to the node containing the dataset that you'd like to export.

2. Under Add destination, choose SageMaker Feature Store.

3. Choose Create Feature Group.

4. In the following dialog box, if your dataset doesn't have an event time column, select Create "EventTime" column.

5. Choose Next.

6. Choose Copy JSON Schema. When you create a feature group, you paste the schema into the feature definitions.

7. Choose Create.

8. For Feature group name, specify a name for your feature group.

9. For Description (optional), specify a description to make your feature group more discoverable.

10. To create a feature group for an online store, do the following.

   a. Select Enable storage online.

   b. For Online store encryption key, specify an AWS managed encryption key or an encryption key of your own.

11. To create a feature group for an offline store, do the following.

   a. Select Enable storage offline. Specify values for the following fields:

      - S3 bucket name – The name of the Amazon S3 bucket that stores the feature group.

      - (Optional) Dataset directory name – The Amazon S3 prefix that you're using to store the feature group.

      - IAM Role ARN – The IAM role that has access to Feature Store.

      - Table Format – Table format of your offline store. You can specify Glue or Iceberg. Glue is the default format.

      - Offline store encryption key – By default, Feature Store uses an AWS Key Management Service managed key, but you can use the field to specify a key of your own.

   b. Specify values for the following fields:

      - S3 bucket name – The name of the bucket storing the feature group.
• **(Optional) Dataset directory name** – The Amazon S3 prefix that you're using to store the feature group.

• **IAM Role ARN** – The IAM role that has access to feature store.

• **Offline store encryption key** – By default, Feature Store uses an AWS managed key, but you can use the field to specify a key of your own.

12. Choose **Continue**.

13. Choose **JSON**.

14. Remove the placeholder brackets in the window.

15. Paste the JSON text from Step 6.

16. Choose **Continue**.

17. For **RECORD IDENTIFIER FEATURE NAME**, choose the column in your dataset that has unique identifiers for each record in your dataset.

18. For **EVENT TIME FEATURE NAME**, choose the column with the timestamp values.

19. Choose **Continue**.

20. (Optional) Add tags to make your feature group more discoverable.

21. Choose **Continue**.

22. Choose **Create feature group**.

23. Navigate back to your Data Wrangler flow and choose the refresh icon next to the **Feature Group** search bar.

---

**Note**

If you've already created a destination node for a feature group within a flow, you can't create another destination node for the same feature group. If you want to create another destination node for the same feature group, you must create another flow file.

Use the following procedure to create a Data Wrangler job.

Create a job from the **Data flow** page and choose the destination nodes that you want to export.

1. Choose **Create job**. The following image shows the pane that appears after you select Create job.
2. For **Job name**, specify the name of the export job.

3. Choose the destination nodes that you want to export.

4. (Optional) For **Output KMS Key**, specify an ARN, ID, or alias of an AWS KMS key. A KMS key is a cryptographic key. You can use the key to encrypt the output data from the job. For more information about AWS KMS keys, see [AWS Key Management Service](https://aws.amazon.com/kms/).

5. The following image shows the **Configure job** page with the **Job configuration** tab open.

![Configure job page](image)

(Optional) Under **Trained parameters**, choose **Refit** if you've done the following:

- Sampled your dataset
- Applied a transform that uses your data to create a new column in the dataset

For more information about refitting the transformations you've made to an entire dataset, see [Refit Transforms to The Entire Dataset and Export Them](https://aws.amazon.com/sagemaker/).
6. Choose **Configure job**.

7. (Optional) Configure the Data Wrangler job. You can make the following configurations:

   - **Job configuration**
   - **Spark memory configuration**
   - **Network configuration**
   - **Tags**
   - **Parameters**
   - **Associate Schedules**

8. Choose **Run**.

**Jupyter notebook**

Use the following procedure to a Jupyter notebook to export to Amazon SageMaker Feature Store.

Use the following procedure to generate a Jupyter notebook and run it to export your Data Wrangler flow to Feature Store.

1. Choose the + next to the node that you want to export.

2. Choose **Export to**.

3. Choose **Amazon SageMaker Feature Store (via Jupyter Notebook)**.

4. Run the Jupyter notebook.
Running a Jupyter notebook runs a Data Wrangler job. Running a Data Wrangler job starts a SageMaker processing job. The processing job ingests the flow into an online and offline feature store.

⚠️ **Important**

The IAM role you use to run this notebook must have the following AWS managed policies attached: `AmazonSageMakerFullAccess` and `AmazonSageMakerFeatureStoreAccess`.

You only need to enable one online or offline feature store when you create a feature group. You can also enable both. To disable online store creation, set `EnableOnlineStore` to `False`:

```python
# Online Store Configuration
online_store_config = {
    "EnableOnlineStore": False
}
```
The notebook uses the column names and types of the dataframe you export to create a feature group schema, which is used to create a feature group. A feature group is a group of features defined in the feature store to describe a record. The feature group defines the schema and features contained in the feature group. A feature group definition is composed of a list of features, a record identifier feature name, an event time feature name, and configurations for its online store and offline store.

Each feature in a feature group can have one of the following types: *String*, *Fractional*, or *Integral*. If a column in your exported dataframe is not one of these types, it defaults to *String*.

The following is an example of a feature group schema.

```python
column_schema = [
    {
        "name": "Height",
        "type": "long"
    },
    {
        "name": "Input",
        "type": "string"
    },
    {
        "name": "Output",
        "type": "string"
    },
    {
        "name": "Sum",
        "type": "string"
    },
    {
        "name": "Time",
        "type": "string"
    }
]
```

Additionally, you must specify a record identifier name and event time feature name:

- The *record identifier name* is the name of the feature whose value uniquely identifies a record defined in the feature store. Only the latest record per identifier value is stored in the online store. The record identifier feature name must be one of feature definitions' names.
The *event time feature name* is the name of the feature that stores the EventTime of a record in a feature group. An EventTime is a point in time when a new event occurs that corresponds to the creation or update of a record in a feature. All records in the feature group must have a corresponding EventTime.

The notebook uses these configurations to create a feature group, process your data at scale, and then ingest the processed data into your online and offline feature stores. To learn more, see [Data Sources and Ingestion](#).

**Refit Transforms to The Entire Dataset and Export Them**

When you import data, Data Wrangler uses a sample of the data to apply the encodings. By default, Data Wrangler uses the first 50,000 rows as a sample, but you can import the entire dataset or use a different sampling method. For more information, see [Import](#).

The following transformations use your data to create a column in the dataset:

- **Encode Categorical**
- **Featurize Text**
- **Handle Outliers**
- **Handle Missing Values**

If you used sampling to import your data, the preceding transforms only use the data from the sample to create the column. The transform might not have used all of the relevant data. For example, if you use the **Encode Categorical** transform, there might have been a category in the entire dataset that wasn't present in the sample.

You can either use a destination node or a Jupyter notebook to refit the transformations to the entire dataset. When Data Wrangler exports the transformations in the flow, it creates a SageMaker processing job. When the processing job finishes, Data Wrangler saves the following files in either the default Amazon S3 location or an S3 location that you specify:

- The Data Wrangler flow file that specifies the transformations that are refit to the dataset
• The dataset with the refit transformations applied to it

You can open a Data Wrangler flow file within Data Wrangler and apply the transformations to a different dataset. For example, if you've applied the transformations to a training dataset, you can open and use the Data Wrangler flow file to apply the transformations to a dataset used for inference.

For a information about using destination nodes to refit transforms and export see the following pages:

• Export to Amazon S3
• Export to Amazon SageMaker Feature Store

Use the following procedure to run a Jupyter notebook to refit the transformations and export the data.

To run a Jupyter notebook and to refit the transformations and export your Data Wrangler flow, do the following.

1. Choose the + next to the node that you want to export.
2. Choose Export to.
3. Choose the location to which you're exporting the data.
4. For the refit_trained_params object, set refit to True.
5. For the output_flow field, specify the name of the output flow file with the refit transformations.
6. Run the Jupyter notebook.

Create a Schedule to Automatically Process New Data

If you're processing data periodically, you can create a schedule to run the processing job automatically. For example, you can create a schedule that runs a processing job automatically when you get new data. For more information about processing jobs, see Export to Amazon S3 and Export to Amazon SageMaker Feature Store.

When you create a job you must specify an IAM role that has permissions to create the job. By default, the IAM role that you use to access Data Wrangler is the SageMakerExecutionRole.
The following permissions allow Data Wrangler to access EventBridge and allow EventBridge to run processing jobs:

- Add the following AWS Managed policy to the Amazon SageMaker Studio Classic execution role that provides Data Wrangler with permissions to use EventBridge:

  arn:aws:iam::aws:policy/AmazonEventBridgeFullAccess

  For more information about the policy, see AWS managed policies for EventBridge.

- Add the following policy to the IAM role that you specify when you create a job in Data Wrangler:

  {
      "Version": "2012-10-17",
      "Statement": [
          {
              "Effect": "Allow",
              "Action": "sagemaker:StartPipelineExecution",
          }
      ]
  }

  If you're using the default IAM role, you add the preceding policy to the Amazon SageMaker Studio Classic execution role.

  Add the following trust policy to the role to allow EventBridge to assume it.

  {
      "Effect": "Allow",
      "Principal": {
          "Service": "events.amazonaws.com"
      },
      "Action": "sts:AssumeRole"
  }
Important

When you create a schedule, Data Wrangler creates an eventRule in EventBridge. You incur charges for both the event rules that you create and the instances used to run the processing job.

For information about EventBridge pricing, see Amazon EventBridge pricing. For information about processing job pricing, see Amazon SageMaker Pricing.

You can set a schedule using one of the following methods:

- **CRON expressions**

  Note
  
  Data Wrangler doesn't support the following expressions:
  
  - LW#
  - Abbreviations for days
  - Abbreviations for months

- **RATE expressions**

  - Recurring – Set an hourly or daily interval to run the job.
  - Specific time – Set specific days and times to run the job.

The following sections provide procedures on creating jobs.

CRON

Use the following procedure to create a schedule with a CRON expression.

To specify a schedule with a CRON expression, do the following.

1. Open your Data Wrangler flow.
2. Choose Create job.
3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.

4. Choose **Next, 2. Configure job**.

5. Select **Associate Schedules**.

6. Choose **Create a new schedule**.

7. For **Schedule Name**, specify the name of the schedule.

8. For **Run Frequency**, choose **CRON**.

9. Specify a valid CRON expression.

10. Choose **Create**.

11. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

   **Note**
   
   You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

12. Choose one of the following:

   - **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.
   - **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

13. Choose **Run**

**RATE**

Use the following procedure to create a schedule with a RATE expression.

To specify a schedule with a RATE expression, do the following.

1. Open your Data Wrangler flow.

2. Choose **Create job**.

3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.

4. Choose **Next, 2. Configure job**.

5. Select **Associate Schedules**.
6. Choose **Create a new schedule**.
7. For **Schedule Name**, specify the name of the schedule.
8. For **Run Frequency**, choose **Rate**.
9. For **Value**, specify an integer.
10. For **Unit**, select one of the following:
    - **Minutes**
    - **Hours**
    - **Days**
11. Choose **Create**.
12. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

**Note**

You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

13. Choose one of the following:
    - **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.
    - **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.
14. Choose **Run**

**Recurring**

Use the following procedure to create a schedule that runs a job on a recurring basis.

To specify a schedule with a CRON expression, do the following.

1. Open your Data Wrangler flow.
2. Choose **Create job**.
3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.
4. Choose **Next, 2. Configure job**.
5. Select **Associate Schedules**.
6. Choose **Create a new schedule**.

7. For **Schedule Name**, specify the name of the schedule.

8. For **Run Frequency**, make sure **Recurring** is selected by default.

9. For **Every x hours**, specify the hourly frequency that the job runs during the day. Valid values are integers in the inclusive range of **1** and **23**.

10. For **On days**, select one of the following options:
    - **Every Day**
    - **Weekends**
    - **Weekdays**
    - **Select Days**

    - (Optional) If you've selected **Select Days**, choose the days of the week to run the job.

    **Note**
    
    The schedule resets every day. If you schedule a job to run every five hours, it runs at the following times during the day:
    
    - 00:00
    - 05:00
    - 10:00
    - 15:00
    - 20:00

11. Choose **Create**.

12. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

    **Note**
    
    You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

13. Choose one of the following:
• **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.

• **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

14. Choose **Run**

**Specific time**

Use the following procedure to create a schedule that runs a job at specific times.

To specify a schedule with a CRON expression, do the following.

1. Open your Data Wrangler flow.
2. Choose **Create job**.
3. (Optional) For **Output KMS key**, specify an AWS KMS key to configure the output of the job.
4. Choose **Next, 2. Configure job**.
5. Select **Associate Schedules**.
6. Choose **Create a new schedule**.
7. For **Schedule Name**, specify the name of the schedule.
8. Choose **Create**.
9. (Optional) Choose **Add another schedule** to run the job on an additional schedule.

**Note**

You can associate a maximum of two schedules. The schedules are independent and don't affect each other unless the times overlap.

10. Choose one of the following:

• **Schedule and run now** – Data Wrangler the job runs immediately and subsequently runs on the schedules.

• **Schedule only** – Data Wrangler the job only runs on the schedules that you specify.

11. Choose **Run**
You can use Amazon SageMaker Studio Classic view the jobs that are scheduled to run. Your processing jobs run within SageMaker Pipelines. Each processing job has its own pipeline. It runs as a processing step within the pipeline. You can view the schedules that you've created within a pipeline. For information about viewing a pipeline, see View a Pipeline.

Use the following procedure to view the jobs that you've scheduled.

To view the jobs you've scheduled, do the following.

1. Open Amazon SageMaker Studio Classic.
2. Open SageMaker Pipelines
3. View the pipelines for the jobs that you've created.

   The pipeline running the job uses the job name as a prefix. For example, if you've created a job named housing-data-feature-enginnering, the name of the pipeline is data-wrangler-housing-data-feature-engineering.

4. Choose the pipeline containing your job.
5. View the status of the pipelines. Pipelines with a Status of Succeeded have run the processing job successfully.

To stop the processing job from running, do the following:

To stop a processing job from running, delete the event rule that specifies the schedule. Deleting an event rule stops all the jobs associated with the schedule from running. For information about deleting a rule, see Disabling or deleting an Amazon EventBridge rule.

You can stop and delete the pipelines associated with the schedules as well. For information about stopping a pipeline, see StopPipelineExecution. For information about deleting a pipeline, see DeletePipeline.

**Use an Interactive Data Preparation Widget in an Amazon SageMaker Studio Classic Notebook to Get Data Insights**

Use the Data Wrangler data preparation widget to interact with your data, get visualizations, explore actionable insights, and fix data quality issues.
You can access the data preparation widget from an Amazon SageMaker Studio Classic notebook. For each column, the widget creates a visualization that helps you better understand its distribution. If a column has data quality issues, a warning appears in its header.

To see the data quality issues, select the column header showing the warning. You can use the information that you get from the insights and the visualizations to apply the widget's built-in transformations to help you fix the issues.

For example, the widget might detect that you have a column that only has one unique value and show you a warning. The warning provides the option to drop the column from the dataset.

**Getting started with running the widget**

Use the following information to help you get started with running a notebook.

Open a notebook in Amazon SageMaker Studio Classic. For information about opening a notebook, see [Create or Open an Amazon SageMaker Studio Classic Notebook](#).

⚠️ **Important**

To run the widget, the notebook must use one of the following images:

- Python 3 (Data Science) with Python 3.7
- Python 3 (Data Science 2.0) with Python 3.8
- Python 3 (Data Science 3.0) with Python 3.10
- SparkAnalytics 1.0
- SparkAnalytics 2.0

For more information about images, see [Available Amazon SageMaker Images](#).

Use the following code to import the data preparation widget and pandas. The widget uses pandas dataframes to analyze your data.

```python
import pandas as pd
import sagemaker_datawrangler
```

The following example code loads a file into the dataframe called df.

```python
### Example code to load a file into the dataframe
def load_data(file_path):
    df = pd.read_csv(file_path)
    return df
```
You can use a dataset in any format that you can load as a pandas dataframe object. For more information about pandas formats, see IO tools (text, CSV, HDF5, ...).

The following cell runs the df variable to start the widget.

```python
df = pd.read_csv("example-dataset.csv")
```

The top of the dataframe has the following options:

- **View the Pandas table** – Switches between the interactive visualization and a pandas table.
- **Use all of the rows in your dataset to compute the insights. Using the entire dataset might increase the time it takes to generate the insights.** – If you don't select the option, Data Wrangler computes the insights for the first 10,000 rows of the dataset.

The dataframe shows the first 1000 rows of the dataset. Each column header has a stacked bar chart that shows the column's characteristics. It shows the proportion of valid values, invalid values, and missing values. You can hover over the different portions of the stacked bar chart to get the calculated percentages.

Each column has a visualization in the header. The following shows the types of visualizations the columns can have:

- Categorical – Bar chart
- Numeric – Histogram
- Datetime – Bar chart
- Text – Bar chart

For each visualization, the data preparation widget highlights outliers in orange.

When you choose a column, it opens a side panel. The side panel shows you the Insights tab. The pane provides a count for the following types of values:

- Invalid values – Values whose type doesn’t match the column type.
- Missing values – Values that are missing, such as NaN or None.
- Valid values – Values that are neither missing nor invalid.
For numeric columns, the **Insights** tab shows the following summary statistics:

- **Minimum** – The smallest value.
- **Maximum** – The largest value.
- **Mean** – The mean of the values.
- **Mode** – The value that appears most frequently.
- **Standard deviation** – The standard deviation of the values.

For categorical columns, the **Insights** tab shows the following summary statistics:

- **Unique values** – The number of unique values in the column.
- **Top** – The value that appears most frequently.

The columns that have warning icons in their headers have data quality issues. Choosing a column opens a **Data quality** tab that you can use to find transforms to help you fix the issue. A warning has one of the following severity levels:

- **Low** – Issues that might not affect your analysis, but can be useful to fix.
- **Medium** – Issues that are likely to affect your analysis, but are likely not critical to fix.
- **High** – Severe issues that we strongly recommend fixing.

**Note**

The widget sorts the column to show the values that have data quality issues at the top of the dataframe. It also highlights the values that are causing the issues. The color of the highlighting corresponds to the severity level.

Under **SUGGESTED TRANSFORMS**, you can choose a transform to fix the data quality issue. The widget can offer multiple transforms that can fix the issue. It can offer recommendations for the transforms that are best suited to the problem. You can move your cursor over the transform to get more information about it.

To apply a transform to the dataset, choose **Apply and export code**. The transform modifies the dataset and updates the visualization with modified values. The code for the transform appears in
the following cell of the notebook. If you apply additional transforms to the dataset, the widget
appends the transforms to the cell. You can use the code that the widget generates to do the
following:

- Customize it to better fit your needs.
- Use it in your own workflows.

You can reproduce all the transforms you've made by rerunning all of the cells in the notebook.

The widget can provide insights and warnings for the target column. The target column is the
column that you're trying to predict. Use the following procedure to get target column insights.

To get target column insights, do the following.

1. Choose the column that you're using as the target column.
2. Choose Select as target column.
3. Choose the problem type. The widget's insights and warnings are tailored to the problem
types. The following are the problem types:

   - Classification – The target column has categorical data.
   - Regression – The target column has numeric data.
5. (Optional) Under Target Column Insights, choose one of the suggested transforms.

Reference for the insights and transforms in the widget

For feature columns (columns that aren't the target column), you can get the following insights to
warn you about issues with your dataset.

- Missing values – The column has missing values such as None, NaN (not a number), or NaT (not
  a timestamp). Many machine learning algorithms don't support missing values in the input data.
  Filling them in or dropping the rows with missing data is therefore a crucial data preparation
  step. If you see the missing values warning, you can use one of the following transforms to
correct the issue.

- Drop missing – Drops rows with missing values. We recommend dropping rows when
  the percentage of rows with missing data is small and imputing the missing values isn't
  appropriate.
• **Replace with new value** – Replaces textual missing values with Other. You can change Other to a different value in the output code. Replaces numeric missing values with 0.

• **Replace with mean** – Replaces missing values with the mean of the column.

• **Replace with median** – Replaces missing values with the median of the column.

• **Drop column** – Drops the column with missing values from the dataset. We recommend dropping the entire column when there's a high percentage of rows with missing data.

• **Disguised missing values** – The column has disguised missing values. A disguised missing value is a value that isn't explicitly encoded as a missing value. For example, instead of using a NaN to indicate a missing value, the value could be Placeholder. You can use one of the following transforms to handle the missing values:
  
  • **Drop missing** – Drops rows with missing values
  
  • **Replace with new value** – Replaces textual missing values with Other. You can change Other to a different value in the output code. Replaces numeric missing values with 0.

• **Constant column** – The column only has one value. It therefore has no predictive power. We strongly recommend using the **Drop column** transform to drop the column from the dataset.

• **ID column** – The column has no repeating values. All of the values in the column are unique. They might be either IDs or database keys. Without additional information, the column has no predictive power. We strongly recommend using the **Drop column** transform to drop the column from the dataset.

• **High cardinality** – The column has a high percentage of unique values. High cardinality limits the predictive power of categorical columns. Examine the importance of the column in your analysis and consider using the **Drop column** transform to drop it.

For the target column, you can get the following insights to warn you about issues with your dataset. You can use the suggested transformation provided with the warning to correct the issue.

• **Mixed data types in target (Regression)** – There are some non-numeric values in the target column. There might be data entry errors. We recommend removing the rows that have the values that can't be converted.

• **Frequent label** – Certain values in the target column appear more frequently than what would be normal in the context of regression. There might be an error in data collection or processing. A frequently appearing category might indicate that either the value is used as a default value or that it's a placeholder for missing values. We recommend using the **Replace with new value** transform to replace the missing values with Other.
• **Too few instances per class** – The target column has categories that appear rarely. Some of the categories don’t have enough rows for the target column to be useful. You can use one of the following transforms:

  • **Drop rare target** – Drops unique values with fewer than ten observations. For example, drops the value `cat` if it appears nine times in the column.

  • **Replace rare target** – Replaces categories that appear rarely in the dataset with the value `Other`.

• **Classes too imbalanced (multi-class classification)** – There are categories in the dataset that appear much more frequently than the other categories. The class imbalance might affect prediction accuracy. For the most accurate predictions possible, we recommend updating the dataset with rows that have the categories that currently appear less frequently.

• **Large amount of classes/too many classes** – There's a large number of classes in the target column. Having many classes might result in longer training times or poor predictive quality. We recommend doing one of the following:

  • Grouping some of the categories into their own category. For example, if six categories are closely related, we recommend using a single category for them.

  • Using an ML algorithm that's resilient to multiple categories.

### Security and Permissions

When you query data from Athena or Amazon Redshift, the queried dataset is automatically stored in the default SageMaker S3 bucket for the AWS Region in which you are using Studio Classic. Additionally, when you export a Jupyter Notebook from Amazon SageMaker Data Wrangler and run it, your data flows, or .flow files, are saved to the same default bucket, under the prefix `data_wrangler_flows`.

For high-level security needs, you can configure a bucket policy that restricts the AWS roles that have access to this default SageMaker S3 bucket. Use the following section to add this type of policy to an S3 bucket. To follow the instructions on this page, use the AWS Command Line Interface (AWS CLI). To learn how, see [Configuring the AWS CLI](#) in the IAM User Guide.

Additionally, you need to grant each IAM role that uses Data Wrangler permissions to access required resources. If you do not require granular permissions for the IAM role you use to access Data Wrangler, you can add the IAM managed policy, `AmazonSageMakerFullAccess`, to an IAM role that you use to create your Studio Classic user. This policy grants you full permission to use...
Data Wrangler. If you require more granular permissions, refer to the section, Grant an IAM Role Permission to Use Data Wrangler.

Add a Bucket Policy To Restrict Access to Datasets Imported to Data Wrangler

You can add a policy to the S3 bucket that contains your Data Wrangler resources using an Amazon S3 bucket policy. Resources that Data Wrangler uploads to your default SageMaker S3 bucket in the AWS Region you are using Studio Classic in include the following:

- Queried Amazon Redshift results. These are stored under the `redshift/` prefix.
- Queried Athena results. These are stored under the `athena/` prefix.
- The .flow files uploaded to Amazon S3 when you run an exported Jupyter Notebook Data Wrangler produces. These are stored under the `data_wrangler_flows/` prefix.

Use the following procedure to create an S3 bucket policy that you can add to restrict IAM role access to that bucket. To learn how to add a policy to an S3 bucket, see How do I add an S3 Bucket policy?

To set up a bucket policy on the S3 bucket that stores your Data Wrangler resources:

1. Configure one or more IAM roles that you want to be able to access Data Wrangler.
2. Open a command prompt or shell. For each role that you create, replace `role-name` with the name of the role and run the following:

   ```
   $ aws iam get-role --role-name role-name
   ```

   In the response, you see a RoleId string which begins with AROA. Copy this string.
3. Add the following policy to the SageMaker default bucket in the AWS Region in which you are using Data Wrangler. Replace `region` with the AWS Region in which the bucket is located, and `account-id` with your AWS account ID. Replace `userId`s starting with `AROAEXAMPLEID` with the IDs of an AWS roles to which you want to grant permission to use Data Wrangler.

   ```
   {
     "Version": "2012-10-17",
     "Statement": [
       {
         "Effect": "Deny",
         "Principal": "*",
   ```
Create an Allowlist for Data Wrangler

Whenever a user starts running Data Wrangler from the Amazon SageMaker Studio Classic user interface, they make call to the SageMaker application programming interface (API) to create a Data Wrangler application.

Your organization might not provide your users with permissions to make those API calls by default. To provide permissions, you must create and attach a policy to the user's IAM roles using the following policy template: Data Wrangler Allow List Example.

Note

The preceding policy example only gives your users access to the Data Wrangler application.

For information about creating a policy, see Creating policies on the JSON tab. When you're creating a policy, copy and paste the JSON policy from Data Wrangler Allow List Example in the JSON tab.
Important
Delete any IAM policies that prevent users from running the following operations:

- CreateApp
- DescribeApp

If you don't delete the policies, your users could still be affected by them.

After you've creating the policy using the template, attach it to the IAM roles of your users. For information about attaching a policy, see Adding IAM identity permissions (console).

Grant an IAM Role Permission to Use Data Wrangler

You can grant an IAM role permission to use Data Wrangler with the general IAM managed policy, AmazonSageMakerFullAccess. This is a general policy that includes permissions required to use all SageMaker services. This policy grants an IAM role full access to Data Wrangler. You should be aware of the following when using AmazonSageMakerFullAccess to grant access to Data Wrangler:

- If you import data from Amazon Redshift, the Database User name must have the prefix sagemaker_access.
- This managed policy only grants permission to access buckets with one of the following in the name: SageMaker, SageMaker, sagemaker, or aws-glue. If want to use Data Wrangler to import from an S3 bucket without these phrases in the name, refer to the last section on this page to learn how to grant permission to an IAM entity to access your S3 buckets.

If you have high-security needs, you can attach the policies in this section to an IAM entity to grant permissions required to use Data Wrangler.

If you have datasets in Amazon Redshift or Athena that an IAM role needs to import from Data Wrangler, you must add a policy to that entity to access these resources. The following policies are the most restrictive policies you can use to give an IAM role permission to import data from Amazon Redshift and Athena.

To learn how to attach a custom policy to an IAM role, refer to Managing IAM policies in the IAM User Guide.
Policy example to grant access to an Athena dataset import

The following policy assumes that the IAM role has permission to access the underlying S3 bucket where data is stored through a separate IAM policy.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "athena:ListDataCatalogs",
                "athena:ListDatabases",
                "athena:ListTableMetadata",
                "athena:GetQueryExecution",
                "athena:GetQueryResults",
                "athena:StartQueryExecution",
                "athena:StopQueryExecution"
            ],
            "Resource": [
                "*
            ]
        },
        {
            "Effect": "Allow",
            "Action": [
                "glue:CreateTable"
            ],
            "Resource": [
                "arn:aws:glue::*:table/*/sagemaker_tmp_*",
                "arn:aws:glue::*:table/sagemaker_featurestore/**",
                "arn:aws:glue::*:catalog",
                "arn:aws:glue::*:database/**
            ]
        },
        {
            "Effect": "Allow",
            "Action": [
                "glue:DeleteTable"
            ],
            "Resource": [
                "arn:aws:glue::*:table/*/sagemaker_tmp_*",
                "arn:aws:glue::*:catalog",
                "arn:aws:glue::*:database/**
            ]
        }
    ]
}
```
Policy example to grant access to an Amazon Redshift dataset import

The following policy grants permission to set up an Amazon Redshift connection to Data Wrangler using database users that have the prefix sagemaker_access in the name. To grant permission to connect using additional database users, add additional entries under "Resources" in the following policy. The following policy assumes that the IAM role has permission to access the underlying S3 bucket where data is stored through a separate IAM policy, if applicable.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "glue:GetDatabases",
                "glue:GetTable",
                "glue:GetTables"
            ],
            "Resource": [
                "arn:aws:glue:*:*:table/*",
                "arn:aws:glue:*:*:catalog",
                "arn:aws:glue:*:*:database/*"
            ]
        },
        {
            "Effect": "Allow",
            "Action": [
                "glue:CreateDatabase",
                "glue:GetDatabase"
            ],
            "Resource": [
                "arn:aws:glue:*:*:catalog",
                "arn:aws:glue:*:*:database/sagemaker_featurestore",
                "arn:aws:glue:*:*:database/sagemaker_processing",
                "arn:aws:glue:*:*:database/default",
                "arn:aws:glue:*:*:database/sagemaker_data_wrangler"
            ]
        }
    ]
}
```
Policy to grant access to an S3 bucket

If your dataset is stored in Amazon S3, you can grant an IAM role permission to access this bucket with a policy similar to the following. This example grants programmatic read-write access to the bucket named `test`.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": ["s3:ListBucket"],
            "Resource": ["arn:aws:s3:::test"]
        },
        {
            "Effect": "Allow",
            "Action": ["redshift-data:ExecuteStatement", "redshift-data:DescribeStatement", "redshift-data:CancelStatement", "redshift-data:GetStatementResult", "redshift-data:ListSchemas", "redshift-data:ListTables"],
            "Resource": ["*"]
        },
        {
            "Effect": "Allow",
            "Action": ["redshift:GetClusterCredentials"],
            "Resource": ["arn:aws:redshift::*:*:sagemaker_access/*", "arn:aws:redshift::*:*:dbname:*"]
        }
    ]
}
```
To import data from Athena and Amazon Redshift, you must grant an IAM role permission to access the following prefixes under the default Amazon S3 bucket in the AWS Region Data Wrangler in which is being used: athena/, redshift/. If a default Amazon S3 bucket does not already exist in the AWS Region, you must also give the IAM role permission to create a bucket in this region.

Additionally, if you want the IAM role to be able to use the Amazon SageMaker Feature Store, SageMaker Pipelines, and Data Wrangler job export options, you must grant access to the prefix data_wrangler_flows/ in this bucket.

Data Wrangler uses the athena/ and redshift/ prefixes to store preview files and imported datasets. To learn more, see Imported Data Storage.

Data Wrangler uses the data_wrangler_flows/ prefix to store .flow files when you run a Jupyter Notebook exported from Data Wrangler. To learn more, see Export.

Use a policy similar to the following to grant the permissions described in the preceding paragraphs.

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "s3:GetObject",
        "s3:PutObject"
      ],
      "Resource": [
        "arn:aws:s3:::sagemaker-region-account-id/data_wrangler_flows/",
        "arn:aws:s3:::sagemaker-region-account-id/data_wrangler_flows/*",
        "arn:aws:s3:::sagemaker-region-account-id/athena",
      ]
    }]
}
```
You can also access data in your Amazon S3 bucket from another AWS account by specifying the Amazon S3 bucket URI. To do this, the IAM policy that grants access to the Amazon S3 bucket in the other account should use a policy similar to the following example, where `BucketFolder` is the specific directory in the user's bucket `UserBucket`. This policy should be added to the user granting access to their bucket for another user.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "s3:GetObject",
        "s3:PutObject",
        "s3:PutObjectAcl"
      ],
      "Resource": "arn:aws:s3:::UserBucket/BucketFolder/*"
    },
    {
      "Effect": "Allow",
      "Action": [
        "s3:GetObject",
        "s3:PutObject",
        "s3:GetObjectLocation"
      ],
      "Resource": "*"
    }
  ]
}
```
The user that is accessing the bucket (not the bucket owner) must add a policy similar to the following example to their user. Note that AccountX and TestUser below refers to the bucket owner and their user respectively.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::AccountX:user/TestUser"
      },
      "Action": [
        "s3:GetObject",
        "s3:PutObject",
        "s3:PutObjectAcl"
      ],
      "Resource": [
        "arn:aws:s3:::UserBucket/BucketFolder/*"
      ]
    },
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::AccountX:user/TestUser"
      },
      "Action": [
      ]
    }
  ]
}
```
Policy example to grant access to use SageMaker Studio

Use a policy like to the following to create an IAM execution role that can be used to set up a Studio Classic instance.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreatePresignedDomainUrl",
                "sagemaker:DescribeDomain",
                "sagemaker:ListDomains",
                "sagemaker:DescribeUserProfile",
                "sagemaker:ListUserProfiles",
                "sagemaker:*App",
                "sagemaker:ListApps"
            ],
            "Resource": "*"
        }
    ]
}
```

Snowflake and Data Wrangler

All permissions for AWS resources are managed via your IAM role attached to your Studio Classic instance. The Snowflake administrator manages Snowflake-specific permissions, as they can grant granular permissions and privileges to each Snowflake user. This includes databases, schemas, tables, warehouses, and storage integration objects. You must ensure that the correct permissions are set up outside of Data Wrangler.
Note that the Snowflake COPY INTO Amazon S3 command moves data from Snowflake to Amazon S3 over the public internet by default, but data in transit is secured using SSL. Data at rest in Amazon S3 is encrypted with SSE-KMS using the default AWS KMS key.

With respect to Snowflake credentials storage, Data Wrangler does not store customer credentials. Data Wrangler uses Secrets Manager to store the credentials in a secret and rotates secrets as part of a best practice security plan. The Snowflake or Studio Classic administrator needs to ensure that the data scientist’s Studio Classic execution role is granted permission to perform GetSecretValue on the secret storing the credentials. If already attached to the Studio Classic execution role, the AmazonSageMakerFullAccess policy has the necessary permissions to read secrets created by Data Wrangler and secrets created by following the naming and tagging convention in the instructions above. Secrets that do not follow the conventions must be separately granted access. We recommend using Secrets Manager to prevent sharing credentials over unsecured channels; however, note that a logged-in user can retrieve the plain-text password by launching a terminal or Python notebook in Studio Classic and then invoking API calls from the Secrets Manager API.

**Data Encryption with AWS KMS**

Within Data Wrangler, you can decrypt encrypted files and add them to your Data Wrangler flow. You can also encrypt the output of the transforms using either a default AWS KMS key or one that you provide.

You can import files if they have the following:

- server-side encryption
- SSE-KMS as the encryption type

To decrypt the file and import to a Data Wrangler flow, you must add the SageMaker Studio Classic user that you're using as a key user.

The following screenshot shows a Studio Classic user role added as a key user. See [IAM Roles](#) to access users under the left panel to make this change.
Amazon S3 customer managed key setup for Data Wrangler imported data storage

By default, Data Wrangler uses Amazon S3 buckets that have the following naming convention: sagemaker-region-account number. For example, if your account number is 111122223333 and you are using Studio Classic in us-east-1, your imported datasets are stored with the following naming convention: sagemaker-us-east-1-111122223333.

The following instructions explain how to set up a customer managed key for your default Amazon S3 bucket.

1. To enable server-side encryption and setup a customer managed key for your default S3 bucket, see Using KMS Encryption.
2. After following step 1, navigate to AWS KMS in your AWS Management Console. Find the customer managed key you selected in step 1 of the previous step and add the Studio Classic role as the key user. To do this, follow the instructions in Allows key users to use a customer managed key.

Encrypting the Data That You Export

You can encrypt the data that you export using one of the following methods:

- Specifying that your Amazon S3 bucket has object use SSE-KMS encryption.
- Specifying an AWS KMS key to encrypt the data that you export from Data Wrangler.

On the Export data page, specify a value for the AWS KMS key ID or ARN.

For more information on using AWS KMS keys, see Protecting Data Using Server-Side Encryption with AWS KMS keys Stored in AWS Key Management Service (SSE-KMS).

Amazon AppFlow Permissions

When you're performing a transfer, you must specify an IAM role that has permissions to perform the transfer. You can use the same IAM role that has permissions to use Data Wrangler. By default, the IAM role that you use to access Data Wrangler is the SageMakerExecutionRole.

The IAM role must have the following permissions:

- Permissions to Amazon AppFlow
- Permissions to the AWS Glue Data Catalog
• Permissions for AWS Glue to discover the data sources that are available

When you run a transfer, Amazon AppFlow stores metadata from the transfer in the AWS Glue Data Catalog. Data Wrangler uses the metadata from the catalog to determine whether it's available for you to query and import.

To add permissions to Amazon AppFlow, add the AmazonAppFlowFullAccess AWS managed policy to the IAM role. For more information about adding policies, see Adding or removing IAM identity permissions.

If you're transferring data to Amazon S3, you must also attach the following policy.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "VisualEditor0",
            "Effect": "Allow",
            "Action": [
                "s3:GetBucketTagging",
                "s3:ListBucketVersions",
                "s3:CreateBucket",
                "s3:ListBucket",
                "s3:GetBucketPolicy",
                "s3:PutEncryptionConfiguration",
                "s3:GetEncryptionConfiguration",
                "s3:PutBucketTagging",
                "s3:GetObjectTagging",
                "s3:GetBucketOwnershipControls",
                "s3:PutObjectTagging",
                "s3:DeleteObject",
                "s3:DeleteBucket",
                "s3:DeleteObjectTagging",
                "s3:GetBucketPublicAccessBlock",
                "s3:GetBucketPolicyStatus",
                "s3:PutBucketPublicAccessBlock",
                "s3:PutAccountPublicAccessBlock",
                "s3:ListAccessPoints",
                "s3:PutBucketOwnershipControls",
                "s3:PutObjectVersionTagging",
                "s3:DeleteObjectVersionTagging",
```
To add AWS Glue permissions, add the AWSGlueConsoleFullAccess managed policy to the IAM role. For more information about AWS Glue permissions with Amazon AppFlow, see [link-to-appflow-page].

Amazon AppFlow needs to access AWS Glue and Data Wrangler for you to import the data that you've transferred. To grant Amazon AppFlow access, add the following trust policy to the IAM role:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::123456789012:root",
        "Service": [
          "appflow.amazonaws.com"
        ]
      },
      "Action": "sts:AssumeRole"
    }
  ]
}
```

To display the Amazon AppFlow data in Data Wrangler, add the following policy to the IAM role:
Using Lifecycle Configurations in Data Wrangler

You might have an Amazon EC2 instance that is configured to run Kernel Gateway applications, but not the Data Wrangler application. Kernel Gateway applications provide access to the environment and the kernels that you use to run Studio Classic notebooks and terminals. The Data Wrangler application is the UI application that runs Data Wrangler. Amazon EC2 instances that aren't Data Wrangler instances require a modification to their lifecycle configurations to run Data Wrangler. Lifecycle configurations are shell scripts that automate the customization of your Amazon SageMaker Studio Classic environment.

For more information about lifecycle configurations, see Use lifecycle configurations with Amazon SageMaker Studio Classic.

The default lifecycle configuration for your instance doesn't support using Data Wrangler. You can make the following modifications to the default configuration to use Data Wrangler with your instance.

```bash
#!/bin/bash
set -eux
STATUS=$(python3 -c "import sagemaker_dataprep; echo $?"
if [ "$STATUS" -eq 0 ]; then
echo 'Instance is of Type Data Wrangler'
else

Security and Permissions
echo 'Instance is not of Type Data Wrangler'

# Replace this with the URL of your git repository
export REPOSITORY_URL="https://github.com/aws-samples/sagemaker-studio-lifecycle-config-examples.git"

git -C /root clone $REPOSITORY_URL

You can save the script as lifecycle_configuration.sh.

You attach the lifecycle configuration to your Studio Classic domain or user profile. For more information about creating and attaching a lifecycle configuration, see Create and associate a lifecycle configuration.

The following instructions show you how to attach a lifecycle configuration to a Studio Classic domain or user profile.

You might run into errors when you're creating or attaching a lifecycle configuration. For information about debugging lifecycle configuration errors, KernelGateway app failure.

Release Notes

Data Wrangler is regularly updated with new features and bug fixes. To upgrade the version of Data Wrangler you are using in Studio Classic, follow the instructions in Shut down and Update Studio Classic Apps.

Release Notes

8/31/2023

New functionality:

You can now create a Data Quality and Insights report on your entire dataset. For more information, see Get Insights On Data and Data Quality.

5/20/2023

New functionality:
You can now import your data from Salesforce Data Cloud. For more information, see Import data from Salesforce Data Cloud.

4/18/2023

New functionality:

You can now get your data in a format that Amazon Personalize can interpret. For more information, see Map Columns for Amazon Personalize.

3/1/2023

New functionality:

You can now use Hive to import your data from Amazon EMR. For more information, see Import data from Amazon EMR.

12/10/2022

New functionality:

You can now export your Data Wrangler flow to an inference endpoint. For more information, see Export to an Inference Endpoint.

New functionality:

You can now use an interactive notebook widget for data preparation. For more information, see Use an Interactive Data Preparation Widget in an Amazon SageMaker Studio Classic Notebook to Get Data Insights.

New functionality:

You can now import data from SaaS platforms. For more information, see Import Data From Software as a Service (SaaS) Platforms.
You can now reuse data flows for different data sets. For more information, see Reusing Data Flows for Different Datasets.

10/05/2022

New functionality:

You can now use Principal Component Analysis (PCA) as a transform. For more information, see Reduce Dimensionality within a Dataset.

10/05/2022

New functionality:

You can now refit parameters in your Data Wrangler flow. For more information, see Export.

10/03/2022

New functionality:

You can now deploy models from your Data Wrangler flow. For more information, see Automatically Train Models on Your Data Flow.

9/20/2022

New functionality:

You can now set data retention periods in Athena. For more information, see Import data from Athena.

6/9/2022

New functionality:

You can now use Amazon SageMaker Autopilot to train a model directly from your Data Wrangler flow. For more information, see Automatically Train Models on Your Data Flow.

5/6/2022

New functionality:
Release Notes

You can now use additional m5 and r5 instances. For more information, see Instances.

4/27/2022

New functionalities:

- You can now get a data quality report. For more information, see Get Insights On Data and Data Quality.
- You can now perform random sampling and stratified sampling. For more information, see Sampling.

4/1/2022

New functionality:

You can now use Databricks as a data source. For more information, see Import data from Databricks (JDBC).

2/2/2022

New functionalities:

- You can now export using destination nodes. For more information, see Export.
- You can import ORC and JSON files. For more information about file types, see Import.
- Data Wrangler now supports using the SMOTE transform. For more information, see Balance Data.
- Data Wrangler now supports similarity encoding for categorical data. For more information, see Similarity encode.
- Data Wrangler now supports unnesting JSON data. For more information, see Unnest JSON Data.
- Data Wrangler now supports expanding the values of an array into separate columns. For more information, see Explode Array.
- Data Wrangler now supports reaching out to the service team when you're having issues. For more information, see Troubleshoot.
Release Notes

- Data Wrangler supports editing and deleting steps in your data flow. For more information, see [Delete a Step from Your Data Flow](#) and [Edit a Step in Your Data Wrangler Flow](#).
- You can now perform transformations on multiple columns. For more information, see [Transform Data](#).
- Data Wrangler now supports cost allocation tags. For more information, see [Using Cost Allocation Tags](#).

10/16/2021

New functionality:

Data Wrangler now supports Athena workgroups. For more information, see [Import data from Athena](#).

10/6/2021

New functionality:

Data Wrangler now supports transforming time series data. For more information, see [Transform Time Series](#).

7/15/2021

New functionalities:

- [Snowflake and Data Wrangler](#) is now supported. You can use Snowflake as a data source in Data Wrangler.
- Added support for custom field delimiter in CSV. Now comma, colon, semicolon, pipe (|) and Tab are supported.
- Now you can export results directly to Amazon S3.
- Added a few new multicollinearity analyzers: Variance Inflation Factors, Principal Component Analysis and Lasso feature selection.

Enhancements:

- The analyze charts can no longer be packed with overlapping labels.
**Release Notes**

**Bug Fixes:**

- One-hot encoder handles empty string gracefully.
- Fixed crashes that occurred when a dataframe column name contained dots.

**4/26/2021**

**Enhancements:**

- Added support for distributed processing Jobs. You can use multiple instances when running a processing job.
- Data Wrangler Processing job now automatically coalesces small outputs when estimated result size is less than 1 gigabytes.
- Feature Store Notebook: Improved feature store ingestion performance
- Data Wrangler Processing jobs now use 1.x as the authoritative container tag for future releases.

**Bug Fixes:**

- Fixed rendering issues for faceted histogram.
- Fixed Export to Processing Job to support vector type columns.
- Fixed Extract using regex operator to return the first captured group if one or more exists in the regular expression or regex.

**2/8/2021**

**New Functionalities:**

- Data Wrangler Flows supports multiple instances.
- Updated Export to Data Wrangler Job Notebook to use SageMaker SDK 2.20.0.
- Updated Export to Pipeline Notebook to use SageMaker SDK 2.20.0.
- Updated Export to Pipeline Notebook to add XGBoost training example as an optional step.

**Enhancements:**
Release Notes

- To improve performance, importing CSV files that contain multiple lines in a single field is no longer supported.

Bug Fixes:

- Fixed type inference issue in Quick model.
- Fixed the bias metric bug in bias reports.
- Fixed the Featurize text transform to work with columns with missing values.
- Fixed Histogram and Scatter plot built-in visualizations to work with datasets that contain array-like columns.
- Athena query now re-runs if the query execution ID has expired.

Troubleshoot

If an issue arises when using Amazon SageMaker Data Wrangler, we recommend you do the following:

- If an error message is provided, read the message and resolve the issue it reports if possible.
- Make sure the IAM role of your Studio Classic user has the required permissions to perform the action. For more information, see Security and Permissions.
- If the issue occurs when you are trying to import from another AWS service, such as Amazon Redshift or Athena, make sure that you have configured the necessary permissions and resources to perform the data import. For more information, see Import.
- If you're still having issues, choose Get help at the top right of your screen to reach out to the Data Wrangler team. For more information, see the following images.
As a last resort, you can try restarting the kernel on which Data Wrangler is running.

1. Save and exit the .flow file for which you want to restart the kernel.
2. Select the **Running Terminals and Kernels** icon, as shown in the following image.

![Running Terminals and Kernels](image)

3. Select the **Stop** icon to the right of the .flow file for which you want to terminate the kernel, as shown in the following image.
4. Refresh the browser.

5. Reopen the .flow file on which you were working.

**Troubleshooting issues with Amazon EMR**

Use the following information to help you troubleshoot errors that might come up when you're using Amazon EMR.

- **Connection failure** – If the connection fails with the following message *The IP address of the EMR cluster isn't private* error message, your Amazon EMR cluster might not have been launched in a private subnet. As a security best practice, Data Wrangler only supports connecting to private Amazon EMR clusters. Choose a private EC2 subnet you launch an EMR cluster.

- **Connection hanging and timing out** – The issue is most likely due to a network connectivity issue. After you start connecting to the cluster, the screen doesn't refresh. After about 2 minutes, you might see the following error *JdbcAddConnectionError: An error occurred when trying to connect to presto: xxx: Connect to xxx failed: Connection timed out (Connection timed out)* will display on top of the screen.
The errors might have two root causes:

- The Amazon EMR and Amazon SageMaker Studio Classic are in different VPCs. We recommend launching both Amazon EMR and Studio Classic in the same VPC. You can also use VPC peering. For more information, see [What is VPC peering?](#).

- The Amazon EMR master security group lacks the inbound traffic rule for the security group of Amazon SageMaker Studio Classic on the port used for Presto. To resolve the issue, allow inbound traffic on port 8889.

Connection fails due to the connection type being misconfigured – You might see the following error message: `Data Wrangler couldn't create a connection to {connection_source} successfully. Try connecting to {connection_source} again. For more information, see Troubleshoot. If you’re still experiencing issues, contact support.`

Check the authentication method. The authentication method that you've specified in Data Wrangler should match the authentication method that you're using on the cluster.

- You don't have HDFS permissions for LDAP authentication – Use the following guidance to resolve the issue [Set up HDFS Permissions using Linux Credentials](#). You can log into the cluster using the following commands:

  ```bash
  hdfs dfs -mkdir /user/USERNAME
  hdfs dfs -chown USERNAME:USERNAME /user/USERNAME
  ```

- LDAP authentication missing connection key error – You might see the following error message: `Data Wrangler couldn't connect to EMR hive successfully. JDBC connection is missing required connection key(s): PWD.`

  For LDAP authentication, you must specify both a username and a password. The JDBC URL stored in Secrets Manager is missing property `PWD`.

- When you're troubleshooting the LDAP configuration: We recommend making sure that the LDAP authenticator (LDAP server) is correctly configured to connect to the Amazon EMR cluster. Use the `ldapwhoami` command to help you resolve the configuration issue. The following are example commands that you can run:

  - For LDAPS: `ldapwhoami -x -H ldaps://ldap-server`
  - For LDAP: `ldapwhoami -x -H ldap://ldap-server`
Either command should return Anonymous if you've configured the authenticator successfully.

**Troubleshooting with Salesforce**

**Lifecycle configuration error**

When your user opens Studio Classic for the first time, they might get an error saying that there's something wrong with their lifecycle configuration. Use Amazon CloudWatch to access the logs written by your lifecycle configuration script. For more information about debugging lifecycle configurations, see [Debug lifecycle configurations](#).

If you aren't able to debug the error, you can create the configuration file manually. You must create the file every time you delete or restart the Jupyter server. Use the following procedure to create the file manually.

**To create a configuration file**

1. Navigate to Studio Classic.
2. Choose **File**, then **New**, then **Terminal**.
4. Open the file in a text editor.
5. Add a JSON object containing the Amazon Resource Name (ARN) of the Secrets Manager secret to the file. You can use the following template to create the object.

   ```json
   {
     "secret_arn": "example-secret-ARN"
   }
   ```

6. Save your changes to the file.

**Unable to access Salesforce Data Cloud from the Data Wrangler flow**

After your user chooses **Salesforce Data Cloud** from your Data Wrangler flow, they might get an error indicating the prerequisites to set up the connection haven't been met. It might be caused by following errors:

- The Salesforce secret in Secrets Manager hasn't been created.
• The Salesforce secret in Secrets Manager has been created, but it's missing the Salesforce tag.

• The Salesforce secret in Secrets Manager has been created in the wrong AWS Region. For example, your user won't be able to access the Salesforce Data Cloud in `ca-central-1` because you've created the secret in `us-east-1`. You can either replicate the secret to `ca-central-1` or create a new secret with the same credentials in `ca-central-1`. For information about replicating secrets, see [Replicate an AWS Secrets Manager secret to other AWS Regions](#).

• The policy that your users are using to access Amazon SageMaker Studio Classic are missing permissions for AWS Secrets Manager

• There's a typo in the Secrets Manager ARN of the JSON object that you've specified through your lifecycle configuration.

• There's a typo in the Secrets Manager secret containing your Salesforce OAuth configuration

**Blank page showing redirect_uri_mismatch**

After your users choose **Save and Connect**, they might get redirected to a page that shows `redirect_uri_mismatch`. The callback URI that you've registered in your Salesforce Connected App settings is either missing or incorrect.

Use the following URL to check that your Studio Classic URL is correctly registered in your Salesforce org's Connected App settings: `https://EXAMPLE_SALESFORCE_ORG/lightning/setup/NavigationMenus/home/`. For more information about using the connected app settings, navigate to the following URL: `https://EXAMPLE_SALESFORCE_ORG/lightning/setup/NavigationMenus/home/`.

ℹ️ **Note**

It takes roughly ten minutes to propagate the URI within Salesforce's systems.

**Shared spaces**

Shared spaces doesn't currently work with the Salesforce Data Cloud integration. You can either delete the shared spaces in the Amazon SageMaker domain that you intend to use, or you can use another domain that doesn't have shared spaces set up.
OAuth Redirect Error

Your users should be able to import their data from the Salesforce Data Cloud after they choose Connect. If they're running into an error, we recommend asking them to do the following:

• Tell them to be patient – When they get redirected back to Amazon SageMaker Studio Classic, it can take up to a minute to complete the authentication process. While they're getting redirected, we recommend telling them to avoid interacting with the browser. For example, they shouldn't close the browser tab, switch to another tab, or interact with the Data Wrangler flow. Interacting with the browser might remove the authorization code required to connect to the data cloud.

• Have your users reconnect to the data cloud – There are transient issues that can cause a connection to the Salesforce Data Cloud to fail. Have your users create a new Data Wrangler flow and try connecting to the Salesforce Data Cloud again.

• Make sure your users close all other tabs with Amazon SageMaker Studio Classic – Having Studio Classic open in multiple tabs can cause the Salesforce Data Cloud connection to fail. Make sure your users only have one Studio Classic tab open.

• Multiple users accessing Studio Classic at the same time – Only one user should access an Amazon SageMaker domain at a time. If multiple users access the same domain, the connection that a user is trying to create to the Salesforce Data Cloud might fail.

Updating both Data Wrangler and Studio Classic might also fix their error. For information about updating Data Wrangler, see Update Data Wrangler. For information about updating Studio Classic, see Shut down and Update SageMaker Studio Classic.

If none of the preceding troubleshooting steps work, you might find an error message from Salesforce with a corresponding description embedded in the Studio Classic URL. The following is an example of a message you could find: error=invalid_client_id&error_description=client%20identifier%20invalid.

You can look at the error message in the URL and try to address the issues it presents. If the error message or description is unclear, we recommend searching the Salesforce Knowledge Base. If searching the knowledge base doesn't work, you can reach out to the Salesforce help desk for more assistance.

Data Wrangler takes a long time to load

When your users are getting redirected back to Data Wrangler from the Salesforce Data Cloud, they might experience long load times.
If this is the user's first time using Data Wrangler or they've deleted the kernel, it might take about 5 minutes to provision the new Amazon EC2 instance to use Data Wrangler.

If this isn't the user's first time using Data Wrangler and they haven't deleted the kernel, you can ask them to refresh the page or close as many browser tabs as possible.

If none of the preceding interventions work, have them set up a new connection to the Salesforce Data Cloud.

**User fails to export their data with an Invalid batch Id error**

When your user exports the transformations that they've made to their Salesforce data, the SageMaker processing job that Data Wrangler uses on the backend might fail. The Salesforce Data Cloud might be temporarily unavailable or there could be a caching issue.

To address the issue, we recommend having your users go back to the step where they're importing the data and changing the order of the columns that they're querying. For example, they can change the following query:

```
SELECT col_A, col_B FROM table
```

To the following query:

```
SELECT col_B, col_A FROM table
```

After they've changed the order of the columns and made sure that the subsequent transformations they've made are still valid, they can start exporting their data again.

**Users can't export a very large dataset**

If your users imported a very large dataset from the Salesforce Data Cloud, they might not be able to export the transformations that they've made. A large dataset might have too many rows, or it can result from a complex query.

We recommend having your users take the following actions:

- Simplifying their SQL query
• Sampling their data

The following are some strategies that they can use to simplify their queries:

• Specify column names instead of using the * operator
• Finding a subset of the data that they'd like to import instead of using a larger subset
• Minimizing joins between very large datasets

They can use sampling to reduce the number of rows in their dataset. For information about sampling methods, your users can refer to Sampling.

Users can't export data due to invalid refresh token

Data Wrangler uses a JDBC driver to integrate with the Salesforce Data Cloud. The method for authentication is OAuth. For OAuth, the refresh token and the access token are two different pieces of data that are used to authorize access to resources within your Salesforce Data Cloud.

The access token, or core token, is what allows you to access your Salesforce data and run queries directly through Data Wrangler. It's short lived and designed to expire quickly. To maintain access to your Salesforce data, Data Wrangler uses the refresh token to get a new access token from Salesforce.

You might have set the refresh to expire too quickly to get a new access token for your users. You might have to revisit your refresh token policy to make sure that it can accommodate queries that take a long time to run for your users. For information about configuring your refresh token policy, see https://EXAMPLE_SALESFORCE_ORG_URL/lightning/setup/ConnectedApplication/home/.

Queries failing or tables not loading

Salesforce experiences service outages. Even if you’ve configured everything correctly, your users might not be able to import their data for periods of time.

Service outages can happen for maintenance reasons. We recommend checking in the following day to see if the issue has been resolved.

If you’re experiencing issues for more than a day, we recommend contacting Salesforce’s help desk for further assistance. For information about contacting Salesforce, see How would you like to contact Salesforce?
**OAUTH_APP_BLOCKED during Studio Classic redirect**

When your user gets redirected back to Amazon SageMaker Studio Classic, they might notice the query parameter error=OAUTH_APP_BLOCKED within the URL. They’re might be experiencing a transient issue that should resolve itself within a day.

It’s possible that you’ve blocked their access to the Connected App as well. For information about resolving the issue, see https://EXAMPLE_SALESFORCE_ORG_URL/lightning/setup/ConnectedApplication/home/.

**OAUTH_APP_DENIED during Studio Classic redirect**

When your user gets redirected back to Amazon SageMaker Studio Classic, they might notice the query parameter error=OAUTH_APP_ACCESS_DENIED within the URL. You haven’t given their profile type permissions to access the Connected App associated with Data Wrangler.

To resolve their access issue, navigate to https://EXAMPLE_SALESFORCE_ORG_URL/lightning/setup/ManageUsers/home/ and check whether the user is assigned to the correct profile.

**Increase Amazon EC2 Instance Limit**

You might see the following error message when you’re using Data Wrangler: The following instance type is not available: ml.m5.4xlarge. Try selecting a different instance below.

The message can indicate that you need to select a different instance type, but it can also indicate that you don’t have enough Amazon EC2 instances to successfully run Data Wrangler on your workflow. You can increase the number of instances by using the following procedure.

To increase the number of instances, do the following.

1. Open the AWS Management Console.
2. In the search bar, specify Services Quotas.
3. Choose Service Quotas.
4. Choose AWS services.
5. In the search bar, specify Amazon SageMaker.
6. Choose Amazon SageMaker.
7. Under **Service quotas**, specify **Studio KernelGateway Apps running on ml.m5.4xlarge instance.**

   **Note**
   
   ml.m5.4xlarge is the default instance type for Data Wrangler. You can use other instance types and request quota increases for them. For more information, see [Instances](#).

8. Select **Studio KernelGateway Apps running on ml.m5.4xlarge instance.**
9. Choose **Request quota increase.**
10. For **Change quota value**, specify a value greater than **Applied quota value.**
11. Choose **Request.**

If your request is approved, AWS sends a notification to the email address associated with your account. You can also check the status of your request by choosing **Quota request history** on the **Service Quotas** page. Processed requests have a **Status** of **Closed**.

**Update Data Wrangler**

To update Data Wrangler to the latest release, first shut down the corresponding KernelGateway app from the Amazon SageMaker Studio Classic control panel. After the KernelGateway app is shut down, restart it by opening a new or existing Data Wrangler flow in Studio Classic. When you open a new or existing Data Wrangler flow, the kernel that starts contains the latest version of Data Wrangler.

**Update your Studio Classic and Data Wrangler instance**

1. Navigate to your [SageMaker Console](#).
2. Choose SageMaker and then Studio Classic.
3. Choose your user name.
4. Under **Apps**, in the row displaying the **App name**, choose **Delete app** for the app that starts with `sagemaker-data-wrangler`, and for the JupyterServer app.
5. Choose **Yes, delete app.**
6. Type delete in the confirmation box.
7. Choose **Delete.**
8. Reopen your Studio Classic instance. When you begin to create a Data Wrangler flow, your instance now uses the latest version of Data Wrangler.

Alternatively, if you are using a Data Wrangler application version that is not the latest version, and you have an existing Data Wrangler flow open, you are prompted to update your Data Wrangler application version in the Studio Classic UI. The following screenshot shows this prompt.

⚠️ **Important**

This updates the Data Wrangler kernel gateway app only. You still need to shut down the JupyterServer app in your user account. To do this, follow the preceding steps.

You can also choose **Remind me later**, in which case an **Update** button appears in the top-right corner of the screen.
Shut Down Data Wrangler

When you are not using Data Wrangler, it is important to shut down the instance on which it runs to avoid incurring additional fees.

To avoid losing work, save your data flow before shutting Data Wrangler down. To save your data flow in Studio Classic, choose File and then choose Save Data Wrangler Flow. Data Wrangler automatically saves your data flow every 60 seconds.

To shut down the Data Wrangler instance in Studio Classic

1. In Studio Classic, select the Running Instances and Kernels icon ( ).
2. Under RUNNING APPS is the sagemaker-data-wrangler-1.0 app. Select the shutdown icon next to this app ( ).
Data Wrangler runs on an ml.m5.4xlarge instance. This instance disappears from **RUNNING INSTANCES** when you shut down the Data Wrangler app.

⚠️ **Important**

If you open Data Wrangler again, an Amazon EC2 instance starts running the application and you will be charged for the compute. In addition to compute, you are also charged for the storage that you use. For example, you're charged for any Amazon S3 buckets that you're using with Data Wrangler.

If you find that you're still getting charged for Data Wrangler after shutting down your applications, there's a Jupyter extension that you can use to automatically shut down idle sessions. For information about the extension, see [SageMaker-Studio-Autoshutdown-Extension](#).

After you shut down the Data Wrangler app, it has to restart the next time you open a Data Wrangler flow file. This can take a few minutes.

### Prepare Data at Scale with Studio Classic using Amazon EMR or AWS Glue

Amazon SageMaker Studio Classic provides data scientists, machine learning (ML) engineers, and general practitioners with tools to perform data analytics and data preparation at scale. Analyzing, transforming, and preparing large amounts of data is a foundational step of any data science and ML workflow. SageMaker Studio Classic comes with built-in integration of Amazon EMR and AWS Glue Interactive Sessions to handle your large-scale interactive data preparation and machine learning workflows, all within your Studio Classic notebook.

**Amazon EMR** is a managed big data platform with resources to help you run petabyte-scale distributed data processing jobs using open-source analytics frameworks on AWS such as Apache Spark, Apache Hive, Presto, HBase, Flink, and Hudi among others. Data engineers and data scientists use Amazon EMR for a wide variety of use cases, including big data analytics, what-if analyses, real-time analytics, and data preparation for machine learning. With Studio Classic integration with Amazon EMR, you can create, browse, discover, and connect to Amazon EMR clusters without leaving your Studio Classic notebook. You can also monitor and debug your Spark workloads with one-click access to the Spark UI from within the notebook. You should consider
Amazon EMR for your data preparation workloads if you want maximum control over hardware and software versions, containers, and big data processing applications.

AWS Glue Interactive Sessions is a serverless service that you can enlist to collect, transform, cleanse, and prepare data for storage in your data lakes and data pipelines. AWS Glue Interactive Sessions provides an on-demand, serverless Apache Spark runtime environment that you can initialize in seconds on a dedicated Data Processing Unit (DPU) without having to worry about provisioning and managing complex compute cluster infrastructure. After initialization, you can quickly browse the AWS Glue data catalog, run large queries, access data governed by AWS Lake Formation, and interactively analyze and prepare data using Spark, right in your Studio Classic notebook. You can then use the prepared data to train, tune, and deploy models using the purpose-built ML tools within SageMaker Studio Classic. You should consider AWS Glue Interactive Sessions for your data preparation workloads when you want a serverless Spark service with moderate control of configurability and flexibility.

Content

- Prepare data using Amazon EMR
- Prepare data using AWS Glue Interactive Sessions

Prepare data using Amazon EMR

Amazon SageMaker Studio Classic comes with built-in integration of Amazon EMR, with which data scientists and data engineers can perform petabyte-scale interactive data preparation and machine learning (ML) right from their Studio Classic notebook. Within a notebook, they can discover and connect to existing Amazon EMR clusters, then interactively explore, visualize, and prepare large-scale data for machine learning using Apache Spark, Apache Hive, Presto. Additionally, users can access Spark UI with a single click to monitor their Spark jobs from their Studio Classic notebooks.

Administrators can use the AWS Service Catalog to define AWS CloudFormation templates of Amazon EMR clusters accessible to Studio Classic users. Data scientists can then choose a predefined template to self-provision an Amazon EMR cluster directly from Amazon SageMaker Studio Classic notebooks. Administrators can further parameterize the templates to let users choose aspects of the cluster to match their workloads within predefined values. For example, a data scientist or data engineer may want to specify the number of core nodes of the cluster up to a predetermined maximum value, or select the instance type of a node from a dropdown menu.
• If you are an administrator, make sure that you have enabled communication between Amazon SageMaker Studio Classic notebooks and Amazon EMR clusters. For instructions, see the Configure networking (for administrators) section. Once this communication is enabled, you have the option to:

• Define cluster templates in AWS Service Catalog and ensure the availability of these templates through Studio Classic’s notebooks: Configure Amazon EMR templates in AWS Service Catalog (for administrators).

• Configure the discoverability of existing Amazon EMR clusters directly from Studio Classic’s notebooks: Configure the discoverability of Amazon EMR clusters (for administrators).

• If you are a data scientist or data engineer looking to self-provision an Amazon EMR cluster, see Launch an Amazon EMR cluster from Studio Classic.

• If you are a data scientist or data engineer looking to discover and connect to existing Amazon EMR clusters from Studio Classic, see Use Amazon EMR clusters from Studio Classic notebooks.

List of topics

• Configure networking (for administrators)
• Create an Amazon EMR cluster from Studio Classic notebooks
• Use Amazon EMR clusters from Studio Classic notebooks
• Access Spark UI from Studio Classic
• Walkthroughs and whitepapers
• Additional Configuration for cross accounts use cases (for administrators)
• Troubleshooting

Configure networking (for administrators)

This section provides information about how administrators can configure their network to allow communication between Amazon SageMaker Studio Classic notebooks and an Amazon EMR cluster.

The networking instructions vary based on whether SageMaker Studio Classic and Amazon EMR are deployed within a private Amazon Virtual Private Cloud (VPC) or communicate over the internet.

By default, SageMaker Studio Classic runs in an AWS managed VPC with internet access. When using an internet connection, Studio Classic accesses AWS resources, such as Amazon S3 buckets,
over the internet. However, if you have security requirements to control access to your data and job containers, we recommend that you configure SageMaker Studio Classic and Amazon EMR so that your data and containers aren’t accessible over the internet. To control access to your resources or run SageMaker Studio Classic without public internet access, you can specify the VPC only network access type when you onboard to Amazon SageMaker domain. In this scenario, SageMaker Studio Classic establishes connections with other AWS services via private VPC endpoints. For information about configuring SageMaker Studio Classic in VPC only mode, see Connect SageMaker Studio Classic notebooks in a VPC to external resources.

The first two sections describe how to ensure communication between SageMaker Studio Classic and an Amazon EMR cluster in VPCs without public internet access. The last section covers how to ensure communication between SageMaker Studio Classic and Amazon EMR using an internet connection. Prior to connecting SageMaker Studio Classic and Amazon EMR without internet access, make sure to establish endpoints for Amazon Simple Storage Service (data storage), Amazon CloudWatch (logging and monitoring), and Amazon SageMaker Runtime (fine-grained role-based access control (RBAC)).

- If your Amazon SageMaker Studio Classic and Amazon EMR cluster are set up in different VPCs in the same AWS account or in different accounts, see Studio Classic and Amazon EMR are deployed in separate VPCs.
- If your Amazon SageMaker Studio Classic and Amazon EMR cluster are set up in the same VPC, see Amazon SageMaker Studio Classic and Amazon EMR are in the same VPC.
- If you chose to connect Amazon SageMaker Studio Classic and Amazon EMR cluster over public internet, see Amazon SageMaker Studio Classic and Amazon EMR communicate over public internet.

**Studio Classic and Amazon EMR are deployed in separate VPCs**

To allow communication between SageMaker Studio Classic and an Amazon EMR cluster when they are deployed in different VPCs:

1. Start by connecting your VPCs through a VPC peering connection.
2. Update your routing tables in each VPC to route the network traffic between Studio Classic subnets and Amazon EMR subnets both ways.
3. Configure your security groups to allow inbound and outbound traffic.
The steps are similar, regardless of whether Amazon SageMaker Studio Classic and the Amazon EMR cluster are deployed within the same AWS account (Single account use case) or different AWS accounts (Cross accounts use case).

1. **VPC peering**

   Create a [VPC peering connection](#) to facilitate the networking between the two VPCs (SageMaker Studio Classic and Amazon EMR).

   a. From your SageMaker Studio Classic account, on the VPC dashboard, choose **Peering connections**, then **Create peering connection**.

   b. Create your request to peer the Studio Classic VPC within the Amazon EMR VPC. When requesting peering in another AWS account, choose **Another account** in **Select another VPC to peer with**.

      For cross accounts peering, the administrator must accept the request from the Amazon EMR account.

      When peering private subnets, you should enable private IP DNS resolution at the VPC peering connection level.

2. **Routing tables**

   Send the network traffic between SageMaker Studio Classic subnets and Amazon EMR subnets both ways.

   After you establish the peering connection, the administrator (on each account for cross-account access) can add routes to the private subnet route tables to route the traffic between the notebooks and the cluster subnets. You can define those routes by going to the **Route Tables** section of each VPC in the VPC dashboard.

   The following illustration of the route table of a Studio Classic VPC subnet shows an example of an outbound route from the Studio Classic account to the Amazon EMR VPC IP range (here 2.0.1.0/24) through the peering connection.
The following illustration of a route table of an Amazon EMR VPC subnet shows an example of return routes from the Amazon EMR VPC to Studio Classic VPC IP range (here 10.0.20.0/24) through the peering connection.

### Security groups

Lastly, the security group of your Studio Classic domain must allow outbound traffic, and the security group of the Amazon EMR primary node must allow inbound traffic on Apache Livy, Hive, or Presto TCP ports (respectively 8998, 10000, and 8889) from the Studio Classic instance security group. Apache Livy is a service that enables interaction with a Amazon EMR cluster over a REST interface.

The following image shows an example of an Amazon VPC setup that enables SageMaker Studio Classic notebooks to provision Amazon EMR clusters from AWS CloudFormation templates and then connect to an Amazon EMR cluster within the same AWS account. The diagram provides an additional illustration of the required endpoints for a direct connection to various AWS services, such as Amazon S3 or Amazon CloudWatch, when the VPCs have no internet access. Alternatively, a NAT gateway must be used to allow instances in private subnets of multiple VPCs to share a single public IP address provided by the internet gateway when accessing the internet.
Amazon SageMaker Studio Classic and Amazon EMR are in the same VPC

If Amazon SageMaker Studio Classic and the cluster are in different subnets, add routes to each private subnet route table to route the traffic between the notebooks and the cluster subnets. You can define those routes by going to the Route Tables section of each VPC in the VPC dashboard. If you deployed Amazon SageMaker Studio Classic and an Amazon EMR cluster in the same VPC and the same subnet, you do not need to route the traffic between the notebooks and the cluster.

Whether or not you needed to update your routing tables, the security group of your Studio Classic domain must allow outbound traffic, and the security group of the Amazon EMR primary node must allow inbound traffic on Apache Livy, Hive, or Presto TCP ports (respectively 8998, 10000, and 8889) from the Studio Classic instance security group. Apache Livy is a service that enables interaction with a Amazon EMR cluster over a REST interface.

Amazon SageMaker Studio Classic and Amazon EMR communicate over public internet

By default, SageMaker Studio Classic provides a network interface that allows communication with the internet through an internet gateway in the VPC associated with the SageMaker domain. If you
choose to connect to Amazon EMR through the public internet, your Amazon EMR cluster needs to accept inbound traffic on Apache Livy, Hive, or Presto TCP ports (respectively 8998, 10000, and 8889) from its internet gateway. *Apache Livy* is a service that enables interaction with an Amazon EMR cluster over a REST interface.

Keep in mind that any port on which you allow inbound traffic represents a potential security vulnerability. Carefully review custom security groups to ensure that you minimize vulnerabilities. For more information, see [Control network traffic with security groups](#).

Alternatively, see [Walkthroughs and whitepapers](#) for a detailed walkthrough of how to enable Kerberos on Amazon EMR, set the cluster in a private subnet, and access the cluster using a [Network Load Balancer (NLB)](#) to expose only specific ports, which are access-controlled via security groups.

**Note**

When connecting to your Apache Livy endpoint through the public internet, we recommend that you secure communications between Amazon SageMaker Studio Classic and your Amazon EMR cluster using TLS. For information on setting up HTTPS with Apache Livy, see [Enabling HTTPS with Apache Livy](#). For information on setting an Amazon EMR cluster with transit encryption enabled, see [Providing certificates for encrypting data in transit with Amazon EMR encryption](#). Additionally, you need to configure Studio Classic to access your certificate key as specified in [Connect to an Amazon EMR cluster over HTTPS](#).

### Create an Amazon EMR cluster from Studio Classic notebooks

Administrators can use [AWS Service Catalog](#) to define [AWS CloudFormation templates](#) of Amazon EMR clusters as products of a portfolio, then make them available to selected users. Using the Service Catalog, administrators can fully control the organizational, security, and networking setup of Amazon EMR clusters. Data scientists and data engineers can then view, select, and customize those templates for their specific workloads to create on-demand Amazon EMR clusters directly from their SageMaker Studio Classic notebooks. This can be done without manually setting up complex configurations. Users can also terminate Amazon EMR clusters from Studio Classic notebooks after use.
• If you are an administrator looking to configure AWS CloudFormation templates as AWS Service Catalog products so users can create Amazon EMR clusters from Studio Classic, see Configure Amazon EMR templates in AWS Service Catalog (for administrators).

• If you are a data scientist or data engineer looking to self-provision an Amazon EMR cluster to process data at scale using open-source frameworks such as Apache Spark, Apache Hive, or Presto, see Launch an Amazon EMR cluster from Studio Classic.

• If you are looking to discover and connect to existing Amazon EMR clusters from Studio Classic, see Use Amazon EMR clusters from Studio Classic notebooks.

Topics

• Configure Amazon EMR templates in AWS Service Catalog (for administrators)

• Launch an Amazon EMR cluster from Studio Classic

Configure Amazon EMR templates in AWS Service Catalog (for administrators)

This section provides details about how administrators can configure an AWS Service Catalog product so users can independently self-provision Amazon EMR clusters from Amazon SageMaker Studio Classic notebooks. Additionally, administrators can configure the Amazon EMR cluster templates in a way so end-users can customize various aspects of the cluster to suit their specific requirements. For example, the administrator can define a list of permissible instance types from which users can choose when creating cluster.

This topic assumes that you are familiar with the creation of portfolios and products in AWS Service Catalog as well as Amazon EMR, and AWS CloudFormation.

Note

You can refer to the AWS CloudFormation templates in aws-samples/sagemaker-studio-emr GitHub repository as examples of CloudFormation stacks to deploy IAM roles, Amazon VPCs, a sandbox Studio Classic domain, a user profile, as well as a CloudFormation template to launch an Amazon EMR cluster. Several options are available depending on your authentication method between Studio Classic and the Amazon EMR cluster. In these examples, a parent CloudFormation template passes the SageMaker VPC ID, security group, and subnet ID parameters to the CloudFormation template of an Amazon EMR cluster.
You can access various examples of CloudFormation Amazon EMR templates in the nested repository [sagemaker-studio-emr/cloudformation/emr_servicecatalog_templates](https://sagemaker-studio-emr/cloudformation/emr_servicecatalog_templates) and further choose from a single account deployment to cross accounts. For more information about the authentication methods available when connecting to an Amazon EMR cluster, see [Use Amazon EMR clusters from Studio Classic notebooks](#).

To simplify the creation of Amazon EMR clusters, administrators can register the [CloudFormation template of an Amazon EMR cluster](#) as a product in the portfolio of the AWS Service Catalog. Then they associate the Service Catalog portfolio with the Studio Classic execution role to ensure the availability of the template in Studio Classic. Furthermore, to make sure that data scientists can discover those templates, provision Amazon EMR clusters, and connect to Amazon EMR clusters from their Studio Classic notebooks, administrators need to set the proper access permissions.

The following list provides the additional settings that administrators need to apply to a baseline CloudFormation stack to enable Studio Classic to access the Service Catalog products and provision Amazon EMR clusters. Those settings must be applied at multiple levels:

- In the Service Catalog portfolio
- In the Service Catalog product
- In the CloudFormation Amazon EMR template declared as the Service Catalog product

Finally, administrators must assign the required permissions to both the Studio Classic execution role accessing the clusters and the account where Amazon EMR is deployed, based on whether Studio Classic and Amazon EMR are in the same or different AWS accounts.

- **Pre-requisites:** **Networking and authentication requirements**
  
  As a prerequisite, ensure that you have reviewed the networking and security requirements in [Configure networking (for administrators)](#) and that you have created a baseline CloudFormation stack supporting the authentication method of your choice. You can find examples of CloudFormation templates in [aws-samples/sagemaker-studio-emr](https://aws-samples/sagemaker-studio-emr).

- **In your Service Catalog portfolio:**

  Add the following section to your portfolio CloudFormation template (see the example in YAML format) to associate your portfolio with the Studio Classic execution role accessing your cluster.

  ```yaml
  SageMakerStudioEMRProductPortfolioPrincipalAssociation:
  ```

Prepare data using Amazon EMR 2388
Type: AWS::ServiceCatalog::PortfolioPrincipalAssociation
Properties:
  PrincipalARN: SageMakerExecutionRole.Arn
  PortfolioId: SageMakerStudioEMRProductPortfolio ID
  PrincipalType: IAM

- In your Service Catalog product:

  Add the following tag key "sagemaker:studio-visibility:emr" and set to the value "true" (here in YAML) to the Service Catalog product referencing the Amazon EMR template resource. This ensures the visibility of the template in Studio Classic.

  SMStudioEMRNoAuthProduct:
    Type: AWS::ServiceCatalog::CloudFormationProduct
    Properties:
      Owner: AWS
      Name: SageMaker Studio Domain No Auth EMR
      ProvisioningArtifactParameters:
        - Name: SageMaker Studio Domain No Auth EMR
          Description: Provisions a SageMaker domain and No Auth EMR Cluster
          Info:
            LoadTemplateFromURL: Link to your CloudFormation template. For example, https://aws-ml-blog.s3.amazonaws.com/artifacts/astra-m4-sagemaker/end-to-end/CFN-EMR-NoStudioNoAuthTemplate-v3.yaml
            Tags:
              - Key: "sagemaker:studio-visibility:emr"
                Value: "true"

- In the CloudFormation template of the Amazon EMR cluster within your Service Catalog product:

  Add the following mandatory stack parameters as a placeholder. This section is populated with the Studio Classic project name and identifier used by the user when provisioning a cluster from Studio Classic.

  SageMakerProjectName:
    Type: String
    Description: Name of the project

  SageMakerProjectId:
    Type: String
    Description: Service generated Id of the project.
Administrators can specify Default and AllowedValues to include choices in the parameters section of a template, letting users input or select custom values when creating a cluster. The following example illustrates additional input parameters that administrators can set when creating an Amazon EMR template.

"Parameters": {
    "EmrClusterName": { 
        "Type": "String",
        "Description": "EMR cluster Name."
    },
    "MasterInstanceType": { 
        "Type": "String",
        "Description": "Instance type of the EMR master node.",
        "Default": "m5.xlarge",
        "AllowedValues": [ 
            "m5.xlarge",
            "m5.2xlarge",
            "m5.4xlarge"
        ]
    },
    "CoreInstanceType": { 
        "Type": "String",
        "Description": "Instance type of the EMR core nodes.",
        "Default": "m5.xlarge",
        "AllowedValues": [ 
            "m5.xlarge",
            "m5.2xlarge",
            "m5.4xlarge",
            "m3.medium",
            "m3.large",
            "m3.xlarge",
            "m3.2xlarge"
        ]
    },
    "CoreInstanceCount": { 
        "Type": "String",
        "Description": "Number of core instances in the EMR cluster.",
        "Default": "2",
        "AllowedValues": [ 
            "2",
            "5",
            "10"
        ]
    }
}
Last, attach the required IAM policies to enable the visibility of CloudFormation Amazon EMR templates and the self-provisioning of Amazon EMR clusters from the Studio Classic notebooks. The role to which you must add those policies depends on whether Studio Classic and Amazon EMR are deployed in the same account (single account) or in different accounts (cross accounts).

- If your Amazon EMR cluster is deployed in the same AWS account as the Studio Classic account, refer to the Single Account tab.
- If your Amazon EMR cluster is deployed in a different AWS account than the Studio Classic account, refer to the Cross Accounts tab.

For more information on cross-account access using roles, see Cross account resource access in IAM or Cross-account policy evaluation logic.

Single account

Attach the following permissions to the Studio Classic execution role accessing your cluster.

The following list provides a breakdown of the permissions required.

- AllowEMRTemplateDiscovery allows the discoverability for Amazon EMR templates.
- AllowSagemakerProjectManagement enables the creation of SageMaker projects. In Studio Classic, access to the AWS Service Catalog is granted through Projects.
- AllowClusterDetailsDiscovery and AllowClusterDiscovery allow the discovery and connection to Amazon EMR clusters.
- AllowPresignedUrl allows the creation of pre-signed URLs to access Spark UI.

The following is a comprehensive JSON that includes these permissions.

```json
{}
```

```json


"Version": "2012-10-17",
"Statement": [
  {
    "Sid": "AllowPresignedUrl",
    "Effect": "Allow",
    "Action": [
      "elasticmapreduce:DescribeCluster",
      "elasticmapreduce:ListInstanceGroups",
      "elasticmapreduce:CreatePersistentAppUI",
      "elasticmapreduce:DescribePersistentAppUI",
      "elasticmapreduce:GetPersistentAppUIPresignedURL",
      "elasticmapreduce:GetOnClusterAppUIPresignedURL"
    ],
    "Resource": [
    ]
  },
  {
    "Sid": "AllowClusterDetailsDiscovery",
    "Effect": "Allow",
    "Action": [
      "elasticmapreduce:DescribeCluster",
      "elasticmapreduce:ListInstances",
      "elasticmapreduce:ListInstanceGroups",
      "elasticmapreduce:DescribeSecurityConfiguration"
    ],
    "Resource": [
    ]
  },
  {
    "Sid": "AllowClusterDiscovery",
    "Effect": "Allow",
    "Action": [
      "elasticmapreduce:ListClusters"
    ],
    "Resource": "*"
  },
  {
    "Sid": "AllowEMRTemplateDiscovery",
    "Effect": "Allow",
    "Action": [
      "servicecatalog:SearchProducts"
    ],
  }
]
Cross accounts

If your Amazon EMR clusters and Studio Classic are deployed in separate AWS accounts, you configure the permissions in multiple steps.

- **On the trusting account** (the account in which Amazon EMR is deployed), create a custom IAM role (referred to as ASSUMABLE-ROLE in this page) with the following permissions and trust relationship.

For information about creating a role on an AWS account, see [Creating an IAM role (console)](creating-an-iam-role-console).

1. Add an IAM policy defining the following permissions.

- **AllowClusterDetailsDiscovery** and **AllowClusterDiscovery** to allow the discovery and connection to Amazon EMR clusters.
- **AllowPresignedUrl** to allow the creation of pre-signed URLs to access Spark UI.

The following is a comprehensive JSON that includes these permissions.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "AllowPresignedUrl",
      "Effect": "Allow",
      "Action": [
        "elasticmapreduce:DescribeCluster",
        "elasticmapreduce:ListInstanceGroups",
      ],
    }
  ]
}
```
2. To grant the trusted account (the account in which Studio Classic is deployed) the permission to assume a role in the trusting account, include the following trust relationship.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "aws:PrincipalId": "arn:aws:sts::emr-region:emr-account:cluster/"
      },
      "Action": "sts:AssumeRole"
    }
  ]
}
```
On the trusted account (the account in which Studio Classic is deployed), add the following permissions and trust relationship to the Studio Classic execution role.

1. Add an IAM policy defining the following permissions.
   - AllowSagemakerProjectManagement to allow the creation of SageMaker projects. In Studio Classic, access to the AWS Service Catalog is granted through Projects.
   - AllowEMRTemplateDiscovery to allow the discoverability of Amazon EMR templates.

The following is a comprehensive JSON that includes these permissions.

```json
{
    "Version": "2012-10-17",
    "Statement": [
    {
        "Sid": "AllowSagemakerProjectManagement",
        "Effect": "Allow",
        "Action": [
            "sagemaker:CreateProject",
            "sagemaker:DeleteProject"
        ],
        "Resource": "arn:aws:sagemaker:::project/*"
    },
    {
        "Sid": "AllowEMRTemplateDiscovery",
        "Effect": "Allow",
        "Action": [
            "servicecatalog:SearchProducts"
        ],
        "Resource": "*"
    }
]
}
```
2. To grant the Studio Classic execution role the permission to assume the ASSUMABLE-ROLE in the **trusting account**, include the following trust relationship.

```
{
   "Version": "2012-10-17",
   "Statement": [
   {
      "Sid": "AllowRoleAssumptionForCrossAccountDiscovery",
      "Effect": "Allow",
      "Action": "sts:AssumeRole",
      "Resource": ["arn:aws:iam::emr-account:role/ASSUMABLE-ROLE"]
   }]
}
```

- Last, see [Additional Configuration for cross accounts use cases (for administrators)](http://example.com) to learn how to provide the ARN of the ASSUMABLE-ROLE to the Studio Classic execution role. The ARN is loaded by the Studio Classic Jupyter server at launch. The Studio Classic execution role assumes that cross-account role to discover and connect to Amazon EMR clusters in the **trusting account**.

Once the CloudFormation templates are available in Amazon SageMaker Studio Classic, data scientists can use them to self-provision Amazon EMR clusters. Each of the "Parameters" specified in the template becomes an input box in the cluster creation form of Studio Classic, with the corresponding "AllowedValues" appearing in a dropdown menu.

The following illustration shows the dynamic form assembled from a CloudFormation Amazon EMR template to create an Amazon EMR cluster in SageMaker Studio Classic.
Visit Launch an Amazon EMR cluster from Studio Classic to learn about how to launch a cluster from Studio Classic using those Amazon EMR templates.

**Launch an Amazon EMR cluster from Studio Classic**

Data scientists and data engineers can self-provision Amazon EMR clusters from Studio Classic using AWS CloudFormation templates configured by their administrators. If you are an administrator looking to configure CloudFormation templates as AWS Service Catalog products so users can create Amazon EMR clusters from Studio Classic, see Configure Amazon EMR templates in AWS Service Catalog (for administrators).

To provision a new Amazon EMR cluster from Studio Classic:

1. Select the **Home** icon in the Studio Classic UI's left-side panel, then select the **Data** node in the navigation menu. Navigate down to the **Clusters** node. This opens up a page listing the Amazon EMR clusters that you can access from SageMaker Studio Classic.

Prepare data using Amazon EMR
2. Choose **Create cluster**. This opens up a page, in the main working area, listing the cluster templates available to you.

3. Select a cluster configuration template by choosing a template name. The selection of a template activates the **Select template** button. Choose **Select template**. This opens up a cluster creation form.

4. Enter the cluster's details, such as a cluster name and any specific configurable parameter set by your administrator, then choose **Create cluster**. The creation of the cluster might take a couple of minutes.

Once the cluster is provisioned, the Studio Classic UI displays a *The cluster has been successfully created* message.

To connect to your cluster, see [Use Amazon EMR clusters from Studio Classic notebooks](#)

**Use Amazon EMR clusters from Studio Classic notebooks**

In this section, you learn about how to discover, connect to, or terminate an Amazon EMR cluster from SageMaker Studio Classic notebooks.
• If you are an administrator, see Configure the discoverability of Amazon EMR clusters (for administrators) to configure the discoverability of Amazon EMR clusters from SageMaker Studio Classic notebooks.

• If you are a data scientist or data engineer looking to discover Amazon EMR clusters from your Studio Classic notebooks, see Discover Amazon EMR clusters from SageMaker Studio Classic.

• If you are a data scientist or data engineer looking to connect to existing Amazon EMR clusters from your Studio Classic notebooks, see Connect to an Amazon EMR cluster from SageMaker Studio Classic.

When connecting to your Amazon EMR cluster from SageMaker Studio Classic, you can authenticate to your cluster with Kerberos, Lightweight Directory Access Protocol (LDAP), or use runtime IAM role authentication. Your authentication method depends on your cluster configuration. You can refer to this example Access Apache Livy using a Network Load Balancer on a Kerberos-enabled Amazon EMR cluster to set up an Amazon EMR cluster that uses Kerberos. Alternatively, you can look at the CloudFormation example templates using Kerberos or LDAP in the aws-samples/sagemaker-studio-emr GitHub repository.

Find the list of available connection commands to an Amazon EMR cluster per authentication method in Enter the connection command to an Amazon EMR cluster manually to connect to your Amazon EMR cluster.

**Supported images and kernels to connect to an Amazon EMR cluster from SageMaker Studio Classic**

SageMaker Studio Classic provides built-in support to connect to Amazon EMR clusters in the following images and kernels:

- DataScience – Python 3 kernel
- DataScience 2.0 – Python 3 kernel
- DataScience 3.0 – Python 3 kernel
- SparkAnalytics 1.0 – SparkMagic and PySpark kernels
- SparkAnalytics 2.0 – SparkMagic and PySpark kernels
- SparkMagic – SparkMagic and PySpark kernels
- PyTorch 1.8 – Python 3 kernels
- TensorFlow 2.6 – Python 3 kernel
- TensorFlow 2.11 – Python 3 kernel
Those images and kernels come with `sagemaker-studio-analytics-extension`, a notebook extension that enables connection to a remote Spark (Amazon EMR) cluster via the `SparkMagic` library using Apache Livy.

To connect to Amazon EMR clusters using another built-in image or your own image, follow the instructions in Bring your own image.

**Bring your own image**

To bring your own image in SageMaker Studio Classic and allow your notebooks to connect to Amazon EMR clusters, install the following `sagemaker-studio-analytics-extension` extension to your kernel. It supports connecting SageMaker Studio Classic notebooks to Spark(Amazon EMR) clusters through the `SparkMagic` library.

```
pip install sparkmagic
pip install sagemaker-studio-sparkmagic-lib
pip install sagemaker-studio-analytics-extension
```

Additionally, to connect to Amazon EMR with Kerberos authentication, you must install the kinit client. Depending on your OS, the command to install the kinit client can vary. To bring an Ubuntu (Debian based) image, use the `apt-get install -y -qq krb5-user` command.

For more information on bringing your own image in SageMaker Studio Classic, see Bring your own SageMaker image.

**Configure the discoverability of Amazon EMR clusters (for administrators)**

This section provides details about how administrators can configure the discoverability of existing Amazon EMR clusters from SageMaker Studio Classic. The clusters can be deployed in the same AWS account as Studio Classic (Single Account tab) or in separate accounts (Cross Accounts tab).

**Single Account**

Attach the following permissions to the SageMaker Studio Classic execution role accessing your cluster.

The following list provides a breakdown of the permissions required.

- **AllowSagemakerProjectManagement** enables the creation of SageMaker projects. In Studio Classic, access to the AWS Service Catalog is granted through Projects.
• AllowClusterDetailsDiscovery and AllowClusterDiscovery allow the discovery and connection to Amazon EMR clusters.

• AllowPresignedUrl allows the creation of pre-signed URLs to access Spark UI.

The following is a comprehensive JSON that includes these permissions.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "AllowPresignedUrl",
         "Effect": "Allow",
         "Action": [
            "elasticmapreduce:DescribeCluster",
            "elasticmapreduce:ListInstanceGroups",
            "elasticmapreduce:CreatePersistentAppUI",
            "elasticmapreduce:DescribePersistentAppUI",
            "elasticmapreduce:GetPersistentAppUIPresignedURL",
            "elasticmapreduce:GetOnClusterAppUIPresignedURL"
         ],
         "Resource": [
            "arn:aws:elasticmapreduce:region:account-id:cluster/*"
         ]
      },
      {
         "Sid": "AllowClusterDetailsDiscovery",
         "Effect": "Allow",
         "Action": [
            "elasticmapreduce:DescribeCluster",
            "elasticmapreduce:ListInstances",
            "elasticmapreduce:ListInstanceGroups",
            "elasticmapreduce:DescribeSecurityConfiguration"
         ],
         "Resource": [
            "arn:aws:elasticmapreduce:region:account-id:cluster/*"
         ]
      },
      {
         "Sid": "AllowClusterDiscovery",
         "Effect": "Allow",
         "Action": [
            "elasticmapreduce:ListClusters"
         ],
         "Resource": [
            "arn:aws:elasticmapreduce:region:account-id:cluster/*"
         ]
      }
   ]
}
```
Cross Accounts

If your Amazon EMR clusters and SageMaker Studio Classic are deployed in separate AWS accounts, you configure the permissions in multiple steps.

- On the trusting account (the account in which Amazon EMR is deployed), create a custom IAM role (referred to as ASSUMABLE-ROLE in this page) with the following permissions and trust relationship.

For information about creating a role on an AWS account, see Creating an IAM role (console).

1. Add a policy defining the following permissions.

   - AllowClusterDetailsDiscovery and AllowClusterDiscovery to allow the discovery and connection to Amazon EMR clusters.
   - AllowPresignedUrl to allow the creation of pre-signed URLs to access Spark UI.

The following is a comprehensive JSON that includes these permissions.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "AllowPresignedUrl",
      "Effect": "Allow",
      "Action": [
        "elasticmapreduce:DescribeCluster",
        "elasticmapreduce:ListInstanceGroups",
        "elasticmapreduce:ListClusters"
      ],
      "Resource": "arn:aws:sagemaker:region:account-id:project/*"
    }
  ]
}
```
2. To grant the trusted account (the account in which SageMaker Studio Classic’s account is deployed) the permission to assume a role in the trusting account, add the following trust relationship.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:elasticmapreduce:emr-region:emr-account:cluster/"
      }
    },
    {
      "Sid": "AllowClusterDetailsDiscovery",
      "Effect": "Allow",
      "Action": [
        "elasticmapreduce:DescribeCluster",
        "elasticmapreduce:ListInstances",
        "elasticmapreduce:ListInstanceGroups",
        "elasticmapreduce:DescribeSecurityConfiguration"
      ],
      "Resource": [
      ]
    },
    {
      "Sid": "AllowClusterDiscovery",
      "Effect": "Allow",
      "Action": [
        "elasticmapreduce:ListClusters"
      ],
      "Resource": "*"
    }
  ]
}
```
On the **trusted account** (the account in which SageMaker Studio Classic is deployed), add the following trust relationship to the Studio Classic execution role.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "AllowRoleAssumptionForCrossAccountDiscovery",
      "Effect": "Allow",
      "Action": "sts:AssumeRole",
      "Resource": ["arn:aws:iam::emr-account:role/ASSUMABLE-ROLE"]
    }
  ]
}
```

Last, see [Additional Configuration for cross accounts use cases (for administrators)](https://docs.aws.amazon.com/sagemaker/latest/dg/configure-aws-account-for-cross-account-discovery.html) to learn how to provide the ARN of the **ASSUMABLE-ROLE** to the Studio Classic execution role. The ARN is loaded by the Studio Classic Jupyter server at launch. The Studio Classic execution role assumes that cross-account role to discover and connect to Amazon EMR clusters in the **trustling account**.

Visit [Discover Amazon EMR clusters from SageMaker Studio Classic](https://docs.aws.amazon.com/sagemaker/latest/dg/discover-aws-account-emr-clusters.html) to learn about how to discover and connect to Amazon EMR clusters from Studio Classic notebooks.

**Discover Amazon EMR clusters from SageMaker Studio Classic**

Data scientists and data engineers can discover, connect to, and manage Amazon EMR clusters from Amazon SageMaker Studio Classic. The Amazon EMR clusters may be in the same AWS account as Amazon SageMaker Studio Classic or in a different AWS account.

If your administrator configured the cross accounts discovery of Amazon EMR clusters, you can see a consolidated list of clusters in the AWS account used by SageMaker Studio Classic as well as in the remote accounts.
If you are an administrator looking to set up the discoverability of Amazon EMR clusters from SageMaker Studio Classic, see Configure the discoverability of Amazon EMR clusters (for administrators).

To view the list of available Amazon EMR clusters from SageMaker Studio Classic:

1. Select the **Home** icon in Studio Classic UI's left-side panel, then select the **Data** node in the navigation menu.

2. Navigate down to the **Clusters** node. This opens up a page listing the Amazon EMR clusters that you can access from SageMaker Studio Classic.

   The list displays the status of each cluster. A cluster status can be **Starting**, **Bootstrapping**, **Running/Walking**, **Terminating**, **Terminated**, and **Terminated with error**. You can filter clusters by status by selecting the filter icon. The following image shows an example of a list of clusters.

   ![Cluster List Example](image)

3. To connect to a particular **Running/Walking** cluster, see Connect to an Amazon EMR cluster from SageMaker Studio Classic.

**Connect to an Amazon EMR cluster from SageMaker Studio Classic**

This section explains how you can connect to an Amazon EMR cluster from a Studio Classic notebook when you use any of the supported kernels.
Connect to an Amazon EMR cluster automatically

To connect to your cluster using the Studio Classic UI, you can either initiate a connection from the list of clusters accessed in Discover Amazon EMR clusters from SageMaker Studio Classic, or from a notebook in SageMaker Studio Classic.

To connect to a particular cluster from your list of clusters

1. Choose the name of the cluster in your list. This activates the Attach to new notebook button.
2. Choose Attach to new notebook. This opens up the images and kernels selection box.
3. Select your image and kernel, then choose Select. For a list of supported images, see Supported images and kernels to connect to an Amazon EMR cluster from SageMaker Studio Classic or refer to Bring your own image.
4. If the cluster you select does not use Kerberos, LDAP, or runtime role authentication, Studio Classic prompts you to select the credential type. Choose from Http basic authentication or No credentials, then enter your credentials, if applicable. A connection command populates the first cell of your notebook and initiates the connection with the Amazon EMR cluster.

Once the connection succeeds, a message confirms the connection and the start of the Spark application.

Alternatively, you can connect to a cluster from a notebook.

1. Choose Cluster at the top of your notebook.

Cluster is only visible when you use a kernel from Supported images and kernels to connect to an Amazon EMR cluster from SageMaker Studio Classic or from Bring your own image. If you cannot see Cluster at the top of your notebook, ensure that your administrator has configured the discoverability of your clusters and switch to a supported kernel.

This opens up a list of available clusters.

2. Select the cluster to which you want to connect, then choose Connect.
3. If you configured your Amazon EMR clusters to support runtime IAM roles and your administrator preloaded your roles in an execution role configuration JSON, you can select
your Amazon EMR access role from the **Amazon EMR execution role** drop down menu. If your roles are not preloaded, Studio Classic uses your Studio Classic execution role by default. For information about using runtime roles with Amazon EMR, see [Connect to an Amazon EMR cluster from Studio Classic using runtime IAM roles](#). When you connect to a cluster, Studio Classic adds a code block to an active cell to establish the connection.

Otherwise, if the cluster you choose does not use Kerberos, LDAP, or runtime role authentication, Studio Classic prompts you to select the credential type. You can choose **HTTP basic authentication** or **No credential**.

4. An active cell populates and runs. This cell contains the connection command to connect to your Amazon EMR cluster.

   Once the connection succeeds, a message confirm the connection and the start of the Spark application.

**Enter the connection command to an Amazon EMR cluster manually**

You can manually connect to your Amazon EMR cluster from a Studio Classic notebook whether or not your Studio Classic application and cluster reside in the same AWS account.

For each of the following authentication types, use the specified command to manually connect to your cluster from your Studio Classic notebook.

- **Kerberos**

  Append the `--assumable-role-arn` argument if you need cross-account Amazon EMR access. Append the `--verify-certificate` argument if you connect to your cluster with HTTPS.

  ```
  %load_ext sagemaker_studio_analytics_extension.magics
  %sm_analytics emr connect --cluster-id cluster_id \  
  --auth-type Kerberos --language python 
  [+--assumable-role-arn EMR_access_role_ARN ] 
  [+--verify-certificate /home/user/certificateKey.pem]
  ```

- **LDAP**

  Append the `--assumable-role-arn` argument if you need cross-account Amazon EMR access. Append the `--verify-certificate` argument if you connect to your cluster with HTTPS.

  ```
  %load_ext sagemaker_studio_analytics_extension.magics
  ```

---

Prepare data using Amazon EMR
%sm_analytics emr connect --cluster-id cluster_id \  
--auth-type Basic_Access --language python \  
[--assumable-role-arn EMR_access_role_ARN ] \  
[--verify-certificate /home/user/certificateKey.pem]

• **NoAuth**

Append the --assumable-role-arn argument if you need cross-account Amazon EMR access. Append the --verify-certificate argument if you connect to your cluster with HTTPS.

%load_ext sagemaker_studio_analytics_extension.magics
%sm_analytics emr connect --cluster-id cluster_id \  
--auth-type None --language python \  
[--assumable-role-arn EMR_access_role_ARN ] \  
[--verify-certificate /home/user/certificateKey.pem]

• **Runtime IAM roles**

Append the --assumable-role-arn argument if you need cross-account Amazon EMR access. Append the --verify-certificate argument if you connect to your cluster with HTTPS.

For more information on connecting to an Amazon EMR cluster using runtime IAM roles, see [Connect to an Amazon EMR cluster from Studio Classic using runtime IAM roles](#).

%load_ext sagemaker_studio_analytics_extension.magics
%sm_analytics emr connect --cluster-id cluster_id \  
--auth-type Basic_Access \  
--emr-execution-role-arn arn:aws:iam::studio_account_id:role/emr-execution-role-name \  
[--assumable-role-arn EMR_access_role_ARN] \  
[--verify-certificate /home/user/certificateKey.pem]

**Connect to an Amazon EMR cluster over HTTPS**

If you have configured your Amazon EMR cluster with transit encryption enabled and Apache Livy server for HTTPS and would like Studio Classic to communicate with Amazon EMR using HTTPS, you need to configure Studio Classic to access your certificate key.

For self-signed or local Certificate Authority (CA) signed certificates, you can do this in two steps:

1. Download the PEM file of your certificate to your local file system using one of the following options:
• Jupyter's built-in file upload function.
• A notebook cell.
• A lifecycle configuration (LCC) script.

For information on how to use an LCC script, see Customize a Notebook Instance Using a Lifecycle Configuration Script.

2. Enable the validation of the certificate by providing the path to your certificate in the --verify-certificate argument of your connection command.

```
%sm_analytics emr connect --cluster-id cluster_id \
--verify-certificate /home/user/certificateKey.pem ...
```

For public CA issued certificates, set the certificate validation by setting the --verify-certificate parameter as true.

Alternatively, you can disable the certificate validation by setting the --verify-certificate parameter as false.

You can find the list of available connection commands to an Amazon EMR cluster in Enter the connection command to an Amazon EMR cluster manually.

## Connect to an Amazon EMR cluster from Studio Classic using runtime IAM roles

When you connect to an Amazon EMR cluster from your Amazon SageMaker Studio Classic notebook, you can visually browse a list of IAM roles, known as runtime roles, and select one on the fly. Subsequently, all your Apache Spark, Apache Hive, or Presto jobs created from your Studio Classic notebook access only the data and resources permitted by policies attached to the runtime role. Also, when data is accessed from data lakes managed with AWS Lake Formation, you can enforce table-level and column-level access using policies attached to the runtime role.

With this capability, you and your teammates can connect to the same cluster, each using a runtime role scoped with permissions matching your individual level of access to data. Your sessions are also isolated from one another on the shared cluster. With this ability to control fine-grained access to data on the same shared cluster, you can simplify provisioning of Amazon EMR clusters, reducing operational overhead and saving costs.

To try out this new feature, see Apply fine-grained data access controls with AWS Lake Formation and Amazon EMR from Amazon SageMaker Studio Classic. This blog post helps you set up a demo
environment where you can try using preconfigured runtime roles to connect to Amazon EMR clusters.

**Prerequisites**

Before you get started, make sure you meet the following prerequisites:

- Use Amazon EMR version 6.9 or above.
- Use JupyterLab version 3 in the Studio Classic Jupyter server application configuration. This version supports Studio Classic connection to Amazon EMR clusters using runtime roles.
- Allow the use of runtime roles in your cluster's security configuration. For more information, see [Runtime roles for Amazon EMR steps](#).
- Create a notebook with any of the kernels listed in [Use Amazon EMR clusters from Studio Classic notebooks](#).
- Make sure you review the instructions in [Set up Studio Classic to use runtime IAM roles](#) to configure runtime roles with Studio Classic.

**Cross-account connection scenarios**

Runtime role authentication supports a variety of cross-account connection scenarios when your data resides outside of your Studio Classic account. The following image shows three different ways you can assign your Amazon EMR cluster, data, and even Amazon EMR execution role between your Studio Classic and data accounts:
In option 1, your Amazon EMR cluster and Amazon EMR execution role are in a separate data account from your Studio Classic account. You define a separate Amazon EMR access role permission policy which grants permission to your Studio Classic execution role to assume the Amazon EMR access role. The Amazon EMR access role then calls the Amazon EMR API GetClusterSessionCredentials on behalf of your Studio Classic execution role, giving you access to the cluster.

In option 2, your Amazon EMR cluster and Amazon EMR execution role are in your Studio Classic account. Your Studio Classic execution role has permission to use the Amazon EMR API GetClusterSessionCredentials to gain access to your cluster. To access the Amazon S3
bucket, give the Amazon EMR execution role cross-account Amazon S3 bucket access permissions — you grant these permissions within your Amazon S3 bucket policy.

In option 3, your Amazon EMR clusters are in your Studio Classic account, and the Amazon EMR execution role is in the data account. Your Studio Classic execution role has permission to use the Amazon EMR API `GetClusterSessionCredentials` to gain access to your cluster. Add the Amazon EMR execution role into the execution role configuration JSON. Then you can select the role in the UI when you choose your cluster. For details about how to set up your execution role configuration JSON file, see Preload your execution roles into Studio Classic.

**Set up Studio Classic to use runtime IAM roles**

To establish runtime role authentication for your Amazon EMR clusters, configure the required IAM policies, network, and usability enhancements. Your setup depends on whether you handle any cross-account arrangements if your Amazon EMR clusters, Amazon EMR execution role, or both, reside outside of your Amazon SageMaker Studio Classic account. The following discussion guides you through the policies to install, how to configure the network to allow traffic between cross-accounts, and the local configuration file to set up to automate your Amazon EMR connection.

**Configure runtime role authentication when your Amazon EMR cluster and Studio Classic are in the same account**

If your Amazon EMR cluster resides in your Studio Classic account, add the basic policy to connect to your Amazon EMR cluster and set permissions to call the Amazon EMR API `GetClusterSessionCredentials`, which gives you access to the cluster. Complete the following steps to add necessary permissions to your Studio Classic execution policy:

1. Add the required IAM policy to connect to Amazon EMR clusters. For details, see Discover Amazon EMR clusters from SageMaker Studio Classic.
2. Grant permission to call the Amazon EMR API `GetClusterSessionCredentials` when you pass one or more permitted Amazon EMR execution roles specified in the policy.
3. (Optional) Grant permission to pass IAM roles that follow any user-defined naming conventions.
4. (Optional) Grant permission to access Amazon EMR clusters that are tagged with specific user-defined strings.
5. If you don’t want to manually call the Amazon EMR connection command, install a SageMaker configuration file in your local Amazon EFS and select the role to use when you select your Amazon EMR cluster. For details about how to preload your IAM roles, see Preload your execution roles into Studio Classic.
The following example policy permits Amazon EMR execution roles belonging to the modeling and training groups to call GetClusterSessionCredentials. In addition, the policyholder can access Amazon EMR clusters tagged with the strings modeling or training.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "VisualEditor0",
         "Effect": "Allow",
         "Action": "elasticmapreduce:GetClusterSessionCredentials",
         "Resource": "*",
         "Condition": {
            "StringLike": {
               "elasticmapreduce:ExecutionRoleArn": [
                  "arn:aws:iam::123456780910:role/emr-execution-role-ml-modeling*",
                  "arn:aws:iam::123456780910:role/emr-execution-role-ml-training*"
               ],
               "elasticmapreduce:ResourceTag/group": [
                  "*modeling*",
                  "*training*"
               ]
            }
         }
      }
   ]
}
```

**Configure runtime role authentication when your cluster and Studio Classic are in different accounts**

If your Amazon EMR cluster is not in your Studio Classic account, allow your Studio Classic execution role to assume the cross-account Amazon EMR access role so you can connect to the cluster. Complete the following steps to set up your cross-account configuration:

1. Create your Studio Classic execution role permission policy so that the execution role can assume the Amazon EMR access role. The following policy is an example:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "VisualEditor0",
         "Effect": "Allow",
         "Action": "elasticmapreduce:GetClusterSessionCredentials",
         "Resource": "*",
         "Condition": {
            "StringLike": {
               "elasticmapreduce:ExecutionRoleArn": [
                  "arn:aws:iam::123456780910:role/emr-execution-role-ml-modeling*",
                  "arn:aws:iam::123456780910:role/emr-execution-role-ml-training*"
               ],
               "elasticmapreduce:ResourceTag/group": [
                  "*modeling*",
                  "*training*"
               ]
            }
         }
      }
   ]
}
```
2. Create the trust policy to specify which Studio Classic account IDs are trusted to assume the Amazon EMR access role. The following policy is an example:

```json
{
  "Version": "2012-10-17",
  "Statement": [  
    {
      "Sid": "AllowCrossAccountSageMakerExecutionRoleToAssumeThisRole",
      "Effect": "Allow",
      "Principal": {
        "AWS": "arn:aws:iam::studio_account_id:role/studio_execution_role"
      },
      "Action": "sts:AssumeRole"
    }
  ]
}
```

3. Create the Amazon EMR access role permission policy, which grants the Amazon EMR execution role the needed permissions to carry out the intended tasks on the cluster. Configure the Amazon EMR access role to call the API `GetClusterSessionCredentials` with the Amazon EMR execution roles specified in the access role permission policy. The following policy is an example:

```json
{
  "Version": "2012-10-17",
  "Statement": [  
    {
      "Sid": "AllowCallingEmrGetClusterSessionCredentialsAPI",
      "Effect": "Allow",
      "Action": "elasticmapreduce:GetClusterSessionCredentials",
      "Resource": "",
      "Condition": {
        "StringLike": {
          "Statement": [  
```
4. Set up the cross-account network so that traffic can move back and forth between your accounts. For guided instruction, see Set up the network in the blog post Create and manage Amazon EMR Clusters from SageMaker Studio Classic to run interactive Spark and ML workloads – Part 2. The steps in the blog post help you complete the following tasks:

   a. VPC-peer your Studio Classic account and your Amazon EMR account to establish a connection.

   b. Manually add routes to the private subnet route tables in both accounts. This permits creation and connection of Amazon EMR clusters from the Studio Classic account to the remote account’s private subnet.

   c. Set up the security group attached to your Studio Classic domain to allow outbound traffic and the security group of the Amazon EMR primary node to allow inbound TCP traffic from the Studio Classic instance security group.

5. If you don’t want to manually call the Amazon EMR connection command, install a SageMaker configuration file in your local Amazon EFS so you can select the role to use when you choose your Amazon EMR cluster. For details about how to preload your IAM roles, see Preload your execution roles into Studio Classic.

Configure Lake Formation access

When you access data from data lakes managed by AWS Lake Formation, you can enforce table-level and column-level access using policies attached to your runtime role. To configure permission for Lake Formation access, see Integrate Amazon EMR with AWS Lake Formation.

Preload your execution roles into Studio Classic

If you don't want to manually call the Amazon EMR connection command, you can install a SageMaker configuration file in your local Amazon EFS so you can select the execution role to use when you choose your Amazon EMR cluster.
To write a configuration file for the Amazon EMR execution roles, associate a Use lifecycle configurations with Amazon SageMaker Studio Classic (LCC) to the Jupyter server application. Alternatively, you can write or update the configuration file and restart the Jupyter server with the command: `restart-jupyter-server`.

The following snippet is an example LCC bash script you can apply if your Studio Classic application and cluster are in the same account:

```bash
#!/bin/bash
set -eux

FILE_DIRECTORY="/home/sagemaker-user/.sagemaker-analytics-configuration-DO_NOT_DELETE"
FILE_NAME="emr-configurations-DO_NOT_DELETE.json"
FILE="$FILE_DIRECTORY/$FILE_NAME"

mkdir -p $FILE_DIRECTORY

cat << 'EOF' > "$FILE"
{
   "emr-execution-role-arns":
   {
      "123456789012": [
         "arn:aws:iam::123456789012:role/emr-execution-role-1",
         "arn:aws:iam::123456789012:role/emr-execution-role-2"
      ]
   }
}
EOF
```

If your Studio Classic application and clusters are in different accounts, specify the Amazon EMR access roles that can use the cluster. In the following example policy, 123456789012 is the ARN for the Amazon EMR cluster account, and 212121212121 and 434343434343 are the ARNs for the permitted Amazon EMR access roles.

```bash
#!/bin/bash
set -eux

FILE_DIRECTORY="/home/sagemaker-user/.sagemaker-analytics-configuration-DO_NOT_DELETE"
FILE_NAME="emr-configurations-DO_NOT_DELETE.json"
FILE="$FILE_DIRECTORY/$FILE_NAME"

```
**Terminate an Amazon EMR cluster from Studio Classic**

The following procedure shows how to terminate an Amazon EMR cluster from a Studio Classic notebook.

**To terminate a cluster in a Running state, navigate to the list of available Amazon EMR clusters.**

1. In SageMaker Studio Classic, select the **Home** icon in Studio Classic UI's left-side panel, then select the **Data** node in the navigation menu.

2. Navigate down to the **Clusters** node. This opens up a page listing the Amazon EMR clusters that you can access from SageMaker Studio Classic.
3. Select the name of the cluster that you want to terminate, then choose **Terminate**.

4. This opens up a confirmation window informing you that any pending work or data on your cluster will be lost permanently after termination. Confirm by choosing **Terminate** again.

**Access Spark UI from Studio Classic**

The following sections give instructions for accessing the Spark UI from SageMaker Studio Classic notebooks. The Spark UI allows you to monitor and debug your Spark Jobs submitted to run on Amazon EMR from Studio Classic notebooks. SSH tunneling and presigned URLs are two ways for accessing the Spark UI.

**Set up SSH tunneling for Spark UI access**

To set up SSH tunneling to access the Spark UI, follow one of the two options in this section.

Options for setting up SSH tunneling:

- **Option 1: Set up an SSH tunnel to the master node using local port forwarding**
- **Option 2, part 1: Set up an SSH tunnel to the master node using dynamic port forwarding**
  - **Option 2, part 2: Configure proxy settings to view websites hosted on the master node**

For information about viewing web interfaces hosted on Amazon EMR clusters, see [View web interfaces hosted on Amazon EMR Clusters](##). You can also visit your Amazon EMR console to get access to the Spark UI.

**Note**

You can set up an SSH tunnel even if presigned URLs are not available to you.

**Presigned URLs**

To create one-click URLs that can access Spark UI on Amazon EMR from SageMaker Studio Classic notebooks, you must enable the following IAM permissions. Choose the option that applies to you:

- **For Amazon EMR clusters that are in the same account as the SageMaker Studio Classic notebook**: Add the following permissions to the SageMaker Studio Classic IAM execution role.
• For Amazon EMR clusters that are in a different account (not SageMaker Studio Classic notebook): Add the following permissions to the cross-account role that you created for Discover Amazon EMR clusters from SageMaker Studio Classic.

**Note**

You can access presigned URLs from the console in the following regions:

- US East (N. Virginia) Region
- US West (N. California) Region
- Canada (Central) Region
- Europe (Frankfurt) Region
- Europe (Stockholm) Region
- Europe (Ireland) Region
- Europe (London) Region
- Europe (Paris) Region
- Asia Pacific (Tokyo) Region
- Asia Pacific (Seoul) Region
- Asia Pacific (Sydney) Region
- Asia Pacific (Mumbai) Region
- Asia Pacific (Singapore) Region
- South America (São Paulo)

The following policy gives access to presigned URLs for your execution role.

```json
{
    "Sid": "AllowPresignedUrl",
    "Effect": "Allow",
    "Action": [
        "elasticmapreduce:DescribeCluster",
        "elasticmapreduce:ListInstanceGroups",
        "elasticmapreduce:CreatePersistentAppUI",
        "elasticmapreduce:DescribePersistentAppUI",
        "elasticmapreduce:GetPersistentAppUIPresignedURL",
        "elasticmapreduce:GetOnClusterAppUIPresignedURL"
    ]
}
```
Walkthroughs and whitepapers

The following blogs use a case study of sentiment prediction for a movie review to illustrate the process of executing a complete machine learning workflow. This includes data preparation, monitoring Spark jobs, and training and deploying a ML model to get predictions directly from your Studio Classic notebook.

- Create and manage Amazon EMR clusters from SageMaker Studio Classic to run interactive Spark and ML workloads.

- To extend the use case to a cross-account configuration where SageMaker Studio Classic and your Amazon EMR cluster are deployed in separate AWS accounts, see Create and manage Amazon EMR clusters from SageMaker Studio Classic to run interactive Spark and ML workloads - Part 2.

See also:

- A walkthrough of the configuration of Access Apache Livy using a Network Load Balancer on a Kerberos-enabled Amazon EMR cluster.

Additional Configuration for cross accounts use cases (for administrators)

To enable cluster discovery across accounts, administrators need to provide the ARN of a cross-account IAM role to the execution role of SageMaker Studio Classic. SageMaker Studio Classic's execution role assumes that remote role to discover and connect to Amazon EMR clusters in the trusting account. The ARN of this role is loaded by the Studio Classic's Jupyter server at launch.

You can specify this information in two ways.

- Write this remote role in a file named emr-discovery-iam-role-arns-DO_NOT_DELETE.json placed in the directory .cross-account-configuration-
DO_NOT_DELETE in your home directory located in the Amazon EFS storage volume used by SageMaker Studio Classic.

- Alternatively, you can automate this process by using Lifecycle Configuration (LCC) scripts. You can attach the LCC to your domain or a specific user profile. The LCC script that you use must be a JupyterServer configuration. For more information on how to create an LCC script, see Use Lifecycle Configurations with Studio Classic.

The following is an example LCC script. To modify the script, replace ASSUMABLE-ROLE and emr-account with your role name and remote account ID, respectively. The number of cross accounts is limited to five.

```bash
#!/bin/bash
set -eux

FILE_DIRECTORY="/home/sagemaker-user/.cross-account-configuration-DO_NOT_DELETE"
FILE_NAME="emr-discovery-iam-role-arns-DO_NOT_DELETE.json"
FILE="${FILE_DIRECTORY}/${FILE_NAME}"

mkdir -p $FILE_DIRECTORY

cat > "$FILE" <<- "EOF"
{
  emr-cross-account1: "arn:aws:iam::emr-cross-account1:role/ASSUMABLE-ROLE",
  emr-cross-account2: "arn:aws:iam::emr-cross-account2:role/ASSUMABLE-ROLE"
}
EOF

After the LCC runs and the files are written, the server reads the file /home/sagemaker-user/.cross-account-configuration-DO_NOT_DELETE/emr-discovery-iam-role-arns-DO_NOT_DELETE.json and stores the cross-account ARN.

Troubleshooting

The following are common errors that might occur while connecting or using Amazon EMR clusters from Studio Classic notebooks.
### Troubleshoot Livy connections hanging or failing

The following are Livy connectivity issues that might occur while using Amazon EMR clusters from Studio Classic notebooks.

- **Your Amazon EMR cluster encountered an out-of-memory error.**

  A possible reason for a Livy connection via `sparkmagic` hanging or failing is if your Amazon EMR cluster encountered an out-of-memory error.

  By default, the Java configuration parameter of the Apache Spark driver, `spark.driver.defaultJavaOptions`, is set to `-XX:OnOutOfMemoryError='kill -9 %p'`. This means that the default action taken when the driver program encounters an `OutOfMemoryError` is to terminate the driver program by sending a SIGKILL signal. When the Apache Spark driver is terminated, any Livy connection via `sparkmagic` that depends on that driver hangs or fails. This is because the Spark driver is responsible for managing the Spark application's resources, including task scheduling and execution. Without the driver, the Spark application cannot function, and any attempts to interact with it fails.

  If you suspect that your Spark cluster is experiencing memory issues, you can check Amazon EMR logs. Containers killed due to out-of-memory errors typically exit with a code of 137. In such cases, you need to restart the Spark application and establish a new Livy connection to resume interaction with the Spark cluster.

  You can refer to the knowledge base article [How do I resolve the error "Container killed by YARN for exceeding memory limits" in Spark on Amazon EMR?](https://aws.amazon.com/premiersupport/knowledgecenter/ask.html?id=KS1CA3476C6986AC2378234187319A0998E037D589F8CBB8A8A81&language=en) on AWS re:Post to learn about various strategies and parameters that can be used to address an out-of-memory issue.

  We recommend reviewing the [Amazon EMR Best Practices Guides](https://aws.amazon.com/sagemaker/en_us/best-practices/) for best practices and tuning guidance on running Apache Spark workloads on your Amazon EMR clusters.

- **Your Livy session times out when connecting to an Amazon EMR cluster for the first time.**

  When you initially connect to an Amazon EMR cluster using `sagemaker-studio-analytics-extension`, which enables connection to a remote Spark (Amazon EMR) cluster via the `SparkMagic` library using `Apache Livy`, you may encounter a connection timeout error:

  An error was encountered: Session 0 did not start up in 60 seconds.

  If your Amazon EMR cluster requires the initialization of a Spark application upon establishing a connection, there is an increased chance of seeing connection timeout errors.
To reduce the chances of getting timeouts when connecting to an Amazon EMR cluster using Livy through the analytics extension, sagemaker-studio-analytics-extension version 0.0.19 and later override the default server session timeout to 120 seconds instead of sparkmagic's default of 60 seconds.

We recommend upgrading your extension 0.0.18 and sooner by running the following upgrade command.

```
pip install --upgrade sagemaker-studio-analytics-extension
```

Note that when providing a custom timeout configuration in sparkmagic, sagemaker-studio-analytics-extension honors this override. However, setting the session timeout to 60 seconds automatically triggers the default server session timeout of 120 seconds in sagemaker-studio-analytics-extension.

Prepare data using AWS Glue Interactive Sessions

AWS Glue Interactive Sessions is an on-demand, serverless, Apache Spark runtime environment that data scientists and engineers can use to rapidly build, test, and run data preparation and analytics applications.

You can initiate an AWS Glue interactive session by starting a SageMaker Studio Classic notebook. When creating your Studio Classic notebook, choose the built-in Glue PySpark or Glue Spark kernel. This automatically starts an interactive, serverless Spark session. You do not need to provision or manage any compute cluster or infrastructure. After initialization, you can explore the AWS Glue Data Catalog, execute complex queries, and interactively analyze and prepare data using Spark within your Studio Classic notebook. You can then use the prepared data to build, train, tune, and deploy models using the purpose-built ML tools within SageMaker Studio Classic.

Before starting your AWS Glue interactive session in SageMaker Studio Classic, you need to set the appropriate roles and policies. Additionally, you may need to provide access to additional resources, such as a storage Amazon S3 bucket, which might require additional policies. For more information about required and additional IAM policies, see Permissions for AWS Glue Interactive Sessions in SageMaker Studio Classic.

SageMaker Studio Classic provides a default configuration for your AWS Glue interactive session, however, you can use AWS Glue's full catalog of Jupyter magic commands to further customize
your environment. For information about the default and additional Jupyter magics that you can use in your AWS Glue interactive session, see Configure your AWS Glue interactive session in SageMaker Studio Classic.

The supported images and kernels for connecting to a AWS Glue interactive session are as follows:

- **Images**: SparkAnalytics 1.0, SparkAnalytics 2.0
- **Kernel**: Glue Python [PySpark and Ray] and Glue Spark

**Prerequisites:**

The SparkAnalytics image that you select to launch your AWS Glue session in Studio Classic is a combination of two frameworks - the SparkMagic framework (used with Amazon EMR), and AWS Glue. For this reason, the prerequisites for both frameworks apply. However, you do not have to set up the Amazon EMR cluster if you only plan to use AWS Glue Interactive Sessions. Before you start your first AWS Glue interactive session in Studio Classic, complete the following:

- Complete the prerequisites required to use the SparkMagic image. For a list of the prerequisites, see the Prerequisites section in Prepare Data at Scale with Studio Classic Notebooks.
- Create an execution role with permissions for both AWS Glue and SageMaker Studio Classic. Add the managed policy AwsGlueSessionUserRestrictedServiceRole, and create a custom policy that includes permissions sts:GetCallerIdentity, iam:GetRole, and IAM:Passrole. For instructions about how to create the necessary permissions, see Permissions for AWS Glue Interactive Sessions in SageMaker Studio Classic.
- Create a SageMaker domain with the execution role you created. For instructions about how to create a domain, see Custom onboarding using IAM.

**Get Started with AWS Glue Interactive Sessions**

In this guide, you learn how to initiate an AWS Glue interactive session in SageMaker Studio Classic, and manage your environment with Jupyter magics.

**Permissions for AWS Glue Interactive Sessions in SageMaker Studio Classic**

This section lists the required policies to run AWS Glue interactive sessions in Studio Classic and explains how to set them up. In particular, it details how to:
• Attach the AwsGlueSessionUserRestrictedServiceRole managed policy to your SageMaker execution role.
• Create an inline custom policy on your SageMaker execution role.
• Modify the trust relationship of your SageMaker execution role.

To attach the AwsGlueSessionUserRestrictedServiceRole managed policy to your execution role

1. Open the IAM console.
2. Select Roles in the left-side panel.
3. Find your Studio Classic execution role. Choose the role name to access the role summary page.
4. Under the Permissions tab, select Attach policies from the Add Permissions dropdown menu.
5. Select the checkbox next to the managed policy AwsGlueSessionUser RestrictedServiceRole.
6. Choose Attach policies.

The summary page shows your newly-added managed policies.

To create the inline custom policy on your execution role

1. Select Create inline policy in the Add Permissions dropdown menu.
2. Select the JSON tab.
3. Copy and paste in the following policy.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "unique_statement_id",
            "Effect": "Allow",
            "Action": [
                "iam:GetRole",
                "iam:PassRole",
                "sts:GetCallerIdentity"
            ]
        }
    ]
}
```
4. Choose **Review policy**.

5. Enter a **Name** and choose **Create policy**.

   The summary page shows your newly-added custom policy.

---

**To modify the trust relationship of your execution role**

1. Select the **Trust relationships** tab.

2. Choose **Edit trust policy**.

3. Copy and paste in the following policy.

   ```json
   {
   "Version": "2012-10-17",
   "Statement": [
   {
   "Effect": "Allow",
   "Principal": {
   "Service": [
   "glue.amazonaws.com",
   "sagemaker.amazonaws.com"
   ],
   "Action": "sts:AssumeRole"
   }
   ]
   }
   }
   ``

4. Choose **Update policy**.

You can add additional roles and policies if you need to access other AWS resources. For a description of the additional roles and policies you can include, see **Interactive sessions with IAM** in the AWS Glue documentation.
Tag propagation

Tags are commonly used to track and allocate costs, control access to your session, isolate your resources, and more. To learn about adding metadata to your AWS resources using tagging, or for details on common use cases, see Additional information.

You can enable the automatic propagation of AWS tags to new AWS Glue interactive sessions created from within the Studio Classic UI. When an AWS Glue interactive session is created from SageMaker Studio Classic, any user-defined tags attached to the user profile or shared space are carried over to the new AWS Glue interactive session. Additionally, SageMaker Studio Classic automatically adds two AWS-generated internal tags ((sagemaker:user-profile-arn and sagemaker:domain-arn) or (sagemaker:shared-space-arn and sagemaker:domain-arn)) to new AWS Glue interactive sessions created from the Studio Classic UI. You can use these tags to aggregate costs across individual domains, user profiles, or spaces.

Enable tag propagation

To enable the automatic propagation of tags to new AWS Glue interactive sessions, set the following permissions for your SageMaker execution role and the IAM role associated with your AWS Glue session:

On your SageMaker execution role:

- Create a new inline policy, and paste the following JSON file. The policy grants the execution role permission to describe (DescribeUserProfile, DescribeSpace, DescribeDomain) and list the tags (ListTag) set on the user profiles, shared spaces, and SageMaker domain.

```json
{
    "Effect": "Allow",
    "Action": [
        "sagemaker:ListTags"
    ]
}
```

Tip

By default, the role associated with the AWS Glue interactive session is the same as the SageMaker execution role. You can specify a different execution role for the AWS Glue interactive session by using the %iam_role magic command. For information on the available Jupyter magic commands to configure AWS Glue interactive sessions, see Configure your AWS Glue interactive session in SageMaker Studio Classic.
On the IAM role of your AWS Glue session: Create a new inline policy, and paste the following JSON file. The policy grants your role permission to attach tags (TagResource) to your session, or retrieve its list of tags (GetTags).

```json
{
    "Effect": "Allow",
    "Action": [
        "glue:TagResource",
        "glue:GetTags"
    ],
    "Resource": ["arn:aws:sagemaker:*:*:user-profile/*",
                  "arn:aws:sagemaker:*:*:space/*"]
}
```
Note

- Failures occurring while applying those permissions do not prevent the creation of AWS Glue interactive sessions. You can find details about the reason of the failure in SageMaker Studio Classic CloudWatch logs.
- You must restart the kernel of your AWS Glue interactive session to propagate the update of a tag’s value.

It is important to note the following points:

- Once a tag is attached to a session, it cannot be removed by propagation.

You can remove tags from an AWS Glue interactive session directly through the AWS CLI, the AWS Glue API, or the https://console.aws.amazon.com/sagemaker/. For example, using the AWS CLI, you can remove a tag by providing the session's ARN and the tag keys you want to remove as follows:

```bash
aws glue untag-resource \
  --resource-arn arn:aws:glue:region:account-id:session:session-name \
  --tags-to-remove tag-key1,tag-key2
```

- SageMaker Studio Classic adds two AWS-generated internal tags ((sagemaker: userProfile-arn and sagemaker:domain-arn) or (sagemaker:shared-space-arn and sagemaker:domain-arn)) to new AWS Glue interactive sessions created from the Studio Classic UI. Those tags count against the limit of 50 tags set on all AWS resources. Both sagemaker:user-profile-arn and sagemaker:shared-space-arn contain the domain ID to which they belong.
- Tags keys starting with aws:, AWS:, or any combination of upper and lowercase letters as a prefix for keys are not propagated and are reserved for AWS use.
Additional information

For more information on tagging, refer to the following resources.

- To learn about adding metadata to your AWS resources with tagging, see Tagging AWS resources.
- For information on tracking costs using tags, see Cost analysis in SageMaker Studio Classic Administration Best Practices.
- For information on controlling access to AWS Glue based on tag keys, see ABAC with AWS Glue.

Launch your AWS Glue interactive session on SageMaker Studio Classic

After you create the roles, policies, and SageMaker domain, you can launch your AWS Glue interactive session in SageMaker Studio Classic.

To launch AWS Glue in SageMaker Studio Classic

1. Create a SageMaker domain. For instructions on how to create a new domain, see Amazon SageMaker domain overview.
3. Select Control Panel in the left-side panel.
4. In the Launch App dropdown menu next to the user name, select Studio.
5. In the Jupyter view, choose File, then New, then Notebook.
6. In the Image dropdown menu, select SparkAnalytics 1.0 or SparkAnalytics 2.0. In the kernel dropdown menu, select Glue Spark or Glue Python [PySpark and Ray]. Choose Select.
7. (optional) Use Jupyter magics to customize your environment. For more information about Jupyter magics, see Configure your AWS Glue interactive session in SageMaker Studio Classic.
8. Start writing your Spark data processing scripts.

Configure your AWS Glue interactive session in SageMaker Studio Classic

Note
All magic configurations are carried over to subsequent sessions for the lifetime of the AWS Glue kernel.
You can use Jupyter magics in your AWS Glue interactive session to modify your session and configuration parameters. Magics are short commands prefixed with % at the start of Jupyter cells that provide a quick and easy way to help you control your environment. In your AWS Glue interactive session, the following magics are set for you by default:

<table>
<thead>
<tr>
<th>Magic</th>
<th>Default value</th>
</tr>
</thead>
<tbody>
<tr>
<td>%glue_version</td>
<td>3.0</td>
</tr>
<tr>
<td>%iam_role</td>
<td>execution role attached to your SageMaker domain</td>
</tr>
<tr>
<td>%region</td>
<td>your region</td>
</tr>
</tbody>
</table>

You can use magics to further customize your environment. For example, if you want to change the number of workers allocated to your job from the default five to 10, you can specify %number_of_workers 10. If you want to configure your session to stop after 10 minutes of idle time instead of the default 2880, you can specify %idle_timeout 10.

All of the Jupyter magics currently available in AWS Glue are also available in SageMaker Studio Classic. For the complete list of AWS Glue magics available, see [Configuring AWS Glue interactive sessions for Jupyter and AWS Glue Studio Classic notebooks](#).

**AWS Glue Interactive Session Pricing**

When you use AWS Glue Interactive Sessions on SageMaker Studio Classic notebooks, you are charged separately for resource usage on AWS Glue and Studio Classic notebooks.

AWS charges for AWS Glue Interactive Sessions based on how long the session is active and the number of Data Processing Units (DPU) used. You are charged an hourly rate for the number of DPUs used to run your workloads, billed in increments of one second. AWS Glue Interactive Sessions assigns a default of five DPUs and requires a minimum of two DPUs. There is also a one-minute minimum billing duration for each interactive session. To see the AWS Glue rates and pricing examples, or to estimate your costs using the AWS Pricing Calculator, see [AWS Glue pricing](#).

Your SageMaker Studio Classic notebook runs on an Amazon EC2 instance and you are charged for the instance type you choose, based on the duration of use. Studio Classic assigns you a default EC2 instance type of ml-t3-medium when you select the SparkAnalytics image and associated...
kernel. You can change the instance type for of your Studio Classic notebook to suit your workload. For information about SageMaker Studio Classic pricing, see Amazon SageMaker Pricing.
Process data

SageMaker Processing refers to SageMaker's capabilities to run data pre and post processing, feature engineering, and model evaluation tasks on SageMaker's fully-managed infrastructure. These tasks are executed as processing jobs. Using SageMaker Processing API, data scientists can run scripts and notebooks to process, transform, and analyze datasets to prepare them for machine learning. When combined with the other critical machine learning tasks provided by SageMaker, such as training and hosting, Processing provides you with the benefits of a fully managed machine learning environment, including all the security and compliance support built into SageMaker. You have the flexibility to use the built-in data processing containers or to bring your own containers for custom processing logic and then submit jobs to run on SageMaker managed infrastructure.

Note

You can create a processing job programmatically by calling the CreateProcessingJob API action in any language supported by SageMaker or by using the AWS CLI. For information on how this API action translates into a function in the language of your choice, see the See Also section of CreateProcessingJob and choose an SDK. As an example, for Python users, refer to the Amazon SageMaker Processing section of SageMaker Python SDK. Alternatively, see the full request syntax of create_processing_job in the AWS SDK for Python (Boto3).

The following diagram shows how Amazon SageMaker spins up a Processing job. Amazon SageMaker takes your script, copies your data from Amazon Simple Storage Service (Amazon S3), and then pulls a processing container. The underlying infrastructure for a Processing job is fully managed by Amazon SageMaker. After you submit a processing job, SageMaker launches the compute instances, processes and analyzes the input data, and releases the resources upon completion. The output of the Processing job is stored in the Amazon S3 bucket you specified.

Note

Your input data must be stored in an Amazon S3 bucket. Alternatively, you can use Amazon Athena or Amazon Redshift as input sources.
Tip
To learn best practices for distributed computing of machine learning (ML) training and processing jobs in general, see Distributed computing with SageMaker best practices.

Use Amazon SageMaker Processing Sample Notebooks

We provide two sample Jupyter notebooks that show how to perform data preprocessing, model evaluation, or both.

For a sample notebook that shows how to run scikit-learn scripts to perform data preprocessing and model training and evaluation with the SageMaker Python SDK for Processing, see scikit-learn Processing. This notebook also shows how to use your own custom container to run processing workloads with your Python libraries and other specific dependencies.

For a sample notebook that shows how to use Amazon SageMaker Processing to perform distributed data preprocessing with Spark, see Distributed Processing (Spark). This notebook also shows how to train a regression model using XGBoost on the preprocessed dataset.

For instructions on how to create and access Jupyter notebook instances that you can use to run these samples in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.
Monitor Amazon SageMaker Processing Jobs with CloudWatch Logs and Metrics

Amazon SageMaker Processing provides Amazon CloudWatch logs and metrics to monitor processing jobs. CloudWatch provides CPU, GPU, memory, GPU memory, and disk metrics, and event logging. For more information, see Monitor Amazon SageMaker with Amazon CloudWatch and Log Amazon SageMaker Events with Amazon CloudWatch.

Data Processing with Apache Spark

Apache Spark is a unified analytics engine for large-scale data processing. Amazon SageMaker provides prebuilt Docker images that include Apache Spark and other dependencies needed to run distributed data processing jobs. With the Amazon SageMaker Python SDK, you can easily apply data transformations and extract features (feature engineering) using the Spark framework. For information about using the SageMaker Python SDK to run Spark processing jobs, see Data Processing with Spark in the Amazon SageMaker Python SDK.

A code repository that contains the source code and Dockerfiles for the Spark images is available on GitHub.

Running a Spark Processing Job

You can use the sagemaker.spark.PySparkProcessor or sagemaker.spark.SparkJarProcessor class to run your Spark application inside of a processing job. Note you can set MaxRuntimeInSeconds to a maximum runtime limit of 5 days. With respect to execution time, and number of instances used, simple spark workloads see a near linear relationship between the number of instances vs. time to completion.

The following code example shows how to run a processing job that invokes your PySpark script preprocess.py.

```python
from sagemaker.spark.processing import PySparkProcessor

spark_processor = PySparkProcessor(
    base_job_name="spark-preprocessor",
    framework_version="2.4",
    role=role,
    instance_count=2,
    ...)"
instance_type="ml.m5.xlarge",
max_runtime_in_seconds=1200,
)

spark_processor.run(
    submit_app="preprocess.py",
    arguments=[
        's3_input_bucket', bucket,
        's3_input_key_prefix', input_prefix,
        's3_output_bucket', bucket,
        's3_output_key_prefix', output_prefix
    ]
)

For an in-depth look, see the Distributed Data Processing with Apache Spark and SageMaker Processing example notebook.

If you are not using the Amazon SageMaker Python SDK and one of its Processor classes to retrieve the pre-built images, you can retrieve these images yourself. The SageMaker prebuilt Docker images are stored in Amazon Elastic Container Registry (Amazon ECR). For a complete list of the available pre-built Docker images, see the available images document.

To learn more about using the SageMaker Python SDK with Processing containers, see Amazon SageMaker Python SDK.

**Data Processing with scikit-learn**

For a sample notebook that shows how to run scikit-learn scripts using a Docker image provided and maintained by SageMaker to preprocess data and evaluate models, see scikit-learn Processing. To use this notebook, you need to install the SageMaker Python SDK for Processing.

This notebook runs a processing job using SKLearnProcessor class from the the SageMaker Python SDK to run a scikit-learn script that you provide. The script preprocesses data, trains a model using a SageMaker training job, and then runs a processing job to evaluate the trained model. The processing job estimates how the model is expected to perform in production.

To learn more about using the SageMaker Python SDK with Processing containers, see the SageMaker Python SDK. For a complete list of pre-built Docker images available for processing jobs, see Docker Registry Paths and Example Code.

The following code example shows how the notebook uses SKLearnProcessor to run your own scikit-learn script using a Docker image provided and maintained by SageMaker, instead of your own Docker image.
from sagemaker.sklearn.processing import SKLearnProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput

sklearn_processor = SKLearnProcessor(framework_version='0.20.0',
    role=role,
    instance_type='ml.m5.xlarge',
    instance_count=1)

sklearn_processor.run(code='preprocessing.py',
    inputs=[ProcessingInput(
        source='s3://path/to/my/input-data.csv',
        destination='/opt/ml/processing/input'),
        outputs=[ProcessingOutput(source='/opt/ml/processing/output/
    train'),
        ProcessingOutput(source='/opt/ml/processing/output/
    validation'),
        ProcessingOutput(source='/opt/ml/processing/output/
    test')]
    )

To process data in parallel using Scikit-Learn on Amazon SageMaker Processing, you can shard input objects by S3 key by setting s3_data_distribution_type='ShardedByS3Key' inside a ProcessingInput so that each instance receives about the same number of input objects.

**Data Processing with Framework Processors**


The FrameworkProcessor class also provides you with customization over the container configuration. The FrameworkProcessor class supports specifying a source directory source_dir for your processing scripts and dependencies. With this capability, you can give the processor access to multiple scripts in a directory instead of only specifying one script. FrameworkProcessor also supports including a requirements.txt file in the source_dir for customizing the Python libraries to install in the container.

For more information on the FrameworkProcessor class and its methods and parameters, see FrameworkProcessor in the *Amazon SageMaker Python SDK*. 
To see examples of using a FrameworkProcessor for each of the supported machine learning frameworks, see the following topics.

**Topics**

- **Hugging Face Framework Processor**
- **MXNet Framework Processor**
- **PyTorch Framework Processor**
- **TensorFlow Framework Processor**
- **XGBoost Framework Processor**

**Hugging Face Framework Processor**

Hugging Face is an open-source provider of natural language processing (NLP) models. The HuggingFaceProcessor in the Amazon SageMaker Python SDK provides you with the ability to run processing jobs with Hugging Face scripts. When you use the HuggingFaceProcessor, you can leverage an Amazon-built Docker container with a managed Hugging Face environment so that you don't need to bring your own container.

The following code example shows how you can use the HuggingFaceProcessor to run your Processing job using a Docker image provided and maintained by SageMaker. Note that when you run the job, you can specify a directory containing your scripts and dependencies in the `source_dir` argument, and you can have a `requirements.txt` file located inside your `source_dir` directory that specifies the dependencies for your processing script(s). SageMaker Processing installs the dependencies in `requirements.txt` in the container for you.

```python
from sagemaker.huggingface import HuggingFaceProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker import get_execution_role

# Initialize the HuggingFaceProcessor
hfp = HuggingFaceProcessor(
    role=get_execution_role(),
    instance_count=1,
    instance_type='ml.g4dn.xlarge',
    transformers_version='4.4.2',
    pytorch_version='1.6.0',
    base_job_name='frameworkprocessor-hf'
)
```
# Run the processing job

```python
hfp.run(
    code='processing-script.py',
    source_dir='scripts',
    inputs=[
        ProcessingInput(
            input_name='data',
            source=f's3://{BUCKET}/{S3_INPUT_PATH}',
            destination='/opt/ml/processing/input/data/'
        )
    ],
    outputs=[
        ProcessingOutput(output_name='train', source='/opt/ml/processing/output/train/', destination=f's3://{BUCKET}/{S3_OUTPUT_PATH}'),
        ProcessingOutput(output_name='test', source='/opt/ml/processing/output/test/', destination=f's3://{BUCKET}/{S3_OUTPUT_PATH}'),
        ProcessingOutput(output_name='val', source='/opt/ml/processing/output/val/', destination=f's3://{BUCKET}/{S3_OUTPUT_PATH}')
    ]
)
```

If you have a `requirements.txt` file, it should be a list of libraries you want to install in the container. The path for `source_dir` can be a relative, absolute, or Amazon S3 URI path. However, if you use an Amazon S3 URI, then it must point to a tar.gz file. You can have multiple scripts in the directory you specify for `source_dir`. To learn more about the HuggingFaceProcessor class, see [Hugging Face Estimator](https://sagemaker.readthedocs.io/en/latest/api/estimator/HuggingFaceEstimator.html) in the [Amazon SageMaker Python SDK](https://github.com/aws/sagemaker-python-sdk).

**MXNet Framework Processor**

Apache MXNet is an open-source deep learning framework commonly used for training and deploying neural networks. The MXNetProcessor in the Amazon SageMaker Python SDK provides you with the ability to run processing jobs with MXNet scripts. When you use the MXNetProcessor, you can leverage an Amazon-built Docker container with a managed MXNet environment so that you don’t need to bring your own container.

The following code example shows how you can use the MXNetProcessor to run your Processing job using a Docker image provided and maintained by SageMaker. Note that when you run the job, you can specify a directory containing your scripts and dependencies in the `source_dir` argument, and you can have a `requirements.txt` file located inside your `source_dir` directory.
that specifies the dependencies for your processing script(s). SageMaker Processing installs the dependencies in `requirements.txt` in the container for you.

```python
from sagemaker.mxnet import MXNetProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker import get_execution_role

#Initialize the MXNetProcessor
mxp = MXNetProcessor(
    framework_version='1.8.0',
    py_version='py37',
    role=get_execution_role(),
    instance_count=1,
    instance_type='ml.c5.xlarge',
    base_job_name='frameworkprocessor-mxnet'
)

#Run the processing job
mxp.run(
    code='processing-script.py',
    source_dir='scripts',
    inputs=[
        ProcessingInput(
            input_name='data',
            source=f's3://{BUCKET}/{S3_INPUT_PATH}',
            destination='/opt/ml/processing/input/data/
        )
    ],
    outputs=[
        ProcessingOutput(
            output_name='processed_data',
            source='/opt/ml/processing/output/',
            destination=f's3://{BUCKET}/{S3_OUTPUT_PATH}'
        )
    ]
)
```

If you have a `requirements.txt` file, it should be a list of libraries you want to install in the container. The path for `source_dir` can be a relative, absolute, or Amazon S3 URI path. However, if you use an Amazon S3 URI, then it must point to a tar.gz file. You can have multiple scripts in the directory you specify for `source_dir`. To learn more about the MXNetProcessor class, see [MXNet Estimator](https://boto3.amazonaws.com/s3dist/index.html) in the [Amazon SageMaker Python SDK](https://docs.aws.amazon.com/sagemaker/latest/dg/).
PyTorch Framework Processor

PyTorch is an open-source machine learning framework. The PyTorchProcessor in the Amazon SageMaker Python SDK provides you with the ability to run processing jobs with PyTorch scripts. When you use the PyTorchProcessor, you can leverage an Amazon-built Docker container with a managed PyTorch environment so that you don’t need to bring your own container.

The following code example shows how you can use the PyTorchProcessor to run your Processing job using a Docker image provided and maintained by SageMaker. Note that when you run the job, you can specify a directory containing your scripts and dependencies in the source_dir argument, and you can have a requirements.txt file located inside your source_dir directory that specifies the dependencies for your processing script(s). SageMaker Processing installs the dependencies in requirements.txt in the container for you.

For the PyTorch versions supported by SageMaker, see the available Deep Learning Container images.

```python
from sagemaker.pytorch.processing import PyTorchProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker import get_execution_role

#Initialize the PyTorchProcessor
pytorch_processor = PyTorchProcessor(
    framework_version='1.8',
    role=get_execution_role(),
    instance_type='ml.m5.xlarge',
    instance_count=1,
    base_job_name='frameworkprocessor-PT'
)

#Run the processing job
pytorch_processor.run(
    code='processing-script.py',
    source_dir='scripts',
    inputs=[
        ProcessingInput(
            input_name='data',
            source=f's3://{BUCKET}/{S3_INPUT_PATH}',
            destination='/opt/ml/processing/input'
        )
    ],
    outputs=[
```
If you have a requirements.txt file, it should be a list of libraries you want to install in the container. The path for source_dir can be a relative, absolute, or Amazon S3 URI path. However, if you use an Amazon S3 URI, then it must point to a tar.gz file. You can have multiple scripts in the directory you specify for source_dir. To learn more about the PyTorchProcessor class, see PyTorch Estimator in the Amazon SageMaker Python SDK.

**TensorFlow Framework Processor**

TensorFlow is an open-source machine learning and artificial intelligence library. The TensorFlowProcessor in the Amazon SageMaker Python SDK provides you with the ability to run processing jobs with TensorFlow scripts. When you use the TensorFlowProcessor, you can leverage an Amazon-built Docker container with a managed TensorFlow environment so that you don’t need to bring your own container.

The following code example shows how you can use the TensorFlowProcessor to run your Processing job using a Docker image provided and maintained by SageMaker. Note that when you run the job, you can specify a directory containing your scripts and dependencies in the source_dir argument, and you can have a requirements.txt file located inside your source_dir directory that specifies the dependencies for your processing script(s). SageMaker Processing installs the dependencies in requirements.txt in the container for you.

```python
from sagemaker.tensorflow import TensorFlowProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker import get_execution_role

#Initialize the TensorFlowProcessor
tp = TensorFlowProcessor(
    framework_version='2.3',
)```
If you have a requirements.txt file, it should be a list of libraries you want to install in the container. The path for source_dir can be a relative, absolute, or Amazon S3 URI path. However, if you use an Amazon S3 URI, then it must point to a tar.gz file. You can have multiple scripts in the directory you specify for source_dir. To learn more about the TensorFlowProcessor class, see TensorFlow Estimator in the Amazon SageMaker Python SDK.

**XGBoost Framework Processor**

XGBoost is an open-source machine learning framework. The XGBoostProcessor in the Amazon SageMaker Python SDK provides you with the ability to run processing jobs with XGBoost scripts.
When you use the XGBoostProcessor, you can leverage an Amazon-built Docker container with a managed XGBoost environment so that you don’t need to bring your own container.

The following code example shows how you can use the XGBoostProcessor to run your Processing job using a Docker image provided and maintained by SageMaker. Note that when you run the job, you can specify a directory containing your scripts and dependencies in the source_dir argument, and you can have a requirements.txt file located inside your source_dir directory that specifies the dependencies for your processing script(s). SageMaker Processing installs the dependencies in requirements.txt in the container for you.

```python
from sagemaker.xgboost import XGBoostProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker import get_execution_role

#Initialize the XGBoostProcessor
xgb = XGBoostProcessor(
    framework_version='1.2-2',
    role=get_execution_role(),
    instance_type='ml.m5.xlarge',
    instance_count=1,
    base_job_name='frameworkprocessor-XGB',
)

#Run the processing job
xgb.run(
    code='processing-script.py',
    source_dir='scripts',
    inputs=[
        ProcessingInput(
            input_name='data',
            source=f's3://{BUCKET}/{S3_INPUT_PATH}',
            destination='/opt/ml/processing/input/data'
        ),
    ],
    outputs=[
        ProcessingOutput(
            output_name='processed_data',
            source='/opt/ml/processing/output/',
            destination=f's3://{BUCKET}/{S3_OUTPUT_PATH}'
        ),
    ],
)
```
If you have a requirements.txt file, it should be a list of libraries you want to install in the container. The path for source_dir can be a relative, absolute, or Amazon S3 URI path. However, if you use an Amazon S3 URI, then it must point to a tar.gz file. You can have multiple scripts in the directory you specify for source_dir. To learn more about the XGBoostProcessor class, see XGBoost Estimator in the Amazon SageMaker Python SDK.

### Use Your Own Processing Code

You can install libraries to run your scripts in your own processing container or, in a more advanced scenario, you can build your own processing container that satisfies the contract to run in Amazon SageMaker. For more information about containers in SageMaker, see Use Docker containers to build models. For a formal specification that defines the contract for an Amazon SageMaker Processing container, see Build Your Own Processing Container (Advanced Scenario).

#### Topics

- Run Scripts with Your Own Processing Container
- Build Your Own Processing Container (Advanced Scenario)

### Run Scripts with Your Own Processing Container

You can use scikit-learn scripts to preprocess data and evaluate your models. To see how to run scikit-learn scripts to perform these tasks, see the scikit-learn Processing sample notebook. This notebook uses the ScriptProcessor class from the Amazon SageMaker Python SDK for Processing.

The following example shows a general workflow for using a ScriptProcessor class with your own processing container. The workflow shows how to create your own image, build your container, and use a ScriptProcessor class to run a Python preprocessing script with the container. The processing job processes your input data and saves the processed data in Amazon Simple Storage Service (Amazon S3).

Before using the following examples, you need to have your own input data and a Python script prepared to process your data. For an end-to-end, guided example of this process, refer back to the scikit-learn Processing sample notebook.

1. Create a Docker directory and add the Dockerfile used to create the processing container. Install pandas and scikit-learn into it. (You could also install your own dependencies with a similar RUN command.)
2. Build the container using the docker command, create an Amazon Elastic Container Registry (Amazon ECR) repository, and push the image to Amazon ECR.

```python
import boto3

account_id = boto3.client('sts').get_caller_identity().get('Account')
region = boto3.Session().region_name
ecr_repository = 'sagemaker-processing-container'
tag = ':latest'
processing_repository_uri = '{}.dkr.ecr.{}.amazonaws.com/{}'.format(account_id, region, ecr_repository + tag)

# Create ECR repository and push docker image
!docker build -t $ecr_repository docker
!aws ecr get-login-password --region {region} | docker login --username AWS --password-stdin {account_id}.dkr.ecr.{region}.amazonaws.com
!aws ecr create-repository --repository-name $ecr_repository
!docker tag {ecr_repository + tag} $processing_repository_uri
!docker push $processing_repository_uri
```

3. Set up the ScriptProcessor from the SageMaker Python SDK to run the script. Replace `image_uri` with the URI for the image you created, and replace `role_arn` with the ARN for an AWS Identity and Access Management role that has access to your target Amazon S3 bucket.

```python
from sagemaker.processing import ScriptProcessor, ProcessingInput, ProcessingOutput

script_processor = ScriptProcessor(command=['python3'],
                                   image_uri='image_uri',
                                   role='role_arn',
                                   instance_count=1,
```

Run Scripts with a Processing Container
4. Run the script. Replace `preprocessing.py` with the name of your own Python processing script, and replace `s3://path/to/my/input-data.csv` with the Amazon S3 path to your input data.

```python
script_processor.run(code='preprocessing.py',
    inputs=[ProcessingInput(
        source='s3://path/to/my/input-data.csv',
        destination='/opt/ml/processing/input')],
    outputs=[ProcessingOutput(source='/opt/ml/processing/output/train'),
        ProcessingOutput(source='/opt/ml/processing/output/validation'),
        ProcessingOutput(source='/opt/ml/processing/output/test')])
```

You can use the same procedure with any other library or system dependencies. You can also use existing Docker images. This includes images that you run on other platforms such as Kubernetes.

**Build Your Own Processing Container (Advanced Scenario)**

You can provide Amazon SageMaker Processing with a Docker image that has your own code and dependencies to run your data processing, feature engineering, and model evaluation workloads.

The following example of a Dockerfile builds a container with the Python libraries scikit-learn and pandas, which you can run as a processing job.

```dockerfile
FROM python:3.7-slim-buster

# Install scikit-learn and pandas
RUN pip3 install pandas==0.25.3 scikit-learn==0.21.3

# Add a Python script and configure Docker to run it
ADD processing_script.py /
ENTRYPOINT ["python3", "/processing_script.py"]
```

For an example of a processing script, see [Get started with SageMaker Processing](#).
Build and push this Docker image to an Amazon Elastic Container Registry (Amazon ECR) repository and ensure that your SageMaker IAM role can pull the image from Amazon ECR. Then you can run this image on Amazon SageMaker Processing.

**How Amazon SageMaker Processing Runs Your Processing Container Image**

Amazon SageMaker Processing runs your processing container image in a similar way as the following command, where `AppSpecification.ImageUri` is the Amazon ECR image URI that you specify in a `CreateProcessingJob` operation.

```
docker run [AppSpecification.ImageUri]
```

This command runs the `ENTRYPOINT` command configured in your Docker image.

You can also override the entrypoint command in the image or give command-line arguments to your entrypoint command using the `AppSpecification.ContainerEntrypoint` and `AppSpecification.ContainerArgument` parameters in your `CreateProcessingJob` request. Specifying these parameters configures Amazon SageMaker Processing to run the container similar to the way that the following command does.

```
```

For example, if you specify the `ContainerEntrypoint` to be `[python3, -v, /processing_script.py]` in your `CreateProcessingJob` request, and `ContainerArguments` to be `[data-format, csv]`, Amazon SageMaker Processing runs your container with the following command.

```
python3 -v /processing_script.py data-format csv
```

When building your processing container, consider the following details:

- Amazon SageMaker Processing decides whether the job completes or fails depending on the exit code of the command run. A processing job completes if all of the processing containers exit successfully with an exit code of 0, and fails if any of the containers exits with a non-zero exit code.
- Amazon SageMaker Processing lets you override the processing container's entrypoint and set command-line arguments just like you can with the Docker API. Docker images can also configure
the entrypoint and command-line arguments using the ENTRYPOINT and CMD instructions. The way CreateProcessingJob's ContainerEntrypoint and ContainerArgument parameters configure a Docker image's entrypoint and arguments mirrors how Docker overrides the entrypoint and arguments through the Docker API:

- If neither ContainerEntrypoint nor ContainerArguments are provided, Processing uses the default ENTRYPOINT or CMD in the image.
- If ContainerEntrypoint is provided, but not ContainerArguments, Processing runs the image with the given entrypoint, and ignores the ENTRYPOINT and CMD in the image.
- If ContainerArguments is provided, but not ContainerEntrypoint, Processing runs the image with the default ENTRYPOINT in the image and with the provided arguments.
- If both ContainerEntrypoint and ContainerArguments are provided, Processing runs the image with the given entrypoint and arguments, and ignores the ENTRYPOINT and CMD in the image.

- You must use the exec form of the ENTRYPOINT instruction in your Dockerfile (ENTRYPOINT ["executable", "param1", "param2"])) instead of the shell form (ENTRYPOINT command param1 param2). This lets your processing container receive SIGINT and SIGKILL signals, which Processing uses to stop processing jobs with the StopProcessingJob API.
- /opt/ml and all its subdirectories are reserved by SageMaker. When building your Processing Docker image, don't place any data required by your processing container in these directories.
- If you plan to use GPU devices, make sure that your containers are nvidia-docker compatible. Include only the CUDA toolkit in containers. Don't bundle NVIDIA drivers with the image. For more information about nvidia-docker, see NVIDIA/nvidia-docker.

How Amazon SageMaker Processing Configures Input and Output For Your Processing Container

When you create a processing job using the CreateProcessingJob operation, you can specify multiple ProcessingInput and ProcessingOutput values.

You use the ProcessingInput parameter to specify an Amazon Simple Storage Service (Amazon S3) URI to download data from, and a path in your processing container to download the data to. The ProcessingOutput parameter configures a path in your processing container from which to upload data, and where in Amazon S3 to upload that data to. For both ProcessingInput and ProcessingOutput, the path in the processing container must begin with /opt/ml/processing/.
For example, you might create a processing job with one ProcessingInput parameter that downloads data from s3://your-data-bucket/path/to/input/csv/data into /opt/ml/processing/csv in your processing container, and a ProcessingOutput parameter that uploads data from /opt/ml/processing/processed_csv to s3://your-data-bucket/path/to/output/csv/data. Your processing job would read the input data, and write output data to /opt/ml/processing/processed_csv. Then it uploads the data written to this path to the specified Amazon S3 output location.

⚠️ **Important**

Symbolic links (symlinks) can not be used to upload output data to Amazon S3. Symlinks are not followed when uploading output data.

### How Amazon SageMaker Processing Provides Logs and Metrics for Your Processing Container

When your processing container writes to stdout or stderr, Amazon SageMaker Processing saves the output from each processing container and puts it in Amazon CloudWatch logs. For information about logging, see [Log Amazon SageMaker Events with Amazon CloudWatch](#).

Amazon SageMaker Processing also provides CloudWatch metrics for each instance running your processing container. For information about metrics, see [Monitor Amazon SageMaker with Amazon CloudWatch](#).

### How Amazon SageMaker Processing Configures Your Processing Container

Amazon SageMaker Processing provides configuration information to your processing container through environment variables and two JSON files—`/opt/ml/config/processingjobconfig.json` and `/opt/ml/config/resourceconfig.json”—at predefined locations in the container.

When a processing job starts, it uses the environment variables that you specified with the Environment map in the CreateProcessingJob request. The `/opt/ml/config/processingjobconfig.json` file contains information about the hostnames of your processing containers, and is also specified in the CreateProcessingJob request.

The following example shows the format of the `/opt/ml/config/processingjobconfig.json` file.
{   "ProcessingJobArn": "<processing_job_arn>",
   "ProcessingJobName": "<processing_job_name>",
   "AppSpecification": {
      "ImageUri": "<image_uri>",
      "ContainerEntrypoint": null,
      "ContainerArguments": null
   },
   "Environment": {
      "KEY": "VALUE"
   },
   "ProcessingInputs": [
      {
         "InputName": "input-1",
         "S3Input": {
            "LocalPath": "/opt/ml/processing/input/dataset",
            "S3Uri": "<s3_uri>",
            "S3DataDistributionType": "FullyReplicated",
            "S3DataType": "S3Prefix",
            "S3InputMode": "File",
            "S3CompressionType": "None",
            "S3DownloadMode": "StartOfJob"
         }
      }
   ],
   "ProcessingOutputConfig": {
      "Outputs": [
         {
            "OutputName": "output-1",
            "S3Output": {
               "LocalPath": "/opt/ml/processing/output/dataset",
               "S3Uri": "<s3_uri>",
               "S3UploadMode": "EndOfJob"
            }
         }
      ],
      "KmsKeyId": null
   },
   "ProcessingResources": {
      "ClusterConfig": {
         "InstanceCount": 1,
         "InstanceType": "ml.m5.xlarge",
         "VolumeSizeInGB": 30,
      }"}
The /opt/ml/config/resourceconfig.json file contains information about the hostnames of your processing containers. Use the following hostnames when creating or running distributed processing code.

```
{
    "current_host": "algo-1",
    "hosts": ["algo-1","algo-2","algo-3"]
}
```

Don't use the information about hostnames contained in /etc/hostname or /etc/hosts because it might be inaccurate.

Hostname information might not be immediately available to the processing container. We recommend adding a retry policy on hostname resolution operations as nodes become available in the cluster.

**Save and Access Metadata Information About Your Processing Job**

To save metadata from the processing container after exiting it, containers can write UTF-8 encoded text to the /opt/ml/output/message file. After the processing job enters any terminal status ("Completed", "Stopped", or "Failed"), the "ExitMessage" field in DescribeProcessingJob contains the first 1 KB of this file. Access that initial part of file with a call to DescribeProcessingJob, which returns it through the ExitMessage parameter. For failed processing jobs, you can use this field to communicate information about why the processing container failed.

⚠️ **Important**
Don't write sensitive data to the /opt/ml/output/message file.
If the data in this file isn't UTF-8 encoded, the job fails and returns a ClientError. If multiple containers exit with an ExitMessage, the content of the ExitMessage from each processing container is concatenated, then truncated to 1 KB.

**Run Your Processing Container Using the SageMaker Python SDK**

You can use the SageMaker Python SDK to run your own processing image by using the Processor class. The following example shows how to run your own processing container with one input from Amazon Simple Storage Service (Amazon S3) and one output to Amazon S3.

```python
from sagemaker.processing import Processor, ProcessingInput, ProcessingOutput

processor = Processor(image_uri='<your_ecr_image_uri>',
                       role=role,
                       instance_count=1,
                       instance_type="ml.m5.xlarge")

processor.run(inputs=[ProcessingInput(
    source='<s3_uri or local path>',
    destination='/opt/ml/processing/input_data')],
               outputs=[ProcessingOutput(
    source='/opt/ml/processing/processed_data',
    destination='<s3_uri>')])
```

Instead of building your processing code into your processing image, you can provide a ScriptProcessor with your image and the command that you want to run, along with the code that you want to run inside that container. For an example, see Run Scripts with Your Own Processing Container.

You can also use the scikit-learn image that Amazon SageMaker Processing provides through SKLearnProcessor to run scikit-learn scripts. For an example, see Data Processing with scikit-learn.
Create, store, and share features with Amazon SageMaker Feature Store

The machine learning (ML) development process often begins with extracting data signals also known as *features* from data to train ML models. Amazon SageMaker Feature Store makes it easy for data scientists, machine learning engineers, and general practitioners to create, share, and manage features for ML development. Feature Store accelerates this process by reducing repetitive data processing and curation work required to convert raw data into features for training an ML algorithm.

Further, the processing logic for your data is authored only once, and features generated are used for both training and inference, reducing the training-serving skew. Feature Store is a centralized store for features and associated metadata so features can be easily discovered and reused. You can create an online or an offline store. The online store is used for low latency real-time inference use cases, and the offline store is used for training and batch inference.

The following diagram shows how you can use Feature Store as part of your machine learning pipeline. First, you read in your raw data and process it. You can ingest data via streaming to the online and offline store, or in batches directly to the offline store. You first create a FeatureGroup and configure it to an online or offline store, or both. Then, you can ingest data into your FeatureGroup and store it in your store. A FeatureGroup is a group of features that is defined via a schema in Feature Store to describe a record.

Online store is primarily designed for supporting real-time predictions that need low millisecond latency reads and high throughput writes. Offline store is primarily intended for batch predictions and model training. Offline store is an append only store and can be used to store and access historical feature data. The offline store can help you store and serve features for exploration and model training. The online store retains only the latest feature data. Feature Groups are mutable and can evolve their schema after creation.
How Feature Store works

In Feature Store, features are stored in a collection called a **feature group**. You can visualize a feature group as a table in which each column is a feature, with a unique identifier for each row. In principle, a feature group is composed of features and values specific to each feature. A **Record** is a collection of values for features that correspond to a unique **RecordIdentifier**. Altogether, a **FeatureGroup** is a group of features defined in your **FeatureStore** to describe a **Record**.

You can use Feature Store in the following modes:

- **Online** – In online mode, features are read with low latency (milliseconds) reads and used for high throughput predictions. This mode requires a feature group to be stored in an online store.
- **Offline** – In offline mode, large streams of data are fed to an offline store, which can be used for training and batch inference. This mode requires a feature group to be stored in an offline store. The offline store uses your S3 bucket for storage and can also fetch data using Athena queries.
- **Online and Offline** – This includes both online and offline modes.

You can ingest data into feature groups in Feature Store in two ways: streaming or in batches. When you ingest data through streaming, a collection of records are pushed to Feature Store by calling a synchronous **PutRecord** API call. This API enables you to maintain the latest feature values in Feature Store and to push new feature values as soon an update is detected.

Alternatively, Feature Store can process and ingest data in batches. You can author features using Amazon SageMaker Data Wrangler, create feature groups in Feature Store and ingest features in batches using a SageMaker Processing job with a notebook exported from Data Wrangler. This mode allows for batch ingestion into the offline store. It also supports ingestion into the online store if the feature group is configured for both online and offline use.
Create feature groups

To ingest features into Feature Store, you must first define the feature group and the feature definitions (feature name and data type) for all features that belong to the feature group. After they are created, feature groups are mutable and can evolve their schema. Feature group names are unique within an AWS Region and AWS account. When creating a feature group, you can also create the metadata for the feature group, such as a short description, storage configuration, features for identifying each record, and the event time, as well as tags to store information such as the author, data source, version, and more.

⚠️ Important

FeatureGroup names or associated metadata such as description or tags should not contain any personal identifiable information (PII) or confidential information.

Find, discover, and share features

After you create a feature group in Feature Store, other authorized users of the feature store can share and discover it. Users can browse through a list of all feature groups in Feature Store or discover existing feature groups by searching by feature group name, description, record identifier name, creation date, and tags.

Real-time inference for features stored in the online store

With Feature Store, you can enrich your features stored in the online store in real time with data from a streaming source (clean stream data from another application) and serve the features with low millisecond latency for real-time inference.

You can also perform joins across different FeatureGroups for real-time inference by querying two different FeatureGroups in the client application.

Offline store for model training and batch inference

Feature Store provides offline storage for feature values in your S3 bucket. Your data is stored in your S3 bucket using a prefixing scheme based on event time. The offline store is an append-only store, enabling Feature Store to maintain a historical record of all feature values. Data is stored in the offline store in Parquet format for optimized storage and query access.
You can query, explore, and visualize features using Data Wrangler from the console. Feature Store supports combining data to produce, train, validate, and test data sets, and allows you to extract data at different points in time.

**Feature data ingestion**

Feature generation pipelines can be created to process large batches (1 million rows of data or more) or small batches, and to write feature data to the offline or online store. Streaming sources such as Amazon Managed Streaming for Apache Kafka or Amazon Kinesis can also be used as data sources from which features are extracted and directly fed to the online store for training, inference, or feature creation.

You can push records to Feature Store by calling the synchronous `PutRecord` API call. Since this is a synchronous API call, it allows small batches of updates to be pushed in a single API call. This enables you to maintain high freshness of the feature values and publish values as soon as an update is detected. These are also called *streaming features*.

When feature data is ingested and updated, Feature Store stores historical data for all features in the offline store. For batch ingest, you can pull feature values from your S3 bucket or use Athena to query. You can also use Data Wrangler to process and engineer new features that can then be exported to a chosen S3 bucket to be accessed by Feature Store. For batch ingestion, you can configure a processing job to batch ingest your data into Feature Store, or you can pull feature values from your S3 bucket using Athena.

To remove a Record from your online store, use the `DeleteRecord` API call. This will also add the deleted record to the offline store.

**Resilience in Feature Store**

Feature Store is distributed across multiple Availability Zones (AZs). An AZ is an isolated location within an AWS Region. If some AZs fail, Feature Store can use other AZs. For more information about AZs, see [Resilience in Amazon SageMaker](#).

**Get started with Amazon SageMaker Feature Store**

The following topics give information about using Amazon SageMaker Feature Store. First learn the Feature Store concepts, then how to manage permissions to use Feature Store, how to create and use feature groups using Studio Classic, Jupyter or JupyterLab notebook, how to use Feature Store
using the User Interface through the console, and how to delete feature groups using the console and AWS SDK for Python (Boto3).

The instructions on using Feature Store through the console depends on if you have enabled Studio or Studio Classic as your default experience. For information on accessing Studio Classic, see Launch Studio Classic Using the Amazon SageMaker Console.

Topics

- Feature Store concepts
- Adding policies to your IAM role
- Use Feature Store with SDK for Python (Boto3)
- Using Amazon SageMaker Feature Store in the console
- Delete a feature group

Feature Store concepts

We list common terms used in Amazon SageMaker Feature Store, followed by example diagrams to visualize a few concepts:

- **Feature Store**: Storage and data management layer for machine learning (ML) features. Serves as the single source of truth to store, retrieve, remove, track, share, discover, and control access to features. In the following example diagram, the Feature Store is a store for your feature groups, which contains your ML data, and provides additional services.

- **Online store**: Low latency, high availability store for a feature group that enables real-time lookup of records. The online store allows quick access to the latest record via the `GetRecord` API.

- **Offline store**: Stores historical data in your Amazon S3 bucket. The offline store is used when low (sub-second) latency reads are not needed. For example, the offline store can be used when you want to store and serve features for exploration, model training, and batch inference.

- **Feature group**: The main resource of Feature Store that contains the data and metadata used for training or predicting with a ML model. A feature group is a logical grouping of features used to describe records. In the following example diagram, a feature group contains your ML data.

- **Feature**: A property that is used as one of the inputs to train or predict using your ML model. In the Feature Store API a feature is an attribute of a record. In the following example diagram, a feature describes a column in your ML data table.
• **Feature definition**: Consists of a name and one of the data types: integral, string or fractional. A feature group contains a list of feature definitions. For more information on Feature Store data types, see Data types.

• **Record**: Collection of values for features for a single record identifier. A combination of record identifier and event time values uniquely identify a record within a feature group. In the following example diagram, a record is a row in your ML data table.

• **Record identifier name**: The record identifier name is the name of the feature that identifies the records. It must refer to one of the names of a feature defined in the feature group's feature definitions. Each feature group is defined with a record identifier name.

• **Event time**: Timestamp that you provide corresponding to when the record event occurred. All records in a feature group must have a corresponding event time. The online store only contains the record corresponding to the latest event time, whereas the offline store contains all historic records. For more information on event time formats, see Data types.

• **Ingestion**: Adding new records to a feature group. Ingestion is typically achieved via the PutRecord API.

Topics

• Concepts overview diagram

• Ingestion diagrams

Concepts overview diagram

The following example diagram conceptualizes a few Feature Store concepts:
The Feature Store contains your feature groups and a feature group contains your ML data. In the example diagram, the original feature group contains a data table that has three features (each describing a column) and two records (rows).

- A feature's definition describes the feature name and data type of the feature values that are associated with records.
- A record contains the feature values and is uniquely identified by its record identifier and must include the event time.

**Ingestion diagrams**

Ingestion is the action of adding a record or records to an existing feature group. The online and offline stores are updated differently for different storage use cases.

**Ingestion to the online store example**

The online store acts as a real-time look-up of records and only keeps the most up-to-date records. Once a record is ingested into an existing online store, the updated online store will only keep the record with the latest event time.
In the following example diagram, the original online store contains a ML data table with one record. A record is ingested with the same record identifier name as the original record, and the ingested record has an earlier event time than the original record. As the updated online store only keeps the record with the latest event time, the updated online store contains the original record.
Ingestion to the offline store example
The offline store acts as a historical look-up of records and keeps all records. After a new record is ingested into an existing offline store, the updated offline store will keep the new record.

In the following example diagram, the original offline store contains a ML data table with one record. A record is ingested with the same record identifier name as the original record, and the ingested record has an event time earlier than the original record. As the updated offline store keeps all of the records, the updated offline store contains both records.
### Original offline store

<table>
<thead>
<tr>
<th>Record identifier name</th>
<th>Feature name</th>
<th>Event time name</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A</td>
<td>Later time</td>
</tr>
</tbody>
</table>

### Ingested record

| 1 | B | Earlier time |

### Updated offline store: keeps all records

<table>
<thead>
<tr>
<th>Record identifier name</th>
<th>Feature name</th>
<th>Event time name</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A</td>
<td>Later time</td>
</tr>
<tr>
<td>1</td>
<td>B</td>
<td>Earlier time</td>
</tr>
</tbody>
</table>
Adding policies to your IAM role

To get started with Amazon SageMaker Feature Store you must have a role and add the required policy to your role, AmazonSageMakerFeatureStoreAccess. The following is a walkthrough on how to view the policies attached to a role and how to add a policy to your role. For information on how to create a role, see SageMaker Roles. For information on how to get your execution role, see Get execution role.

1. Open the IAM console at https://console.aws.amazon.com/iam/.
2. In the navigation pane on the left of the IAM console, choose Roles.
3. In the search bar enter the role you are using for Amazon SageMaker Feature Store.
   For examples on how to find your execution role ARN for a notebook within SageMaker, see Get execution role. The role is at the end of the execution role ARN.
4. After you enter the role in the search bar, choose the role.
   Under Permissions policies you can view the policies attached to the role.
5. After you choose the role, choose Add permissions, then choose Attach policies.
6. In the search bar under Other permissions policies enter AmazonSageMakerFeatureStoreAccess and press enter. If the policy does not show, you may already have the policy attached, listed under your Current permissions policies.
7. After you press enter, select the check box next to the policy and then choose Add permissions.
8. After you have attached the policy to your role, the policy will appear under Permissions policies for your IAM role.

Use Feature Store with SDK for Python (Boto3)

The feature group is the main Feature Store resource that contains your machine learning (ML) data and metadata stored in Amazon SageMaker Feature Store. A feature group is a logical grouping of features and records. A feature group's definition is composed of a configurations for its online and offline store and a list of feature definitions that are used to describe the values of your records. The feature definitions must include a record identifier name and an event time name. For more information on feature store concepts, see Feature Store concepts.
Prior to using a feature store you typically load your dataset, run transformations, and set up your features for ingestion. This process has a lot of variation and is highly dependent on your data. The example code in the following topics refer to the [Introduction to Feature Store](#) and [Fraud Detection with Amazon SageMaker Feature Store](#) example notebooks, respectively. Both use the AWS SDK for Python (Boto3). For more Feature Store examples and resources, see [Amazon SageMaker Feature Store resources](#).

Feature Store supports the following feature types: `String`, `Fractional` (IEEE 64-bit floating point value), and `Integral` (Int64 - 64 bit signed integral value). The default type is set to `String`. This means that, if a column in your dataset is not of a `float` or `long` feature type, it defaults to `String` in your feature store.

You may use a schema to describe your data’s columns and data types. You pass this schema into `FeatureDefinitions`, a required parameter for a `FeatureGroup`. You can use the SDK for Python (Boto3), which has automatic data type detection when you use the `load_feature_definitions` function.

The default behavior when a new feature record is added with an already existing record ID is as follows. In the offline store, the new record will be appended. In the online store, if the event time of the new record is less than the existing event time then nothing will happen, but if the event time of the new record is greater than or equal to the existing event time, the record will be overwritten.

When you create a new feature group you can choose one of the following table formats:

- AWS Glue (Default)
- Apache Iceberg

Ingesting data, especially when streaming, can result in a large number of small files deposited into the offline store. This can negatively impact query performance due the higher number of file operations required. To avoid potential performance issues, use the Apache Iceberg table format when creating new feature groups. With Iceberg you can compact the small data files into fewer large files in the partition, resulting in significantly faster queries. This compaction operation is concurrent and does not affect ongoing read and write operations on the feature group. If you choose the Iceberg option when creating new feature groups, Amazon SageMaker Feature Store will create the Iceberg tables using Parquet file format, and register the tables with the AWS Glue Data Catalog.
Important

Note that for feature groups in Iceberg table format, you must specify `String` as the value for the event time. If you specify any other type, you can't create the feature group successfully.

In the following we list some available Feature Store managed resources.

Topics

- Introduction to Feature Store example notebook
- Fraud detection with Feature Store example notebook

Introduction to Feature Store example notebook

The example code on this page refers to the Introduction to Feature Store example notebook. We recommend that you run this notebook in Studio Classic, notebook instances, or JupyterLab because the code in this guide is conceptual and not fully functional if copied.

Use the following to clone the `aws/amazon-sagemaker-examples` GitHub repository, containing the example notebook:

- **For Studio Classic**

  Launch Studio Classic. You can open Studio Classic if Studio or Studio Classic is enabled as your default experience. For instructions on how to open Studio Classic, see Launch Studio Classic Using the Amazon SageMaker Console.

  Clone the `aws/amazon-sagemaker-examples` GitHub repository to Studio Classic by following the steps in Clone a Git Repository in SageMaker Studio Classic.

- **For Amazon SageMaker notebook instances**

  Launch SageMaker notebook instance by following the instructions in Access Notebook Instances.

  Check if the examples are already in your notebooks by following the instructions in Example Notebooks. If not, follow the instructions in Add a Git Repository to Your Amazon SageMaker Account.
Now that you have the SageMaker example notebooks, navigate to the `amazon-sagemaker-examples/sagemaker-featurestore` directory and open the [Introduction to Feature Store](#) example notebook.

**Step 1: Set up your SageMaker session**

To start using Feature Store, create a SageMaker session. Then, set up the Amazon Simple Storage Service (Amazon S3) bucket that you want to use for your features. The Amazon S3 bucket is your offline store. The following code uses the SageMaker default bucket and adds a custom prefix to it.

```python
# SageMaker Python SDK version 2.x is required
import sagemaker
import sys
import boto3
import pandas as pd
import numpy as np
import io
from sagemaker.session import Session
from sagemaker import get_execution_role

prefix = 'sagemaker-featurestore-introduction'
role = get_execution_role()

sagemaker_session = sagemaker.Session()
region = sagemaker_session.boto_region_name
s3_bucket_name = sagemaker_session.default_bucket()
```

**Note**

The role that you use to run the notebook must have the following managed policies attached to it: `AmazonS3FullAccess` and `AmazonSageMakerFeatureStoreAccess`. For information about adding policies to your IAM role, see [Adding policies to your IAM role](#).

**Step 2: Inspect your data**

In this notebook example, we ingest synthetic data from the [GitHub repository](#) that hosts the full notebook.
customer_data = pd.read_csv("data/feature_store_introduction_customer.csv")
orders_data = pd.read_csv("data/feature_store_introduction_orders.csv")

print(customer_data.head())
print(orders_data.head())

The following diagram illustrates the steps that data goes through before Feature Store ingests it. In this notebook, we illustrate the use case where you have data from multiple sources and want to store them independently in a Feature Store. Our example considers data from a data warehouse (customer data), and data from a real-time streaming service (order data).

Step 3: Create feature groups

We first start by creating feature group names for customer_data and orders_data. Following this, we create two feature groups, one for customer_data and another for orders_data:

```python
import time
from time import strftime, gmtime

customers_feature_group_name = 'customers-feature-group-' + strftime('%d-%H-%M-%S',
                                                                  gmtime())
```
orders_feature_group_name = 'orders-feature-group-' + strftime('%d-%H-%M-%S', gmtime())

Instantiate a FeatureGroup object for customers_data and orders_data:

```python
from sagemaker.feature_store.feature_group import FeatureGroup

customers_feature_group = FeatureGroup(
    name=customers_feature_group_name, sagemaker_session=sagemaker_session
)
orders_feature_group = FeatureGroup(
    name=orders_feature_group_name, sagemaker_session=sagemaker_session
)
```

```python
import time
current_time_sec = int(round(time.time()))
record_identifier_feature_name = "customer_id"
```

Append EventTime feature to your data frame. This parameter is required, and timestamps each data point:

```python
customer_data["EventTime"] = pd.Series([current_time_sec]*len(customer_data),
        dtype="float64")
orders_data["EventTime"] = pd.Series([current_time_sec]*len(orders_data),
        dtype="float64")
```

Load feature definitions to your feature group:

```python
customers_feature_group.load_feature_definitions(data_frame=customer_data)
orders_feature_group.load_feature_definitions(data_frame=orders_data)
```

The following calls create to create two feature groups, customers_feature_group and orders_feature_group, respectively:

```python
customers_feature_group.create(
    s3_uri=f"s3://{s3_bucket_name}/{prefix}",
    record_identifier_name=record_identifier_feature_name,
    event_time_feature_name="EventTime",
    role_arn=role,
    enable_online_store=True
)
```
orders_feature_group.create(
    s3_uri=f"s3://{s3_bucket_name}/{prefix}",
    record_identifier_name=record_identifier_feature_name,
    event_time_feature_name="EventTime",
    role_arn=role,
    enable_online_store=True
)

To confirm that your feature group was created, we display it by using DescribeFeatureGroup and ListFeatureGroups APIs:

customers_feature_group.describe()

orders_feature_group.describe()

sagemaker_session.boto_session.client('sagemaker',
    region_name=region).list_feature_groups() # We use the boto client to list FeatureGroups

Step 4: Ingest data into a feature group

After feature groups are created, we can put data into them. If you're using the SageMaker AWS SDK for Python (Boto3), use the ingest API call. If you're using SDK for Python (Boto3), then use the PutRecord API. It will take less than 1 minute to ingest data both of these options. This example uses the SageMaker SDK for Python (Boto3), so it uses the ingest API call:

def check_feature_group_status(feature_group):
    status = feature_group.describe().get("FeatureGroupStatus")
    while status == "Creating":
        print("Waiting for Feature Group to be Created")
        time.sleep(5)
        status = feature_group.describe().get("FeatureGroupStatus")
    print(f"FeatureGroup {feature_group.name} successfully created.")

check_feature_group_status(customers_feature_group)
check_feature_group_status(orders_feature_group)

customers_feature_group.ingest()}
Using an arbitrary customer record id, 573291 we use get_record to check that the data has been ingested into the feature group.

```python
customer_id = 573291
sample_record = sagemaker_session.boto_session.client('sagemaker-featurestore-runtime', region_name=region).get_record(FeatureGroupName=customers_feature_group_name, RecordIdentifierValueAsString=str(customer_id))
print(sample_record)
```

The following demonstrates how to use the batch_get_record to get a batch of records.

```python
all_records = sagemaker_session.boto_session.client('sagemaker-featurestore-runtime', region_name=region).batch_get_record(
    Identifiers=[
        {
            "FeatureGroupName": customers_feature_group_name,
            "RecordIdentifiersValueAsString": ["573291", "109382", "828400", "124013"],
        },
        {
            "FeatureGroupName": orders_feature_group_name,
            "RecordIdentifiersValueAsString": ["573291", "109382", "828400", "124013"],
        },
    ]
)
print(all_records)
```

**Step 5: Clean up**

Here we remove the Feature Groups that we created.
customers_feature_group.delete()
orders_feature_group.delete()

**Step 6: Next steps**

In this example notebook, you learned how to get started with Feature Store, create feature groups, and ingest data into them.

For an advanced example on how to use Feature Store for a fraud detection use case, see [Fraud Detection with Feature Store](#).

**Step 7: Code examples for programmers**

In this notebook we used a variety of different API calls. Most of them are accessible through the SageMaker Python SDK, however some only exist within Boto3. You can invoke the SageMaker Python SDK API calls directly on your Feature Store objects, whereas to invoke API calls that exist within Boto3, you must first access a Boto3 client through your Boto3 and SageMaker sessions: for example, `sagemaker_session.boto_session.client()`.

The following is a list of API calls for this notebook. These calls exist within the SDK for Python and exist in Boto3, for your reference:

**SDK for Python (Boto3) API Calls**

- describe()
- ingest()
- delete()
- create()
- load_feature_definitions()

**Boto3 API Calls**

- list_feature_groups()
- get_record()

**Fraud detection with Feature Store example notebook**

The example code on this page refers to the example notebook: [Fraud Detection with Amazon SageMaker Feature Store](#). We recommend that you run this notebook in Studio Classic, notebook...
instances, or JupyterLab because the code in this guide is conceptual and not fully functional if copied.

Use the following to clone the `aws/amazon-sagemaker-examples` GitHub repository, containing the example notebook.

- **For Studio Classic**

  First launch Studio Classic. You can open Studio Classic if Studio or Studio Classic is enabled as your default experience. To open Studio Classic, see [Launch Studio Classic Using the Amazon SageMaker Console](#).

  Clone the `aws/amazon-sagemaker-examples` GitHub repository to Studio Classic by following the steps in [Clone a Git Repository in SageMaker Studio Classic](#).

- **For Amazon SageMaker notebook instances**

  First launch SageMaker notebook instance by following the instructions in [Access Notebook Instances](#).

  Check if the examples are already in your notebooks by following the instructions in [Example Notebooks](#). If not, follow the instructions in [Add a Git Repository to Your Amazon SageMaker Account](#).

Now that you have the SageMaker example notebooks, navigate to the `amazon-sagemaker-examples/sagemaker-featurestore` directory and open the [Fraud Detection with Amazon SageMaker Feature Store](#) example notebook.

**Step 1: Set up your Feature Store session**

To start using Feature Store, create a SageMaker session, Boto3 session, and a Feature Store session. Also, set up the Amazon S3 bucket you want to use for your features. This is your offline store. The following code uses the SageMaker default bucket and adds a custom prefix to it.

```
Note
The role that you use to run the notebook must have the following managed policies attached to it: AmazonSageMakerFullAccess and AmazonSageMakerFeatureStoreAccess. For information about adding policies to your IAM role, see [Adding policies to your IAM role](#).
```
import boto3
import sagemaker
from sagemaker.session import Session

sagemaker_session = sagemaker.Session()
region = sagemaker_session.boto_region_name
boto_session = boto3.Session(region_name=region)
role = sagemaker.get_execution_role()
default_bucket = sagemaker_session.default_bucket()
prefix = 'sagemaker-featurestore'
offline_feature_store_bucket = 's3://{}/{}'.format(default_bucket, prefix)

sagemaker_client = boto_session.client(service_name='sagemaker', region_name=region)
featurestore_runtime = boto_session.client(service_name='sagemaker-featurestore-runtime', region_name=region)

feature_store_session = Session(
    boto_session=boto_session,
    sagemaker_client=sagemaker_client,
    sagemaker_featurestore_runtime_client=featurestore_runtime
)

**Step 2: Load datasets and partition data into feature groups**

Load your data into data frames for each of your features. You use these data frames after you set up the feature group. In the fraud detection example, you can see these steps in the following code.

```python
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import io

s3_client = boto3.client(service_name='s3', region_name=region)

fraud_detection_bucket_name = 'sagemaker-featurestore-fraud-detection'
identity_file_key = 'sampled_identity.csv'
transaction_file_key = 'sampled_transactions.csv'

identity_data_object = s3_client.get_object(Bucket=fraud_detection_bucket_name,
                                           Key=identity_file_key)
transaction_data_object = s3_client.get_object(Bucket=fraud_detection_bucket_name,
                                            Key=transaction_file_key)
```
Amazon SageMaker

Developer Guide

identity_data = pd.read_csv(io.BytesIO(identity_data_object['Body'].read()))
transaction_data = pd.read_csv(io.BytesIO(transaction_data_object['Body'].read()))
identity_data = identity_data.round(5)
transaction_data = transaction_data.round(5)
identity_data = identity_data.fillna(0)
transaction_data = transaction_data.fillna(0)
# Feature transformations for this dataset are applied before ingestion into
FeatureStore.
# One hot encode card4, card6
encoded_card_bank = pd.get_dummies(transaction_data['card4'], prefix = 'card_bank')
encoded_card_type = pd.get_dummies(transaction_data['card6'], prefix = 'card_type')
transformed_transaction_data = pd.concat([transaction_data, encoded_card_type,
encoded_card_bank], axis=1)
transformed_transaction_data =
transformed_transaction_data.rename(columns={"card_bank_american express":
"card_bank_american_express"})

Step 3: Set up feature groups
When you set up your feature groups, you need to customize the feature names with a unique
name and set up each feature group with the FeatureGroup class.
from sagemaker.feature_store.feature_group import FeatureGroup
feature_group_name = "some string for a name"
feature_group = FeatureGroup(name=feature_group_name,
sagemaker_session=feature_store_session)

For example, in the fraud detection example, the two feature groups are identity and
transaction. In the following code you can see how the names are customized with a timestamp,
and then each group is set up by passing in the name and the session.
import time
from time import gmtime, strftime, sleep
from sagemaker.feature_store.feature_group import FeatureGroup
identity_feature_group_name = 'identity-feature-group-' + strftime('%d-%H-%M-%S',
gmtime())
Use Feature Store with SDK for Python (Boto3)

2476


Step 4: Set up record identifier and event time features

In this step, you specify a record identifier name and an event time feature name. This name maps to the column of the corresponding features in your data. For example, in the fraud detection example, the column of interest is TransactionID. EventTime can be appended to your data when no timestamp is available. In the following code, you can see how these variables are set, and then EventTime is appended to both feature’s data.

```python
record_identifier_name = "TransactionID"
event_time_feature_name = "EventTime"
current_time_sec = int(round(time.time()))
identity_data[event_time_feature_name] = pd.Series([current_time_sec]*len(identity_data), dtype="float64")
transformed_transaction_data[event_time_feature_name] = pd.Series([current_time_sec]*len(transaction_data), dtype="float64")
```

Step 5: Load feature definitions

You can now load the feature definitions by passing a data frame containing the feature data. In the following code for the fraud detection example, the identity feature and transaction feature are each loaded by using load_feature_definitions, and this function automatically detects the data type of each column of data. For developers using a schema rather than automatic detection, see the Export Feature Groups from Data Wrangler example for code that shows how to load the schema, map it, and add it as a FeatureDefinition that you can use to create the FeatureGroup. This example also covers a AWS SDK for Python (Boto3) implementation, which you can use instead of the SageMaker Python SDK.

```python
identity_feature_group.load_feature_definitions(data_frame=identity_data); # output is suppressed
transaction_feature_group.load_feature_definitions(data_frame=transformed_transaction_data); # output is suppressed
```
**Step 6: Create a feature group**

In this step, you use the `create` function to create the feature group. The following code shows all of the available parameters. The online store is not created by default, so you must set this as `True` if you want to enable it. The `s3_uri` is the S3 bucket location of your offline store.

```python
# create a FeatureGroup
feature_group.create(
    description = "Some info about the feature group",
    feature_group_name = feature_group_name,
    record_identifier_name = record_identifier_name,
    event_time_feature_name = event_time_feature_name,
    feature_definitions = feature_definitions,
    role_arn = role,
    s3_uri = offline_feature_store_bucket,
    enable_online_store = True,
    online_store_kms_key_id = None,
    offline_store_kms_key_id = None,
    disable_glue_table_creation = False,
    data_catalog_config = None,
    tags = ["tag1","tag2"])
```

The following code from the fraud detection example shows a minimal `create` call for each of the two features groups being created.

```python
identity_feature_group.create(
    s3_uri=offline_feature_store_bucket,
    record_identifier_name=record_identifier_name,
    event_time_feature_name=event_time_feature_name,
    role_arn=role,
    enable_online_store=True
)

transaction_feature_group.create(
    s3_uri=offline_feature_store_bucket,
    record_identifier_name=record_identifier_name,
    event_time_feature_name=event_time_feature_name,
    role_arn=role,
    enable_online_store=True
)
```
When you create a feature group, it takes time to load the data, and you need to wait until the feature group is created before you can use it. You can check status using the following method.

```python
status = feature_group.describe().get("FeatureGroupStatus")
```

While the feature group is being created, you receive Creating as a response. When this step has finished successfully, the response is Created. Other possible statuses are CreateFailed, Deleting, or DeleteFailed.

**Step 7: Work with feature groups**

Now that you've set up your feature group, you can perform any of the following tasks:

**Topics**
- Describe a feature group
- List feature groups
- Put records in a feature group
- Get records from a feature group
- Generate hive DDL commands
- Build a training dataset
- Write and execute an Athena query
- Delete a feature group

**Describe a feature group**

You can retrieve information about your feature group with the `describe` function.

```python
feature_group.describe()
```

**List feature groups**

You can list all of your feature groups with the `list_feature_groups` function.

```python
sagemaker_client.list_feature_groups()
```
Put records in a feature group

You can use the `ingest` function to load your feature data. You pass in a data frame of feature data, set the number of workers, and choose to wait for it to return or not. The following example demonstrates using the `ingest` function.

```python
feature_group.ingest(
    data_frame=feature_data, max_workers=3, wait=True
)
```

For each feature group you have, run the `ingest` function on the feature data you want to load.

Get records from a feature group

You can use the `get_record` function to retrieve the data for a specific feature by its record identifier. The following example uses an example identifier to retrieve the record.

```python
record_identifier_value = str(2990130)
featurestore_runtime.get_record(FeatureGroupName=transaction_feature_group_name,
    RecordIdentifierValueAsString=record_identifier_value)
```

An example response from the fraud detection example:

```json
...
'Record': [{'FeatureName': 'TransactionID', 'ValueAsString': '2990130'},
    {'FeatureName': 'isFraud', 'ValueAsString': '0'},
    {'FeatureName': 'TransactionDT', 'ValueAsString': '152647'},
    {'FeatureName': 'TransactionAmt', 'ValueAsString': '75.0'},
    {'FeatureName': 'ProductCD', 'ValueAsString': 'H'},
    {'FeatureName': 'card1', 'ValueAsString': '4577'},
    ...
```

Generate hive DDL commands

The SageMaker Python SDK's `FeatureStore` class also provides the functionality to generate Hive DDL commands. The schema of the table is generated based on the feature definitions. Columns are named after feature name and data-type are inferred based on feature type.

```python
print(feature_group.as_hive_ddl())
```

Example output:
CREATE EXTERNAL TABLE IF NOT EXISTS sagemaker_featurestore.identity-feature-group-27-19-33-00 (  TransactionID INT  id_01 FLOAT  id_02 FLOAT  id_03 FLOAT  id_04 FLOAT  ...)

**Build a training dataset**

Feature Store automatically builds an AWS Glue data catalog when you create feature groups and you can turn this off if you want. The following describes how to create a single training dataset with feature values from both identity and transaction feature groups created earlier in this topic. Also, the following describes how to run an Amazon Athena query to join data stored in the offline store from both identity and transaction feature groups.

To start, create an Athena query using `athena_query()` for both identity and transaction feature groups. The `table_name` is the AWS Glue table that is autogenerated by Feature Store.

```python
identity_query = identity_feature_group.athena_query()
transaction_query = transaction_feature_group.athena_query()

identity_table = identity_query.table_name
transaction_table = transaction_query.table_name
```

**Write and execute an Athena query**

You write your query using SQL on these feature groups, and then execute the query with the `.run()` command and specify your Amazon S3 bucket location for the data set to be saved there.

```python
# Athena query
query_string = 'SELECT * FROM "'+transaction_table+'" LEFT JOIN "'+identity_table+'" ON "'+transaction_table+'".transactionid = "'+identity_table+'".transactionid'

# run Athena query. The output is loaded to a Pandas dataframe.
dataset = pd.DataFrame()
identity_query.run(query_string=query_string,
                  output_location='s3://'+default_s3_bucket_name+'/query_results/)
identity_query.wait()
dataset = identity_query.as_dataframe()
```
From here you can train a model using this data set and then perform inference.

**Delete a feature group**

You can delete a feature group with the `delete` function.

```python
feature_group.delete()
```

The following code example is from the fraud detection example.

```python
identity_feature_group.delete()
transaction_feature_group.delete()
```

For more information, see the [Delete a feature group API](https://docs.aws.amazon.com/sagemaker/latest/dg/API_DeleteFeatureGroup.html).

**Using Amazon SageMaker Feature Store in the console**

You can use Amazon SageMaker Feature Store on the console to create, view, update, and monitor your feature groups. Monitoring in this guide includes viewing pipeline executions and lineage of your feature groups. This guide provides instructions on how to achieve these tasks from the console.

For Feature Store examples and resources using the Amazon SageMaker APIs and AWS SDK for Python (Boto3), see [Amazon SageMaker Feature Store resources](https://docs.aws.amazon.com/sagemaker/latest/dg/feature-store-resources.html).

**Topics**

- [Create a feature group from the console](#)
- [View feature group details from the console](#)
- [Update a feature group from the console](#)
- [View pipeline executions from the console](#)
- [View lineage from the console](#)

**Create a feature group from the console**

The create feature group process has four steps:

1. Enter feature group information.
2. Enter feature definitions.
3. Enter required features.
4. Enter feature group tags.

Consider which of the following options fits your use case:

- Create an online store, an offline store, or both. For more information about the differences between online and offline stores, see Feature Store concepts.

- Use a default AWS Key Management Service key or your own KMS key. The default key is AWS KMS key (SSE-KMS). You can reduce AWS KMS request costs by configuring use of Amazon S3 Bucket Keys on the offline store Amazon S3 bucket. The Amazon S3 Bucket Key must be enabled before using the bucket for your feature groups. For more information about reducing the cost by using Amazon S3 Bucket Keys, see Reducing the cost of SSE-KMS with Amazon S3 Bucket Keys.

You can use the same key for both online and offline stores, or have a unique key for each. For more information about AWS KMS, see AWS Key Management Service.

- If you create an offline store:
  - Decide if you want to create an Amazon S3 bucket or use an existing one. When using an existing one, you must know the Amazon S3 bucket URL or Amazon S3 bucket name and dataset directory name, if applicable.
  - Choose which Amazon Resource Name (ARN) to use to specify the IAM role. For more information about how to find your role and attached policies, see Adding policies to your IAM role.
  - Decide whether to use the AWS Glue (default) or Apache Iceberg table format. In most use cases, you use the Apache Iceberg table format. For more information about table formats, see Use Feature Store with SDK for Python (Boto3).

You can use the console to view the lineage of a feature group. The instructions for using Feature Store on the console vary depending on whether you enabled Amazon SageMaker Studio or Amazon SageMaker Studio Classic as your default experience.

Create feature groups if Studio is your default experience (console)

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose Data from the left navigation pane to expand the dropdown list.
3. From the dropdown list, choose **Feature Store**.

4. Choose **Create feature group**.

5. Under **Feature group details**, enter a feature group name.

6. (Optional) Enter a description of the feature group.

7. Under **Feature group storage configuration**, choose a storage configuration from the dropdown list. For information about storage configurations, see [Feature Store storage configurations](#).

8. If you have chosen to enable the online storage:
   a. If you only enable the online storage, you can choose a **Storage type** from the dropdown list. For information about online store storage types, see [Online store](#).
   b. (Optional) Apply **Time to Live (TTL)** by toggling the switch to **On** and specifying the **Time to Live duration** value and unit. This will update the default TTL duration for all records added to the feature group *after the feature group is created*. For more information about TTL, see [Time to live (TTL) duration for records](#).

9. If you have chosen to enable the offline storage:
   a. Under the **Amazon S3 bucket name**, enter a new bucket name, or enter an existing bucket URL, manually.
   b. From the **Table format** dropdown list, choose the table format. In most use cases, you should use the Apache Iceberg table format. For more information about table formats, see [Use Feature Store with SDK for Python (Boto3)](#).
   c. Under **IAM role ARN**, choose the IAM role ARN you want to attach to this feature group. For more information about how to find your role and attached policies, see [Adding policies to your IAM role](#).
   d. If you have chosen to enable the offline storage **Table format** and **AWS Glue (default)** **Table format**, under **Data catalog**, you can choose one of the following two options:
      - Use default values for your AWS Glue Data Catalog.
      - Provide your existing Data Catalog name, table name, and database name to extend your existing AWS Glue Data Catalog.

10. Under the **Online store encryption key** or **Offline store encryption key** dropdown list, choose one of the following options:
    - Use **AWS managed AWS KMS key** (default)
• Enter an AWS KMS key ARN and enter your AWS KMS key ARN under Offline store encryption key ARN. For more information about AWS KMS, see AWS Key Management Service.

11. If applicable, you will have the option to choose your throughput mode, which impacts how you are charged. Under Throughput mode, choose a mode from the dropdown list and input the read and write capacities when available. For information about throughput modes, like when the mode can be applied and capacity units, see Throughput modes.

12. After you specify all of the required information, the Continue button appears available. Choose Continue.

13. Under Specify feature definitions, you have two options for providing a schema for your features: a JSON editor, or a table editor.

   • JSON editor: In the JSON tab, enter or copy and paste your feature definitions in the JSON format.

   • Table editor: In the Table tab, enter the feature feature name and choose the corresponding data type for each feature in your feature group. Choose + Add feature definitions to include more features. Be aware that you cannot remove feature definitions from your feature groups. However, you can add and update feature definitions after the feature group is created.

There must be at least two features in a feature group that represent the record identifier and event time:

   • The record Feature type can be a string, fractional, or an integral.

   • The event time Feature type must be a string or a fractional. However, if you chose the Iceberg table format, the event time must be a string.

14. After all of the features are included, choose Continue.

15. Under Select required features, you must specify the record identifier and event time features. Do this by choosing the feature name under Record identifier feature name and Event time feature name dropdown lists, respectively.

16. After you choose the record identifier and event time features, choose Continue.

17. (Optional) To add tags for the feature group, choose Add new tag. Then enter a tag key and the corresponding value under Key and Value, respectively.

18. Choose Continue.
19. Under **Review feature group**, review the feature group information. To edit any step, choose the **Edit** button that corresponds to that step. This brings you to the corresponding step for editing. To return to step 5, choose **Continue** until you return to step 5.

20. After you finalize the setup for your feature group, choose **Create feature group**.

If an issue occurs during setup, a pop-up alert message appears at the bottom of the page with tips for solving the issue. You can return to previous steps to fix the issues by choosing **Edit** for the step with conflicts.

After the feature group has been successfully created, a green pop-up message appears at the bottom of the page. The new feature group also appears in your feature groups catalog.

**Create feature groups if Studio Classic is your default experience (console)**

1. Open the Studio Classic console by following the instructions in [Launch Amazon SageMaker Studio Classic](#).
2. Choose the **Home** icon on the left navigation pane.
3. Choose **Data**.
4. From the dropdown list, choose **Feature Store**.
5. Choose **Create feature group**.
6. Under **Feature group details**, enter a feature group name.
7. (Optional) Enter a description of the feature group.
8. Under **Feature group storage configuration**, choose a storage configuration from the dropdown list. For information about storage configurations, see [Feature Store storage configurations](#).
9. If you have chosen to enable the online storage:
   a. If you only enable the online storage, you may choose a **Storage type** from the dropdown list. For information about online store storage types, see [Online store](#).
   b. (Optional) Apply **Time to Live (TTL)** by toggling the switch to **On** and specifying the **Time to Live duration** value and unit. This will update the default TTL duration for all records added to the feature group after the feature group is created. For more information about TTL, see [Time to live (TTL) duration for records](#).
10. If you have chosen to enable the offline storage:
   
a. Under the **Amazon S3 bucket name**, enter a new bucket name or enter an existing bucket URL manually.

b. From the **Table format** dropdown list, choose the table format. In most use cases, you should use the Apache Iceberg table format. For more information about table formats, see [Use Feature Store with SDK for Python (Boto3)](https://docs.aws.amazon.com/sagemaker/latest/dg/feature-store-sdk-python-boto3.html).

c. Under **IAM role ARN**, choose the IAM role ARN you want to attach to this feature group. For more information about how to find your role and attached policies, see [Adding policies to your IAM role](https://docs.aws.amazon.com/sagemaker/latest/dg/feature-store-sdk-python-boto3.html).

d. If you have chosen to enable the offline storage **Table format** and AWS Glue (default) **Table format**, under **Data catalog**, you can choose one of the following two options:
   
   - **Use default values for your AWS Glue Data Catalog.**
   - Provide your existing Data Catalog name, table name, and database name to extend your existing AWS Glue Data Catalog.

11. Under the **Online store encryption key** or **Offline store encryption key** dropdown list, choose one of the following options:
   
   - **Use AWS managed AWS KMS key (default)**
   - **Enter an AWS KMS key ARN** and enter your AWS KMS key ARN under **Offline store encryption key ARN**. For more information about AWS KMS, see [AWS Key Management Service](https://docs.aws.amazon.com/kms/latest/devguide/)

12. After you specify all of the required information, the **Continue** button appears available. Choose **Continue**.

13. Under **Specify feature definitions**, you have two options for providing a schema for your features: a JSON editor, or a table editor.

   - **JSON editor**: In the **JSON** tab, enter or copy and paste your feature definitions in the JSON format.
   - **Table editor**: In the **Table** tab, enter the feature feature name and choose the corresponding data type for each feature in your feature group. Choose **Add feature definitions** to include more features. Be aware that you cannot remove feature definitions from your feature groups. However, you can add and update feature definitions after the feature group is created.
There must be at least two features in a feature group that represent the record identifier and event time:

- The record **Feature type** can be a string, fractional, or an integral.
- The event time **Feature type** must be a string or a fractional. However, if you chose the Iceberg table format, the event time must be a string.

14. After all of the features are included, choose **Continue**.

15. Under **Select required features**, you must specify the record identifier and event time features. Do this by choosing the feature name under **Record identifier feature name** and **Event time feature name** dropdown lists, respectively.

16. After you choose the record identifier and event time features, choose **Continue**.

17. (Optional) To add tags for the feature group, choose **Add new tag**. Then enter a tag key and the corresponding value under **Key** and **Value**, respectively.

18. Choose **Continue**.

19. Under **Review feature group**, review the feature group information. To edit any step, choose the **Edit** button that corresponds to that step. This brings you to the corresponding step for editing. To return to step 5, choose **Continue** until you return to step 5.

20. After you finalize the setup for your feature group, choose **Create feature group**.

   If an issue occurs during setup, a pop-up alert message appears at the bottom of the page with tips for solving the issue. You can return to previous steps to fix the issues by choosing **Edit** for the step with conflicts.

   After the feature group has been successfully created, a green pop-up message appears at the bottom of the page. The new feature group also appears in your feature groups catalog.

**View feature group details from the console**

You can view details of your feature groups after a feature group has successfully been created in the Feature Store.

You can use the console or the Amazon SageMaker Feature Store API to view your feature group details. The instructions for using Feature Store through the console depends on if you have enabled Amazon SageMaker Studio or Amazon SageMaker Studio Classic as your default experience.
View feature group details if Studio is your default experience (console)

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose Data in the left navigation pane, to expand the dropdown list.
3. From the dropdown list, choose Feature Store.
4. (Optional) To view your feature groups, choose My account. To view shared feature groups, choose Cross account.
5. Under the Feature group catalog tab, choose your feature group name from the list. This opens the feature group page.
6. On the Features tab, you can find a list of all of the features. Use the filter to refine your list. Choose a feature to view its details.
7. Under the Details tab and the Information subtab, you can review your feature group information. This includes Latest execution, Offline storage settings, Online storage settings, and more.
8. Under the Details tab and the Tags subtab, you can review your feature group tags. Choose Add new tag to add a new tag or Remove to remove a tag.
9. Under the Pipeline Executions tab, you can view the associated pipelines or pipeline executions for your feature group.
10. Under the Lineage tab, you can view the lineage of your feature group.

View feature group details if Studio Classic is your default experience (console)

1. Open the Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.
2. Choose the Home icon
   ( in the left navigation pane.
3. Choose Data.
4. From the dropdown list, choose Feature Store.
5. (Optional) To view your feature groups, choose My account. To view shared feature groups, choose Cross account.
6. Under the Feature group catalog tab, choose your feature group name from the list. This opens the feature group page.
On the **Features** tab, you can find a list of all of the features. Use the filter to refine your list. Choose a feature to view its details.

Under the **Details** tab and the **Information** subtab, you can review your feature group information, including **Latest execution**, **Offline storage settings**, **Online storage settings**, and more.

Under the **Details** tab and the **Tags** subtab, you can review your feature group tags. Choose **Add new tag** to add a new tag or **Remove** to remove a tag.

Under the **Pipeline Executions** tab, you can view the associated pipelines or pipeline executions for your feature group.

Under the **Lineage** tab, you can view the lineage of your feature group.

### Update a feature group from the console

You can update your feature groups after a feature group has successfully been created in the Feature Store.

You can use the console or the Amazon SageMaker Feature Store API to update a feature group. The instructions for using Feature Store through the console depends on if you have enabled [Amazon SageMaker Studio](http://aws.amazon.com) or [Amazon SageMaker Studio Classic](http://aws.amazon.com) as your default experience.

#### Update a feature group if Studio is your default experience (console)

1. Open the Studio console by following the instructions in [Launch Amazon SageMaker Studio](http://aws.amazon.com).
2. Choose **Data** in the left navigation pane, to expand the dropdown list.
3. From the dropdown list, choose **Feature Store**.
4. (Optional) To view your feature groups, choose **My account**. To view shared feature groups, choose **Cross account**.
5. Under the **Feature group catalog** tab, search for and choose your feature group name from the list. This opens the feature group page.
6. Choose **Update feature group**.
7. (Optional) If applicable, you can change your throughput mode, which impacts how you are charged. Under **Throughput mode**, choose a mode from the dropdown list and input the read and write capacities when available. For information about throughput modes, like when the mode can be applied and capacity units, see [Throughput modes](http://aws.amazon.com).
8. (Optional) If your feature group uses the online store, you can update the default **Time to Live (TTL)**. If TTL hasn't been enabled for the feature group, toggle the switch button under
Time to Live (TTL) to On. You can specify the TTL value and unit under Time to Live duration. This will update the default TTL duration for all records added to the feature group after the feature group is updated.

9. (Optional) You can add feature definitions to your feature group but be aware that you cannot remove feature definitions from your feature groups. To add a feature definition, choose + Add feature definition and then specify the new feature definition name under the Name column and select the feature type under the Feature type column.

10. Choose Save changes.

11. To confirm your changes, choose Confirm.

Update a feature group if Studio Classic is your default experience (console)

1. Open the Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.

2. Choose the Home icon

3. Choose Data.

4. From the dropdown list, choose Feature Store.

5. (Optional) To view your feature groups, choose My account. To view shared feature groups, choose Cross account.

6. Under the Feature group catalog tab, search for and choose your feature group name from the list. This opens the feature group page.

7. Choose Update feature group.

8. (Optional) If your feature group uses the online store, you can update the default Time to Live (TTL). If TTL hasn't been enabled for the feature group, toggle the switch button under Time to Live (TTL) to On. You can specify the TTL value and unit under Time to Live duration. This will update the default TTL duration for all records added to the feature group after the feature group is updated.

9. (Optional) You can add feature definitions to your feature group but be aware that you cannot remove feature definitions from your feature groups. To add a feature definition, choose + Add feature definition and then specify the new feature definition name under the Name column and select the feature type under the Feature type column.

10. Choose Save changes.
11. To confirm your changes, choose **Confirm**.

**View pipeline executions from the console**

You can view the latest pipeline execution information for a feature or feature group under **Pipeline executions**. You can also get links to pipelines, executions, code, and other useful execution information.

You can use the console to view your pipeline executions. The instructions for using Feature Store through the console depends on if you have enabled Amazon SageMaker Studio or Amazon SageMaker Studio Classic as your default experience.

**View pipeline executions if Studio is your default experience (console)**

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose **Data** in the left navigation pane, to expand the dropdown list.
3. From the dropdown list, choose **Feature Store**.
4. (Optional) To view your feature groups, choose **My account**. To view shared feature groups, choose **Cross account**.
5. Choose a feature group or feature to view their pipeline executions.
6. Choose the **Pipeline executions** tab.
7. Search for a pipeline from the **Select a pipeline** dropdown list.
8. You can view the links for the pipeline, execution, and code details. You can also view the execution owner, status, date, and duration.

**View pipeline executions if Studio Classic is your default experience (console)**

1. Open the Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.
2. Choose the **Home** icon
   
   ( )
   
   in the left navigation pane.
3. Choose **Data**.
4. From the dropdown list, choose **Feature Store**.
5. (Optional) To view your feature groups, choose **My account**. To view shared feature groups, choose **Cross account**.

6. Choose a feature group or feature to view their pipeline executions.

7. Choose the **Pipeline executions** tab.

8. Search for a pipeline from the **Select a pipeline** dropdown list.

9. You can view the links for the pipeline, execution, and code details. You can also view the execution owner, status, date, and duration.

### View lineage from the console

You can view the lineage of a feature group. The lineage includes the information about the execution code of your feature processing workflow, what data sources were used, and how they are ingested to the feature group or feature.

You can use the console to view the lineage of a feature group. The instructions on using Feature Store through the console depends on if you have enabled [Amazon SageMaker Studio](https://aws.amazon.com/sagemaker/studio) or [Amazon SageMaker Studio Classic](https://aws.amazon.com/sagemaker/studio-classic) as your default experience.

### View lineage if Studio is your default experience (console)

1. Open the Studio console by following the instructions in [Launch Amazon SageMaker Studio](https://aws.amazon.com/sagemaker/studio).

2. Choose **Data** from the left navigation pane to expand the dropdown list.

3. From the dropdown list, choose **Feature Store**.

4. (Optional) To view your feature groups, choose **My account**. To view shared feature groups, choose **Cross account**.

5. Choose a feature group or feature to view its lineage details.

6. Choose the **Lineage** tab.

7. Choose a feature group or pipeline node to expand the node. This contains more information about a feature group or pipeline.

8. You can zoom in, zoom out, or recenter the lineage graph by using the buttons on the bottom left of the screen.

9. You can move through the lineage map when you choose and drag the screen. To move your lineage maps using nodes as the focal point, you can press **Tab** or **Shift+Tab** to switch between nodes.
10. If applicable, you can navigate the lineage upstream (left, earlier) or downstream (right, most recent). Do this by choosing a node and then choosing **Query upstream lineage** or **Query downstream lineage**.

**View lineage if Studio Classic is your default experience (console)**

1. Open Studio Classic by following the instructions in [Launch Amazon SageMaker Studio Classic](#).
2. Choose the **Home** icon (in the left navigation pane).
3. Choose **Data**.
4. From the dropdown list, choose **Feature Store**.
5. (Optional) To view your feature groups, choose **My account**. To view shared feature groups, choose **Cross account**.
6. Choose a feature group or feature to view its lineage details.
7. Choose the **Lineage** tab.
8. Choose a feature group or pipeline node to expand the node. This contains more information about a feature group or pipeline.
9. You can zoom in, zoom out, or recenter the lineage graph by using the buttons on the bottom left of the screen.
10. You can move through the lineage map when you choose and drag the screen. To move your lineage maps using nodes as the focal point, you can press **Tab** or **Shift+Tab** to switch between nodes.
11. If applicable, you can navigate the lineage upstream (left, earlier) or downstream (right, most recent). Do this by choosing a node and then choosing **Query upstream lineage** or **Query downstream lineage**.

**Delete a feature group**

You can use the console or the Amazon SageMaker Feature Store API to delete your feature group. The instructions on using Feature Store through the console depends on if you have enabled Studio or Studio Classic as your default experience. For more information about the differences between the two, or how to change your default, see [Amazon SageMaker Studio](#).
The following sections provide an overview on how to delete a feature group.

Topics

- Delete a feature group using the console
- Delete feature group example Python code

Delete a feature group using the console

This section shows two ways to delete a feature group in the console, depending on your default experience: Studio or Studio Classic.

Delete feature group if Studio is your default experience (console)

1. Open the Studio console by following instructions in [Launch Amazon SageMaker Studio Classic](#).
2. Choose Data in the left navigation pane to expand the dropdown list.
3. From the dropdown list, choose Feature Store.
4. (Optional) To view your feature groups, choose My account. To view shared feature groups, choose Cross account.
5. In the Feature Group Catalog tab, choose the feature group to delete under Feature group name.
6. Choose Delete feature group.
7. In the pop-up window, confirm deletion by entering delete in the field, then choose Delete.

Delete feature group if Studio Classic is your default experience (console)

1. Open the Studio Classic console by following the instructions in [Launch Amazon SageMaker Studio Classic](#).
2. In the left navigation pane, choose the Home icon (🏠).
3. Choose Data.
4. From the dropdown list, choose Feature Store.
5. (Optional) To view your feature groups, choose My account. To view shared feature groups, choose Cross account.
6. In the **Feature Group Catalog** tab, choose the feature group to delete under **Feature group name**.

7. Choose **Delete feature group**.

8. In the pop-up window, confirm deletion by typing **delete** in the field, then choose **Delete**.

**Delete feature group example Python code**

The following code uses the [DeleteFeatureGroup] API operation to delete your feature group using the AWS SDK for Python (Boto3). It assumes that you've set up Feature Store and created a feature group. For more information about getting started, see [Introduction to Feature Store example notebook].

```python
import sagemaker
from sagemaker.feature_store.feature_group import FeatureGroup

sagemaker_session = sagemaker.Session()
fg_name = 'your-feature-group-name'
my_fg = FeatureGroup(name=fg_name, sagemaker_session=sagemaker_session)
my_fg.delete()
```

**Data sources and ingestion**

Records are added to your feature groups through ingestion. Depending on your desired use case, the ingested records may be kept within the feature group or not. This depends on the storage configuration, if your feature group uses the offline or online store. The offline store is used as a historical database, that is typically used for data exploration, machine learning (ML) model training, and batch inference. The online store is used as a real-time lookup of records, that is typically used for ML model serving. For more information on Feature Store concepts and ingestion, see [Feature Store concepts].

There are multiple ways to bring your data into Amazon SageMaker Feature Store. Feature Store offers a single API call for data ingestion called **PutRecord** that enables you to ingest data in batches or from streaming sources. You can use Amazon SageMaker Data Wrangler to engineer features and then ingest your features into your Feature Store. You can also use Amazon EMR for batch data ingestion through a Spark connector.

In the following topics we will discuss the difference between
Topics

- Stream ingestion
- Data Wrangler with Feature Store
- Batch ingestion with Amazon SageMaker Feature Store Spark

Stream ingestion

You can use streaming sources such as Kafka or Kinesis as a data source, where records are extracted from, and directly feed records to the online store for training, inference or feature creation. Records can be ingested into your feature group by using the synchronous PutRecord API call. Since this is a synchronous API call it allows small batches of updates to be pushed in a single API call. This enables you to maintain high freshness of the feature values and publish values as soon an update is detected. These are also called streaming features.

Data Wrangler with Feature Store

Data Wrangler is a feature of Studio Classic that provides an end-to-end solution to import, prepare, transform, featurize, and analyze data. Data Wrangler enables you to engineer your features and ingest them into your online or offline store feature groups.

The following instructions exports a Jupyter notebook that contains all of the source code needed to create a Feature Store feature group that adds your features from Data Wrangler to an online or offline store.

The instructions on exporting your Data Wrangler data flow to Feature Store on the console vary depending on whether you enabled enabled Amazon SageMaker Studio or Amazon SageMaker Studio Classic as your default experience.

Export your Data Wrangler data flow to Feature Store if Studio is your default experience (console)

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose Data from the left panel, to expand the dropdown list.
3. From the dropdown list, choose Data Wrangler.
4. If you have an instance of Amazon SageMaker Canvas already running, choose Open Canvas.
   
   If you don't have an instance of SageMaker Canvas running, choose Run in Canvas.
5. On the SageMaker Canvas console, choose **Data Wrangler** in the left navigation pane.
6. Choose **Data flows** to view your data flows.
7. Choose + to expand the dropdown list.
8. Choose **Export data flow** to expand the dropdown list.
9. Choose **Save to SageMaker Feature Store (via JupyterLab Notebook)**.
10. Under **Export data flow as notebook**, choose one of the following options:
   - **Download a local copy** to download the dataflow to your local machine.
   - **Export to S3 location** to download the dataflow to an Amazon Simple Storage Service location and enter the Amazon S3 location or choose **Browse** to find your Amazon S3 location.
11. Choose **Export**.

**Export your Data Wrangler data flow to Feature Store if Studio Classic is your default experience (console)**

1. Open the Studio Classic console by following the instructions in [Launch Amazon SageMaker Studio Classic](#).
2. Choose the **Home** icon
   - (in the left navigation pane.
3. Choose **Data**.
4. From the dropdown list, choose **Data Wrangler**.
5. Choose your workflow.
6. Choose the **Export** tab.
7. Choose **Export Step**.
8. Choose **Feature Store**.

After the feature group has been created, you can also select and join data across multiple feature groups to create new engineered features in Data Wrangler and then export your data set to an Amazon S3 bucket.

For more information on how to export to Feature Store, see [Export to SageMaker Feature Store](#).
Batch ingestion with Amazon SageMaker Feature Store Spark

Amazon SageMaker Feature Store Spark is a Spark connector that connects the Spark library to Feature Store. Feature Store Spark simplifies data ingestion from Spark DataFrames to feature groups. Feature Store supports batch data ingestion with Spark, using your existing ETL pipeline, on Amazon EMR, GIS, an AWS Glue job, an Amazon SageMaker Processing job, or a SageMaker notebook.

Methods for installing and implementing batch data ingestion are provided for Python and Scala developers. Python developers can use the open-source sagemaker-feature-store-pyspark Python library for local development, installation on Amazon EMR, and for Jupyter Notebooks by following the instructions in the Amazon SageMaker Feature Store Spark GitHub repository. Scala developers can use the Feature Store Spark connector available in the Amazon SageMaker Feature Store Spark SDK Maven central repository.

You can use the Spark connector to ingest data in the following ways, depending on if the online store, offline store, or both are enabled.

1. Ingest by default – If the online store is enabled, Spark connector first ingests your dataframe into the online store using the PutRecord API. Only the record with the largest event time remains in the online store. If the offline store is enabled, within 15 minutes Feature Store ingests your dataframe into the offline store. For more information about how the online and offline stores work, see Feature Store concepts.

   You can accomplish this by not specifying target_stores in the .ingest_data(...) method.

2. Offline store direct ingestion – If offline store is enabled, Spark connector batch ingests your dataframe directly into the offline store. Ingesting the dataframe directly into the offline store doesn't update the online store.

   You can accomplish this by setting target_stores=\"OfflineStore\" in the .ingest_data(...) method.

3. Online store only – If online store is enabled, Spark connector ingests your dataframe into the online store using the PutRecord API. Ingesting the dataframe directly into the online store doesn't update the offline store.

   You can accomplish this by setting target_stores=\"OnlineStore\" in the .ingest_data(...) method.
For information about using the different ingestion methods, see [Example implementations](#).

**Topics**
- [Feature Store Spark installation](#)
- [Retrieving the JAR for Feature Store Spark](#)
- [Example implementations](#)

**Feature Store Spark installation**

**Scala users**

The Feature Store Spark SDK is available in the [Amazon SageMaker Feature Store Spark SDK Maven central repository](#) for Scala users.

**Requirements**

- Spark >= 3.0.0 and <= 3.3.0
- iceberg-spark-runtime >= 0.14.0
- Scala >= 2.12.x
- Amazon EMR >= 6.1.0 (only if you are using Amazon EMR)

**Declare the dependency in POM.xml**

The Feature Store Spark connector has a dependency on the iceberg-spark-runtime library. You must therefore add corresponding version of the iceberg-spark-runtime library to the dependency if you're ingesting data into a feature group that you've auto-created with the Iceberg table format. For example, if you're using Spark 3.1, you must declare the following in your project's POM.xml:

```xml
<dependency>
  <groupId>software.amazon.sagemaker.featurestore</groupId>
  <artifactId>sagemaker-feature-store-spark-sdk_2.12</artifactId>
  <version>1.0.0</version>
</dependency>

<dependency>
  <groupId>org.apache.iceberg</groupId>
  <artifactId>iceberg-spark-runtime-3.1_2.12</artifactId>
</dependency>
```
Python users

The Feature Store Spark SDK is available in the open-source Amazon SageMaker Feature Store Spark GitHub repository.

Requirements

• Spark >= 3.0.0 and <= 3.3.0
• Amazon EMR >= 6.1.0 (only if you are using Amazon EMR)
• Kernel = conda_python3

We recommend setting the $SPARK_HOME to the directory where you have Spark installed. During installation, Feature Store uploads the required JAR to SPARK_HOME, so that the dependencies load automatically. Spark starting a JVM is required to make this PySpark library work.

Local installation

To find more info about the installation, enable verbose mode by appending --verbose to the following installation command.

```
pip3 install sagemaker-feature-store-pyspark-3.1 --no-binary :all:
```

Installation on Amazon EMR

Create an Amazon EMR cluster with the release version 6.1.0 or later. Enable SSH to help you troubleshoot any issues.

You can do one of the following to install the library:

• Create a custom step within Amazon EMR.
• Connect to your cluster using SSH and install the library from there.
Note
The following information uses Spark version 3.1, but you can specify any version that meets the requirements.

```
export SPARK_HOME=/usr/lib/spark
sudo -E pip3 install sagemaker-feature-store-pyspark-3.1 --no-binary :all: --verbose
```

Note
If you want to install the dependent JARs automatically to SPARK_HOME, do not use the bootstrap step.

Installation on a SageMaker notebook instance

Install a version of PySpark that's compatible with the Spark connector using the following commands:

```
!pip3 install pyspark==3.1.1
!pip3 install sagemaker-feature-store-pyspark-3.1 --no-binary :all:
```

If you're performing batch ingestion to the offline store, the dependencies aren't within the notebook instance environment.

```
from pyspark.sql import SparkSession
import feature_store_pyspark

extra_jars = ",".join(feature_store_pyspark.classpath_jars())

spark = SparkSession.builder \
  .config("spark.jars", extra_jars) \
  .config("spark.jars.packages", "org.apache.hadoop:hadoop-aws:3.2.1,org.apache.hadoop:hadoop-common:3.2.1") \
  .getOrCreate()
```
Installation on notebooks with GIS

⚠️ Important
You must use AWS Glue Version 2.0 or later.

Use the following information to help you install the PySpark connector in an AWS Glue Interactive Session (GIS).

Amazon SageMaker Feature Store Spark requires a specific Spark connector JAR during the initialization of the session to be uploaded to your Amazon S3 bucket. For more information on uploading the required JAR to your S3 bucket, see [Retrieving the JAR for Feature Store Spark](#).

After you’ve uploaded the JAR, you must provide the GIS sessions with the JAR using the following command.

```
%extra_jars s3://<YOUR_BUCKET>/spark-connector-jars/sagemaker-feature-store-spark-sdk.jar
```

To install Feature Store Spark in the AWS Glue runtime, use the `%additional_python_modules` magic command within the GIS notebook. AWS Glue runs `pip` to the modules that you’ve specified under `%additional_python_modules`.

```
%additional_python_modules sagemaker-feature-store-pyspark-3.1
```

Before you start the AWS Glue session, you must use both of the preceding magic commands.

Installation on an AWS Glue job

⚠️ Important
You must use AWS Glue Version 2.0 or later.

To install the Spark connector on a AWS Glue job, use the `--extra-jars` argument to provide the necessary JARs and `--additional-python-modules` to install the Spark Connector as job parameters when you create the AWS Glue job as shown in the following example. For more
information on uploading the required JAR to your S3 bucket, see [Retrieving the JAR for Feature Store Spark](#).

```python
glue_client = boto3.client('glue', region_name=region)  
response = glue_client.create_job(  
    Name=pipeline_id,  
    Description='Feature Store Compute Job',  
    Role=glue_role_arn,  
    ExecutionProperty={'MaxConcurrentRuns': max_concurrent_run},  
    Command={  
        'Name': 'glueetl',  
        'ScriptLocation': script_location_uri,  
        'PythonVersion': '3'  
    },  
    DefaultArguments={  
        '--TempDir': temp_dir_location_uri,  
        '--additional-python-modules': 'sagemaker-feature-store-pyspark-3.1',  
        '--extra-jars': 's3:/<YOUR_BUCKET>/spark-connector-jars/sagemaker-feature-store-spark-sdk.jar',  
        ...  
    },  
    MaxRetries=3,  
    NumberOfWorkers=149,  
    Timeout=2880,  
    GlueVersion='3.0',  
    WorkerType='G.2X'  
)
```

**Installation on an Amazon SageMaker Processing job**

To use Feature Store Spark with Amazon SageMaker Processing jobs, bring your own image. For more information about bringing your image, see [Bring your own SageMaker image](#). Add the installation step to a Dockerfile. After you’ve pushed the Docker image to an Amazon ECR repository, you can use the PySparkProcessor to create the processing job. For more information about creating a processing job with the PySpark processor, see [Data Processing with Apache Spark](#).

The following is an example of adding an installation step to the Dockerfile.

```bash
FROM <ACCOUNT_ID>.dkr.ecr.<AWS_REGION>.amazonaws.com/sagemaker-spark-processing:3.1-cpu-py38-v1.0
```
**Retrieving the JAR for Feature Store Spark**

To retrieve the Feature Store Spark dependency JAR, you must install the Spark connector from the Python Package Index (PyPI) repository using `pip` in any Python environment with network access. A SageMaker Jupyter Notebook is an example of a Python environment with network access.

The following command installs the Spark connector.

```bash
!pip install sagemaker-feature-store-pyspark-3.1
```

After you've installed Feature Store Spark, you can retrieve the JAR location and upload the JAR to Amazon S3.

The `feature-store-pyspark-dependency-jars` command provides the location of the necessary dependency JAR that Feature Store Spark added. You can use the command to retrieve the JAR and upload it to Amazon S3.

```python
jar_location = !feature-store-pyspark-dependency-jars
jar_location = jar_location[0]

s3_client = boto3.client("s3")
s3_client.upload_file(jar_location, "<YOUR_BUCKET>" ,"spark-connector-jars/sagemaker-feature-store-spark-sdk.jar")
```

**Example implementations**

Example Python script

`FeatureStoreBatchIngestion.py`

```python
from pyspark.sql import SparkSession
from feature_store_pyspark.FeatureStoreManager import FeatureStoreManager
import feature_store_pyspark
```
spark = SparkSession.builder.getOrCreate()

# Construct test DataFrame
columns = ["RecordIdentifier", "EventTime"]

df = spark.createDataFrame(data).toDF(*columns)

# Initialize FeatureStoreManager with a role arn if your feature group is created by
# another account
feature_store_manager = FeatureStoreManager("arn:aws:iam::111122223333:role/role-arn")

# Load the feature definitions from input schema. The feature definitions can be
# used to create a feature group
feature_definitions = feature_store_manager.load_feature_definitions_from_schema(df)

feature_group_arn = "arn:aws:sagemaker:<AWS_REGION>:<ACCOUNT_ID>:feature-group/<YOUR_FEATURE_GROUP_NAME>"

# Ingest by default. The connector will leverage PutRecord API to ingest your data
# in stream
# https://docs.aws.amazon.com/sagemaker/latest/APIReference/
# API_feature_store_PutRecord.html
feature_store_manager.ingest_data(input_data_frame=df,
   feature_group_arn=feature_group_arn)

# To select the target stores for ingestion, you can specify the target store as the
# parameter
# If OnlineStore is selected, the connector will leverage PutRecord API to ingest
# your data in stream
feature_store_manager.ingest_data(input_data_frame=df,
   feature_group_arn=feature_group_arn, target_stores=["OfflineStore", "OnlineStore"])

# If only OfflineStore is selected, the connector will batch write the data to
# offline store directly
feature_store_manager.ingest_data(input_data_frame=df,
   feature_group_arn=feature_group_arn, target_stores=["OfflineStore"])

# To retrieve the records failed to be ingested by spark connector
failed_records_df = feature_store_manager.get_failed_stream_ingestion_data_frame()
Submit a Spark job with example Python script

The PySpark version requires an extra dependent JAR to be imported, so extra steps are needed to run the Spark application.

If you did not specify `SPARK_HOME` during installation, then you have to load required JARs in JVM when running `spark-submit feature-store-pyspark-dependency-jars` is a Python script installed by the Spark library to automatically fetch the path to all JARs for you.

```
spark-submit --jars `feature-store-pyspark-dependency-jars` FeatureStoreBatchIngestion.py
```

If you are running this application on Amazon EMR, we recommended that you run the application in client mode, so that you do not need to distribute the dependent JARs to other task nodes. Add one more step in Amazon EMR cluster with Spark argument similar to the following:

```
spark-submit --deploy-mode client --master yarn s3://<PATH_TO_SCRIPT>/ FeatureStoreBatchIngestion.py
```

Example Scala script

`FeatureStoreBatchIngestion.scala`

```scala
import software.amazon.sagemaker.featurestore.sparksdk.FeatureStoreManager
import org.apache.spark.sql.types.{StringType, StructField, StructType}
import org.apache.spark.sql.{Row, SparkSession}

object TestSparkApp {
  def main(args: Array[String]): Unit = {
    val spark = SparkSession.builder().getOrCreate()

    // Construct test DataFrame
    val data = List(
      Row("1", "2021-07-01T12:20:12Z"),
      Row("2", "2021-07-02T13:30:00Z"),
      Row("3", "2021-07-03T14:45:15Z"),
      Row("4", "2021-07-04T15:55:30Z"),
      Row("5", "2021-07-05T16:05:45Z")
    )

    // Create DataFrame from test data
    val df = spark.createDataFrame(data).
    .withColumn("id", new ColumnString("id", StringType)).
    .withColumn("timestamp", new ColumnString("timestamp", StringType)).

    // Save DataFrame to S3
    df.write.saveAsTable("feature-store-batch-ingestion")
  }
}
```
Row("2", "2021-07-02T12:20:13Z"),
Row("3", "2021-07-03T12:20:14Z")
)

val schema = StructType(
    List(StructField("RecordIdentifier", StringType), StructField("EventTime", StringType))
)

val df = spark.createDataFrame(spark.sparkContext.parallelize(data), schema)

// Initialize FeatureStoreManager with a role arn if your feature group is created by another account
val featureStoreManager = new FeatureStoreManager("arn:aws:iam::111122223333:role/role-arn")

// Load the feature definitions from input schema. The feature definitions can be used to create a feature group
val featureDefinitions = featureStoreManager.loadFeatureDefinitionsFromSchema(df)


// Ingest by default. The connector will leverage PutRecord API to ingest your data in stream
featureStoreManager.ingestData(df, featureGroupArn)

// To select the target stores for ingestion, you can specify the target store as the parameter
// If OnlineStore is selected, the connector will leverage PutRecord API to ingest your data in stream
featureStoreManager.ingestData(df, featureGroupArn, List("OfflineStore", "OnlineStore"))

// If only OfflineStore is selected, the connector will batch write the data to offline store directly
featureStoreManager.ingestData(df, featureGroupArn, ["OfflineStore"])

// To retrieve the records failed to be ingested by spark connector
val failedRecordsDf = featureStoreManager.getFailedStreamIngestionDataFrame()
Submit a Spark job

Scala

You should be able to use Feature Store Spark as a normal dependency. No extra instruction is needed to run the application on all platforms.

Feature Processing

Amazon SageMaker Feature Store Feature Processing is a capability with which you can transform raw data into machine learning (ML) features. It provides you with a Feature Processor SDK with which you can transform and ingest data from batch data sources into your feature groups. With this capability, Feature Store takes care of the underlying infrastructure including provisioning the compute environments and creating and maintaining SageMaker Pipelines to load and ingest data. This way you can focus on your feature processor definitions that includes a transformation function (for example, count of product views, mean of transaction value), sources (where to apply this transformation on), and sinks (where to write the computed feature values to).

Feature Processor pipeline is a SageMaker Pipelines pipeline. As a SageMaker Pipelines, you can also track scheduled Feature Processor pipelines with SageMaker lineage in the console. For more information on SageMaker Lineage, see Amazon SageMaker ML Lineage Tracking This includes tracking scheduled executions, visualizing lineage to trace features back to their data sources, and viewing shared feature processors in a single environment. For information on using Feature Store with the console, see View pipeline executions from the console.

Topics

- Feature Store Feature Processor SDK
- Running Feature Store Feature Processor remotely
- Creating and running Feature Store Feature Processor pipelines
- Scheduled and event based executions for Feature Processor pipelines
- Monitor Amazon SageMaker Feature Store Feature Processor pipelines
- IAM permissions and execution roles
- Feature Processor restrictions, limits, and quotas
- Data sources
Feature Store Feature Processor SDK

Declare a Feature Store Feature Processor definition by decorating your transformation functions with the @feature_processor decorator. The SageMaker SDK for Python (Boto3) automatically loads data from the configured input data sources, applies the decorated transformation function, and then ingests the transformed data to a target feature group. Decorated transformation functions must conform to the expected signature of the @feature_processor decorator. For more information about the @feature_processor decorator, see @feature_processor Decorator in the Amazon SageMaker Feature Store Read the Docs.

With the @feature_processor decorator, your transformation function runs in a Spark runtime environment where the input arguments provided to your function and its return value are Spark DataFrames. The number of input parameters in your transformation function must match the number of inputs configured in the @feature_processor decorator.

For more information on the @feature_processor decorator, see the Feature Processor Feature Store SDK for Python (Boto3).

The following code are basic examples on how to use the @feature_processor decorator. For more specific example usage cases, see Example Feature Processing code for common use cases.

The Feature Processor SDK can be installed from the SageMaker Python SDK and its extras using the following command.

```
pip install sagemaker[feature-processor]
```

In the following examples, us-east-1 is the region of the resource, 111122223333 is the resource owner account ID, and your-feature-group-name is the feature group name.

The following is a basic feature processor definition, where the @feature_processor decorator configures a CSV input from Amazon S3 to be loaded and provided to your transformation function (for example, transform), and prepares it for ingestion to a feature group. The last line runs it.

```
from sagemaker.feature_store.feature_processor import CSVDataSource, feature_processor

CSV_DATA_SOURCE = CSVDataSource('s3://your-bucket/prefix-to-csv/')
```
@feature_processor(inputs=[CSV_DATA_SOURCE], output=OUTPUT_FG)
def transform(csv_input_df):
    return csv_input_df
transform()

The @feature_processor parameters include:

- **inputs** (List[str]): A list of data sources that are used in your Feature Store Feature Processor. If your data sources are feature groups or stored in Amazon S3 you may be able to use Feature Store provided data source definitions for feature processor. For a full list of Feature Store provided data source definitions, see the Feature Processor Data Source in the Amazon SageMaker Feature Store Read the Docs.

- **output** (str): The ARN of the feature group to ingest the output of the decorated function.

- **target_stores** (Optional[List[str]]): A list of stores (for example, OnlineStore or OfflineStore) to ingest to the output. If unspecified, data is ingested to all of the output feature group's enabled stores.

- **parameters** (Dict[str, Any]): A dictionary to be provided to your transformation function.

- **enable_ingestion** (bool): A flag to indicate whether the transformation function's outputs are ingested to the output feature group. This flag is useful during the development phase. If unspecified, ingestion is enabled.

Optional wrapped function parameters (provided as an argument if provided in the function signature) include:

- **params** (Dict[str, Any]): The dictionary defined in the @feature_processor parameters. It also contains system configured parameters that can be referenced with the key system, such as the scheduled_time parameter.

- **spark** (SparkSession): A reference to the SparkSession instance initialized for the Spark Application.

The following code is an example of using the params and spark parameters.

```python
from sagemaker.feature_store.feature_processor import CSVDataSource, feature_processor
CSV_DATA_SOURCE = CSVDataSource('s3://your-bucket/prefix-to-csv/')
```
The scheduled_time system parameter (provided in the params argument to your function) is an important value to support retrying each execution. The value can help to uniquely identify the Feature Processor's execution and can be used as a reference point for daterange–based inputs (for example, only loading the last 24 hours worth of data) to guarantee the input range independent of the code's actual execution time. If the Feature Processor runs on a schedule (see Scheduled and event based executions for Feature Processor pipelines) then its value is fixed to the time it is scheduled to run. The argument can be overridden during synchronous execution using the SDK's execute API to support use cases such as data backfills or re-running a missed past execution. Its value is the current time if the Feature Processor runs any other way.

For information about authoring Spark code, see the Spark SQL Programming Guide.

For more code samples for common use-cases, see the Example Feature Processing code for common use cases.

Note that transformation functions decorated with @feature_processor do not return a value. To programmatically test your function, you can remove or monkey patch the @feature_processor decorator such that it acts as a pass-through to the wrapped function.

For more details on the @feature_processor decorator, see Amazon SageMaker Feature Store Python SDK.

## Running Feature Store Feature Processor remotely

To run your Feature Processors on large data sets that require hardware more powerful than what is locally available, you can decorate your code with the @remote decorator to run your local
Python code as a single or multi-node distributed SageMaker training job. For more information on running your code as a SageMaker training job, see Run your local code as a SageMaker training job.

The following is a usage example of the @remote decorator along with the @feature_processor decorator.

```python
from sagemaker.remote_function.spark_config import SparkConfig
from sagemaker.remote_function import remote
from sagemaker.feature_store.feature_processor import CSVDataSource, feature_processor

CSV_DATA_SOURCE = CSVDataSource('s3://bucket/prefix-to-csv/)

@remote(
    spark_config=SparkConfig(),
    instance_type="ml.m5.2xlarge",
    dependencies="/local/requirements.txt"
)
@feature_processor(
    inputs=[CSV_DATA_SOURCE],
    output=OUTPUT_FG,
)
def transform(csv_input_df):
    return csv_input_df

transform()
```

The `spark_config` parameter indicates that the remote job runs as a Spark application. The `SparkConfig` instance can be used to configure the Spark Configuration and provide additional dependencies to the Spark application such as Python files, JARs, and files.

For faster iterations when developing your feature processing code, you can specify the `keep_alive_period_in_seconds` argument in the `@remote` decorator to retain configured resources in a warm pool for subsequent training jobs. For more information on warm pools, see `KeepAlivePeriodInSeconds` in the API Reference guide.

The following code is an example of local requirements.txt:

```
sagemaker>=2.167.0
```
This will install the corresponding SageMaker SDK version in remote job which is required for executing the method annotated by @feature-processor.

Creating and running Feature Store Feature Processor pipelines

The Feature Processor SDK provides APIs to promote your Feature Processor Definitions into a fully managed SageMaker Pipeline. For more information on SageMaker Pipelines, see SageMaker Pipelines Overview. To convert your Feature Processor Definitions in to a SageMaker Pipeline, use the to_pipeline API with your Feature Processor definition. You can schedule executions of your Feature Processor Definition can be scheduled, operationally monitor them with CloudWatch metrics, and integrate them with EventBridge to act as event sources or subscribers. For more information about monitoring pipelines created with SageMaker Pipelines, see Monitor Amazon SageMaker Feature Store Feature Processor pipelines.

To view your Feature Processor pipelines, see View pipeline executions from the console.

If your function is also decorated with the @remote decorator, then its configurations is carried over to the Feature Processor pipeline. You can specify advanced configurations such as compute instance type and count, runtime dependencies, network and security configurations using the @remote decorator.

The following example uses the to_pipeline and execute APIs.

```python
from sagemaker.feature_store.feature_processor import (  
    execute, to_pipeline, describe, TransformationCode
)

pipeline_name="feature-processor-pipeline"

pipeline_arn = to_pipeline(  
    pipeline_name=pipeline_name,  
    step=transform,  
    transformation_code=TransformationCode(s3_uri="s3://bucket/prefix"),
)

pipeline_execution_arn = execute(  
    pipeline_name=pipeline_name
)
```

The to_pipeline API is semantically an upsert operation. It updates the pipeline if it already exists; otherwise, it creates a pipeline.
The `to_pipeline` API optionally accepts an Amazon S3 URI that references a file containing the Feature Processor definition to associate it with the Feature Processor pipeline to track the transformation function and its versions in its SageMaker machine learning lineage.

To retrieve a list of every Feature Processor pipeline in your account, you can use the `list_pipelines` API. A subsequent request to the `describe` API returns details related to the Feature Processor pipeline including, but not limited to, SageMaker Pipelines and schedule details.

The following example uses the `list_pipelines` and `describe` APIs.

```python
from sagemaker.feature_store.feature_processor import list_pipelines, describe

feature_processor_pipelines = list_pipelines()

pipeline_description = describe(
    pipeline_name = feature_processor_pipelines[0]
)
```

## Scheduled and event based executions for Feature Processor pipelines

Amazon SageMaker Feature Store Feature Processing pipeline executions can be configured to start automatically and asynchronously based on a preconfigured schedule or as a result of another AWS service event. For example, you can schedule Feature Processing pipelines to execute on the first of every month or chain two pipelines together so that a target pipeline is executed automatically after a source pipeline execution completes.

### Topics

- [Schedule based executions](#)
- [Event based executions](#)

### Schedule based executions

The Feature Processor SDK provides a `schedule` API to run Feature Processor pipelines on a recurring basis with Amazon EventBridge Scheduler integration. The schedule can be specified with an `at`, `rate`, or `cron` expression using the `ScheduleExpression` parameter with the same expressions supported by Amazon EventBridge. The schedule API is semantically an upsert operation in that it updates the schedule if it already exists; otherwise, it creates it. For more information on the EventBridge expressions and examples, see [Schedule types on EventBridge Scheduler](#) in the EventBridge Scheduler User Guide.
The following examples use the Feature Processor schedule API, using the at, rate, and cron expressions.

```python
from sagemaker.feature_store.feature_processor import schedule
pipeline_name='feature-processor-pipeline'

event_bridge_schedule_arn = schedule(
    pipeline_name=pipeline_name,
    schedule_expression="at(2020-11-30T00:00:00)"
)

event_bridge_schedule_arn = schedule(
    pipeline_name=pipeline_name,
    schedule_expression="rate(24 hours)"
)

event_bridge_schedule_arn = schedule(
    pipeline_name=pipeline_name,
    schedule_expression="cron(0 0-23/1 ? * * 2023-2024)"
)
```

The default timezone for date and time inputs in the schedule API are in UTC. For more information about EventBridge Scheduler schedule expressions, see ScheduleExpression in the EventBridge Scheduler API Reference documentation.

Scheduled Feature Processor pipeline executions provide your transformation function with the scheduled execution time, to be used as an idempotency token or a fixed reference point for date range–based inputs. To disable (i.e., pause) or re-enable a schedule, use the state parameter of the schedule API with ‘DISABLED’ or ‘ENABLED’, respectively.

For information about Feature Processor, see Feature Processor SDK data sources.

**Event based executions**

A Feature Processing pipeline can be configured to automatically execute when an AWS event occurs. The Feature Processing SDK provides a put_trigger function that accepts a list of source events and a target pipeline. The source events must be instances of FeatureProcessorPipelineEvent, that specifies a pipeline and execution status events.
The `put_trigger` function configures an Amazon EventBridge rule and target to route events and allows you to specify an EventBridge event pattern to respond to any AWS event. For information on these concepts, see Amazon EventBridge rules, targets, and event patterns.

Triggers can be enabled or disabled. EventBridge will start a target pipeline execution using the role provided in the `role_arn` parameter of the `put_trigger` API. The execution role is used by default if the SDK is used in a Amazon SageMaker Studio Classic or Notebook environment. For information on how to get your execution role, see Get execution role.

The following example sets up:

- A SageMaker Pipeline using the `to_pipeline` API, that takes in your target pipeline name (target-pipeline) and your transformation function (transform). For information on your Feature Processor and transform function, see Feature Processor SDK data sources.

- A trigger using the `put_trigger` API, that takes in FeatureProcessorPipelineEvent for the event and your target pipeline name (target-pipeline).

The `FeatureProcessorPipelineEvent` defines the trigger for when the status of your source pipeline (source-pipeline) becomes Succeeded. For information on the Feature Processor Pipeline event function, see FeatureProcessorPipelineEvent in the Feature Store Read the Docs.

```python
from sagemaker.feature_store.feature_processor import put_trigger, to_pipeline, FeatureProcessorPipelineEvent
to_pipeline(pipeline_name="target-pipeline", step=transform)
put_trigger(
    source_pipeline_events=[
        FeatureProcessorPipelineEvent(
            pipeline_name="source-pipeline",
            status=["Succeeded"]
        )
    ],
    target_pipeline="target-pipeline"
)
```
For an example of using event based triggers to create continuous executions and automatic retries for your Feature Processor pipeline, see Continuous executions and automatic retries using event based triggers.

For an example of using event based triggers to create continuous streaming and automatic retries using event based triggers, see Streaming custom data source examples.

**Monitor Amazon SageMaker Feature Store Feature Processor pipelines**

AWS provides monitoring tools to watch your Amazon SageMaker resources and applications in real time, report when something goes wrong, and take automatic actions when appropriate. Feature Store Feature Processor pipelines are SageMaker Pipelines, so the standard monitoring mechanisms and integrations are available. Operational metrics such as execution failures can be monitored via Amazon CloudWatch metrics and Amazon EventBridge events.

For more information on how to monitor and operationalize Feature Store Feature Processor, see the following resources:

- Monitor AWS resources provisioned while using Amazon SageMaker - General guidance on monitoring and auditing activity for SageMaker resources.
- SageMaker Pipelines Metrics - CloudWatch Metrics emitted by SageMaker Pipelines.
- Pipeline execution state change - EventBridge events emitted for SageMaker Pipelines and executions.
- Troubleshooting Amazon SageMaker Model Building Pipelines - General debugging and troubleshooting tips for SageMaker Pipelines.

Feature Store Feature Processor execution logs can be found in Amazon CloudWatch Logs under the /aws/sagemaker/TrainingJobs log group, where you can find the execution log streams using lookup conventions. For executions created by directly invoking the @feature_processor decorated function, you can find logs in your local execution environment’s console. For @remote decorated executions, the CloudWatch Logs stream name contains the name of the function and the execution timestamp. For Feature Processor pipeline executions, the CloudWatch Logs stream for the step contains the feature-processor string and the pipeline execution ID.

Feature Store Feature Processor pipelines and recent execution statuses can be found in Amazon SageMaker Studio Classic for a given feature group in the Feature Store UI. Feature groups related to the Feature Processor pipelines as either inputs or outputs are displayed in the UI. In addition,
the lineage view can provide context into upstream executions, such as data producing Feature Processor pipelines and data sources, for further debugging. For more information on using the lineage view using Studio Classic, see View lineage from the console.

**IAM permissions and execution roles**

To use the The Amazon SageMaker Python SDK requires permissions to interact with AWS services. The following policies are required for full Feature Processor functionality. You can attach the AmazonSageMakerFullAccess and AmazonEventBridgeSchedulerFullAccess AWS Managed Policies attached to your IAM role. For information on attaching policies to your IAM role, see Adding policies to your IAM role. See the following examples for details.

The trust policy of the role to which this policy is applied must allow the "scheduler.amazonaws.com", "sagemaker.amazonaws.com", and "glue.amazonaws.com" principles.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "",
            "Effect": "Allow",
            "Principal": {
                "Service": [
                    "scheduler.amazonaws.com",
                    "sagemaker.amazonaws.com",
                    "glue.amazonaws.com"
                ],
            },
            "Action": "sts:AssumeRole"
        }
    ]
}
```

**Feature Processor restrictions, limits, and quotas**

Amazon SageMaker Feature Store Feature Processing relies on SageMaker machine learning (ML) lineage tracking. The Feature Store Feature Processor uses lineage contexts to represent and track Feature Processing Pipelines and Pipeline versions. Each Feature Store Feature Processor consumes at least two lineage contexts (one for the Feature Processing Pipeline and another for the version). If the input or output data source of a Feature Processing Pipeline changes, an additional lineage
context is created. You can update SageMaker ML lineage limits by reaching out to AWS support for a limit increase. Default limits for resources used by Feature Store Feature Processor are as follows. For information on SageMaker ML lineage tracking, see Amazon SageMaker ML Lineage Tracking.

For more information on SageMaker quotas, see Amazon SageMaker endpoints and quotas.

Lineage limits per Region

- Contexts – 500 (soft limit)
- Artifacts – 6,000 (soft limit)
- Associations – 6,000 (soft limit)

Training Limits per Region

- Longest run time for a training job – 432,000 seconds
- Maximum number of instances per training job – 20
- The maximum number of CreateTrainingJob requests that you can make, per second, in this account in the current Region – 1 TPS
- Keep alive period for cluster reuse – 3,600 seconds

Maximum number of Pipelines and concurrent pipeline executions per Region

- Maximum number of pipelines allowed per account – 500
- Maximum number of concurrent pipeline executions allowed per account – 20
- Time at which pipeline executions time out – 672 hours

Data sources

Amazon SageMaker Feature Store Feature Processing supports multiple data sources. The Feature Processor SDK for Python (Boto3) provides constructs to load data from feature groups or objects stored in Amazon S3. In addition, you can author custom data sources to load data from other data sources. For information about Feature Store provided data sources, see Feature Processor data source Feature Store Python SDK.

Topics

- Feature Processor SDK data sources
Feature Processor SDK data sources

The Amazon SageMaker Feature Store Feature Processor SDK for Python (Boto3) provides constructs to load data from feature groups or objects stored in Amazon S3. For a full list of Feature Store provided data source definitions, see the Feature Processor data source Feature Store Python SDK.

For examples on how to use the Feature Store Python SDK data source definitions, see Example Feature Processing code for common use cases.

FeatureGroupDataSource

The FeatureGroupDataSource is used to specify a feature group as an input data source for a Feature Processor. Data can be loaded from an offline store feature group. Attempting to load your data from an online store feature group will result in a validation error. You can specify start and end offsets to limit the data that is loaded to a specific time range. For example, you can specify a start offset of ‘14 days’ to load only the last two weeks of data, and you can additionally specify an end offset of ‘7 days’ to limit the input to the previous week of data.

Feature Store provided data source definitions

The Feature Store Python SDK contain data source definitions that can be used to specify various input data sources for a Feature Processor. These include CSV, Parquet, and Iceberg table sources. For a full list of Feature Store provided data source definitions, see the Feature Processor data source Feature Store Python SDK.

Custom data sources

On this page we will describe how to create a custom data source class and show some usage examples. With custom data sources, you can use the SageMaker SDK for Python (Boto3) provided APIs in the same way as if you are using Amazon SageMaker Feature Store provided data sources.

To use a custom data source to transform and ingest data into a feature group using Feature Processing, you will need to extend the PySparkDataSource class with the following class members and function.
• **data_source_name** (str): an arbitrary name for the data source. For example, Amazon Redshift, Snowflake, or a Glue Catalog ARN.

• **data_source_unique_id** (str): a unique identifier that refers to the specific resource being accessed. For example, table name, DDB Table ARN, Amazon S3 prefix. All usage of the same **data_source_unique_id** in custom data sources will be associated to the same data source in the lineage view. Lineage includes information about the execution code of a feature processing workflow, what data sources were used, and how they are ingested into the feature group or feature. For information about viewing lineage of a feature group in **Studio**, see [View lineage from the console](#).

• **read_data** (func): a method used to connect with the feature processor. Returns a Spark data frame. For examples, see [Custom data source examples](#).

Both **data_source_name** and **data_source_unique_id** are used to uniquely identify your lineage entity. The following is an example for a custom data source class named CustomDataSource.

```python
from sagemaker.feature_store.feature_processor import PySparkDataSource
from pyspark.sql import DataFrame

class CustomDataSource(PySparkDataSource):
    data_source_name = "custom-data-source-name"
    data_source_unique_id = "custom-data-source-id"

    def read_data(self, parameter, spark) -> DataFrame:
        your own code here to read data into a Spark dataframe
        return dataframe
```

### Custom data source examples

This section provides examples of custom data sources implementations for Feature Processors. For more information on custom data sources, see [Custom data sources](#).

Security is a shared responsibility between AWS and our customers. AWS is responsible for protecting the infrastructure that runs the services in the AWS Cloud. Customers are responsible for all of their necessary security configuration and management tasks. For example, secrets such as access credentials to data stores should not be hard coded in your custom data sources. You can use AWS Secrets Manager to manage these credentials. For information about Secrets Manager, see...
What is AWS Secrets Manager? in the AWS Secrets Manager user guide. The following examples will use Secrets Manager for your credentials.

Topics

- Amazon Redshift Clusters (JDBC) custom data source examples
- Snowflake custom data source examples
- Databricks (JDBC) custom data source examples
- Streaming custom data source examples

Amazon Redshift Clusters (JDBC) custom data source examples

Amazon Redshift offers a JDBC driver that can be used to read data with Spark. For information about how to download the Amazon Redshift JDBC driver, see Download the Amazon Redshift JDBC driver, version 2.1.

To create the custom Amazon Redshift data source class, you will need to overwrite the read_data method from the Custom data sources.

To connect with an Amazon Redshift cluster you need your:

- Amazon Redshift JDBC URL (jdbc-url)

  For information about obtaining your Amazon Redshift JDBC URL, see Getting the JDBC URL in the Amazon Redshift Database Developer Guide.

- Amazon Redshift user name (redshift-user) and password (redshift-password)

  For information about how to create and manage database users using the Amazon Redshift SQL commands, see Users in the Amazon Redshift Database Developer Guide.

- Amazon Redshift table name (redshift-table-name)

  For information about how to create a table with some examples, see CREATE TABLE in the Amazon Redshift Database Developer Guide.

- (Optional) If using Secrets Manager, you'll need the secret name (secret-redshift-account-info) where you store your Amazon Redshift access username and password on Secrets Manager.

  For information about Secrets Manager, see Find secrets in AWS Secrets Manager in the AWS Secrets Manager User Guide.
• AWS Region (*your-region*)

For information about obtaining your current session's region name using SDK for Python (Boto3), see `region_name` in the Boto3 documentation.

The following example demonstrates how to retrieve the JDBC URL and personal access token from Secrets Manager and override the `read_data` for your custom data source class, `DatabricksDataSource`.

```python
from sagemaker.feature_store.feature_processor import PySparkDataSource
import json
import boto3

class RedshiftDataSource(PySparkDataSource):
    data_source_name = "Redshift"
    data_source_unique_id = "redshift-resource-arn"

    def read_data(self, spark, params):
        url = "jdbc-url?user=redshift-user&password=redshift-password"
        aws_iam_role_arn = "redshift-command-access-role"
        secret_name = "secret-redshift-account-info"
        region_name = "your-region"

        session = boto3.session.Session()
        sm_client = session.client(
            service_name='secretsmanager',
            region_name=region_name,
        )

        secrets = json.loads(sm_client.get_secret_value(SecretId=secret_name)['SecretString'])
        jdbc_url = url.replace("jdbc-url", secrets['jdbcurl']).replace("redshift-user", secrets['username']).replace("redshift-password", secrets['password'])

        return spark.read \
            .format("jdbc") \
            .option("url", url) \
            .option("driver", "com.amazon.redshift.Driver") \
            .option("dbtable", "redshift-table-name") \
            .option("tempdir", "s3a://your-bucket-name/your-bucket-prefix") \
```

Data sources
The following example shows how to connect RedshiftDataSource to your feature_processor decorator.

```python
from sagemaker.feature_store.feature_processor import feature_processor

@feature_processor(
    inputs=[RedshiftDataSource()],
    output="feature-group-arn",
    target_stores=["OfflineStore"],
    spark_config={"spark.jars.packages": "com.amazon.redshift:redshift-jdbc42:2.1.0.16"}
)
def transform(input_df):
    return input_df
```

To run the feature processor job remotely, you need to provide the jdbc driver by defining SparkConfig and pass it to the @remote decorator.

```python
from sagemaker.remote_function import remote
from sagemaker.remote_function.spark_config import SparkConfig

config = {
    "Classification": "spark-defaults",
    "Properties": {
        "spark.jars.packages": "com.amazon.redshift:redshift-jdbc42:2.1.0.16"
    }
}

@remote(
    spark_config=SparkConfig(configuration=config),
    instance_type="ml.m5.2xlarge",
)
@feature_processor(
    inputs=[RedshiftDataSource()],
    output="feature-group-arn",
    target_stores=["OfflineStore"],
)
def transform(input_df):
    return input_df
```
Snowflake custom data source examples

Snowflake provides a Spark connector that can be used for your `feature_processor` decorator. For information about Snowflake connector for Spark, see [Snowflake Connector for Spark](#) in the Snowflake documentation.

To create the custom Snowflake data source class, you will need to override the `read_data` method from the [Custom data sources](#) and add the Spark connector packages to the Spark classpath.

To connect with a Snowflake data source you need:

- **Snowflake URL** *(sf-url)*
  
  For information about URLs for accessing Snowflake web interfaces, see [Account Identifiers](#) in the Snowflake documentation.

- **Snowflake database** *(sf-database)*
  
  For information about obtaining the name of your database using Snowflake, see `CURRENT_DATABASE` in the Snowflake documentation.

- **Snowflake database schema** *(sf-schema)*
  
  For information about obtaining the name of your schema using Snowflake, see `CURRENT_SCHEMA` in the Snowflake documentation.

- **Snowflake warehouse** *(sf-warehouse)*
  
  For information about obtaining the name of your warehouse using Snowflake, see `CURRENT_WAREHOUSE` in the Snowflake documentation.

- **Snowflake table name** *(sf-table-name)*

- **(Optional) If using Secrets Manager, you'll need the secret name** *(secret-snowflake-account-info)* where you store your Snowflake access username and password on Secrets Manager.

  For information about Secrets Manager, see [Find secrets in AWS Secrets Manager](#) in the AWS Secrets Manager User Guide.

- **AWS Region** *(your-region)*

  For information about obtaining your current session’s region name using SDK for Python (Boto3), see `region_name` in the Boto3 documentation.
The following example demonstrates how to retrieve the Snowflake user name and password from Secrets Manager and override the read_data function for your custom data source class SnowflakeDataSource.

class SnowflakeDataSource(PySparkDataSource):
    sf_options = {
        "sfUrl" : "sf-url",
        "sfDatabase" : "sf-database",
        "sfSchema" : "sf-schema",
        "sfWarehouse" : "sf-warehouse",
    }
    data_source_name = "Snowflake"
    data_source_unique_id = "sf-url"

    def read_data(self, spark, params):
        secret_name = "secret-snowflake-account-info"
        region_name = "your-region"

        session = boto3.session.Session()
        sm_client = session.client(
            service_name='secretsmanager',
            region_name=region_name,
        )

        secrets = json.loads(sm_client.get_secret_value(SecretId=secret_name)["SecretString"])
        self.sf_options["sfUser"] = secrets.get("username")
        self.sf_options["sfPassword"] = secrets.get("password")

        return spark.read.format("net.snowflake.spark.snowflake") \\.
            .options(**self.sf_options) \\.
            .option("dbtable", "sf-table-name") \\.
            .load()
The following example shows how to connect SnowflakeDataSource to your feature_processor decorator.

```python
from sagemaker.feature_store.feature_processor import feature_processor

@feature_processor(
    inputs=[SnowflakeDataSource()],
    output="feature-group-arn",
    target_stores=["OfflineStore"],
    spark_config={"spark.jars.packages": "net.snowflake:spark-snowflake_2.12:2.12.0-spark_3.3"}
)
def transform(input_df):
    return input_df
```

To run the feature processor job remotely, you need to provide the packages via defining SparkConfig and pass it to @remote decorator. The Spark packages in the following example are such that spark-snowflake_2.12 is the Feature Processor Scala version, 2.12.0 is the Snowflake version you wish to use, and spark_3.3 is the Feature Processor Spark version.

```python
from sagemaker.remote_function import remote
from sagemaker.remote_function.spark_config import SparkConfig

config = {
    "Classification": "spark-defaults",
    "Properties": {
        "spark.jars.packages": "net.snowflake:spark-snowflake_2.12:2.12.0-spark_3.3"
    }
}

@remote(
    spark_config=SparkConfig(configuration=config),
    instance_type="ml.m5.2xlarge",
)
@feature_processor(
    inputs=[SnowflakeDataSource()],
    output="feature-group-arn",
    target_stores=["OfflineStore"],
)
def transform(input_df):
    return input_df
```
Databricks (JDBC) custom data source examples

Spark can read data from Databricks by using the Databricks JDBC driver. For information about the Databricks JDBC driver, see Configure the Databricks ODBC and JDBC drivers in the Databricks documentation.

Note
You can read data from any other database by including the corresponding JDBC driver in Spark classpath. For more information, see JDBC To Other Databases in the Spark SQL Guide.

To create the custom Databricks data source class, you will need to override the read_data method from the Custom data sources and add the JDBC jar to the Spark classpath.

To connect with a Databricks data source you need:

- Databricks URL (databricks-url)
  For information about your Databricks URL, see Building the connection URL for the Databricks driver in the Databricks documentation.

- Databricks personal access token (personal-access-token)
  For information about your Databricks access token, see Databricks personal access token authentication in the Databricks documentation.

- Data catalog name (db-catalog)
  For information about your Databricks catalog name, see Catalog name in the Databricks documentation.

- Schema name (db-schema)
  For information about your Databricks schema name, see Schema name in the Databricks documentation.

- Table name (db-table-name)
  For information about your Databricks table name, see Table name in the Databricks documentation.
• (Optional) If using Secrets Manager, you’ll need the secret name `secret-databricks-account-info` where you store your Databricks access username and password on Secrets Manager.

For information about Secrets Manager, see [Find secrets in AWS Secrets Manager](https://docs.aws.amazon.com/secretsmanager/latest/userguide/) in the AWS Secrets Manager User Guide.

• AWS Region *(your-region)*

For information about obtaining your current session’s region name using SDK for Python (Boto3), see `region_name` in the Boto3 documentation.

The following example demonstrates how to retrieve the JDBC URL and personal access token from Secrets Manager and overwrite the `read_data` for your custom data source class, DatabricksDataSource.

```python
from sagemaker.feature_store.feature_processor import PySparkDataSource
import json
import boto3

class DatabricksDataSource(PySparkDataSource):
    data_source_name = "Databricks"
    data_source_unique_id = "databricks-url"

    def read_data(self, spark, params):
        secret_name = "secret-databricks-account-info"
        region_name = "your-region"

        session = boto3.session.Session()
        sm_client = session.client(
            service_name='secretsmanager',
            region_name=region_name,
        )

        secrets = json.loads(sm_client.get_secret_value(SecretId=secret_name)["SecretString"])
        jdbc_url = secrets["jdbcurl"].replace("personal-access-token", secrets['pwd'])

        return spark.read.format("jdbc") \
            .option("url", jdbc_url) \
"
The following example shows how to upload the JDBC driver jar, `jdbc-jar-file-name.jar`, to Amazon S3 in order to add it to the Spark classpath. For information about downloading the Spark JDBC driver (`jdbc-jar-file-name.jar`) from Databricks, see [Download JDBC Driver](#) in the Databricks website.

```python
from sagemaker.feature_store.feature_processor import feature_processor

@feature_processor(
    inputs=[DatabricksDataSource()],
    output="feature-group-arn",
    target_stores="OfflineStore",
    spark_config={"spark.jars": "s3://your-bucket-name/your-bucket-prefix/jdbc-jar-file-name.jar"}
)
def transform(input_df):
    return input_df
```

To run the feature processor job remotely, you need to provide the jars by defining `SparkConfig` and pass it to the `@remote` decorator.

```python
from sagemaker.remote_function import remote
from sagemaker.remote_function.spark_config import SparkConfig

config = {
    "Classification": "spark-defaults",
    "Properties": {
        "spark.jars": "s3://your-bucket-name/your-bucket-prefix/jdbc-jar-file-name.jar"
    }
}

@remote(
    spark_config=SparkConfig(configuration=config),
    instance_type="ml.m5.2xlarge",
)
@feature_processor(
    inputs=[DatabricksDataSource()],
    output="feature-group-arn",
    target_stores="OfflineStore",
)
def transform(input_df):
    return input_df

Streaming custom data source examples

You can connect to streaming data sources like Amazon Kinesis, and author transforms with Spark Structured Streaming to read from streaming data sources. For information about the Kinesis connector, see Kinesis Connector for Spark Structured Streaming in GitHub. For information about Amazon Kinesis, see What Is Amazon Kinesis Data Streams? in the Amazon Kinesis Developer Guide.

To create the custom Amazon Kinesis data source class, you will need to extend the BaseDataSource class and override the read_data method from Custom data sources.

To connect to an Amazon Kinesis data stream you need:

- Kinesis ARN (kinesis-resource-arn)
  
  For information on Kinesis data stream ARNs, see Amazon Resource Names (ARNs) for Kinesis Data Streams in the Amazon Kinesis Developer Guide.

- Kinesis data stream name (kinesis-stream-name)

- AWS Region (your-region)

  For information about obtaining your current session's region name using SDK for Python (Boto3), see region_name in the Boto3 documentation.

from sagemaker.feature_store.feature_processor import BaseDataSource
from sagemaker.feature_store.feature_processor import feature_processor

class KinesisDataSource(BaseDataSource):
    
    data_source_name = "Kinesis"
    data_source_unique_id = "kinesis-resource-arn"

    def read_data(self, spark, params):
        return spark.readStream.format("kinesis") \
        .option("streamName", "kinesis-stream-name") \
        .option("awsUseInstanceProfile", "false") \
        .option("endpointUrl", "https://kinesis.your-region.amazonaws.com")
The following example demonstrates how to connect KinesisDataSource to your feature_processor decorator.

```python
from sagemaker.remote_function import remote
from sagemaker.remote_function.spark_config import SparkConfig
import feature_store_pyspark.FeatureStoreManager as fsm

def ingest_micro_batch_into_fg(input_df, epoch_id):
    feature_group_arn = "feature-group-arn"
    fsm.FeatureStoreManager().ingest_data(
        input_data_frame = input_df,
        feature_group_arn = feature_group_arn
    )

@remote(
    spark_config=SparkConfig(
        configuration={
            "Classification": "spark-defaults",
            "Properties":{
                "spark.sql.streaming.schemaInference": "true",
                "spark.jars.packages": "com.roncemer.spark/spark-sql-kinesis_2.13/1.2.2_spark-3.2"
            }
        }
    ),
    instance_type="ml.m5.2xlarge",
    max_runtime_in_seconds=2419200 # 28 days
)
@feature_processor(
    inputs=[KinesisDataSource()],
    output="feature-group-arn"
)

def transform(input_df):
    output_stream = (
        input_df.selectExpr("CAST(rand() AS STRING) as partitionKey", "CAST(data AS STRING)"
    ).writeStream.foreachBatch(ingest_micro_batch_into_fg)
    .trigger(processingTime="1 minute")
    .option("checkpointLocation", "s3a://checkpoint-path")
    .start()
)"
In the example code above we use a few Spark Structured Streaming options while streaming micro-batches into your feature group. For a full list of options, see the Structured Streaming Programming Guide in the Apache Spark documentation.

- The `foreachBatch` sink mode is a feature that allows you to apply operations and write logic on the output data of each micro-batch of a streaming query.

  For information on `foreachBatch`, see Using Foreach and ForeachBatch in the Apache Spark Structured Streaming Programming Guide.

- The `checkpointLocation` option periodically saves the state of the streaming application. The streaming log is saved in checkpoint location `s3a://checkpoint-path`.

  For information on the `checkpointLocation` option, see Recovering from Failures with Checkpointing in the Apache Spark Structured Streaming Programming Guide.

- The `trigger` setting defines how often the micro-batch processing is triggered in a streaming application. In the example, the processing time trigger type is used with one-minute micro-batch intervals, specified by `trigger(processingTime="1 minute")`. To backfill from a stream source, you can use the available-now trigger type, specified by `trigger(availableNow=True)`.

  For a full list of `trigger` types, see Triggers in the Apache Spark Structured Streaming Programming Guide.

Continuous streaming and automatic retries using event based triggers

The Feature Processor uses SageMaker Training as compute infrastructure and it has a maximum runtime limit of 28 days. You can use event based triggers to extend your continuous streaming for a longer period of time and recover from transient failures. For more information on schedule and event based executions, see Scheduled and event based executions for Feature Processor pipelines.

The following is an example of setting up an event based trigger to keep the streaming Feature Processor pipeline running continuously. This uses the streaming transform function defined in the previous example. A target pipeline can be configured to be triggered when a STOPPED or FAILED event occurs for a source pipeline execution. Note that the same pipeline is used as the source and target so that it run continuously.
import sagemaker.feature_store.feature_processor as fp
from sagemaker.feature_store.feature_processor import FeatureProcessorPipelineEvent
from sagemaker.feature_store.feature_processor import FeatureProcessorPipelineExecutionStatus

streaming_pipeline_name = "streaming-pipeline"
streaming_pipeline_arn = fp.to_pipeline(
    pipeline_name = streaming_pipeline_name,
    step = transform # defined in previous section
)

fp.put_trigger(
    source_pipeline_events=FeatureProcessorPipelineEvents(
        pipeline_name=source_pipeline_name,
        pipeline_execution_status=[
            FeatureProcessorPipelineExecutionStatus.STOPPED,
            FeatureProcessorPipelineExecutionStatus.FAILED
        ],
        target_pipeline=target_pipeline_name
    )
)

Example Feature Processing code for common use cases

The following examples provide sample Feature Processing code for common use cases. For a more detailed example notebook showcasing specific use cases, see Amazon SageMaker Feature Store Feature Processing notebook.

In the following examples, us-east-1 is the region of the resource, 111122223333 is the resource owner account ID, and your-feature-group-name is the feature group name.

The transactions data set used in the following examples has the following schema:

```
'FeatureDefinitions': [
    {'FeatureName': 'txn_id', 'FeatureType': 'String'},
    {'FeatureName': 'txn_time', 'FeatureType': 'String'},
    {'FeatureName': 'credit_card_num', 'FeatureType': 'String'},
    {'FeatureName': 'txn_amount', 'FeatureType': 'Fractional'}
]
```

Topics

- Joining data from multiple data sources
• **Sliding window aggregates**
• **Tumbling window aggregates**
• **Promotion from the offline store to online store**
• **Transformations with the Pandas library**
• **Continuous executions and automatic retries using event based triggers**

### Joining data from multiple data sources

```python
@feature_processor(
    inputs=[
        CSVDataSource('s3://bucket/customer'),
        FeatureGroupDataSource('transactions')
    ],
    output='arn:aws:sagemaker:us-east-1:111122223333:feature-group/your-feature-group-name'
)
def join(transactions_df, customer_df):
    '''Combine two data sources with an inner join on a common column'''

    return transactions_df.join(
        customer_df, transactions_df.customer_id == customer_df.customer_id, "inner"
    )
```

### Sliding window aggregates

```python
@feature_processor(
    inputs=[FeatureGroupDataSource('transactions')],
    output='arn:aws:sagemaker:us-east-1:111122223333:feature-group/your-feature-group-name'
)
def sliding_window_aggregates(transactions_df):
    '''Aggregates over 1-week windows, across 1-day sliding windows.''

    from pyspark.sql.functions import window, avg, count

    return (transactions_df
        .groupBy("credit_card_num", window("txn_time", "1 week", "1 day"))
        .agg(avg("txn_amount").alias("avg_week"), count("*").alias("count_week"))
        .orderBy("window.start")
        .select("credit_card_num", "window.start", "avg_week", "count_week")
```
Tumbling window aggregates

```python
@feature_processor(
    inputs=[FeatureGroupDataSource('transactions')],
    output='arn:aws:sagemaker:us-east-1:111122223333:feature-group/your-feature-group-name'
)
def tumbling_window_aggregates(transactions_df, spark):
    '''Aggregates over 1-week windows, across 1-day tumbling windows, as a SQL query.'''

    transactions_df.createOrReplaceTempView('transactions')
    return spark.sql(f'''SELECT credit_card_num, window.start, AVG(amount) AS avg, COUNT(*) AS count
                        FROM transactions
                        GROUP BY credit_card_num, window(txn_time, "1 week")
                        ORDER BY window.start
                        ''')
```

Promotion from the offline store to online store

```python
@feature_processor(
    inputs=[FeatureGroupDataSource('transactions')],
    target_stores=['OnlineStore'],
    output='arn:aws:sagemaker:us-east-1:111122223333:feature-group/transactions'
)
def offline_to_online():
    '''Move data from the offline store to the online store of the same feature group.'''

    transactions_df.createOrReplaceTempView('transactions')
    return spark.sql(f'''SELECT txn_id, txn_time, credit_card_num, amount
                        FROM (SELECT *,
                                row_number() OVER (PARTITION BY txn_id
                                                            ORDER BY "txn_time" DESC, Api_Invocation_Time DESC, write_time DESC)
                                AS row_number
                        FROM (SELECT *
                                row_number()
                        OVER
                                )
                        AS over
                        ORDER BY "txn_time" DESC, Api_Invocation_Time DESC, write_time DESC)
```

Example Feature Processing code for common use cases

Example Feature Processing code for common use cases
FROM transactions
    WHERE row_number = 1

Transformations with the Pandas library

Transformations with the Pandas library

@feature_processor(
    inputs=[FeatureGroupDataSource('transactions')],
    target_stores=['OnlineStore'],
    output='arn:aws:sagemaker:us-east-1:111122223333:feature-group/transactions'
)
def pandas(transactions_df):
    '''Author transformations using the Pandas interface.

    Requires PyArrow to be installed via pip.
    For more details: https://spark.apache.org/docs/latest/api/python/user_guide/pandas_on_spark
    
    ..
    ..
    import pyspark.pandas as ps

    # PySpark DF to Pandas-On-Spark DF (Distributed DF with Pandas interface).
    pandas_on_spark_df = transactions_df.pandas_api()
    # Pandas-On-Spark DF to Pandas DF (Single Machine Only).
    pandas_df = pandas_on_spark_df.to_pandas()

    # Reverse: Pandas DF to Pandas-On-Spark DF
    pandas_on_spark_df = ps.from_pandas(pandas_df)
    # Reverse: Pandas-On-Spark DF to PySpark DF
    spark_df = pandas_on_spark_df.to_spark()

    return spark_df

Continuous executions and automatic retries using event based triggers

from sagemaker.feature_store.feature_processor import put_trigger, to_pipeline,
    FeatureProcessorPipelineEvent
from sagemaker.feature_store.feature_processor import import
    FeatureProcessorPipelineExecutionStatus

streaming_pipeline_name = "target-pipeline"
Time to live (TTL) duration for records

Amazon SageMaker Feature Store provides the option for records to be hard deleted from the online store after a time duration is reached, with time to live (TTL) duration (TtlDuration). The record will expire after the record's EventTime plus the TtlDuration is reached, or ExpiresAt = EventTime + TtlDuration. The TtlDuration can be applied at a feature group level, where all records within the feature group will have the TtlDuration by default, or at an individual record level. If TtlDuration is unspecified, the default value is null and the record will remain in the online store until it is overwritten.

A record deleted using TtlDuration is hard deleted, or completely removed from the online store, and the deleted record is added to the offline store. For more information on hard delete and deletion modes, see DeleteRecord in the Amazon SageMaker API Reference guide. When a record is hard deleted it immediately becomes inaccessible using Feature Store APIs.

⚠️ Important

TTL typically deletes expired items within a few days. Depending on the size and activity level of a table, the actual delete operation of an expired item can vary. Because TTL is meant to be a background process, the nature of the capacity used to expire and delete items via TTL is variable (but free of charge). For more information on how items are deleted from a DynamoDB table, see How it works: DynamoDB Time to Live (TTL).
**TtlDuration** must be a dictionary containing a **Unit** and a **Value**, where the **Unit** must be a string with values "Seconds", "Minutes", "Hours", "Days", or "Weeks" and **Value** must be an integer greater than or equal to 1. **TtlDuration** can be applied while using the **CreateFeatureGroup**, **UpdateFeatureGroup**, and **PutRecord** APIs. See the request and response syntax in the SDK for Python (Boto3) documentation for **CreateFeatureGroup**, **UpdateFeatureGroup**, and **PutRecord** APIs.

- When **TtlDuration** is applied at a feature group level (using the **CreateFeatureGroup** or **UpdateFeatureGroup** APIs), the applied **TtlDuration** becomes the default **TtlDuration** for all records that are added to the feature group from the point in time that the API is called. When applying **TtlDuration** with the **UpdateFeatureGroup** API, this will **not** become the default **TtlDuration** for records that were created **before** the API is called.

  To remove the default **TtlDuration** from an existing feature group, use the **UpdateFeatureGroup** API and set the **TtlDuration** **Unit** and **Value** to **null**.

- When **TtlDuration** is applied at a record level (for example, using **PutRecord** API), the **TtlDuration** duration applies to that record and is used instead of the feature group level default **TtlDuration**.

- When **TtlDuration** is applied on a feature group level it may take a few minutes for **TtlDuration** to come into effect.

- If **TtlDuration** is used when there is no online store, you will receive a **Validation Exception (400)** error.

The following example code shows how to apply **TtlDuration** while updating a feature group, such that the records added to the feature group **after running the API** will by default expire four weeks after their event times.

```python
import boto3

sagemaker_client = boto3.client("sagemaker")
feature_group_name = '<YOUR_FEATURE_GROUP_NAME>'

sagemaker_client.update_feature_group(
    FeatureGroupName=feature_group_name,
    OnlineStoreConfig={
        TtlDuration:{
            Unit: "Weeks",
            Value: 4
        }
    }
)
```

**Time to live (TTL) duration for records**
You can use the DescribeFeatureGroup API to view the default TtlDuration.

To view the expiration times, ExpiresAt (in UTC time ISO-8601 format), while using the GetRecord or BatchGetRecord APIs you must set ExpirationTimeResponse to ENABLED. See the request and response syntax in the SDK for Python (Boto3) documentation for DescribeFeatureGroup, GetRecord, and BatchGetRecord APIs.

Cross account feature group discoverability and access

Data scientists and data engineers can benefit from exploring and accessing features that span multiple accounts, in order to promote data consistency, streamline collaboration, and reduce duplication of effort.

With Amazon SageMaker Feature Store, you can share feature group resources across accounts. The resources that can be shared in Feature Store are feature group entities or the feature group catalog, where the feature group catalog contains all of the feature group entities on your account. The resource owner account shares resources with the resource consumer accounts. There are two distinct categories of permissions associated with sharing resources:

- **Discoverability permission**: Discoverability means being able to see feature group names and metadata. When you share the feature group catalog and grant the discoverability permission, all feature group entities in the account that you share from (resource owner account) become discoverable by the accounts that you are sharing with (resource consumer account). For example, if you make the feature group catalog in the resource owner account discoverable to a resource consumer account, then principals of the resource consumer account can see all feature groups contained in the resource owner account. It means discoverability is “all or nothing” at the account level (regionalized). This permission is granted to resource consumer accounts by using the feature group catalog resource type.

- **Access permissions**: When you grant an access permission, you do so at a feature group resource level (not at account level). This gives you more granular control over granting access to data. The type of access permissions that can be granted are: read-only, read-write, and admin. For example, you can select only certain feature groups from the resource owner account to be accessible by principals of the resource consumer account, depending on your business needs.
This permission is granted to resource consumer accounts by using the feature group resource type and specifying feature group entities.

The distinction between discoverability and access is important to keep in mind when you set up cross account sharing. Also, the methods of sharing resources differ depending on whether you are sharing online or offline feature groups. For information about online and offline feature groups, see Feature Store concepts. In the following topics, you can learn how to apply discoverability and access permissions to your shared resources.

The following example diagram visualizes the feature group catalog resource versus a feature group resource entity. The feature group catalog contains all of your feature group entities and can be shared using the discoverability permission. When granted a discoverability permission, the resource consumer account can search and discover all feature group entities within the resource owner account. A feature group entity contains your machine learning data and can be shared using the access permission. When granted an access permission, the resource consumer account can access the feature group data, with access determined by the relevant access permission.
Enabling cross account discoverability

With AWS Resource Access Manager (AWS RAM) you can securely share the feature group catalog, which contains all of your feature group and feature resources, with other AWS accounts. This lets members of your team search and discover feature groups and features that span multiple accounts, promoting data consistency, streamlining collaboration, and reducing duplication of effort.
The resource owner account can share resources with other individual AWS accounts by granting permissions using AWS RAM. The resource consumer account is the AWS account with whom a resource is shared, limited by the permissions granted from the resource owner account. If you are an organization, you may want to take advantage of AWS Organizations, with which you can share resources with individual AWS accounts, with all accounts in your organization, or in an Organization Unit (OU), without having to apply permissions to each account. For instructional videos and more information about AWS RAM concepts and benefits, see What is AWS Resource Access Manager? in the AWS RAM User Guide.

This section covers how the resource owner account can choose the feature group catalog and grant discoverability privilege to resource consumer accounts, and then how the resource consumer accounts with the discoverability privilege can use search and discover the feature groups within the resource owner account. The discoverability permission does not grant access permissions (read-only, read-write, or admin). Access permissions are granted at a resource level and not at the account level. For information about granting access permissions, see Enabling cross account access.

The following topics discuss how to share the feature group catalog and how to search for shared resources with discoverability permissions applied.

Topics

• Share your feature group catalog
• Search discoverable resources

Share your feature group catalog

The feature group catalog, DefaultFeatureGroupCatalog, contains all feature group entities owned by the resource owner account. The catalog can be shared by the resource owner account to grant discoverability to a single or multiple resource consumer accounts, by creating a resource share in AWS Resource Access Manager (AWS RAM). A feature group is the main resource in Amazon SageMaker Feature Store and is composed of feature definitions and records that are managed by Feature Store. For more information about feature groups, see Feature Store concepts.

Discoverability means that the resource consumer accounts can search for the discoverable resources and view them as if they were in their own account (excluding tags). When allowing the feature group catalog to be discoverable, the resource consumer accounts by default are not granted access permissions (read-only, read-write, or admin). Access permissions are granted at a
resource level and not at the account level. For information about granting access permissions, see Enabling cross account access.

In order to enable cross account discoverability you will need to specify the SageMaker Resource Catalog and the feature group catalog while using the AWS RAM Create a resources share instructions in the AWS RAM developer guide. In the following we give the specifications for using the AWS RAM console instructions.

1. **Specify resource share details:**
   - Resource type: Choose SageMaker Resource Catalogs.
   - ARN: Choose the feature group catalog ARN with the format: arn:aws:sagemaker:us-east-1:111122223333:sagemaker-catalog/DefaultFeatureGroupCatalog
     
     **us-east-1** is the region of the resource and **111122223333** is the resource owner account ID.
   - Resource ID: Choose DefaultFeatureGroupCatalog.

2. **Associate managed permissions:**
   - Managed permission: Choose AWSRAMPermissionSageMakerCatalogResourceSearch.

3. **Grant access to principals:**
   - Choose the principal types (AWS account, Organization, or Organizational unit) and enter the appropriate ID.

   If you are an organization, you may want to take advantage of AWS Organizations, with which you can share resources with individual AWS accounts, with all accounts in your organization, or with an Organization Unit (OU), without having to apply permissions to each account. For more information about sharing your resources and granting permissions within AWS, see Enable resource sharing within AWS Organizations in the AWS Resource Access Manager Developer Guide.

4. **Review and create:**
   - Review then choose Create resource share.

It may take a few minutes for the resource share and principal, or resource consumer account, associations to complete. Once the resource share and principal associations are set, the specified resource consumer accounts receive an invitation to join the resource share. The resource consumer
accounts can view and accept the invitations by opening the [Shared with me: Resource shares](https://aws.amazon.com/about-aws/whats-new/) page in the AWS RAM console. For more information on accepting and viewing resources in AWS RAM, see [Access AWS resources shared with you](https://docs.aws.amazon.com/ram/latest/userguide/access-invitations.html). Invitations are not sent in these cases:

- If you are part of an organization in AWS Organizations and sharing in your organization is enabled, then principals in the organization automatically get access to the shared resources without invitations.
- If you share with the AWS account that owns the resource, then the principals in that account automatically get access to the shared resources without invitations.

For more information about accepting and using a resource share, see [Search discoverable resources](https://docs.aws.amazon.com/ram/latest/userguide/search-discoverable-resources.html).

### Share the feature group catalog using the AWS SDK for Python (Boto3)

You can use the AWS SDK for Python (Boto3) for AWS RAM APIs to create a resource share. The following code is an example of a resource owner account ID `111122223333` within the region `us-east-1` creating a resource share named `test-cross-account-catalog`, sharing the feature group catalog with the resource consumer account ID `444455556666`. To use the Python SDK for AWS RAM APIs, attach the `AWSRAMPermissionSageMakerCatalogResourceSearch` policy with the execution role. See [AWS RAM APIs](https://docs.aws.amazon.com/ram/latest/userguide/ram-api.html) for more details.

```python
# Call list resource catalogs as a prerequisite for RAM share
sagemaker_client.list_resource_catalogs()

# Share DefaultFeatureGroupCatalog with other account
ram_client = boto3.client("ram")
response = ram_client.create_resource_share(
    name='test-cross-account-catalog', # Change to your custom resource share name
    resourceArns=[
        'arn:aws:sagemaker:us-east-1:111122223333:sagemaker-catalog/' +
        'DefaultFeatureGroupCatalog', # Change 111122223333 to the resource owner account ID
    ],
    principals=[
        '444455556666', # Change 444455556666 to the resource consumer account ID
    ],
    permissionArns = ["arn:aws:ram::aws:permission/
AWSRAMPermissionSageMakerCatalogResourceSearch"] # AWSRAMPermissionSageMakerCatalogResourceSearch is the only policy allowed for SageMaker Catalog
```

---

Enabling cross account discoverability

2546
Principals are actors in a security system. In a resource-based policy, the allowed principals are IAM users, IAM roles, the root account, or another AWS service.

**Search discoverable resources**

The resource owner account must grant permissions to resource consumer accounts to allow for discoverability or access (read-only, read-write, or admin) privileges with a shared resource. In the following sections, we provide instructions on how to accept an invitation to shared resources and examples showing how to search for discoverable feature groups.

**Accept an invitation to shared resources**

As the resource consumer account, you receive an invitation to join a resource share once the resource owner account has granted permission. To accept the invitation to any shared resources, open the [Shared with me: Resource shares](https://console.aws.amazon.com/rms/home#resource-shares:resourceowneraccount) page in the AWS RAM console to view and respond to invitations. Invitations are not sent in these cases:

- If you are part of an organization in AWS Organizations and sharing in your organization is enabled, then principals in the organization automatically get access to the shared resources without invitations.
- If you share with the AWS account that owns the resource, then the principals in that account automatically get access to the shared resources without invitations.

For more information about accepting and using a resource share in AWS RAM, see [Respond to the resource share invitation](https).

**Search discoverable feature groups example**

Once resources are shared with a resource consumer account with the discoverability permission applied, the resource consumer account can search for and discover the shared resources in Amazon SageMaker Feature Store using the console UI and the Feature Store SDK. Note that you cannot search on tags for cross account resources. The maximum number of feature group catalogs viewable is 1000. For more information about granting discoverability permissions, see [Enabling cross account discoverability](https).

For details about viewing shared feature groups in the console, see [Find feature groups in your Feature Store](https).
In the following example, the resource consumer account uses SageMaker search to search for resources made discoverable to them when `CrossAccountFilterOption` is set to "CrossAccount":

```python
from sagemaker.session import Session

sagemaker_session = Session(boto_session=boto_session)

sagemaker_session.search(
    resource="FeatureGroup",
    search_expression={
        "Filters": [
            {
                "Name": "FeatureGroupName",
                "Value": "MyFeatureGroup",
                "Operator": "Contains",
            },
        ],
        "Operator": "And",
    },
    sort_by="Name",
    sort_order="Ascending",
    next_token="token",
    max_results=50,
    CrossAccountFilterOption="CrossAccount"
)
```

For more information about SageMaker search and the request parameters, see [Search](#) in the Amazon SageMaker API Reference.

### Enabling cross account access

The access permissions are read-only, read-write, and admin permissions. The permission name, description, and list of specific APIs available for each permission are listed in the following:

- **Read-only permission** (*AWSRAMPermissionFeatureGroupReadOnly*): The read privilege allows resource consumer accounts to read records in the shared feature groups and view details and metadata.
  - `DescribeFeatureGroup`: Retrieves details about a feature group and its configuration
  - `DescribeFeatureMetadata`: Shows the metadata for a feature within a feature group
  - `BatchGetRecord`: Retrieves a batch of records from a feature group
- **GetRecord**: Retrieves a record from a feature group
- **Read-write permission** ([AWSRAMPermissionSagemakerFeatureGroupReadWrite](#)): The read-write privilege allows resource consumer accounts to write records to, and delete records from, the shared feature groups, in addition to read permissions.
- **PutRecord**: Writes a record to a feature group
- **DeleteRecord**: Removes a record from a feature group
- **APIs listed in AWSRAMPermissionFeatureGroupReadOnly**
- **Admin permission** ([AWSRAMPermissionSagemakerFeatureGroupAdmin](#)): The admin privilege allows the resource consumer accounts to update the description and parameters of features within the shared feature groups, update the configuration of the shared feature groups, in addition to read-write permissions.
- **DescribeFeatureMetadata**: Shows the metadata for a feature within a feature group
- **UpdateFeatureGroup**: Updates a feature group configuration
- **UpdateFeatureMetadata**: Updates description and parameters of a feature in the feature group
- **APIs listed in AWSRAMPermissionSagemakerFeatureGroupReadWrite**

In the following topics you can learn how to share online store and offline feature groups—there are differences between the two when it comes to sharing.

**Topics**

- [Share online feature groups with AWS Resource Access Manager](#)
- [Cross account offline store access](#)

**Share online feature groups with AWS Resource Access Manager**

With AWS Resource Access Manager (AWS RAM) you can securely share Amazon SageMaker Feature Store online feature groups with other AWS accounts. Members of your team can explore and access feature groups that span multiple accounts, promoting data consistency, streamlining collaboration, and reducing duplication of effort.

The resource owner account can share resources with other individual AWS accounts by granting permissions using AWS RAM. The resource consumer account is the AWS account with whom a resource is shared, limited by the permissions granted from the resource owner account. If you
are an organization, you may want to take advantage of AWS Organizations, with which you can share resources with individual AWS accounts, with all accounts in your organization, or in an Organization Unit (OU), without having to apply permissions to each account. For instructional videos and more information about AWS RAM concepts and benefits, see What is AWS Resource Access Manager? in the AWS RAM User Guide.

Note that there is a soft maximum limit to the transactions per second (TPS) per API per AWS account. The maximum TPS limit applies to all transactions on the resources within the resource owner account, so transactions from the resource consumer accounts also count towards this maximum limit. For information about service quotas and how to request a quota increase, see AWS service quotas.

This section covers how the resource owner account can choose feature groups and grant access privileges (read-only, read-write, and admin) to resource consumer accounts, and then how the resource consumer accounts with access privileges can use those feature groups. The access permissions do not allow for the resource consumer accounts to search and discover feature groups. To allow for resource consumer accounts to search and discover feature groups from the resource owner account, the resource owner account must grant discoverability permission to the resource consumer accounts, where all of the feature groups within the resource owner account are discoverable by the resource consumer accounts. For more information about granting the discoverability permission, see Enabling cross account discoverability.

The following topics show how to share Feature Store online store resources using the AWS RAM console. For information about sharing your resources and granting permissions within AWS using the AWS RAM console or AWS Command Line Interface (AWS CLI), see Sharing your AWS resources.

Topics

• Share your feature group entities
• Use online store shared resources with access permissions

Share your feature group entities

As the resource owner account you can use the feature group resource type for Amazon SageMaker Feature Store to share feature group entities, by creating a resource share in AWS Resource Access Manager (AWS RAM).

Use the following instructions along with the Sharing your AWS resources instructions in the AWS RAM User Guide.
When sharing the feature group resource type using the AWS RAM console, you need to make the following choices.

1. **Specify resource share details:**
   - Resource type: Choose **SageMaker Feature Groups**.
     - `us-east-1` is the region of the resource, 111122223333 is the resource owner account ID, and `your-feature-group-name` is the feature group you are sharing.
   - Resource ID: Choose the feature group, `your-feature-group-name`, to which you want to grant access permissions.

2. **Associate managed permissions:**
   - Managed permission: Choose the access permission. For more information about access permissions, see [Enabling cross account access](#).

3. **Grant access to principals:**
   - Choose the principal type (AWS account, Organization, Organizational unit, IAM role, or IAM user) and enter the appropriate ID or ARN.

4. **Review and create:**
   - Review then choose **Create resource share**.

Granting any access permission does not grant resource consumer accounts the discoverability permission, so the resource consumer accounts with access permissions cannot search and discover those feature groups. To allow for resource consumer accounts to search and discover feature groups from the resource owner account, the resource owner account must grant the discoverability permission to the resource consumer accounts, where all of the feature groups within the resource owner account are discoverable by the resource consumer accounts. For more information about granting the discoverability permission, see [Enabling cross account discoverability](#).

If the resource consumer accounts are only granted access permissions, the feature group entities can still be viewed on AWS RAM. To view resources on AWS RAM, see [Access AWS resources shared with you](#) in the AWS RAM User Guide.
It may take a few minutes for the resource share and principal, or resource consumer account, associations to complete. Once the resource share and principal associations are set, the specified resource consumer accounts receive an invitation to join the resource share. The resource consumer accounts can view and accept the invitations by opening the Shared with me: Resource shares page in the AWS RAM console. Invitations are not sent in these cases:

- If you are part of an organization in AWS Organizations and sharing in your organization is enabled, then principals in the organization automatically get access to the shared resources without invitations.
- If you share with the AWS account that owns the resource, then the principals in that account automatically get access to the shared resources without invitations.

For more information about accepting and using a resource share in AWS RAM, see Using shared AWS resources in the AWS RAM User Guide.

Share online store feature groups using the AWS SDK for Python (Boto3)

You can use the AWS SDK for Python (Boto3) for AWS RAM APIs to create a resource share. The following code is an example of a resource owner account ID 111122223333 creating a resource share named 'test-cross-account-fg', sharing the feature group named 'my-feature-group' with the resource consumer account ID 444455556666 while granting the AWSRAMPermissionSageMakerFeatureGroupReadOnly permission. For more information about access permissions, see Enabling cross account access. To use the Python SDK for AWS RAM APIs, you need to attach AWS RAM full access managed policy with execution role. See create_resource_share AWS RAM API for more details.

```python
import boto3

# Choose feature group name
feature_group_name = 'my-feature-group'  # Change to your feature group name

# Share 'my-feature-group' with other account
ram_client = boto3.client("ram")
response = ram_client.create_resource_share(
    name='test-cross-account-fg',  # Change to your custom resource share name
    resourceArns=[
        'arn:aws:sagemaker:us-east-1:111122223333:feature-group/' + feature_group_name,
        # Change 111122223333 to the resource owner account ID
    ],
    principals=[
        # Change to your custom resource share principal
    ]
)
```

Enabling cross account access
Principals are actors in a security system. In a resource-based policy, the allowed principals are IAM users, IAM roles, the root account, or another AWS service.

Use online store shared resources with access permissions

The resource owner account must grant permissions to resource consumer accounts to allow for discoverability, read-only, write, or admin privileges with a shared resource. In the following sections, we provide instructions on how to accept an invitation to access shared resources and provide examples showing how to view and interact with shared feature groups.

Accept an invitation to access shared resources using AWS RAM

As the resource consumer account, you will receive an invitation to join a resource share once the resource owner account has granted permission. To accept the invitation to any shared resources, open the Shared with me: Resource shares page in the AWS RAM console to view and respond to invitations. Invitations are not sent in these cases:

- If you are part of an organization in AWS Organizations and sharing in your organization is enabled, then principals in the organization automatically get access to the shared resources without invitations.
- If you share with the AWS account that owns the resource, then the principals in that account automatically get access to the shared resources without invitations.

For more information about accepting and using a resource share in AWS RAM, see Using shared AWS resources in the AWS RAM User Guide.

View shared resources on the AWS RAM console

Granting any access permissions does not grant resource consumer accounts the discoverability permission, so the resource consumer accounts with access permissions cannot search and discover those feature groups. To allow for resource consumer accounts to search and discover feature groups from the resource owner account, the resource owner account must grant the discoverability permission to the resource consumer accounts, where all of the feature groups within the resource owner account are discoverable by the resource consumer accounts. For
more information about granting the discoverability permission, see [Enabling cross account discoverability](#).

To view the shared resources on the AWS RAM console, open the [Shared with me: Resource shares](#) page in the AWS RAM console.

### Read and write actions with a shared feature groups example

Once your resource consumer account is granted the appropriate permissions by the resource owner account, you can perform actions on the shared resources using the Feature Store SDK. You can do this by providing the resource ARN as the FeatureGroupName. To obtain the Feature Group ARN, you can use the AWS SDK for Python (Boto3) `DescribeFeatureGroup` function or use the console UI. For information about using the console UI to view feature group details, see [View feature group details from the console](#).

The following examples use `PutRecord` and `GetRecord` with a shared feature group entity. See the request and response syntax in the AWS SDK for Python (Boto3) documentation for `PutRecord` and `GetRecordAPIs`.

```python
import boto3

sagemaker_featurestore_runtime = boto3.client('sagemaker-featurestore-runtime')

# Put record into feature group named 'test-fg' within the resource owner account ID 111122223333
featurestore_runtime.put_record(
    Record=[value.to_dict() for value in record] # You will need to define record prior to calling PutRecord
)
```

```python
import boto3

sagemaker_featurestore_runtime = boto3.client('sagemaker-featurestore-runtime')

# Choose record identifier
record_identifier_value = str(2990130)

# Get record from feature group named 'test-fg' within the resource owner account ID 111122223333
featurestore_runtime.get_record(
)
```
Cross account offline store access

Amazon SageMaker Feature Store allows users to create a feature group in one account (Account A) and configure it with an offline store using an Amazon S3 bucket in another account (Account B). You can set this up using the steps in the following section.

Topics

- **Step 1: Set up the offline store access role in Account A**
- **Step 2: Set up an offline store Amazon S3 bucket in Account B**
- **Step 3: Set up an offline store AWS KMS encryption key in Account A**
- **Step 4: Create a feature group in Account A**

**Step 1: Set up the offline store access role in Account A**

First, set up a role for Amazon SageMaker Feature Store to write the data into the offline store. The simplest way to accomplish this is to create a new role using the AmazonSageMakerFeatureStoreAccess policy or to use an existing role that already has the AmazonSageMakerFeatureStoreAccess policy attached. This document refers to this policy as `Account-A-Offline-Feature-Store-Role-ARN`.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "s3:PutObject",
            "s3:GetBucketAcl",
            "s3:PutObjectAcl"
         ],
         "Resource": [
            "arn:aws:s3:::*SageMaker*",
            "arn:aws:s3:::*Sagemaker*",
```
The preceding code snippet shows the AmazonSageMakerFeatureStoreAccess policy. The Resource section of the policy is scoped down by default to S3 buckets with names that contain SageMaker, Sagemaker, or sagemaker. This means the offline store Amazon S3 bucket being used must follow this naming convention. If this is not your case, or if you want to further scope down the resource, you can copy and paste the policy to your Amazon S3 bucket policy in the console, customize the Resource section to be arn:aws:s3:::your-offline-store-bucket-name, and then attach to the role.

Additionally, this role must have AWS KMS permissions attached. At a minimum, it requires the kms:GenerateDataKey permission to be able to write to the offline store using your customer managed key. See Step 3 to learn about why a customer managed key is needed for the cross account scenario and how to set it up. The following example shows an inline policy:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "VisualEditor0",
      "Effect": "Allow",
      "Action": [
        "kms:GenerateDataKey"
      ],
    }
  ]
}
```

The Resource section of this policy is scoped to any key in Account A. To further scope this down, after setting up the offline store KMS key in Step 3, return to this policy and replace it with the key ARN.

**Step 2: Set up an offline store Amazon S3 bucket in Account B**

Create an Amazon S3 bucket in Account B. If you are using the default AmazonSageMakerFeatureStoreAccess policy, the bucket name must include SageMaker,
Sagemaker, or sagemaker. Edit the bucket policy as shown in the following example to allow Account A to read and write objects.

This document refers to the following example bucket policy as Account-B-Offline-Feature-Store-Bucket.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "S3CrossAccountBucketAccess",
         "Effect": "Allow",
         "Action": [
            "s3:PutObject",
            "s3:PutObjectAcl",
            "s3:GetBucketAcl"
         ],
         "Principal": {
            "AWS": [
               "*Account-A-Offline-Feature-Store-Role-ARN*"
            ],
         },
         "Resource": [
            "arn:aws:s3:::offline-store-bucket-name/*",
            "arn:aws:s3:::offline-store-bucket-name"
         ]
      }
   ]
}
```

In the preceding policy, the principal is Account-A-Offline-Feature-Store-Role-ARN, which is the role created in Account A in Step 1 and provided to Amazon SageMaker Feature Store to write to the offline store. You can provide multiple ARN roles under Principal.

**Step 3: Set up an offline store AWS KMS encryption key in Account A**

Amazon SageMaker Feature Store ensures that server-side encryption is always enabled for Amazon S3 objects in the offline store. For cross account use cases, you must provide a customer managed key so that you are in control of who can write to the offline store (in this case, Account-A-Offline-Feature-Store-Role-ARN from Account A) and who can read from the offline store (in this case, identities from Account B).
This document refers to the following example key policy as Account-A-Offline-Feature-Store-KMS-Key-ARN.

```json
{
    "Version": "2012-10-17",
    "Id": "key-consolepolicy-3",
    "Statement": [
        {
            "Sid": "Enable IAM User Permissions",
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::Account-A-Account-Id:root"
            },
            "Action": "kms:*",
            "Resource": "*"
        },
        {
            "Sid": "Allow access for Key Administrators",
            "Effect": "Allow",
            "Principal": {
                "AWS": [
                    "arn:aws:iam::Account-A-Account-Id:role/Administrator"
                ]
            },
            "Action": [
                "kms:Create*",
                "kms:Describe*",
                "kms:Enable*",
                "kms:List*",
                "kms:Put*",
                "kms:Update*",
                "kms:Revoke*",
                "kms:Disable*",
                "kms:Get*",
                "kms:Delete*",
                "kms:TagResource",
                "kms:UntagResource",
                "kms:ScheduleKeyDeletion",
                "kms:CancelKeyDeletion"
            ],
            "Resource": "*"
        }
    ]
}
```
"Sid": "Allow Feature Store to get information about the customer managed key",
"Effect": "Allow",
"Principal": {
    "Service": "sagemaker.amazonaws.com"
},
"Action": [
    "kms:Describe*",
    "kms:Get*",
    "kms:List*"
],
"Resource": "*"
},
{
    "Sid": "Allow use of the key",
    "Effect": "Allow",
    "Principal": {
        "AWS": [
            "*Account-A-Offline-Feature-Store-Role-ARN*",
            "*arn:aws:iam::Account-B-Account-Id:root*"
        ]
    },
    "Action": [
        "kms:Encrypt",
        "kms:Decrypt",
        "kms:DescribeKey",
        "kms:CreateGrant",
        "kms:RetireGrant",
        "kms:ReEncryptFrom",
        "kms:ReEncryptTo",
        "kms:GenerateDataKey",
        "kms:ListAliases",
        "kms:ListGrants"
    ],
    "Resource": "*",
}
}

**Step 4: Create a feature group in Account A**

Next, create the feature group in Account A, with an offline store Amazon S3 bucket in Account B. To do this, provide the following parameters for
RoleArn, OfflineStoreConfig.S3StorageConfig.KmsKeyId, and OfflineStoreConfig.S3StorageConfig.S3Uri, respectively:

- Provide Account-A-Offline-Feature-Store-Role-ARN as the RoleArn.
- Provide Account-A-Offline-Feature-Store-KMS-Key-ARN for OfflineStoreConfig.S3StorageConfig.KmsKeyId.
- Provide Account-B-Offline-Feature-Store-Bucket for OfflineStoreConfig.S3StorageConfig.S3Uri.

**Feature Store storage configurations**

Amazon SageMaker Feature Store consists of an online store and an offline store. The online store enables real-time lookup of features for inference, while the offline store contains historical data for model training and batch inference. When creating a feature group, you have the option of enabling either the online store, offline store, or both. When you enable both, they sync to avoid discrepancies between training and serving data. For more information about the online and offline stores and other Feature Store concepts, see [Feature Store concepts](#).

The following topics discuss online store storage types and offline store table formats.

**Topics**

- [Online store](#)
- [Offline store](#)
- [Throughput modes](#)

**Online store**

The online store is a low-latency, high-availability data store that provides real-time lookup of features. It is typically used for machine learning (ML) model serving. You can chose between the standard online store (Standard) or an in-memory tier online store (InMemory), at the point when you create a feature group. In this way, you can select the storage type that best matches the read and write patterns for a particular application, while considering performance and cost. For more details about pricing, see [Amazon SageMaker Pricing](#).

The online store contains the following StorageType options. For more information about the online store contents, see [OnlineStoreConfig](#).
Standard tier storage type

The Standard tier is a managed low-latency data store for online store feature groups. It provides fast data retrieval for ML model service for your applications. Standard is the default storage type.

In-memory tier storage type

The InMemory tier is a managed data store for online store feature groups that supports very low-latency retrieval. It provides large-scale real-time data retrieval for ML model serving used for high throughput applications. The InMemory tier is powered by Amazon ElastiCache for Redis. For more information, see [What is Amazon ElastiCache for Redis?](#).

The online store InMemory tier supports collection types, namely list, set, and vector. For more information about the InMemory collection types, see [Collection types](#).

Feature Store provides low latency read and writes to the online store. The application latency is primarily made up of two primary components: infrastructure or network latency and Feature Store API latency. Reduction of network latency helps with getting the lowest latency reads and writes to Feature Store. You can reduce the network latency to Feature Store by deploying AWS PrivateLink to Feature Store Runtime endpoint. With AWS PrivateLink, you can privately access all Feature Store Runtime API operations from your Amazon Virtual Private Cloud (VPC) in a scalable manner by using interface VPC endpoints. An AWS PrivateLink deployment with the privateDNSEnabled option set as true:

- It keeps all Feature Store read/write traffic within your VPC.
- It keeps traffic in the same AZ as the client that originated it when using Feature Store. This avoids the “hops” between AZs reducing the network latency.

Follow the steps in [Access an AWS service using an interface VPC endpoint](#) to setup AWS PrivateLink to Feature Store. The service name for Feature Store Runtime in AWS PrivateLink is com.amazonaws.region.sagemaker.featurestore-runtime.

The InMemory tier online store scales automatically based about storage usage and requests. The automated scaling can take a few minutes to adapt to a new usage pattern if it changes rapidly. During automated scaling:

- Write operations to the feature group may receive throttling errors. You should retry your requests a few minutes later.
• Read operations to the feature group may receive throttling errors. Standard retry strategies are suitable in this case.

• Read operations may see elevated latency.

The default InMemory tier feature group maximum size is 50 GiB.

Note that the InMemory tier currently supports online feature groups only, not online+offline feature groups, so there is not replication between online and offline stores for the InMemory tier. Also, the InMemory tier does not currently support customer managed KMS keys.

**Offline store**

The offline store is used for historical data when sub-second retrieval is not needed. It is typically used for data exploration, model training, and batch inference.

When you enable both the online and offline stores for your feature group, both stores sync to avoid discrepancies between training and serving data. Please note that an online store feature group with the InMemory storage type enabled does not currently support a corresponding feature group in the offline store (no online to offline replication). For more information about ML model serving in Amazon SageMaker Feature Store, see [Online store](#).

The offline store contains the following TableFormat options. For information about the offline store contents, see [OfflineStoreConfig](#) in the Amazon SageMaker API Reference.

**Glue table format**

The Glue format (default) is a standard Hive type table format for AWS Glue. With AWS Glue, you can discover, prepare, move, and integrate data from multiple sources. It also includes additional productivity and data ops tooling for authoring, running jobs, and implementing business workflows. For more information about AWS Glue, see [What is AWS Glue?](#).

**Iceberg table format**

The Iceberg format (recommended) is an open table format for very large analytic tables. With Iceberg, you can compact the small data files into fewer large files in the partition, resulting in significantly faster queries. This compaction operation is concurrent and does not affect ongoing read and write operations on the feature group. For more information about optimizing Iceberg tables, see the [Amazon Athena](#) and [AWS Lake Formation](#) user guides.
Iceberg manages large collections of files as tables and supports modern analytical data lake operations. If you choose the Iceberg option when creating new feature groups, Amazon SageMaker Feature Store creates the Iceberg tables using Parquet file format, and registers the tables with the AWS Glue Data Catalog. For more information about Iceberg table formats, see Using Apache Iceberg tables.

⚠️ Important

Note that for feature groups in Iceberg table format, you must specify String as the feature type for the event time. If you specify any other type, you can't create the feature group successfully.

### Throughput modes

Amazon SageMaker Feature Store provides two pricing models to choose from: on-demand (On-demand) and provisioned (Provisioned) throughput modes. On-demand works best for less predictable traffic, while Provisioned works best for consistent and predictable traffic.

You have the option to switch between On-demand and Provisioned throughput modes for a given feature group, to accommodate periods in which application traffic patterns are changing or less predictable. You can only update your feature group throughput mode to On-demand once in a 24 hour period. The throughput mode can be updated programmatically using the UpdateFeatureGroup API or through the console UI. For more information about using the console, see Using Amazon SageMaker Feature Store in the console.

You can use the Provisioned throughput mode with offline-only feature groups or feature groups with the Standard storage type. For other storage configurations, the On-demand throughput mode is used. For information about the online and offline storage configurations, see Online store and Offline store, respectively.

For more details about pricing, see Amazon SageMaker Pricing.

**Topics**

- On-demand throughput mode
- Provisioned throughput mode
- Throughput mode metrics
- Throughput mode limits
On-demand throughput mode

The On-demand (default) throughput mode works best when you are using feature groups with unknown workload, unpredictable application traffic, and you cannot forecast the capacity requirements.

The On-demand mode charges you for the reads and writes that your application performs on your feature groups. You do not need to specify how much read and write throughput you expect your application to perform because Feature Store instantly accommodates your workloads as they ramp up or down. You pay only for what you use, which is measured in ReadRequestsUnits and WriteRequestsUnits.

You can enable the On-demand throughput mode using the CreateFeatureGroup or UpdateFeatureGroup APIs or through the console UI. For more information about using the console UI, see Using Amazon SageMaker Feature Store in the console.

Important

You can only update your feature group throughput mode to On-demand once in a 24 hour period.

Provisioned throughput mode

The Provisioned throughput mode works best when you are using feature groups with predictable workloads and you can forecast the capacity requirements to control costs. This can make it more cost effective for certain workloads where you can anticipate throughput requirements in advance.

When you set a feature group to Provisioned mode, you specify capacity units which are the maximum amount of capacity that an application can consume from a feature group. If your application exceeds this Provisioned throughput capacity, it is subject to request throttling.

The following includes information about the read and write capacity units.

- Retrieving a single record of up to 4 KB using the GetRecord API will consume at least 1 RCU (read capacity unit). Retrieving larger payloads may take more. The total number of read capacity units required depends on the item size, including a small per record metadata added by the Feature Store service.
A single write request with a payload of 1 KB using the PutRecord API will consume at least 1 WCU (write capacity unit), with fractional payloads rounded up to nearest KB. It may consume more depending on the event time, deletion status of the record, and time to live (TTL) status. For more information about TTL, see Time to live (TTL) duration for records.

### Important

When setting your capacity units please consider the following:

- You will be charged for the read and write capacities you provision for your feature group, even if you do not fully utilize the Provisioned capacity.
- If you set a read or write capacity too low, your requests may experience throttling.
- In some cases, records may consume an extra capacity unit due to record level metadata that is added by the Feature Store service to enable various features.
- Retrieving only a subset of features using GetRecord or BatchGetRecord APIs will still consume RCU corresponding to the entire record.
- For write capacity, you should provision 2x the recent peak capacity to avoid throttling when performing backfills or bulk ingestion that may result in a large number of historical record writes. This is because writing historical records consumes additional write capacity.
- Feature Store does not currently support auto scaling for Provisioned mode.

You can enable the On-demand throughput mode using the CreateFeatureGroup or UpdateFeatureGroup APIs or through the console UI. For more information about using the console UI, see Using Amazon SageMaker Feature Store in the console.

The following describes how you can increase or decrease the RCU and WCU throughput for your feature groups when Provisioned mode is enabled.

### Increasing provisioned throughput

You can increase RCU or WCU as often as needed using the UpdateFeatureGroup API or the console UI.

### Decreasing provisioned throughput
You can decrease RCU and WCU (or both) for feature groups using `UpdateFeatureGroup` API or the console UI.

There is a default quota on the number of Provisioned capacity decreases you can perform on your feature group per day. A day is defined according to Universal Time Coordinated (UTC). On a given day, you can start by performing up to four decreases within one hour as long as you have not performed any other decreases yet during that day. Subsequently, you can perform one additional decrease per hour as long as there were no decreases in the preceding hour. This effectively brings the maximum number of decreases in a day to 27 times (4 decreases in the first hour, and 1 decrease for each of the subsequent 1-hour windows in a day).

**Throughput mode metrics**

A feature group in On-demand mode will emit `ConsumedReadRequestsUnits` and `ConsumedWriteRequestsUnits` metrics. A feature group in Provisioned mode will emit `ConsumedReadCapacityUnits` and `ConsumedWriteCapacityUnits` metrics. For more information about Feature Store metrics, see [Amazon SageMaker Feature Store Metrics](#).

**Throughput mode limits**

Each AWS account has default service quotas or limits that are applied to help ensure availability and manage billing risks. For information about the default quotas and limits, see [Quotas, naming rules and data types](#).

In some cases, these limits may be lower than what is stated in the documentation. If you need higher limits, you can submit a request for an increase. It's a good idea to do so before reaching current limits to avoid interruptions to your work. For information about service quotas and how to request a quota increase, see [AWS service quotas](#).

**Collection types**

Collection types provide a way to organize and structure data for efficient retrieval and analysis. They are used in ML databases to define the schema of a dataset and its elements. In Amazon SageMaker Feature Store, the supported collection types include list, set, and vector.

Collections are a grouping of elements in which each element within the collection must have the same feature type (String, Integral, or Fractional). For example, a collection can contain elements with all of the element feature types as Fractional, but a collection cannot contain elements with some feature types as Fractional and some feature types as String.
Only InMemory online store feature groups currently support collection types. The following list describes the collection type options.

**List:** An ordered collection of elements.

- The length of the list is determined by how many elements are in the collection.
- Example: You can have a list such as ['a', 'b', 'a'], because the list preserves the order and can have repeat elements.

**Set:** An unordered collection of unique elements.

- The length of the set is determined by how many unique elements are in the collection.
- Example: You cannot have a set such as ['a', 'b', 'a'], because it contains a repeat element. The set will instead have the elements ['a', 'b'], because the set only contains unique elements.

**Vector:** A specialized list that represents a fixed-size array of elements. The order of the elements hold significance, such that the positions of the elements represent certain properties of the data.

- The elements in the vector collection type must have the Fractional feature type.
- You may only have one vector collection type per online store InMemory tier feature group.
- The dimension (number of elements in the vector) of the vector is predetermined by you and is specified using VectorDimension. The max dimension limit is 8192.
- Example: You can have a vector such as [4.2, -6.3, 4.2], where the first, second, and third elements can represent the x, y, and z positions in physical space.

There are no limits on the length of the collections, as long as they don’t exceed the maximum size of a record. For the maximum size of a record, see [Quotas, naming rules and data types](#).

### Add features and records to a feature group

You can use the Amazon SageMaker Feature Store API or the console to update and describe your feature group as well as add features and records to your feature group. A feature group is an object that contains your data and a feature describes a column in the table. When you add a feature to the feature group you are effectively adding a column to the table. When you add a new record to the feature group you are filling in values for features associated with a specific record identifier. For more information on Feature Store concepts, see [Feature Store concepts](#).
After you successfully add features to a feature group, you cannot remove those features. The features that you have added do not add any data to your records. You can add new records to the feature group or overwrite them using the `PutRecord` API. For examples on updating, describing, and putting records into a feature group, see [Example code](#).

You can use the console to add features to a feature group. For more information on how to update your feature groups using the console, see [Update a feature group from the console](#).

The following sections provide an overview of using Feature Store APIs to add features to a feature group followed by examples. With the API, you can also add or overwrite records after you've updated the feature group.

### Topics
- [API](#)
- [Example code](#)

### API

Use the `UpdateFeatureGroup` operation to add features to a feature group.

You can use the `DescribeFeatureGroup` operation to see if you've added the features successfully.

To add or overwrite records, use the `PutRecord` operation.

To see the updates that you've made to a record, use the `GetRecord` operation. To see the updates that you've made to multiple records, use the `BatchGetRecord` operation. It can take up to five minutes for the updates that you've made to appear.

You can use the example code in the following section to walk through adding features and records using the AWS SDK for Python (Boto3).

### Example code

The example code walks you through the following process:

1. Adding features to the feature group
2. Verifying that you've added them successfully
3. Adding a record to the feature group
4. Verifying that you've added it successfully

**Step 1: Add features to a feature group**

The following code uses the UpdateFeatureGroup operation to add new features to the feature group. It assumes that you've set up Feature Store and created a feature group. For more information about getting started, see [Introduction to Feature Store example notebook](#).

```python
import boto3

sagemaker_client = boto3.client("sagemaker")

sagemaker_client.update_feature_group(
    FeatureGroupName=feature_group_name,
    FeatureAdditions=[
        {"FeatureName": "new-feature-1", "FeatureType": "Integral"},
        {"FeatureName": "new-feature-2", "FeatureType": "Fractional"},
        {"FeatureName": "new-feature-3", "FeatureType": "String"}
    ]
)
```

The following code uses the DescribeFeatureGroup operation to check the status of the update. If the LastUpdateStatus field is Successful, you've added the features successfully.

```python
sagemaker_client.describe_feature_group(
    FeatureGroupName=feature_group_name
)
```

**Step 2: Add a new record to the feature group**

The following code uses the PutRecord operation to add records to the feature group that you've created.

```python
record_identifier_value = 'new_record'

sagemaker_featurestore_runtime_client = boto3.client("sagemaker-featurestore-runtime")
```
Use the **GetRecord** operation to see which records in your feature group don't have data for the features that you've added. You can use the **PutRecord** operation to overwrite the records that don't have data for the features that you've added.

### Find features in your feature groups

With Amazon SageMaker Feature Store, you can search for the features that you created in your feature groups. You can search through all of your features without needing to select a feature group first. The search functionality helps find the features that are relevant to your use case.

**Note**

The feature groups where you're searching for features must be within your AWS Region and AWS account. For shared feature groups, the feature groups must be made
discoverable to your AWS account. For more instructions on how to share the feature group catalog and grant discoverability, see [Share your feature group catalog](#).

If you're on a team, and teammates are looking for features to use in their models, they can search through the features in all of the feature groups.

You can add searchable parameters and descriptions to make your features more discoverable. For more information, see [Adding searchable metadata to your features](#).

You can search for features using either the console or by using the **Search** API operation in SageMaker. The following table lists all of the searchable metadata and whether you can search for it in the console or with the API.

<table>
<thead>
<tr>
<th>Searchable metadata</th>
<th>API field name</th>
<th>Searchable in the console?</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Parameters</td>
<td>AllParameters</td>
<td>Yes</td>
</tr>
<tr>
<td>Creation time</td>
<td>CreationTime</td>
<td>Yes</td>
</tr>
<tr>
<td>Description</td>
<td>Description</td>
<td>Yes</td>
</tr>
<tr>
<td>Feature group name</td>
<td>FeatureGroupName</td>
<td>No</td>
</tr>
<tr>
<td>Feature name</td>
<td>FeatureName</td>
<td>Yes</td>
</tr>
<tr>
<td>Feature type</td>
<td>FeatureType</td>
<td>No</td>
</tr>
<tr>
<td>Last modified time</td>
<td>LastModifiedTime</td>
<td>No</td>
</tr>
<tr>
<td>Parameters</td>
<td>Parameters.<em>key</em></td>
<td>Yes</td>
</tr>
</tbody>
</table>

**How to search for your features**

The instructions for using Feature Store through the console depends on whether you have enabled [Amazon SageMaker Studio](#) or [Amazon SageMaker Studio Classic](#) as your default experience. Choose one of the following instructions based on your use case.
Search for features if Studio is your default experience (console)

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose Data in the left navigation pane to expand the dropdown list.
3. From the dropdown list, choose Feature Store.
4. (Optional) To view your features, choose My account. To view shared features, choose Cross account.
5. Under the Feature Catalog tab, choose My account to view your feature groups.
6. Under the Feature Catalog tab, choose Cross account to view feature groups that others made discoverable to you. Under Created by, you can view the resource owner account ID.
7. You can search for your feature in the Search dropdown list:
   - (Optional) To filter your search, choose the filter icon next to the Search dropdown list. You can use filters to specify parameters or date ranges in your search results. If you search for a parameter, specify both its key and value. To find your features, specify time ranges, or clear (deselect) columns that you don't want to query.
   - For shared resources, you can only edit feature group metadata or feature definitions if you have the proper access permission granted from the resource owner account. The discoverability permission alone won't allow you to edit metadata or feature definitions. For more information about granting access permissions, see Enabling cross account access.

Search for features if Studio Classic is your default experience (console)

Use the latest version of Amazon SageMaker Studio Classic so that you have the most recent version of the search functionality. For information about updating Studio Classic, see Shut down and Update SageMaker Studio Classic.

1. Open the Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.
2. Choose the Home icon
   (in the left navigation pane.
3. Choose Data.
4. From the dropdown list, choose Feature Store.
5. (Optional) To view your features, choose **My account**. To view shared features, choose **Cross account**.

6. Under the **Feature Catalog** tab, choose **My account** to view your feature groups.

7. Under the **Feature Catalog** tab, choose **Cross account** to view feature groups that others made discoverable to you. Under **Created by**, you can view the resource owner account ID.

8. You can search for your feature in the **Search** dropdown list:

   • (Optional) To filter your search, choose the filter icon next to the **Search** dropdown list. You can use filters to specify parameters or date ranges in your search results. If you search for a parameter, specify both its key and value. To find your features, specify time ranges, or clear (deselect) columns that you don't want to query.

   • For shared resources, you can only edit feature group metadata or feature definitions if you have the proper access permission granted from the resource owner account. The discoverability permission alone won't allow you to edit metadata or feature definitions. For more information about granting access permissions, see [Enabling cross account access](#).

**Search for your features using SDK for Python (Boto3)**

The code in this section uses the **Search** operation in the AWS SDK for Python (Boto3) to run the search query to find features in your feature groups. For information about the other languages to submit a query, see [See Also](#) in the *Amazon SageMaker API Reference*.

For more Feature Store examples and resources, see [Amazon SageMaker Feature Store resources](#).

The following code shows different example search queries using the API:

```python
# Return all features in your feature groups
sagemaker_client.search(
    Resource="FeatureMetadata",
)

# Search for all features that belong to a feature group that contain the "ver" substring
sagemaker_client.search(
    Resource="FeatureMetadata",
    SearchExpression={
        'Filters': [
            {
```
How to search for your features

# Search for all features that belong to a feature group that have the EXACT name "airport"
sagemaker_client.search(
    Resource="FeatureMetadata",
    SearchExpression={
        'Filters': [
            {
                'Name': 'FeatureGroupName',
                'Operator': 'Equals',
                'Value': 'airport'
            }
        ]
    }
)

# Search for all features that belong to a feature group that contains the name "ver"
AND have a name that contains "wha"
AND have a parameter (key or value) that contains "hea"

sagemaker_client.search(
    Resource="FeatureMetadata",
    SearchExpression={
        'Filters': [
            {
                'Name': 'FeatureGroupName',
                'Operator': 'Contains',
                'Value': 'ver'
            },
            {
                'Name': 'FeatureName',
                'Operator': 'Contains',
                'Value': 'wha'
            },
            {
                'Name': 'AllParameters',
                'Operator': 'Contains',
                'Value': 'hea'
            }
        ]
    }
)
# Search for all features that belong to a feature group with substring "ver" in its name
OR features that have a name that contain "wha"
OR features that have a parameter (key or value) that contains "hea"

```python
sagemaker_client.search(
    Resource="FeatureMetadata",
    SearchExpression={
        'Filters': [
            {'Name': 'FeatureGroupName', 'Operator': 'Contains', 'Value': 'ver'},
            {'Name': 'FeatureName', 'Operator': 'Contains', 'Value': 'wha'},
            {'Name': 'AllParameters', 'Operator': 'Contains', 'Value': 'hea'},
        ],
        'Operator': 'Or'  # note that this is explicitly set to "Or"- the default is "And"
    }
)
```

# Search for all features that belong to a feature group with substring "ver" in its name
OR features that have a name that contain "wha"
OR parameters with the value 'Sage' for the 'org' key

```python
sagemaker_client.search(
    Resource="FeatureMetadata",
    SearchExpression={
        'Filters': [
            {'Name': 'FeatureGroupName', 'Operator': 'Contains', 'Value': 'ver'},
            {'Name': 'FeatureName', 'Operator': 'Contains', 'Value': 'wha'},
            {'Name': 'AllParameters', 'Operator': 'Contains', 'Value': 'hea'},
        ],
        'Operator': 'Or'  # note that this is explicitly set to "Or"- the default is "And"
    }
)
```
Find feature groups in your Feature Store

With Amazon SageMaker Feature Store, you can search for the feature groups using either the console or the Search operation. You can use the search functionality to find features and feature groups that are relevant to the models that you're creating. You can use the search functionality to quickly find the feature groups that are relevant to your use case.

Note

The feature groups that you're searching for must be within your AWS Region and AWS account, or shared with and made discoverable to your AWS account. For more information about how to share the feature group catalog and grant discoverability, see Share your feature group catalog.

The following table shows the searchable fields and whether you can use the console to search for a specific field.
You can search for features using either Amazon SageMaker Studio Classic or the Search operation in the SageMaker API. The following table lists all of the searchable metadata and whether you can search for it in the console. Tags are searchable for your own feature groups but are not searchable for feature groups made discoverable to you.

<table>
<thead>
<tr>
<th>Searchable metadata</th>
<th>API field name</th>
<th>Searchable in the console?</th>
<th>Searchable with cross account?</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Tags</td>
<td>AllTags</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Creation Failure Reason</td>
<td>FailureReason</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Creation Status</td>
<td>FeatureGroupStatus</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Creation time</td>
<td>CreationTime</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Description</td>
<td>Description</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Event Time Feature Name</td>
<td>EventTimeFeatureName</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Feature Definitions</td>
<td>FeatureDefinitions</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Feature Group ARN</td>
<td>FeatureGroupARN</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Feature Group Name</td>
<td>FeatureGroupName</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Offline Store Configuration</td>
<td>OfflineStoreConfig</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Offline Store Status</td>
<td>OfflineStoreStatus</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Last Update Status</td>
<td>LastUpdateStatus</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Record Identifier Feature Name</td>
<td>RecordIdentifierFeatureName</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Tags</td>
<td>Tags.key</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>
How to find feature groups

You can use the console or the Amazon SageMaker Feature Store API to find your feature groups. The instructions for using Feature Store through the console depends on if you have enabled Amazon SageMaker Studio or Amazon SageMaker Studio Classic as your default experience.

Find feature groups if Studio is your default experience (console)

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose Data in the left navigation pane to expand the dropdown list.
3. From the dropdown list, choose Feature Store.
4. (Optional) To view your feature groups, choose My account. To view shared feature groups, choose Cross account.
5. Under the Feature Group Catalog tab, choose My account to view your feature groups.
6. Under the Feature Group Catalog tab, choose Cross account to view feature groups that others made discoverable to you. Under Created by, you can view the resource owner account ID.
7. You can search for your feature groups in the Search dropdown list:
   • (Optional) To filter your search, choose the filter icon next to the Search dropdown list. You can use filters to specify parameters or date ranges in your search results. If you search for a parameter, specify both its key and value. To find your feature groups, you can specify time ranges, clear (deselect) columns that you don't want to query, choose stores to search, or search by status.
   • For shared resources, you can only edit feature group metadata or feature definitions if you have the proper access permission granted from the resource owner account. The discoverability permission alone won't allow you to edit metadata or feature definitions. For more information about granting access permissions, see Enabling cross account access.

Find feature groups if Studio Classic is your default experience (console)

Use the latest version of Amazon SageMaker Studio Classic to get the most recent version of the search functionality if you are accessing Feature Store through the Studio Classic application. For information about updating Studio Classic, see Shut down and Update SageMaker Studio Classic.

1. Open the Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.
2. Choose the **Home** icon
   in the left navigation pane.

3. Choose **Data**.

4. From the dropdown list, choose **Feature Store**.

5. (Optional) To view your feature groups, choose **My account**. To view shared feature groups, choose **Cross account**.

6. Under the **Feature Group Catalog** tab, choose **My account** to view your feature groups.

7. Under the **Feature Group Catalog** tab, choose **Cross account** to view feature groups that others made discoverable to you. Under **Created by**, you can view the resource owner account ID.

8. You can search for your feature groups in the **Search** dropdown list:
   - (Optional) To filter your search, choose the filter icon next to the **Search** dropdown list. You can use filters to specify parameters or date ranges in your search results. If you search for a parameter, specify both its key and value. To find your feature groups more easily, you can specify time ranges, clear (deselect) columns that you don't want to query, choose stores to search, or search by status.
   - For shared resources, you can only edit feature group metadata or feature definitions if you have the proper access permission granted from the resource owner account. The discoverability permission alone won't allow you to edit metadata or feature definitions. For more information about granting access permissions, see [Enabling cross account access](#).

### Find feature groups using SDK for Python (Boto3)

The code in this section uses the **Search** operation in the AWS SDK for Python (Boto3) to run the search query to find feature groups. For information about the other languages to submit a query, see [See Also](#) in the [Amazon SageMaker API Reference](#).

For more Feature Store examples and resources, see [Amazon SageMaker Feature Store resources](#).

The following code shows different example search queries using the API:

```python
# Return all feature groups
sagemaker_client.search(
    Resource="FeatureGroups",
)
```
# Search for feature groups that are shared with your account
```
sagemaker_session.search(
    resource="FeatureGroup",
    search_expression={
        "Filters": [
        {
            "Name": "FeatureGroupName",
            "Value": "MyFeatureGroup",
            "Operator": "Contains",
        }
        ],
        "Operator": "And",
    },
    sort_by="Name",
    sort_order="Ascending",
    next_token="token",
    max_results=50,
    CrossAccountFilterOption="SameAccount"
)
```

# Search for all feature groups with a name that contains the "ver" substring
```
sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
        {
            'Name': 'FeatureGroupName',
            'Operator': 'Contains',
            'Value': 'ver'
        }
        ]
    }
)
```

# Search for all feature groups that have the EXACT name "airport"
```
sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
        {
            'Name': 'FeatureGroupName',
            'Operator': 'Equals',
            'Value': 'airport'
        }
        ]
    }
)
```
# Search for all feature groups that contains the name "ver"
# AND have a record identifier feature name that contains "wha"
# AND have a tag (key or value) that contains "hea"

sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
            {
                'Name': 'FeatureGroupName',
                'Operator': 'Contains',
                'Value': 'ver'
            },
            {
                'Name': 'RecordIdentifierFeatureName',
                'Operator': 'Contains',
                'Value': 'wha'
            },
            {
                'Name': 'AllTags',
                'Operator': 'Contains',
                'Value': 'hea'
            }
        ]
    }
)

# Search for all feature groups with substring "ver" in its name
# OR feature groups that have a record identifier feature name that contains "wha"
# OR feature groups that have a tag (key or value) that contains "hea"

sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
            {
                'Name': 'FeatureGroupName',
                'Operator': 'Contains',
                'Value': 'ver'
            }
        ]
    }
)
# Search for all feature groups with substring "ver" in its name
# OR feature groups that have a record identifier feature name that contains "wha"
# OR tags with the value 'Sage' for the 'org' key

```python
sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
            { 'Name': 'FeatureGroupName', 'Operator': 'Contains', 'Value': 'ver' },
            { 'Name': 'RecordIdentifierFeatureName', 'Operator': 'Contains', 'Value': 'wha' },
            { 'Name': 'Tags.org', 'Operator': 'Contains', 'Value': 'Sage' }],
        'Operator': 'Or' # note that this is explicitly set to "Or"- the default is "And"
    })
```

How to find feature groups
# Search for all offline only feature groups
sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
            {
                'Name': 'OnlineStoreConfig.EnableOnlineStore',
                'Operator': 'NotEquals',
                'Value': 'true'
            },
            {
                'Name': 'OfflineStoreConfig.S3StorageConfig.S3Uri',
                'Operator': 'Exists'
            }
        ]
    }
)

# Search for all online only feature groups
sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
            {
                'Name': 'OnlineStoreConfig.EnableOnlineStore',
                'Operator': 'Equals',
                'Value': 'true'
            },
            {
                'Name': 'OfflineStoreConfig.S3StorageConfig.S3Uri',
                'Operator': 'NotExists'
            }
        ]
    }
)

# Search for all feature groups that are BOTH online and offline
sagemaker_client.search(
    Resource="FeatureGroups",
    SearchExpression={
        'Filters': [
            {
                'Name': 'OnlineStoreConfig.EnableOnlineStore',
                'Operator': 'Equals',
                'Value': 'true'
            }
        ]
    }
)
You can also use python SDK of AWS RAM APIs to create resource share. The API signature is given below. To use python SDK of AWS RAM API, you need attach AWS RAM full access managed policy with execution Role.

```
response = client.create_resource_share(
    name='string',
    resourceArns=[
        'string',
    ],
    principals=[
        'string',
    ],
    tags=[
        {
            'key': 'string',
            'value': 'string'
        },
    ],
    allowExternalPrincipals=True|False,
    clientToken='string',
    permissionArns=[
        'string',
    ]
)
```

**Adding searchable metadata to your features**

In Amazon SageMaker Feature Store, you can search through all of your features. To make your features more discoverable, you can add metadata to them. You can add the following types of metadata:
• Description – A searchable description of the feature.

• Parameters – Searchable key-value pairs.

The description can have up to 255 characters. For parameters, you must specify a key-value pair in your search. You can add up to 25 parameters.

To update the metadata of a feature, you can use either the console or the `UpdateFeatureMetadata` operation.

How to add searchable metadata to your features

You can use the console or the Amazon SageMaker Feature Store API to add searchable metadata to your features. Instructions for using Feature Store through the console depend on whether you have enabled Amazon SageMaker Studio or Amazon SageMaker Studio Classic as your default experience.

Add searchable metadata to features if Studio is your default experience (console)

1. Open the Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. Choose Data in the left navigation pane, to expand the dropdown list.
3. From the dropdown list, choose Feature Store.
4. (Optional) To view your features, choose My account. To view shared features, choose Cross account.
5. To view your feature groups, under the Feature Catalog tab, choose My account.
6. Under the Feature Catalog tab, choose Cross account to view feature groups that others make discoverable to you. Under Created by, you can view the resource owner account ID of the feature group.
7. You can search for your feature in the Search dropdown list.

• (Optional) To filter your search, choose the filter icon next to the Search dropdown list. You can use filters to specify parameters or date ranges in your search results. If you search for a parameter, specify both its key and value. To find your features more easily, you can specify time ranges or deselect columns that you don't want to query.

• For shared resources, you can only edit feature group metadata or feature definitions if you have the proper access permission granted from the resource owner account. Having the discoverability permission alone doesn't allow you to edit metadata or feature definitions. For more information about granting access permissions, see Enabling cross account access.
8. Choose your feature.

9. Choose **Edit metadata**.

10. In the **Description** field, add or update the description.

11. In the **Parameters** field under **Parameters**, specify a key-value pair for the parameter.

12. (Optional) Choose **Add new parameter** to add another parameter.

13. Choose **Save changes**.

14. Choose **Confirm**.

---

**Add searchable metadata to your features if Studio Classic is your default experience (console)**

1. Open the Studio Classic console by following the instructions in [Launch Studio Classic Using the Amazon SageMaker Console](#).

2. In the left navigation pane, choose the **Home** icon (🏠).

3. Choose **Data**.

4. From the dropdown list, choose **Feature Store**.

5. (Optional) To view your features, choose **My account**. To view shared features, choose **Cross account**.

6. To view your feature groups, under the **Feature Catalog** tab, choose **My account**.

7. Under the **Feature Catalog** tab, choose **Cross account** to view feature groups that other accounts made discoverable to you. Under **Created by**, you can view the resource owner account ID of the feature group.

8. You can search for your feature in the **Search** dropdown list.

   - (Optional) To filter your search, choose the filter icon next to the **Search** dropdown list. You can use filters to specify parameters or date ranges in your search results. If you search for a parameter, specify both its key and value. To find your features more easily, you can specify time ranges or deselect columns that you don't want to query.

   - For shared resources, you can only edit feature group metadata or feature definitions if you have the proper access permission granted from the resource owner account. Having the discoverability permission alone doesn't allow you to edit metadata or feature definitions. For more information about granting access permissions, see [Enabling cross account access](#).

9. Choose your feature.
10. Choose **Edit metadata**.

11. In the **Description** field, add or update the description.

12. In the **Parameters** field under **Parameters**, specify a key-value pair for the parameter.

13. (Optional) Choose **Add new parameter** to add another parameter.

14. Choose **Save changes**.

15. Choose **Confirm**.

**Add searchable metadata to your features using SDK for Python (Boto3)**

The code in this section uses the `UpdateFeatureMetadata` operation in the AWS SDK for Python (Boto3) to add searchable metadata to your features for different scenarios. For information about the other languages to submit a query, see **See Also** in the *Amazon SageMaker API Reference*.

For more Feature Store examples and resources, see [Amazon SageMaker Feature Store resources](https://docs.aws.amazon.com/sagemaker/latest/dg/feature-store.html).

### Add a list of parameters to a feature

To add a list of parameters to a feature, specify values for the following fields:

- FeatureGroupName
- Feature
- Parameters

The following example code uses the AWS SDK for Python (Boto3) to add two parameters.

```python
sagemaker_client.update_feature_metadata(
    FeatureGroupName="feature_group_name",
    FeatureName="feature-name",
    ParameterAdditions=[
        {"Key": "example-key-0", "Value": "example-value-0"},
        {"Key": "example-key-1", "Value": "example-value-1"},
    ]
)
```

### Add a description to a feature

To add a description to a feature, specify values for the following fields:
- FeatureGroupName
- Feature
- Description

```python
sagemaker_client.update_feature_metadata(
    FeatureGroupName="feature-group-name",
    FeatureName="feature-name",
    Description="description"
)
```

**Remove parameters for a feature**

To remove all parameters for a feature, do the following.

Specify values for the following fields:

- FeatureGroupName
- Feature

Specify the keys for the parameters that you're removing under ParameterRemovals.

```python
sagemaker_client.update_feature_metadata(
    FeatureGroupName="feature-group-name",
    FeatureName="feature-name",
    ParameterRemovals=[
    {"Key": "example-key-0"},
    {"Key": "example-key-1"},
    ]
)
```

**Remove the description for a feature**

To remove the description for a feature, do the following.

Specify values for the following fields:

- FeatureGroupName
• Feature

Specify an empty string for Description.

```python
sagemaker_client.update_feature_metadata(
    FeatureGroupName="feature-group-name",
    FeatureName="feature-name",
    Description=""
)
```

**Example code**

After you've updated the metadata for a feature, you can use the `DescribeFeatureMetadata` operation to see the updates that you've made.

The following code goes through an example workflow using the AWS SDK for Python (Boto3). The example code does the following:

1. Sets up your SageMaker environment.
2. Creates a feature group.
3. Adds features to the group.
4. Adds metadata to the features.

For more Feature Store examples and resources, see [Amazon SageMaker Feature Store resources](#).

**Step 1: Setup**

To start using Feature Store, create SageMaker, boto3 and Feature Store sessions. Then set up the S3 bucket you want to use for your features. This is your offline store. The following code uses the SageMaker default bucket and adds a custom prefix to it.

**Note**

The role that you use must have the following managed policies attached to it: `AmazonS3FullAccess` and `AmazonSageMakerFeatureStoreAccess`. 
# SageMaker Python SDK version 2.x is required
%pip install 'sagemaker>=2.0.0'
import sagemaker
import sys
import boto3
import pandas as pd
import numpy as np
import io
from sagemaker.session import Session
from sagemaker import get_execution_role
from botocore.exceptions import ClientError

prefix = 'sagemaker-featurestore-introduction'
role = get_execution_role()
sagemaker_session = sagemaker.Session()
region = sagemaker_session.boto_region_name
s3_bucket_name = sagemaker_session.default_bucket()
sagemaker_client = boto_session.client(service_name='sagemaker', region_name=region)

Step 2: Create a feature group and add features

The following code is an example of creating a feature group with feature definitions.

feature_group_name = "test-for-feature-metadata"
feature_definitions = [
    {"FeatureName": "feature-1", "FeatureType": "String"},
    {"FeatureName": "feature-2", "FeatureType": "String"},
    {"FeatureName": "feature-3", "FeatureType": "String"},
    {"FeatureName": "feature-4", "FeatureType": "String"},
    {"FeatureName": "feature-5", "FeatureType": "String"}
]
try:
    sagemaker_client.create_feature_group(
        FeatureGroupName=feature_group_name,
        RecordIdentifierFeatureName="feature-1",
        EventTimeFeatureName="feature-2",
    )

How to add searchable metadata to your features
Step 3: Add metadata

Before you add metadata, use the `DescribeFeatureGroup` operation to make sure that the status of the feature group is `Created`.

```python
sagemaker_client.describe_feature_group(
    FeatureGroupName=feature_group_name
)
```

Add a description to the feature.

```python
sagemaker_client.update_feature_metadata(
    FeatureGroupName=feature_group_name,
    FeatureName="feature-1",
    Description="new description"
)
```

You can use the `DescribeFeatureMetadata` operation to see if you successfully updated the description for the feature group.

```python
sagemaker_client.describe_feature_metadata(
    FeatureGroupName=feature_group_name,
    FeatureName="feature-1"
)
```

You can also use it to add parameters to the feature group.
You can use the DescribeFeatureMetadata operation again to see if you have successfully added the parameters.

Create a dataset from your feature groups

After a Feature Store feature group has been created in an offline store, you can choose to use the following methods to get your data:

- Using the Amazon SageMaker Python SDK
- Running SQL queries in the Amazon Athena

Important

Feature Store requires data to be registered in a AWS Glue data catalog. By default, Feature Store automatically builds an AWS Glue data catalog when you create a feature group.

After you've created feature groups for your offline store and populated them with data, you can create a dataset by running queries or using the SDK to join data stored in the offline store from different feature groups. You can also join the feature groups to a single pandas dataframe. You can use Amazon Athena to write and execute SQL queries.
Note

To make sure that your data is up to date, you can set up a AWS Glue crawler to run on a schedule.

To set up a AWS Glue crawler, specify an IAM role that the crawler is using to access the offline store's Amazon S3 buckets. For more information, see Create an IAM role. For more information on how to use AWS Glue and Athena to build a training dataset for model training and inference, see Use Feature Store with SDK for Python (Boto3).

Using the Amazon SageMaker Python SDK to get your data from your feature groups

You can use the Feature Store APIs to create a dataset from your feature groups. Data scientists create ML datasets for training by retrieving ML feature data from one or more feature groups in the offline store. Use the create_dataset() function to create the dataset. You can use the SDK to do the following:

- Create a dataset from multiple feature groups.
- Create a dataset from the feature groups and a pandas data frame.

By default, Feature Store doesn't include records that you've deleted from the dataset. It also doesn't include duplicated records. A duplicate record has the record ID and timestamp value in the event time column.

Before you use the SDK to create a dataset, you must start a SageMaker session. Use the following code to start the session.

```python
import boto3
from sagemaker.session import Session
from sagemaker.feature_store.feature_store import FeatureStore

region = boto3.Session().region_name
boto_session = boto3.Session(region_name=region)

sagemaker_client = boto_session.client(
    service_name="sagemaker", region_name=region
)

featurestore_runtime = boto_session.client(
```

```python
The following code shows an example of creating a dataset from multiple feature groups. The following code snippet uses the example feature groups "base_fg_name", "first_fg_name", and "second_fg_name", which may not exist or have the same schema within your Feature Store. It is recommended to replace these feature groups with feature groups that exist within your Feature Store. For information on how to create a feature group, see Step 3: Create feature groups.

```python
from sagemaker.feature_store.feature_group import FeatureGroup

s3_bucket_name = "offline-store-sdk-test"
base_fg_name = "base_fg_name"
based_fg = FeatureGroup(name=base_fg_name, sagemaker_session=feature_store_session)

first_fg_name = "first_fg_name"
first_fg = FeatureGroup(name=first_fg_name, sagemaker_session=feature_store_session)

second_fg_name = "second_fg_name"
second_fg = FeatureGroup(name=second_fg_name, sagemaker_session=feature_store_session)

feature_store = FeatureStore(feature_store_session)
builder = feature_store.create_dataset(
    base=base_fg,
    output_path=f"s3://{DOC-EXAMPLE-BUCKET1}"
).with_feature_group(first_fg)
.with_feature_group(second_fg, "base_id", ["base_feature_1"])
```

The following code shows an example of creating a dataset from multiple feature groups and a pandas dataframe.

```python
base_data = [[1, 187512346.0, 123, 128],
```
The Feature Store APIs provide you with helper methods for the `create_dataset` function. You can use them to do the following:

- Create a dataset from multiple feature groups.
- Create a dataset from multiple feature groups and a pandas dataframe.
- Create a dataset from a single feature group and a pandas dataframe.
- Create a dataset using a point in time accurate join where records in the joined feature group follow sequentially.
- Create a dataset with the duplicated records, instead of following the default behavior of the function.
- Create a dataset with the deleted records, instead of following the default behavior of the function.
- Create a dataset for time periods that you specify.
- Save the dataset as a CSV file.
- Save the dataset as a pandas dataframe.

The `base` feature group is an important concept for joins. The base feature group is the feature group that has other feature groups or the pandas dataframe joined to it. For each dataset you can add the following optional methods to the `create_dataset` function to configure how you're creating dataset:

```python
data = [[2, 187512347.0, 168, 258],
        [3, 187512348.0, 125, 184],
        [1, 187512349.0, 195, 206]]
base_data_df = pd.DataFrame(
    data,
    columns=['base_id', 'base_time', 'base_feature_1', 'base_feature_2'])

builder = feature_store.create_dataset(
    base=base_data_df,
    event_time_identifier_feature_name='base_time',
    record_identifier_feature_name='base_id',
    output_path=f's3://{s3_bucket_name}'
).with_feature_group(first_fg)
.with_feature_group(second_fg, 'base_id', ['base_feature_1'])
```
with_feature_group – Performs an inner join between the base feature group and another feature group using the record identifier and the target feature name in the base feature group. The following provides information about the parameters that you specify:

- feature_group – The feature group that you're joining.
- target_feature_name_in_base – The name of the feature in the base feature group that you're using as a key in the join. The record identifier in the other feature groups are the other keys that Feature Store uses in the join.
- included_feature_names – A list of strings representing the feature names of the base feature group. You can use the field to specify the features that you want to include in the dataset.
- feature_name_in_target – Optional string representing the feature in the target feature group that will be compared to the target feature in the base feature group.
- join_comparator – Optional JoinComparatorEnum representing the comparator used when joining the target feature in the base feature group and the feature in the target feature group. These JoinComparatorEnum values can be GREATER_THAN, GREATER_THAN_OR_EQUAL_TO, LESS_THAN, LESS_THAN_OR_EQUAL_TO, NOT_EQUAL_TO or EQUALS by default.
- join_type – Optional JoinTypeEnum representing the type of join between the base and target feature groups. These JoinTypeEnum values can be LEFT_JOIN, RIGHT_JOIN, FULL_JOIN, CROSS_JOIN or INNER_JOIN by default.
- with_event_time_range – Creates a dataset using the event time range that you specify.
- as_of – Creates a dataset up to a timestamp that you specify. For example, if you specify datetime(2021, 11, 28, 23, 55, 59, 342380) as the value, creates a dataset up to November 28th, 2021.
- point_time_accurate_join – Creates a dataset where all of the event time values of the base feature group is less than all the event time values of the feature group or pandas dataframe that you're joining.
- include_duplicated_records – Keeps duplicated values in the feature groups.
- include_deleted_records – Keeps deleted values in the feature groups.
- with_number_of_recent_records_by_record_identifier – An integer that you specify to determine how many of the most recent records appear in the dataset.
- with_number_of_records_by_record_identifier – An integer that represents how many records appear in the dataset.
After you've configured the dataset, you can specify the output using one of the following methods:

- **to_csv_file** – Saves the dataset as a CSV file.
- **to_dataframe** – Saves the dataset as a pandas dataframe.

You can retrieve data that comes after a specific period in time. The following code retrieves data after a timestamp.

```python
g1 = FeatureGroup("example-feature-group-1")
feature_store.create_dataset(
    base=g1,
    output_path="s3://example-S3-path"
).with_number_of_records_from_query_results(5).to_csv_file()
```

You can also retrieve data from a specific time period. You can use the following code to get data for a specific time range:

```python
g1 = FeatureGroup("fg1")
feature_store.create_dataset(
    base=g1,
    output_path="example-S3-path"
).with_event_time_range(
    datetime(2021, 11, 28, 23, 55, 59, 342380),
    datetime(2020, 11, 28, 23, 55, 59, 342380)
).to_csv_file() #example time range specified in datetime functions
```

You might want to join multiple feature groups to a pandas dataframe where the event time values of the feature group happen no later than the event time of the data frame. Use the following code as a template to help you perform the join.

```python
g1 = FeatureGroup("fg1")
g2 = FeatureGroup("fg2")
events = [['2020-02-01T08:30:00Z', 6, 1],
    ['2020-02-02T10:15:30Z', 5, 2],
    ['2020-02-03T13:20:59Z', 1, 3],
    ['2021-01-01T00:00:00Z', 1, 4]]
df = pd.DataFrame(events, columns=['event_time', 'customer-id', 'title-id'])
feature_store.create_dataset(
    base=df,
    output_path="s3://example-S3-path"
).with_event_time_range(
    datetime(2021, 11, 28, 23, 55, 59, 342380),
    datetime(2020, 11, 28, 23, 55, 59, 342380)
).to_csv_file() #example time range specified in datetime functions
```
You can also retrieve data that comes after a specific period in time. The following code retrieves data after the time specified by the timestamp in the `as_of` method.

```python
fg1 = FeatureGroup("fg1")
feature_store.create_dataset(
    base=fg1,
    output_path="s3://example-s3-file-path"
).as_of(datetime(2021, 11, 28, 23, 55, 59, 342380)
).to_csv_file() # example datetime values
```

**Sample Amazon Athena queries**

You can write queries in Amazon Athena to create a dataset from your feature groups. You can also write queries that create a dataset from feature groups and a single pandas dataframe.

**Interactive Exploration**

This query selects the first 1000 records.

```sql
SELECT *
FROM <FeatureGroup.DataCatalogConfig.DatabaseName>.<FeatureGroup.DataCatalogConfig.TableName>
LIMIT 1000
```

**Latest snapshot without duplicates**

This query selects the latest non-duplicate records.

```sql
SELECT *
FROM
    (SELECT *,
        row_number()
    OVER (PARTITION BY <RecordIdentifierFeatureName>
```
Latest snapshot without duplicates and deleted records in the offline store

This query filters out any deleted records and selects non-duplicate records from the offline store.

```
SELECT *
FROM
    (SELECT *
     , row_number()
     OVER (PARTITION BY <RecordIdentifierFeatureName>
     ORDER BY <EventTimeFeatureName> desc, Api_Invocation_Time DESC, write_time DESC)
     AS row_num
     FROM
     <FeatureGroup.DataCatalogConfig.DatabaseName>.<FeatureGroup.DataCatalogConfig.TableName>
WHERE row_num = 1 AND
NOT is_deleted;
```

Time Travel without duplicates and deleted records in the offline store

This query filters out any deleted records and selects non-duplicate records from a particular point in time.

```
SELECT *
FROM
    (SELECT *
     , row_number()
     OVER (PARTITION BY <RecordIdentifierFeatureName>
     ORDER BY <EventTimeFeatureName> desc, Api_Invocation_Time DESC, write_time DESC)
     AS row_num
     FROM
     <FeatureGroup.DataCatalogConfig.DatabaseName>.<FeatureGroup.DataCatalogConfig.TableName>
WHERE <EventTimeFeatureName> <= timestamp '<timestamp>'
-- replace timestamp '<timestamp>' with just <timestamp> if EventTimeFeature is of type fractional
WHERE row_num = 1 AND
NOT is_deleted;
```
Delete records from your feature groups

You can use the Amazon SageMaker Feature Store API to delete records from your feature groups. A feature group is an object that contains your machine learning (ML) data, where the columns of your data are described by features and your data are contained in records. A record contains values for features that are associated with a specific record identifier.

There are two storage configurations for your feature groups: online store and offline store. The online store only keeps the record with the latest event time and is typically used for real-time lookup for ML inference. The offline store keeps all records and acts as a historical database and is typically used for feature exploration, ML training, and batch inference.

For more information on Feature Store concepts, see Ingestion diagrams.

There are two ways to delete records from your feature groups, and the behavior is different depending on the storage configuration. In the following topics we will describe how to soft and hard delete records from the online and offline stores and provide examples.

Topics

- Delete records from the online store
- Delete records from the offline store

Delete records from the online store

You can soft or hard delete a record from the online store using the DeleteRecord API by using the DeletionMode request parameter to specify SoftDelete (default) or HardDelete. For more information on the DeleteRecord API, see DeleteRecord in the Amazon SageMaker API Reference.

With the online store:

- When you soft delete (default), the record is no longer retrievable by GetRecord or BatchGetRecord and the feature column values are set to null, except for the RecordIdentifier and EventTime feature values.
- When you hard delete, the record is completely removed from the online store.

In both cases Feature Store appends the deleted record marker to the OfflineStore. The deleted record marker is a record with the same RecordIdentifier as the original, but with is_deleted
value set to True, EventTime set to the delete input EventTime, and other feature values set to null.

Note that the EventTime specified in DeleteRecord should be set later than the EventTime of the existing record in the OnlineStore for that same RecordIdentifier. If it is not, the deletion does not occur:

- For SoftDelete, the existing (not deleted) record remains in the OnlineStore, though the delete record marker is still written to the OfflineStore.
- HardDelete returns EventTime: 400 ValidationException to indicate that the delete operation failed. No delete record marker is written to the OfflineStore.

The following examples use the SDK for Python (Boto3) delete_record operation to delete a record from a feature group. To delete a record from a feature group, you will need:

- Feature group name (feature-group-name)
- Record identifier value as a string (record-identifier-value)
- Deletion event time (deletion-event-time)

The deletion event time should be later than the event time of the record you wish to delete.

**Online store soft delete example**

For soft delete you will need use the DeleteRecord API and can use the default DeletionMode or set the DeletionMode to SoftDelete.

```python
import boto3
client = boto3.client('sagemaker-featurestore-runtime')

client.delete_record(
    FeatureGroupName='feature-group-name',
    RecordIdentifierValueAsString='record-identifier-value',
    EventTime='deletion-event-time',
    TargetStores=['OnlineStore'],
    DeletionMode='SoftDelete'
)
```

Delete records from the online store
Online store hard delete example

For hard delete you will need use the DeleteRecord API and set the DeletionMode to HardDelete.

```python
import boto3
client = boto3.client('sagemaker-featurestore-runtime')

client.delete_record(
    FeatureGroupName='feature-group-name',
    RecordIdentifierValueAsString='record-identifier-value',
    EventTime='deletion-event-timestamp',
    TargetStores=['OnlineStore'],
    DeletionMode='HardDelete'
)
```

Delete records from the offline store

With Amazon SageMaker Feature Store you can soft and hard delete a record from the OfflineStore Iceberg table format. With the OfflineStore Iceberg table format:

- When you soft delete a record the latest version of the Iceberg table file will not contain the record, but previous versions will still contain the record and can be accessed using time travel. For information on time travel, see [Querying Iceberg table data and performing time travel](#) in the Athena user guide.
- When you hard delete a record you are removing previous versions of the Iceberg table that contain the record. In this case you should specify which versions of the Iceberg table you wish to delete.

Obtain your Iceberg table name

To soft and hard delete from your OfflineStore Iceberg table, you will need to obtain your Iceberg table name, `iceberg-table-name`. The following instructions assumes you have already used Feature Store to create a feature group using the offline store storage configuration using the Iceberg table format, with `DisableGlueTableCreation = False` (default). For more information on creating feature groups, see [Get started with Amazon SageMaker Feature Store](#).
To obtain your `iceberg-table-name`, use the DescribeFeatureGroup API to obtain DataCatalogConfig. This contains the metadata of the Glue table which serves as data catalog for the OfflineStore. The TableName within the DataCatalogConfig is your `iceberg-table-name`.

**Amazon Athena offline store soft and hard delete example**

The following instructions use Amazon Athena to soft delete then hard delete a record from the OfflineStore Iceberg table. This assumes that the record you intend to delete in your OfflineStore is a deleted record marker. For information on the deleted record marker in your OfflineStore, see Delete records from the online store.

1. Obtain your Iceberg table name, `iceberg-table-name`. For information on how to obtain your Iceberg table name, see Obtain your Iceberg table name.

2. Run the DELETE command to soft delete the records on the OfflineStore, such that the latest version (or snapshot) of the Iceberg table will not contain the records. The following example deletes the records where is_deleted is 'True' and the previous event-time versions of the those records. You may add additional conditions based on other features to restrict the deletion. For more information on using DELETE with Athena, see DELETE in the Athena user guide.

```sql
DELETE FROM iceberg-table-name WHERE record-id-feature-name IS IN ( SELECT record-id-feature-name FROM iceberg-table-name WHERE is_deleted = 'True')
```

The soft deleted records are still viewable on previous file versions by performing time travel. For information on performing time travel, see Querying Iceberg table data and performing time travel in the Athena user guide.

3. Remove the record from previous versions of your Iceberg tables to hard delete the record from OfflineStore:

   a. Run the OPTIMIZE command to rewrite the data files into a more optimized layout, based on their size and number of associated delete files. For more information on optimizing Iceberg tables and the syntax, see Optimizing Iceberg tables in the Athena user guide.

```sql
OPTIMIZE iceberg-table-name REWRITE DATA USING BIN_PACK
```

   b. (Optional, only need to run once) Run the ALTER TABLE command to alter the Iceberg table set values, and set when previous file versions are to be hard
deleted according to your specifications. This can be done by assigning values to vacuum_min_snapshots_to_keep and vacuum_max_snapshot_age_seconds properties. For more information on altering your Iceberg table set properties, see ALTER TABLE SET PROPERTIES in the Athena user guide. For more information on Iceberg table property key-value pairs, see Table properties in the Athena user guide.

```
ALTER TABLE iceberg-table-name SET TBLPROPERTIES (  
    'vacuum_min_snapshots_to_keep'='your-specified-value',  
    'vacuum_max_snapshot_age_seconds'='your-specified-value'  
)
```

c. Run the VACUUM command to remove no longer needed data files for your Iceberg tables, not referenced by the current version. The VACUUM command should run after the deleted record is no longer referenced in the current snapshot. For example, vacuum_max_snapshot_age_seconds after the deletion. For more information on VACUUM with Athena and the syntax, see VACUUM.

```
VACUUM iceberg-table-name
```

Apache Spark offline store soft and hard delete example

To soft and then hard delete a record from the OfflineStore Iceberg table using Apache Spark, you can follow the same instructions as in the Amazon Athena offline store soft and hard delete example above, but using Spark procedures. For a full list of procedures, see Spark Procedures in the Apache Iceberg documentation.

- When soft deleting from the OfflineStore: instead of using the DELETE command in Athena, use the DELETE_FROM command in Apache Spark.
- To remove the record from previous versions of your Iceberg tables to hard delete the record from OfflineStore:
  - When changing your Iceberg table configuration: instead of using the ALTER TABLE command from Athena, use expire_snapshots procedure.
  - To remove no longer needed data files from your Iceberg tables: instead of using the VACUUM command in Athena, use the remove_orphan_files procedure.
Logging Feature Store operations by using AWS CloudTrail

Amazon SageMaker Feature Store is integrated with AWS CloudTrail, a service that provides a record of actions taken by a user, role, or an AWS service in Feature Store. CloudTrail captures all of the API calls for Feature Store listed on this page. The logged events include API calls from Feature Store resource management and data operations. When you create a trail, you activate continuous delivery of CloudTrail events from Feature Store to an Amazon S3 bucket. Using the information collected by CloudTrail, you can determine the request that was made to Feature Store, the IP address from which the request was made, who made the request, when it was made, and additional details.

To learn more about CloudTrail, see the [AWS CloudTrail User Guide](#).

**Management events**

Management events capture operations performed on Feature Store resources in your AWS account. For example, the log generated from the management events provides visibility if a user creates or deletes a Feature Store. The following APIs log management events with Amazon SageMaker Feature Store.

- `CreateFeatureGroup`
- `DeleteFeatureGroup`
- `DescribeFeatureGroup`
- `UpdateFeatureGroup`

Amazon SageMaker API calls and management events are logged by default when you create the account, as described in [Log Amazon SageMaker API Calls with AWS CloudTrail](#). For more information, see [Logging management events for trails](#).

**Data events**

Data events capture data plane operations performed using the Feature Store resources in your AWS account. For example, the log generated from the data events provides visibility if a user adds or deletes a record within a feature group. The following APIs log data events with Amazon SageMaker Feature Store.

- `BatchGetRecord`
Data events are not logged by CloudTrail trails by default. To activate logging of data events, turn on logging of data plane API activity in CloudTrail. For more information, see CloudTrail's [Logging data events for trails](#).

The following is an example CloudTrail event for a PutRecord API call:

```json
{
  "eventVersion": "1.08",
  "userIdentity": {
    "type": "IAMUser",
    "principalId": "USERPRINCIPALID",
    "arn": "arn:aws:iam::123456789012:user/user",
    "accountId": "123456789012",
    "accessKeyId": "USERACCESSKEYID",
    "userName": "your-user-name"
  },
  "eventTime": "2023-01-01T01:00:00Z",
  "eventSource": "sagemaker.amazonaws.com",
  "eventName": "PutRecord",
  "awsRegion": "us-east-1",
  "sourceIPAddress": "192.0.2.0",
  "userAgent": "your-user-agent",
  "requestParameters": {
    "featureGroupName": "your-feature-group-name"
  },
  "responseElements": null,
  "requestID": "request-id",
  "eventID": "event-id",
  "readOnly": false,
  "resources": [
    {
      "accountId": "123456789012",
      "type": "AWS::SageMaker::FeatureGroup",
      "ARN": "arn:aws:sagemaker:us-east-1:123456789012:feature-group/your-feature-group-name"
    }
  ],
  "eventType": "AwsApiCall",
```
Security and access control

Amazon SageMaker Feature Store enables you to create two types of stores: an online store or offline store. The online store is used for low latency real-time inference use cases whereas the offline store is used for training and batch inference use cases. When you create a feature group for online or offline use you can provide a AWS Key Management Service customer managed key to encrypt all your data at rest. In case you do not provide a AWS KMS key then we ensure that your data is encrypted on the server side using an AWS owned AWS KMS key or AWS managed AWS KMS key. While creating a feature group, you can select storage type and optionally provide a AWS KMS key for encrypting data, then you can call various APIs for data management such as PutRecord, GetRecord, DeleteRecord.

Feature Store allows you to grant or deny access to individuals at the feature group-level and enables cross-account access to Feature Store. For example, you can set up developer accounts to access the offline store for model training and exploration that do not have write access to production accounts. You can set up production accounts to access both online and offline stores. Feature Store uses unique customer AWS KMS keys for offline and online store data at-rest encryption. Access control is enabled through both API and AWS KMS key access. You can also create feature group-level access control.

For more information about customer managed key, see customer managed keys. For more information about AWS KMS, see AWS KMS.

Using AWS KMS permissions for Amazon SageMaker Feature Store

Encryption at rest protects Feature Store under an AWS KMS customer managed key. By default, it uses an AWS owned customer managed key for OnlineStore and AWS managed customer managed key for OfflineStore. Feature Store supports an option to encrypt your online or offline store under customer managed key. You can select the customer managed key for Feature Store when you create your online or offline store, and they can be different for each store.
Feature Store supports only symmetric customer managed keys. You cannot use an asymmetric customer managed key to encrypt your data in your online or offline store. For help determining whether a customer managed key is symmetric or asymmetric, see Identifying symmetric and asymmetric customer managed keys.

When you use a customer managed key, you can take advantage of the following features:

- You create and manage the customer managed key, including setting the key policies, IAM policies and grants to control access to the customer managed key. You can enable and disable the customer managed key, enable and disable automatic key rotation, and delete the customer managed key when it is no longer in use.
- You can use a customer managed key with imported key material or a customer managed key in a custom key store that you own and manage.
- You can audit the encryption and decryption of your online or offline store by examining the API calls to AWS KMS in AWS CloudTrail logs.

You do not pay a monthly fee for AWS owned customer managed keys. Customer managed keys will incur a charge for each API call and AWS Key Management Service quotas apply to each customer managed key.

Authorizing use of a customer managed Key for your online store

If you use a customer managed key to protect your online store, the policies on that customer managed key must give Feature Store permission to use it on your behalf. You have full control over the policies and grants on a customer managed key.

Feature Store does not need additional authorization to use the default AWS owned KMS key to protect your online or offline stores in your AWS account.

Customer managed key policy

When you select a customer managed key to protect your Online Store, Feature Store must have permission to use the customer managed key on behalf of the principal who makes the selection. That principal, a user or role, must have the permissions on the customer managed key that Feature Store requires. You can provide these permissions in a key policy, an IAM policy, or a grant. At a minimum, Feature Store requires the following permissions on a customer managed key:
"kms:Encrypt", "kms:Decrypt", "kms:DescribeKey", "kms:CreateGrant", "kms:RetireGrant",
"kms:ReEncryptFrom", "kms:ReEncryptTo", "kms:GenerateDataKey", "kms:ListAliases",
"kms:ListGrants", "kms:RevokeGrant"

For example, the following example key policy provides only the required permissions. The policy has the following effects:

- Allows Feature Store to use the customer managed key in cryptographic operations and create grants, but only when it is acting on behalf of principals in the account who have permission to use your Feature Store. If the principals specified in the policy statement don't have permission to use your Feature Store, the call fails, even when it comes from the Feature Store service.

- The kms:ViaService condition key allows the permissions only when the request comes from FeatureStore on behalf of the principals listed in the policy statement. These principals can't call these operations directly. The value for kms:ViaService should be sagemaker.*.amazonaws.com.

Before using an example key policy, replace the example principals with actual principals from your AWS account.

```json
{"Id": "key-policy-feature-store",
 "Version":"2012-10-17",
 "Statement": [
  {"Sid": "Allow access through Amazon SageMaker Feature Store for all principals in the account that are authorized to use Amazon SageMaker Feature Store",
   "Effect": "Allow",
   "Principal": {"AWS": "arn:aws:iam::111122223333:user/featurestore-user"},
   "Action": [
```
"kms:Encrypt",
"kms:Decrypt",
"kms:DescribeKey",
"kms:CreateGrant",
"kms:RetireGrant",
"kms:ReEncryptFrom",
"kms:ReEncryptTo",
"kms:GenerateDataKey",
"kms:ListAliases",
"kms:ListGrants"
],
"Resource": "*",
"Condition": {"StringLike": {"kms:ViaService": "sagemaker.*.amazonaws.com"
}
},
{"Sid": "Allow administrators to view the customer managed key and revoke grants",
"Effect": "Allow",
"Principal": {"AWS": "arn:aws:iam::111122223333:role/featurestore-admin"
},
"Action": [
"kms:Describe*",
"kms:Get*",
"kms:List*",
"kms:RevokeGrant"
],
"Resource": "*
",
{"Sid": "Enable IAM User Permissions",
"Effect": "Allow",
"Principal": {"AWS": "arn:aws:iam::123456789:root"
},
"Action": "kms:*",
"Resource": "*
"}
]
Using grants to authorize Feature Store

In addition to key policies, Feature Store uses grants to set permissions on the customer managed key. To view the grants on a customer managed key in your account, use the `ListGrants` operation. Feature Store does not need grants, or any additional permissions, to use the AWS owned customer managed key to protect your online store.

Feature Store uses the grant permissions when it performs background system maintenance and continuous data protection tasks.

Each grant is specific to an online store. If the account includes multiple stores encrypted under the same customer managed key, there will be unique grants per FeatureGroup using the same customer managed key.

The key policy can also allow the account to revoke the grant on the customer managed key. However, if you revoke the grant on an active encrypted online store, Feature Store won't be able to protect and maintain the store.

Monitoring Feature Store interaction with AWS KMS

If you use a customer managed key to protect your online or offline store, you can use AWS CloudTrail logs to track the requests that Feature Store sends to AWS KMS on your behalf.

Accessing data in your online store

The caller (either user or role) to ALL DataPlane operations (Put, Get, DeleteRecord) must have below permissions on the customer managed key:

"kms:Decrypt"

Authorizing use of a customer managed key for your offline store

The roleArn that is passed as a parameter to `createFeatureGroup` must have below permissions to the OfflineStore KmsKeyId:

"kms:GenerateDataKey"
Note
The key policy for the online store also works for the offline store, only when the kms:ViaService condition is not specified.

Important
You can specify an AWS KMS encryption key to encrypt the Amazon S3 location used for your offline feature store when you create a feature group. If AWS KMS encryption key is not specified, by default we encrypt all data at rest using AWS KMS key. By defining your bucket-level key for SSE, you can reduce AWS KMS requests costs by up to 99 percent.

Quotas, naming rules and data types

Quota terminologies

- **Read Request Unit (RRU):** Measure of read throughput, where the number of RRUs per read request is equal to the ceiling of read record's size divided into 4KB chunks. The minimum RRU per request is 0.
- **Write Request Unit (WRU):** Measure of write throughput, where the number of WRUs per write request is equal to the ceiling of the written record's size divided into 1KB chunks. The minimum WRU per request is 1 (including delete operations).

Limits and quotas

Note
Soft limits can be increased based on your needs.

- **Maximum number of feature groups per AWS account:** Soft limit of 100.
- **Maximum number of feature definitions per feature group:** 2500.
- **Maximum number of RRU per record identifier:** 2400 RRU per second.
- **Maximum number of WRU per record identifier:** 500 WRU per second.
• **Max Read Capacity Units (RCU) that can be provisioned on a single feature group:** 40000 RCU.
• **Max Write Capacity Units (WCU) that can be provisioned on a single feature group:** 40000 WCU.
• **Max Read Capacity Units that can be provisioned across all feature groups in a region:** 80000 RCU.
• **Max Write Capacity Units that can be provisioned across all feature groups in a region:** 80000 WCU.
• **Maximum Transactions per second (TPS) per API per AWS account:** Soft limit of 10000 TPS per API excluding the BatchGetRecord API call, which has a soft limit of 500 TPS.
• **Maximum size of a record:** 350KB.
• **Maximum size of a record identifier:** 2KB.
• **Maximum size of a feature value:** 350KB.
• **Maximum number of concurrent feature group creation workflows:** 4.
• **BatchGetRecord API:** Can contain as many as 100 records and can query up to 100 feature groups.

For information about service quotas and how to request a quota increase, see [AWS service quotas](https://aws.amazon.com/).  

**Naming rules**

• **Reserved Words:** The following are reserved words and cannot be used as feature names in feature definitions: `is_deleted`, `write_time`, and `api_invocation_time`.

**Data types**

• **String Feature Type:** Strings are Unicode with UTF-8 binary encoding. The minimum length of a string can be zero, the maximum length is constrained by the maximum size of a record.
• **Fractional Feature Type:** Fractional feature values must conform to a double precision floating point number as defined by the [IEEE 754 standard](https://en.wikipedia.org/wiki/IEEE_754).
• **Integral Feature Type:** Feature Store supports integral values in the range of a 64-bit signed integer. Minimum value of $-2^{63}$ and a maximum value: $2^{63} - 1$.
• **Event Time Features:** All feature groups have an event time feature with nanosecond precision. Any event time with lower than nanosecond precision will lead to backwards incompatibility. The feature can have a feature type of either String or Fractional.
A string event time is accepted in ISO-8601 format, in UTC time, conforming to the pattern(s): [yyyy-MM-dd'T'HH:mm:ssZ, yyyy-MM-dd'T'HH:mm:ss.SSSSSSSSSSZ].

A fractional event time value is accepted as seconds from unix epoch. Event times must be in the range of [0000-01-01T00:00:00.000000000Z, 9999-12-31T23:59:59.999999999Z]. For feature groups in the Iceberg table format, you can only use String type for the event time.

**Amazon SageMaker Feature Store offline store data format**

Amazon SageMaker Feature Store supports the AWS Glue and Apache Iceberg table formats for the offline store. You can choose the table format when you're creating a new feature group. AWS Glue is the default format.

Amazon SageMaker Feature Store offline store data is stored in an Amazon S3 bucket within your account. When you call `PutRecord`, your data is buffered, batched, and written into Amazon S3 within 15 minutes. Feature Store only supports the Parquet file format when writing your data to your offline store. Specifically, when your data is written to your offline store, the data can be retrieved from your Amazon S3 bucket in Parquet format. Each file can contain multiple Records.

For the Iceberg format, Feature Store saves the table's metadata in the same Amazon S3 bucket that you're using to store the offline store data. You can find it under the `metadata` prefix.

Feature Store also exposes the `OfflineStoreConfig.S3StorageConfig.ResolvedOutputS3Uri` field, which can be found from in the `DescribeFeatureGroup` API call. This is the S3 path under which the files for the specific feature group are written.

The following additional fields are added by Feature Store to each record when they persist in the offline store:

- **api_invocation_time** – The timestamp when the service receives the `PutRecord` or `DeleteRecord` call. If using managed ingestion (e.g. Data Wrangler), this is the timestamp when data was written into the offline store.

- **write_time** – The timestamp when data was written into the offline store. Can be used for constructing time-travel related queries.

- **is_deleted** – False by default. If `DeleteRecord` is called, a new `Record` is inserted into `RecordIdentifierValue` and set to `True` in the offline store.
Amazon SageMaker Feature Store offline store URI structures

In the following examples `DOC-EXAMPLE-BUCKET` is the Amazon S3 bucket within your account, `example-prefix` is your example prefix, `111122223333` is your account ID, `AWS Region` is your region, `feature-group-name` is the name of your feature group.

**AWS Glue table format**

Records in the offline store stored using the AWS Glue table format are partitioned by event time into hourly partitions. You can’t configure the partitioning scheme. The following URI structure shows the organization of a Parquet file using the AWS Glue format:

```
s3://DOC-EXAMPLE-BUCKET/example-prefix/111122223333/sagemaker/AWS Region/offline-store/feature-group-name-feature-group-creation-time/data/year=year/month=month/day=day/hour=hour/timestamp_of_latest_event_time_in_file_16-random-alphanumeric-digits.parquet
```

The following example is the output location of a Parquet file for a file with `feature-group-name` as `customer-purchase-history-patterns`:

```
s3://DOC-EXAMPLE-BUCKET/example-prefix/111122223333/sagemaker/AWS Region/offline-store/customer-purchase-history-patterns-1593511200/data/year=2020/month=06/day=31/hour=00/20200631T064401Z_108934320012Az11.parquet
```

**Iceberg table format**

Records in the offline store stored in the Iceberg table format are partitioned by event time into daily partitions. You can’t configure the partitioning scheme. The following URI structure shows the organization of the data files saved in the Iceberg table format:

```
```

The following example is the output location of a Parquet file for a file with `feature-group-name` as `customer-purchase-history-patterns`, and the `event-time-feature-name` is `EventTime`:
The following example is the location of a metadata file for data files saved in the Iceberg table format.

```
s3://DOC-EXAMPLE-BUCKET/example-prefix/111122223333/sagemaker/AWS Region/offline-store/feature-group-name-feature-group-creation-time/metadata/
```

## Amazon SageMaker Feature Store resources

The following lists the available resources for Amazon SageMaker Feature Store users. For the Feature Store main page, see [Amazon SageMaker Feature Store](#).

### Feature Store example notebooks and workshops

To get started using Amazon SageMaker Feature Store, you can choose from a variety of example Jupyter notebooks from the following table. If this is your first time using Feature Store, try out the Introduction to Feature Store notebook. To run any these notebooks, you must attach this policy to your IAM execution role: AmazonSageMakerFeatureStoreAccess.

See [IAM Roles](#) to access your role and attach this policy. For a walkthrough on how to view the policies attached to a role and how to add a policy to your role, see [Adding policies to your IAM role](#).

The following table lists a variety of resources to help you get started with Feature Store. This table contains examples, instructions, and example notebooks to guide you in how to use Feature Store for the first time to specific use cases. The code in these resources use the SageMaker SDK for Python (Boto3).

<table>
<thead>
<tr>
<th>Page</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Get started with Amazon SageMaker Feature Store in Read the Docs.</td>
<td>A list of example notebooks to introduce you to Feature Store and its features to help you get started.</td>
</tr>
</tbody>
</table>
Feature Store Python SDK and API

Python Software Development Kit (SDK) and Application Programming Interface (API) are tools used for creating software applications. The Feature Store SDK for Python (Boto3) and API are listed in the following table.

<table>
<thead>
<tr>
<th>Page</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feature Store APIs in the Amazon SageMaker Python SDK Read the Docs</td>
<td>The Feature Store APIs in Read the Docs.</td>
</tr>
<tr>
<td>Feature Store Python SDK in the Amazon SageMaker Python SDK Github repository</td>
<td>The Feature Store Python SDK Github repository.</td>
</tr>
<tr>
<td>Feature Store Runtime operations and data types in the SDK for Python (Boto3) documentation</td>
<td>Feature Store Runtime client that contains all data plane API operations and data types for Feature Store.</td>
</tr>
<tr>
<td>Amazon SageMaker Feature Store Runtime in the Amazon SageMaker API Reference</td>
<td>Some feature group level actions supported by Feature Store. If the API operation or data type you are looking for is not listed here, please use search in the guide.</td>
</tr>
<tr>
<td>Page</td>
<td>Description</td>
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</tr>
<tr>
<td><strong>Amazon SageMaker Feature Store Runtime</strong> in the Amazon SageMaker API Reference</td>
<td>Record level actions supported by Feature Store. If the API operation or data type you are looking for is not listed here, please use search in the guide.</td>
</tr>
</tbody>
</table>
Train machine learning models

The training stage of the full machine learning (ML) lifecycle spans from accessing your training dataset to generating a final model and selecting the best performing model for deployment. The following sections provide an overview of available SageMaker training features and resources with in-depth technical information for each.

The simplest training workflow in SageMaker

If you’re using SageMaker for the first time and want to find a quick ML solution to train a model on your dataset, consider using a no-code or low-code solution such as SageMaker Canvas, SageMaker JumpStart within SageMaker Studio Classic, or SageMaker Autopilot.

For intermediate coding experiences, consider using a SageMaker Studio Classic notebook or SageMaker Notebook Instances. To get started, follow the instructions at the section called “Step 4: Train a Model” of the SageMaker Getting Started guide. We recommend this for use cases in which you create your own model and training script using an ML framework.

The following architecture diagram shows how SageMaker manages ML training jobs and provisions Amazon EC2 instances on behalf of SageMaker users. You as a SageMaker user can bring your own training dataset, saving it to Amazon S3. You can choose an ML model training from available SageMaker built-in algorithms, or bring your own training script with a model built with popular machine learning frameworks.
Full view of the SageMaker Training workflow and features

The full journey of ML training involves tasks beyond data ingestion to ML models, training models on compute instances, and obtaining model artifacts and outputs. You need to evaluate every phase of before, during, and after training to make sure your model is trained well to meet the target accuracy for your objectives.

The following flow chart shows a high-level overview of your actions (in blue boxes) and available SageMaker Training features (in light blue boxes) throughout the training phase of the ML lifecycle.
The following sections walk you through each phase of training depicted in the previous flow chart and useful features offered by SageMaker throughout the three sub-stages of the ML training.

**Topics**

- Before training
- During training
- After training
Before training

There are a number of scenarios of setting up data resources and access you need to consider before training. Refer to the following diagram and details of each before-training stage to get a sense of what decisions you need to make.

- **Prepare data**: Before training, you must have finished data cleaning and feature engineering during the data preparation stage. SageMaker has several labeling and feature engineering tools
to help you. See Label Data, Prepare and Analyze Datasets, Process Data, and Create, Store, and Share Features for more information.

- **Choose an algorithm or framework:** Depending on how much customization you need, there are different options for algorithms and frameworks.
  - If you prefer a low-code implementation of a pre-built algorithm, use one of the built-in algorithms offered by SageMaker. For more information, see Choose an Algorithm.
  - If you need more flexibility to customize your model, run your training script using your preferred frameworks and toolkits within SageMaker. For more information, see ML Frameworks and Toolkits.
  - To extend pre-built SageMaker Docker images as the base image of your own container, see Use Pre-built SageMaker Docker images.
  - To bring your custom Docker container to SageMaker, see Adapting your own Docker container to work with SageMaker. You need to install the sagemaker-training-toolkit to your container.

- **Manage data storage:** Understand mapping between the data storage (such as Amazon S3, Amazon EFS, or Amazon FSx) and the training container that runs in the Amazon EC2 compute instance. SageMaker helps map the storage paths and local paths in the training container. You can also manually specify them. After mapping is done, consider using one of the data transmission modes: File, Pipe, and FastFile mode. To learn how SageMaker maps storage paths, see Training Storage Folders.

- **Set up access to training data:** Use Amazon SageMaker domain, a domain user profile, IAM, Amazon VPC, and AWS KMS to meet the requirements of the most security-sensitive organizations.
  - For account administration, see Amazon SageMaker domain.
  - For a complete reference about IAM policies and security, see Security in Amazon SageMaker.

- **Stream your input data:** SageMaker provides three data input modes, File, Pipe, and FastFile. The default input mode is File mode, which loads the entire dataset during initializing the training job. To learn about general best practices for streaming data from your data storage to the training container, see Access Training Data.

  In case of Pipe mode, you can also consider using an augmented manifest file to stream your data directly from Amazon Simple Storage Service (Amazon S3) and train your model. Using pipe mode reduces disk space because Amazon Elastic Block Store only needs to store your final model artifacts, rather than storing your full training dataset. For more information, see Provide Dataset Metadata to Training Jobs with an Augmented Manifest File.
**Analyze your data for bias:** Before training, you can analyze your dataset and model for bias against a disfavored group so that you can check that your model learns an unbiased dataset using [SageMaker Clarify](#).

**Choose which SageMaker SDK to use:** There are two ways to launch a training job in SageMaker: using the high-level SageMaker Python SDK, or using the low-level SageMaker APIs for the SDK for Python (Boto3) or the AWS CLI. The SageMaker Python SDK abstracts the low-level SageMaker API to provide convenient tools. As aforementioned in the section called “The simplest training workflow in SageMaker”, you can also pursue no-code or minimal-code options using [SageMaker Canvas](#), [SageMaker JumpStart within SageMaker Studio Classic](#), or [SageMaker Autopilot](#).

### During training

During training, you need to continuously improve training stability, training speed, training efficiency while scaling compute resources, cost optimization, and, most importantly, model performance. Read on for more information about during-training stages and relevant SageMaker Training features.

*Diagram of decisions to make during training*
• **Set up infrastructure**: Choose the right instance type and infrastructure management tools for your use case. You can start from a small instance and scale up depending on your workload. For training a model on a tabular dataset, start with the smallest CPU instance of the C4 or C5 instance families. For training a large model for computer vision or natural language processing, start with the smallest GPU instance of the P2, P3, G4dn or G5 instance families. You can also mix different instance types in a cluster, or keep instances in warm pools using the following instance management tools offered by SageMaker. You can also use persistent cache to reduce latency and billable time on iterative training jobs over the latency reduction from warm pools alone. To learn more, see the following topics.

  - [Train Using a Heterogeneous Cluster](#)
  - [Train Using SageMaker Managed Warm Pools](#)
  - [Using persistent cache](#)

You must have sufficient quota to run a training job. If you run your training job on an instance where you have insufficient quota, you will receive a `ResourceLimitExceeded` error. To check the currently available quotas in your account, use your [Service Quotas console](#). To learn how to request a quota increase, see [Supported Regions and Quotas](#). Also, to find pricing information and available instance types depending on the AWS Regions, look up the tables in the [Amazon SageMaker Pricing](#) page.

• **Run a training job from a local code**: You can annotate your local code with a remote decorator to run your code as a SageMaker training job from inside Amazon SageMaker Studio Classic, an Amazon SageMaker notebook or from your local integrated development environment. For more information, see [Run your local code as a SageMaker training job](#).

• **Track training jobs**: Monitor and track your training jobs using SageMaker Experiments, SageMaker Debugger, or Amazon CloudWatch. You can watch the model performance in terms of accuracy and convergence, and run comparative analysis of metrics between multiple training jobs by using SageMaker Experiments. You can watch the compute resource utilization rate by using SageMaker Debugger's profiling tools or Amazon CloudWatch. To learn more, see the following topics.

  - [Manage Machine Learning with Amazon SageMaker Experiments](#)
  - [Profile Training Jobs Using Amazon SageMaker Debugger](#)
  - [Monitor and Analyze Using CloudWatch Metrics](#)

Additionally, for deep learning tasks, use the [Amazon SageMaker Debugger model debugging tools](#) and [built-in rules](#) to identify more complex issues in model convergence and weight update processes.
• **Distributed training:** If your training job is going into a stable stage without breaking due to misconfiguration of the training infrastructure or out-of-memory issues, you might want to find more options to scale your job and run over an extended period of time for days and even months. When you’re ready to scale up, consider distributed training. SageMaker provides various options for distributed computation from light ML workloads to heavy deep learning workloads.

For deep learning tasks that involve training very large models on very large datasets, consider using one of the [SageMaker distributed training strategies](https://docs.aws.amazon.com/sagemaker/latest/dg/distributed-training-strategies.html) to scale up and achieve data parallelism, model parallelism, or a combination of the two. You can also use [SageMaker Training Compiler](https://docs.aws.amazon.com/sagemaker/latest/dg/compiling-models.html) for compiling and optimizing model graphs on GPU instances. These SageMaker features support deep learning frameworks such as PyTorch, TensorFlow, and Hugging Face Transformers.

• **Model hyperparameter tuning:** Tune your model hyperparameters using [Automatic Model Tuning with SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/automatic-model-tuning.html). SageMaker provides hyperparameter tuning methods such as grid search and Bayesian search, launching parallel hyperparameter tuning jobs with early-stopping functionality for non-improving hyperparameter tuning jobs.

• **Checkpointing and cost saving with Spot instances:** If training time is not a big concern, you might consider optimizing model training costs with managed Spot instances. Note that you must activate checkpointing for Spot training to keep restoring from intermittent job pauses due to Spot instance replacements. You can also use the checkpointing functionality to back up your models in case of unexpected training job termination. To learn more, see the following topics.

  • [Managed Spot Training](https://docs.aws.amazon.com/sagemaker/latest/dg/managed-spot-training.html)
  • [Use Checkpoints](https://docs.aws.amazon.com/sagemaker/latest/dg/checkpoints.html)

**After training**

After training, you obtain a final model artifact to use for model deployment and inference. There are additional actions involved in the after-training phase as shown in the following diagram.
• **Obtain baseline model:** After you have the model artifact, you can set it as a baseline model. Consider the following post-training actions and using SageMaker features before moving on to model deployment to production.

• **Examine model performance and check for bias:** Use Amazon CloudWatch Metrics and SageMaker Clarify for post-training bias to detect any bias in incoming data and model over time against the baseline. You need to evaluate your new data and model predictions against the new data regularly or in real time. Using these features, you can receive alerts about any acute changes or anomalies, as well as gradual changes or drifts in data and model.

• You can also use the Incremental Training functionality of SageMaker to load and update your model (or fine-tune) with an expanded dataset.

• You can register model training as a step in your SageMaker Pipeline or as part of other Workflow features offered by SageMaker in order to orchestrate the full ML lifecycle.
Train a Model with Amazon SageMaker

The following diagram shows how you train and deploy a model with Amazon SageMaker. Your training code accesses your training data and outputs model artifacts from an S3 bucket. Then you can make requests to a model endpoint to run inference. You can store both the training and inference container images in an Amazon Elastic Container Registry (ECR).

The following guide highlights two components of SageMaker: model training and model deployment.

To train a model in SageMaker, you create a training job. The training job includes the following information:
• The URL of the Amazon Simple Storage Service (Amazon S3) bucket where you've stored the training data.

• The compute resources that you want SageMaker to use for model training. Compute resources are machine learning (ML) compute instances that are managed by SageMaker.

• The URL of the S3 bucket where you want to store the output of the job.

• The Amazon Elastic Container Registry path where the training code is stored. For more information, see Docker Registry Paths and Example Code.

**Note**

Your input dataset must be in the same AWS Region as your training job.

You have the following options for a training algorithm:

• **Use an algorithm provided by SageMaker**—SageMaker provides dozens of built-in training algorithms and hundreds of pre-trained models. If one of these meets your needs, it's a great out-of-the-box solution for quick model training. For a list of algorithms provided by SageMaker, see Use Amazon SageMaker Built-in Algorithms or Pre-trained Models. To try an exercise that uses an algorithm provided by SageMaker, see Get started. You can also use SageMaker JumpStart to use algorithms and models through the Studio Classic UI.

• **Use SageMaker Debugger**—to inspect training parameters and data throughout the training process when working with the TensorFlow, PyTorch, and Apache MXNet learning frameworks or the XGBoost algorithm. Debugger automatically detects and alerts users to commonly occurring errors such as parameter values getting too large or small. For more information about using Debugger, see Use Amazon SageMaker Debugger to debug and improve model performance. Debugger sample notebooks are available at Amazon SageMaker Debugger Samples.

• **Use Apache Spark with SageMaker**—SageMaker provides a library that you can use in Apache Spark to train models with SageMaker. Using the library provided by SageMaker is similar to using Apache Spark MLLib. For more information, see Use Apache Spark with Amazon SageMaker.

• **Submit custom code to train with deep learning frameworks**—You can submit custom Python code that uses TensorFlow, PyTorch, or Apache MXNet for model training. For more information, see Use TensorFlow with Amazon SageMaker, Use PyTorch with Amazon SageMaker, and Use Apache MXNet with Amazon SageMaker.
• **Use your own custom algorithms**—Put your code together as a Docker image and specify the registry path of the image in a SageMaker CreateTrainingJob API call. For more information, see [Use Docker containers to build models](#).

• **Use an algorithm that you subscribe to from AWS Marketplace**—For information, see [Find and Subscribe to Algorithms and Model Packages on AWS Marketplace](#).

After you create the training job, SageMaker launches the ML compute instances and uses the training code and the training dataset to train the model. It saves the resulting model artifacts and other output in the S3 bucket you specified for that purpose.

You can create a training job with the SageMaker console or the API. For information about creating a training job with the API, see the [CreateTrainingJob](#) API.

When you create a training job with the API, SageMaker replicates the entire dataset on ML compute instances by default. To make SageMaker replicate a subset of the data on each ML compute instance, you must set the `S3DataDistributionType` field to `ShardedByS3Key`. You can set this field using the low-level SDK. For more information, see `S3DataDistributionType` in `S3DataSource`.

⚠️ **Important**

To prevent your algorithm container from contending for memory, we reserve memory for our SageMaker critical system processes on your ML compute instances and therefore you cannot expect to see all the memory for your instance type.

---

### Choose an Algorithm

Machine learning can help you accomplish empirical tasks that require some sort of inductive inference. This task involves induction as it uses data to train algorithms to make generalizable inferences. This means that the algorithms can make statistically reliable predictions or decisions, or complete other tasks when applied to new data that was not used to train them.

To help you select the best algorithm for your task, we classify these tasks on various levels of abstraction. At the highest level of abstraction, machine learning attempts to find patterns or relationships between features or less structured items, such as text in a data set. Pattern recognition techniques can be classified into distinct machine learning paradigms, each of which
address specific problem types. There are currently three basic paradigms for machine learning used to address various problem types:

- **Supervised learning**
- **Unsupervised learning**
- **Reinforcement learning**

The types of problems that each learning paradigm can address are identified by considering the inferences (or predictions, decisions, or other tasks) you want to make from the type of data that you have or could collect. Machine learning paradigms use algorithmic methods to address their various problem types. The algorithms provide recipes for solving these problems.

However, many algorithms, such as neural networks, can be deployed with different learning paradigms and on different types of problems. Multiple algorithms can also address a specific problem type. Some algorithms are more generally applicable and others are quite specific for certain kinds of objectives and data. So the mapping between machine learning algorithms and problem types is many-to-many. Also, there are various implementation options available for algorithms.

The following sections provide guidance concerning implementation options, machine learning paradigms, and algorithms appropriate for different problem types.

**Topics**

- [Choose an algorithm implementation](#)
- [Problem types for the basic machine learning paradigms](#)
- [Use Amazon SageMaker Built-in Algorithms or Pre-trained Models](#)
- [Use Reinforcement Learning with Amazon SageMaker](#)

**Choose an algorithm implementation**

After choosing an algorithm, you must decide which implementation of it you want to use. Amazon SageMaker supports three implementation options that require increasing levels of effort.

- **Pre-trained models** require the least effort and are models ready to deploy or to fine-tune and deploy using SageMaker JumpStart.
• **Built-in algorithms** require more effort and scale if the data set is large and significant resources are needed to train and deploy the model.

• If there is no built-in solution that works, try to develop one that uses **pre-made images for machine and deep learning frameworks** for supported frameworks such as Scikit-Learn, TensorFlow, PyTorch, MXNet, or Chainer.

• If you need to run custom packages or use any code which isn’t a part of a supported framework or available via PyPi, then you need to build **your own custom Docker image** that is configured to install the necessary packages or software. The custom image must also be pushed to an online repository like the Amazon Elastic Container Registry.

**Topics**

• [Use a built-in algorithm](#)

• [Use script mode in a supported framework](#)

• [Use a custom Docker image](#)

**Algorithm implementation guidance**

<table>
<thead>
<tr>
<th>Implementation</th>
<th>Requires code</th>
<th>Pre-coded algorithms</th>
<th>Support for third party packages</th>
<th>Support for custom code</th>
<th>Level of effort</th>
</tr>
</thead>
<tbody>
<tr>
<td>Built-in</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>Low</td>
</tr>
<tr>
<td>Scikit-learn</td>
<td>Yes</td>
<td>Yes</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium</td>
</tr>
<tr>
<td>Spark ML</td>
<td>Yes</td>
<td>Yes</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium</td>
</tr>
<tr>
<td>XGBoost (open source)</td>
<td>Yes</td>
<td>Yes</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium</td>
</tr>
<tr>
<td>TensorFlow</td>
<td>Yes</td>
<td>No</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium-high</td>
</tr>
<tr>
<td>PyTorch</td>
<td>Yes</td>
<td>No</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium-high</td>
</tr>
<tr>
<td>MXNet</td>
<td>Yes</td>
<td>No</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium-high</td>
</tr>
<tr>
<td>Chainer</td>
<td>Yes</td>
<td>No</td>
<td>PyPi only</td>
<td>Yes</td>
<td>Medium-high</td>
</tr>
</tbody>
</table>
Use a built-in algorithm

When choosing an algorithm for your type of problem and data, the easiest option is to use one of Amazon SageMaker's built-in algorithms. These built-in algorithms come with two major benefits.

- The built-in algorithms require no coding to start running experiments. The only inputs you need to provide are the data, hyperparameters, and compute resources. This allows you to run experiments more quickly, with less overhead for tracking results and code changes.
- The built-in algorithms come with parallelization across multiple compute instances and GPU support right out of the box for all applicable algorithms (some algorithms may not be included due to inherent limitations). If you have a lot of data with which to train your model, most built-in algorithms can easily scale to meet the demand. Even if you already have a pre-trained model, it may still be easier to use its corollary in SageMaker and input the hyper-parameters you already know than to port it over, using script mode on a supported framework.

For more information on the built-in algorithms provided by SageMaker, see Use Amazon SageMaker Built-in Algorithms or Pre-trained Models.

For important information about docker registry paths, data formats, recommended EC2 instance types, and CloudWatch logs common to all of the built-in algorithms provided by SageMaker, see Common Information About Built-in Algorithms.

Use script mode in a supported framework

If the algorithm you want to use for your model is not supported by a built-in choice and you are comfortable coding your own solution, then you should consider using an Amazon SageMaker supported framework. This is referred to as "script mode" because you write your custom code (script) in a text file with a .py extension. As the table above indicates, SageMaker supports most of the popular machine learning frameworks. These frameworks come preloaded with the corresponding framework and some additional Python packages, such as Pandas and NumPy,
so you can write your own code for training an algorithm. These frameworks also allow you to install any Python package hosted on PyPi by including a requirements.txt file with your training code or to include your own code directories. R is also supported natively in SageMaker notebook kernels. Some frameworks, like scikit-learn and Spark ML, have pre-coded algorithms you can use easily, while other frameworks like TensorFlow and PyTorch may require you to implement the algorithm yourself. The only limitation when using a supported framework image is that you cannot import any software packages that are not hosted on PyPi or that are not already included with the framework’s image.

For more information on the frameworks supported by SageMaker, see Machine Learning Frameworks and Languages.

Use a custom Docker image

Amazon SageMaker's built-in algorithms and supported frameworks should cover most use cases, but there are times when you may need to use an algorithm from a package not included in any of the supported frameworks. You might also have a pre-trained model picked or persisted somewhere which you need to deploy. SageMaker uses Docker images to host the training and serving of all models, so you can supply your own custom Docker image if the package or software you need is not included in a supported framework. This may be your own Python package or an algorithm coded in a language like Stan or Julia. For these images you must also configure the training of the algorithm and serving of the model properly in your Dockerfile. This requires intermediate knowledge of Docker and is not recommended unless you are comfortable writing your own machine learning algorithm. Your Docker image must be uploaded to an online repository, such as the Amazon Elastic Container Registry (ECR) before you can train and serve your model properly.

For more information on custom Docker images in SageMaker, see Use Docker containers to build models.

Problem types for the basic machine learning paradigms

The following three sections describe the main problem types addressed by the three basic paradigms for machine learning. For a list of the built-in algorithms that SageMaker provides to address these problem types, see Use Amazon SageMaker Built-in Algorithms or Pre-trained Models.

Topics

• Supervised learning
• **Unsupervised learning**

• **Reinforcement learning**

**Supervised learning**

If your data set consists of features or attributes (inputs) that contain target values (outputs), then you have a supervised learning problem. If your target values are categorical (mathematically discrete), then you have a **classification problem**. It is a standard practice to distinguish binary from multiclass classification.

• **Binary classification** is a type of supervised learning that assigns an individual to one of two predefined and mutually exclusive classes based on the individual's attributes. It is supervised because the models are trained using examples in which the attributes are provided with correctly labeled objects. A medical diagnosis for whether an individual has a disease or not based on the results of diagnostic tests is an example of binary classification.

• **Multiclass classification** is a type of supervised learning that assigns an individual to one of several classes based on the individual's attributes. It is supervised because the models are trained using examples in which the attributes are provided with correctly labeled objects. An example is the prediction of the topic most relevant to a text document. A document may be classified as being about religion, politics, or finance, or as about one of several other predefined topic classes.

If the target values you are trying to predict are mathematically continuous, then you have a **regression** problem. Regression estimates the values of a dependent target variable based on one or more other variables or attributes that are correlated with it. An example is the prediction of house prices using features like the number of bathrooms and bedrooms and the square footage of the house and garden. Regression analysis can create a model that takes one or more of these features as an input and predicts the price of a house.

For more information on the built-in supervised learning algorithms provided by SageMaker, see **Supervised Learning**.

**Unsupervised learning**

If your data set consists of features or attributes (inputs) that do not contain labels or target values (outputs), then you have an unsupervised learning problem. In this type of problem, the output must be predicted based on the pattern discovered in the input data. The goal in
unsupervised learning problems is to discover patterns such as groupings within the data. There are a large variety of tasks or problem types to which unsupervised learning can be applied. Principal component and cluster analyses are two of the main methods commonly deployed for preprocessing data. Here is a short list of problem types that can be addressed by unsupervised learning:

• **Dimension reduction** is typically part of a data exploration step used to determine the most relevant features to use for model construction. The idea is to transform data from a high-dimensional, sparsely populated space into a low-dimensional space that retains most significant properties of the original data. This provides relief for the curse of dimensionality that can arise with sparsely populated, high-dimensional data on which statistical analysis becomes problematic. It can also be used to help understand data, reducing high-dimensional data to a lower dimensionality that can be visualized.

• **Cluster analysis** is a class of techniques that are used to classify objects or cases into groups called clusters. It attempts to find discrete groupings within data, where members of a group are as similar as possible to one another and as different as possible from members of other groups. You define the features or attributes that you want the algorithm to use to determine similarity, select a distance function to measure similarity, and specify the number of clusters to use in the analysis.

• **Anomaly detection** is the identification of rare items, events, or observations in a data set which raise suspicions because they differ significantly from the rest of the data. The identification of anomalous items can be used, for example, to detect bank fraud or medical errors. Anomalies are also referred to as outliers, novelties, noise, deviations, and exceptions.

• **Density estimation** is the construction of estimates of unobservable underlying probability density functions based on observed data. A natural use of density estimates is for data exploration. Density estimates can discover features such as skewness and multimodality in the data. The most basic form of density estimation is a rescaled histogram.

SageMaker provides several built-in machine learning algorithms that you can use for these unsupervised learning tasks. For more information on the built-in unsupervised algorithms provided by SageMaker, see [Unsupervised Learning](#).

**Reinforcement learning**

Reinforcement learning is a type of learning that is based on interaction with the environment. This type of learning is used by an agent that must learn behavior through trial-and-error interactions with a dynamic environment in which the goal is to maximize the long-term rewards that the agent
receives as a result of its actions. Rewards are maximized by trading off exploring actions that have uncertain rewards with exploiting actions that have known rewards.

For more information on SageMaker's frameworks, toolkits, and environments for reinforcement learning, see Use Reinforcement Learning with Amazon SageMaker.

**Use Amazon SageMaker Built-in Algorithms or Pre-trained Models**

Amazon SageMaker provides a suite of built-in algorithms, pre-trained models, and pre-built solution templates to help data scientists and machine learning practitioners get started on training and deploying machine learning models quickly. For someone who is new to SageMaker, choosing the right algorithm for your particular use case can be a challenging task. The following table provides a quick cheat sheet that shows how you can start with an example problem or use case and find an appropriate built-in algorithm offered by SageMaker that is valid for that problem type. Additional guidance organized by learning paradigms (supervised and unsupervised) and important data domains (text and images) is provided in the sections following the table.

**Table: Mapping use cases to built-in algorithms**

<table>
<thead>
<tr>
<th>Example problems and use cases</th>
<th>Learning paradigm or domain</th>
<th>Problem types</th>
<th>Data input format</th>
<th>Built-in algorithms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Here a few examples out of the 15 problem types that can be addressed by the pre-trained models and pre-built solution templates provided by SageMaker JumpStart:</td>
<td>Pre-trained models and pre-built solution templates</td>
<td>Image Classification</td>
<td>Image, Text, Tabular</td>
<td>Popular models, including MobileNet, YOLO, Faster R-CNN, BERT, lightGBM, and CatBoost</td>
</tr>
<tr>
<td>Question answering: chatbot that</td>
<td></td>
<td>Tabular Classification</td>
<td></td>
<td>For a list of pre-trained models available, see JumpStart Models.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tabular Regression</td>
<td></td>
<td>For a list of pre-built solution</td>
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<tr>
<td>Example problems and use cases</td>
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<tr>
<td>outputs an answer for a given question.</td>
<td>Question Answering</td>
<td>Sentence Pair Classification</td>
<td>Image Embedding</td>
<td>templates available, see <a href="#">JumpStart Solutions</a></td>
</tr>
<tr>
<td>Text analysis: analyze texts from models specific to an industry domain such as finance.</td>
<td>Named Entity Recognition</td>
<td>Instance Segmentation</td>
<td>Text Generation</td>
<td></td>
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<tr>
<td></td>
<td>Text Summarization</td>
<td>Semantic Segmentation</td>
<td>Machine Translation</td>
<td></td>
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<tr>
<td>Example problems and use cases</td>
<td>Learning paradigm or domain</td>
<td>Problem types</td>
<td>Data input format</td>
<td>Built-in algorithms</td>
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<tr>
<td>Predict if an item belongs to a category: an email spam filter</td>
<td>Supervised Learning</td>
<td>Binary/multi-class classification</td>
<td>Tabular</td>
<td>AutoGluon-Tabular, CatBoost, Factorization Machines Algorithm, K-Nearest Neighbors (k-NN) Algorithm, LightGBM, Linear Learner Algorithm, TabTransformer, XGBoost Algorithm</td>
</tr>
<tr>
<td>Predict a numeric/continuous value: estimate the value of a house</td>
<td></td>
<td>Regression</td>
<td>Tabular</td>
<td>AutoGluon-Tabular, CatBoost, Factorization Machines Algorithm, K-Nearest Neighbors (k-NN) Algorithm, LightGBM, Linear Learner Algorithm, TabTransformer, XGBoost Algorithm</td>
</tr>
<tr>
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<tr>
<td>Based on historical data for a behavior, predict future behavior: predict sales on a new product based on previous sales data.</td>
<td>Time-series forecasting</td>
<td>Tabular</td>
<td>DeepAR Forecasting Algorithm</td>
<td></td>
</tr>
<tr>
<td>Improve the data embedding s of the high-dimensional objects: identify duplicate support tickets or find the correct routing based on similarity of text in the tickets</td>
<td>Embeddings: convert high-dimensional objects into low-dimensional space.</td>
<td>Tabular</td>
<td>Object2Vec Algorithm</td>
<td></td>
</tr>
<tr>
<td>Drop those columns from a dataset that have a weak relation with the label/target variable: the color of a car when predicting its mileage.</td>
<td>Feature engineering: dimensionality reduction</td>
<td>Tabular</td>
<td>Principal Component Analysis (PCA) Algorithm</td>
<td></td>
</tr>
<tr>
<td>Example problems and use cases</td>
<td>Learning paradigm or domain</td>
<td>Problem types</td>
<td>Data input format</td>
<td>Built-in algorithms</td>
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<tr>
<td>Detect abnormal behavior in application: spot when an IoT sensor is sending abnormal readings</td>
<td></td>
<td>Anomaly detection</td>
<td>Tabular</td>
<td>Random Cut Forest (RCF) Algorithm</td>
</tr>
<tr>
<td>Protect your application from suspicious users: detect if an IP address accessing a service might be from a bad actor</td>
<td></td>
<td>IP anomaly detection</td>
<td>Tabular</td>
<td>IP Insights</td>
</tr>
<tr>
<td>Group similar objects/data together: find high-, medium-, and low-spend ing customers from their transaction histories</td>
<td></td>
<td>Clustering or grouping</td>
<td>Tabular</td>
<td>K-Means Algorithm</td>
</tr>
<tr>
<td>Example problems and use cases</td>
<td>Learning paradigm or domain</td>
<td>Problem types</td>
<td>Data input format</td>
<td>Built-in algorithms</td>
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<td>------------------------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>Organize a set of documents into topics (not known in advance): tag a document as belonging to a medical category based on the terms used in the document.</td>
<td></td>
<td>Topic modeling</td>
<td>Text</td>
<td>Latent Dirichlet Allocation (LDA) Algorithm, Neural Topic Model (NTM) Algorithm</td>
</tr>
<tr>
<td>Assign pre-defined categories to documents in a corpus: categorize books in a library into academic disciplines</td>
<td>Textual Analysis</td>
<td>Text classification</td>
<td>Text</td>
<td>BlazingText algorithm, Text Classification - TensorFlow</td>
</tr>
<tr>
<td>Convert text from one language to other: Spanish to English</td>
<td></td>
<td>Machine translation algorithm</td>
<td>Text</td>
<td>Sequence-to-Sequence Algorithm</td>
</tr>
<tr>
<td>Summarize a long text corpus: an abstract for a research paper</td>
<td></td>
<td>Text summarization</td>
<td>Text</td>
<td>Sequence-to-Sequence Algorithm</td>
</tr>
<tr>
<td>Example problems and use cases</td>
<td>Learning paradigm or domain</td>
<td>Problem types</td>
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<td>Built-in algorithms</td>
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</tr>
<tr>
<td>Convert audio files to text: transcribe call center conversations for further analysis</td>
<td></td>
<td>Speech-to-text</td>
<td>Text</td>
<td>Sequence-to-Sequence Algorithm</td>
</tr>
<tr>
<td>Label/tag an image based on the content of the image: alerts about adult content in an image</td>
<td>Image Processing</td>
<td>Image and multi-label classification</td>
<td>Image</td>
<td>Image Classification - MXNet</td>
</tr>
<tr>
<td>Classify something in an image using transfer learning.</td>
<td></td>
<td>Image classification</td>
<td>Image</td>
<td>Image Classification - TensorFlow</td>
</tr>
<tr>
<td>Detect people and objects in an image: police review a large photo gallery for a missing person</td>
<td></td>
<td>Object detection and classification</td>
<td>Image</td>
<td>Object Detection - MXNet, Object Detection - TensorFlow</td>
</tr>
<tr>
<td>Example problems and use cases</td>
<td>Learning paradigm or domain</td>
<td>Problem types</td>
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</tr>
<tr>
<td>Tag every pixel of an image individually with a category: self-driving cars prepare to identify objects in their way</td>
<td>Computer vision</td>
<td>Image</td>
<td>Semantic Segmentation Algorithm</td>
<td></td>
</tr>
</tbody>
</table>

For important information about Docker registry paths, data formats, recommended Amazon EC2 instance types, and CloudWatch logs common to all of the built-in algorithms provided by SageMaker, see Common Information About Built-in Algorithms.

The following sections provide additional guidance for the Amazon SageMaker built-in algorithms grouped by the supervised and unsupervised learning paradigms to which they belong. For descriptions of these learning paradigms and their associated problem types, see Choose an Algorithm. Sections are also provided for the SageMaker built-in algorithms available to address two important machine learning domains: textual analysis and image processing.

- **Pre-trained Models and Solution Templates**
- **Supervised Learning**
- **Unsupervised Learning**
- **Textual Analysis**
- **Image Processing**

**Pre-trained Models and Solution Templates**

SageMaker JumpStart provides a wide range of pre-trained models, pre-built solution templates, and examples for popular problem types that use the SageMaker SDK as well as Studio Classic. For more information about these models, solutions, and the example notebooks provided by SageMaker JumpStart, see SageMaker JumpStart.
Supervised Learning

Amazon SageMaker provides several built-in general purpose algorithms that can be used for either classification or regression problems.

- **AutoGluon-Tabular**—an open-source AutoML framework that succeeds by ensembling models and stacking them in multiple layers.
- **CatBoost**—an implementation of the gradient-boosted trees algorithm that introduces ordered boosting and an innovative algorithm for processing categorical features.
- **Factorization Machines Algorithm**—an extension of a linear model that is designed to economically capture interactions between features within high-dimensional sparse datasets.
- **K-Nearest Neighbors (k-NN) Algorithm**—a non-parametric method that uses the k nearest labeled points to assign a label to a new data point for classification or a predicted target value from the average of the k nearest points for regression.
- **LightGBM**—an implementation of the gradient-boosted trees algorithm that adds two novel techniques for improved efficiency and scalability: Gradient-based One-Side Sampling (GOSS) and Exclusive Feature Bundling (EFB).
- **Linear Learner Algorithm**—learns a linear function for regression or a linear threshold function for classification.
- **TabTransformer**—a novel deep tabular data modeling architecture built on self-attention-based Transformers.
- **XGBoost Algorithm**—an implementation of the gradient-boosted trees algorithm that combines an ensemble of estimates from a set of simpler and weaker models.

Amazon SageMaker also provides several built-in supervised learning algorithms that are used for more specialized tasks during feature engineering and forecasting from time series data.

- **Object2Vec Algorithm**—a new highly customizable multi-purpose algorithm used for feature engineering. It can learn low-dimensional dense embeddings of high-dimensional objects to produce features that improve training efficiencies for downstream models. While this is a supervised algorithm, as it requires labeled data for training, there are many scenarios in which the relationship labels can be obtained purely from natural clusterings in data, without any explicit human annotation.
- **DeepAR Forecasting Algorithm**—a supervised learning algorithm for forecasting scalar (one-dimensional) time series using recurrent neural networks (RNN).
Unsupervised Learning

Amazon SageMaker provides several built-in algorithms that can be used for a variety of unsupervised learning tasks such as clustering, dimension reduction, pattern recognition, and anomaly detection.

- **Principal Component Analysis (PCA) Algorithm**—reduces the dimensionality (number of features) within a dataset by projecting data points onto the first few principal components. The objective is to retain as much information or variation as possible. For mathematicians, principal components are eigenvectors of the data's covariance matrix.

- **K-Means Algorithm**—finds discrete groupings within data, where members of a group are as similar as possible to one another and as different as possible from members of other groups.

- **IP Insights**—learns the usage patterns for IPv4 addresses. It is designed to capture associations between IPv4 addresses and various entities, such as user IDs or account numbers.

- **Random Cut Forest (RCF) Algorithm**—detects anomalous data points within a data set that diverge from otherwise well-structured or patterned data.

Textual Analysis

SageMaker provides algorithms that are tailored to the analysis of textual documents used in natural language processing, document classification or summarization, topic modeling or classification, and language transcription or translation.

- **BlazingText algorithm**—a highly optimized implementation of the Word2vec and text classification algorithms that scale to large datasets easily. It is useful for many downstream natural language processing (NLP) tasks.

- **Sequence-to-Sequence Algorithm**—a supervised algorithm commonly used for neural machine translation.

- **Latent Dirichlet Allocation (LDA) Algorithm**—an algorithm suitable for determining topics in a set of documents. It is an *unsupervised algorithm*, which means that it doesn't use example data with answers during training.

- **Neural Topic Model (NTM) Algorithm**—another unsupervised technique for determining topics in a set of documents, using a neural network approach.

- **Text Classification - TensorFlow**—a supervised algorithm that supports transfer learning with available pretrained models for text classification.
Image Processing

SageMaker also provides image processing algorithms that are used for image classification, object detection, and computer vision.

- **Image Classification - MXNet**—uses example data with answers (referred to as a *supervised algorithm*). Use this algorithm to classify images.

- **Image Classification - TensorFlow**—uses pretrained TensorFlow Hub models to fine-tune for specific tasks (referred to as a *supervised algorithm*). Use this algorithm to classify images.

- **Semantic Segmentation Algorithm**—provides a fine-grained, pixel-level approach to developing computer vision applications.

- **Object Detection - MXNet**—detects and classifies objects in images using a single deep neural network. It is a supervised learning algorithm that takes images as input and identifies all instances of objects within the image scene.

- **Object Detection - TensorFlow**—detects bounding boxes and object labels in an image. It is a supervised learning algorithm that supports transfer learning with available pretrained TensorFlow models.

Topics

- **Common Information About Built-in Algorithms**
- **Built-in SageMaker Algorithms for Tabular Data**
- **Built-in SageMaker Algorithms for Text Data**
- **Built-in SageMaker Algorithms for Time-Series Data**
- **Unsupervised Built-in SageMaker Algorithms**
- **Built-in SageMaker Algorithms for Computer Vision**

Common Information About Built-in Algorithms

The following table lists parameters for each of the algorithms provided by Amazon SageMaker.
<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>AutoGluon-Tabular</td>
<td>training and (optional ly) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>BlazingText</td>
<td>train</td>
<td>File or Pipe</td>
<td>Text file (one sentence per line with space-separated tokens)</td>
<td>CPU or GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>CatBoost</td>
<td>training and (optional ly) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>DeepAR Forecasting</td>
<td>train and (optional ly) test</td>
<td>File</td>
<td>JSON Lines or Parquet</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Factorization Machines</td>
<td>train and (optional ly) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf</td>
<td>CPU (GPU for dense data)</td>
<td>Yes</td>
</tr>
<tr>
<td>Image Classification - MXNet</td>
<td>train and validation, (optional ly) train_lst, validation</td>
<td>File or Pipe</td>
<td>recordIO or image files (.jpg or .png)</td>
<td>GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
<td>Instance class</td>
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<tr>
<td>n_lst, and model</td>
<td>File</td>
<td>image files (.jpg, jpeg, or .png)</td>
<td>CPU or GPU</td>
<td>Yes (only across multiple GPUs on a single instance)</td>
<td></td>
</tr>
<tr>
<td>IP Insights</td>
<td>training and validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>K-Means</td>
<td>train and (optionally) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU (single GPU device on one or more instances)</td>
<td>No</td>
</tr>
<tr>
<td>K-Nearest-Neighbors (k-NN)</td>
<td>train and (optionally) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU (single GPU device on one or more instances)</td>
<td>Yes</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
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<tr>
<td>LDA</td>
<td>train and (optionally) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>LightGBM</td>
<td>train/training and (optionally) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Linear Learner</td>
<td>train and (optionally) validation, test, or both</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Neural Topic Model</td>
<td>train and (optionally) validation, test, or both</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Object2Vec</td>
<td>train and (optionally) validation, test, or both</td>
<td>File</td>
<td>JSON Lines</td>
<td>CPU or GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
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</tr>
<tr>
<td>Object Detection - MXNet</td>
<td>train and validation, (optional) train_annotation, validation_annotation, and model</td>
<td>File or Pipe</td>
<td>recordIO or image files (.jpg or .png)</td>
<td>GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Object Detection - TensorFlow</td>
<td>training and validation</td>
<td>File</td>
<td>image files (.jpg, .jpeg, or .png)</td>
<td>GPU</td>
<td>Yes (only across multiple GPUs on a single instance)</td>
</tr>
<tr>
<td>PCA</td>
<td>train and (optional) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Random Cut Forest</td>
<td>train and (optional) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
<td>Instance class</td>
<td>Parallelizable</td>
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</tr>
<tr>
<td>Semantic Segmentation</td>
<td>train and validation, train_annotation, validation_annotation, and (optionally) label_map and model</td>
<td>File or Pipe</td>
<td>Image files</td>
<td>GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Seq2Seq Modeling</td>
<td>train, validation, and vocab</td>
<td>File</td>
<td>recordIO-protobuf</td>
<td>GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>TabTransformer</td>
<td>training and (optionally) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Text Classification - TensorFlow</td>
<td>training and validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU</td>
<td>Yes (only across multiple GPUs on a single instance)</td>
</tr>
</tbody>
</table>

<p>| Use Built-in Algorithms | 2652 |</p>
<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>XGBoost (0.90-1, 0.90-2, 1.0-1, 1.2-1, 1.2-21)</td>
<td>train and (optional) validation</td>
<td>File or Pipe</td>
<td>CSV, LibSVM, or Parquet</td>
<td>CPU (or GPU for 1.2-1)</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Algorithms that are *parallelizable* can be deployed on multiple compute instances for distributed training.

The following topics provide information about data formats, recommended Amazon EC2 instance types, and CloudWatch logs common to all of the built-in algorithms provided by Amazon SageMaker.

**Note**

To look up the Docker image URIs of the built-in algorithms managed by SageMaker, see [Docker Registry Paths and Example Code](#).

**Topics**

- [Common Data Formats for Built-in Algorithms](#)
- [Instance Types for Built-in Algorithms](#)
- [Logs for Built-in Algorithms](#)

**Common Data Formats for Built-in Algorithms**

The following topics explain the data formats for the algorithms provided by Amazon SageMaker.

**Topics**

- [Common Data Formats for Training](#)
- [Common Data Formats for Inference](#)
Common Data Formats for Training

To prepare for training, you can preprocess your data using a variety of AWS services, including AWS Glue, Amazon EMR, Amazon Redshift, Amazon Relational Database Service, and Amazon Athena. After preprocessing, publish the data to an Amazon S3 bucket. For training, the data need to go through a series of conversions and transformations, including:

- Training data serialization (handled by you)
- Training data deserialization (handled by the algorithm)
- Training model serialization (handled by the algorithm)
- Trained model deserialization (optional, handled by you)

When using Amazon SageMaker in the training portion of the algorithm, make sure to upload all data at once. If more data is added to that location, a new training call would need to be made to construct a brand new model.

Topics

- Content Types Supported by Built-In Algorithms
- Using Pipe Mode
- Using CSV Format
- Using RecordIO Format
- Trained Model Deserialization

Content Types Supported by Built-In Algorithms

The following table lists some of the commonly supported `ContentType` values and the algorithms that use them:

<table>
<thead>
<tr>
<th>ContentType</th>
<th>Algorithm</th>
</tr>
</thead>
<tbody>
<tr>
<td>application/x-image</td>
<td>Object Detection Algorithm, Semantic Segmentation</td>
</tr>
<tr>
<td>application/x-recordio</td>
<td>Object Detection Algorithm</td>
</tr>
<tr>
<td>ContentType</td>
<td>Algorithm</td>
</tr>
<tr>
<td>-------------</td>
<td>-----------</td>
</tr>
<tr>
<td>application/x-recordio-protobuf</td>
<td>Factorization Machines, K-Means, k-NN, Latent Dirichlet Allocation, Linear Learner, NTM, PCA, RCF, Sequence-to-Sequence</td>
</tr>
<tr>
<td>application/jsonlines</td>
<td>BlazingText, DeepAR</td>
</tr>
<tr>
<td>image/jpeg</td>
<td>Object Detection Algorithm, Semantic Segmentation</td>
</tr>
<tr>
<td>image/png</td>
<td>Object Detection Algorithm, Semantic Segmentation</td>
</tr>
<tr>
<td>text/libsvm</td>
<td>XGBoost</td>
</tr>
</tbody>
</table>

For a summary of the parameters used by each algorithm, see the documentation for the individual algorithms or this [table](#).

**Using Pipe Mode**

In *Pipe mode*, your training job streams data directly from Amazon Simple Storage Service (Amazon S3). Streaming can provide faster start times for training jobs and better throughput. This is in contrast to *File mode*, in which your data from Amazon S3 is stored on the training instance volumes. File mode uses disk space to store both your final model artifacts and your full training dataset. By streaming in your data directly from Amazon S3 in Pipe mode, you reduce the size of Amazon Elastic Block Store volumes of your training instances. Pipe mode needs only enough disk space to store your final model artifacts. See the [AlgorithmSpecification](#) for additional details on the training input mode.

**Using CSV Format**

Many Amazon SageMaker algorithms support training with data in CSV format. To use data in CSV format for training, in the input data channel specification, specify `text/csv` as the **ContentType**. Amazon SageMaker requires that a CSV file does not have a header record and that the target variable is in the first column. To run unsupervised learning algorithms that don't have a target, specify the number of label columns in the content type. For example, in this case `'content_type=text/csv;label_size=0'`. For a notebook example that uses CSV format,
see Breast Cancer Prediction. For more information, see Now use Pipe mode with CSV datasets for faster training on Amazon SageMaker built-in algorithms.

**Using RecordIO Format**

In the protobuf recordIO format, SageMaker converts each observation in the dataset into a binary representation as a set of 4-byte floats, then loads it in the protobuf values field. If you are using Python for your data preparation, we strongly recommend that you use these existing transformations. However, if you are using another language, the protobuf definition file below provides the schema that you use to convert your data into SageMaker protobuf format.

---

### Note

For an example that shows how to convert the commonly used numPy array into the protobuf recordIO format, see *An Introduction to Factorization Machines with MNIST*.

```
syntax = "proto2";

package aialgs.data;

option java_package = "com.amazonaws.aialgorithms.proto";
option java_outer_classname = "RecordProtos";

// A sparse or dense rank-R tensor that stores data as doubles (float64).
message Float32Tensor {
  // Each value in the vector. If keys is empty, this is treated as a dense vector.
  repeated float values = 1 [packed = true];

  // If key is not empty, the vector is treated as sparse, with each key specifying the location of the value in the sparse vector.
  repeated uint64 keys = 2 [packed = true];

  // An optional shape that allows the vector to represent a matrix.
  // For example, if shape = [10, 20], floor(keys[i] / 20) gives the row, and keys[i] % 20 gives the column.
  // This also supports n-dimensional tensors.
  // Note: If the tensor is sparse, you must specify this value.
  repeated uint64 shape = 3 [packed = true];
}
```
// A sparse or dense rank-R tensor that stores data as doubles (float64).
message Float64Tensor {
  // Each value in the vector. If keys is empty, this is treated as a
  // dense vector.
  repeated double values = 1 [packed = true];

  // If this is not empty, the vector is treated as sparse, with
  // each key specifying the location of the value in the sparse vector.
  repeated uint64 keys = 2 [packed = true];

  // An optional shape that allows the vector to represent a matrix.
  // For example, if shape = [ 10, 20 ], floor(keys[i] / 10) gives the row,
  // and keys[i] % 20 gives the column.
  // This also supports n-dimensonal tensors.
  // Note: If the tensor is sparse, you must specify this value.
  repeated uint64 shape = 3 [packed = true];
}

// A sparse or dense rank-R tensor that stores data as 32-bit ints (int32).
message Int32Tensor {
  // Each value in the vector. If keys is empty, this is treated as a
  // dense vector.
  repeated int32 values = 1 [packed = true];

  // If this is not empty, the vector is treated as sparse with
  // each key specifying the location of the value in the sparse vector.
  repeated uint64 keys = 2 [packed = true];

  // An optional shape that allows the vector to represent a matrix.
  // For Example, if shape = [ 10, 20 ], floor(keys[i] / 10) gives the row,
  // and keys[i] % 20 gives the column.
  // This also supports n-dimensonal tensors.
  // Note: If the tensor is sparse, you must specify this value.
  repeated uint64 shape = 3 [packed = true];
}

// Support for storing binary data for parsing in other ways (such as JPEG/etc).
// This is an example of another type of value and may not immediately be supported.
message Bytes {
  repeated bytes value = 1;

  // If the content type of the data is known, stores it.
  // This allows for the possibility of using decoders for common formats
  // in the future.
optional string content_type = 2;
}

message Value {
  oneof value {
    // The numbering assumes the possible use of:
    // - float16, float128
    // - int8, int16, int32
    Float32Tensor float32_tensor = 2;
    Float64Tensor float64_tensor = 3;
    Int32Tensor int32_tensor = 7;
    Bytes bytes = 9;
  }
}

message Record {
  // Map from the name of the feature to the value.
  //
  // For vectors and libsvm-like datasets,
  // a single feature with the name `values`
  // should be specified.
  map<string, Value> features = 1;

  // An optional set of labels for this record.
  // Similar to the features field above, the key used for
  // generic scalar / vector labels should be 'values'.
  map<string, Value> label = 2;

  // A unique identifier for this record in the dataset.
  //
  // Whilst not necessary, this allows better
  // debugging where there are data issues.
  //
  // This is not used by the algorithm directly.
  optional string uid = 3;

  // Textual metadata describing the record.
  //
  // This may include JSON-serialized information
  // about the source of the record.
  //
  // This is not used by the algorithm directly.
  optional string metadata = 4;
// An optional serialized JSON object that allows per-record
// hyper-parameters/configuration/other information to be set.
//
// The meaning/interpretation of this field is defined by
// the algorithm author and may not be supported.
//
// This is used to pass additional inference configuration
// when batch inference is used (e.g. types of scores to return).
optional string configuration = 5;
}

After creating the protocol buffer, store it in an Amazon S3 location that Amazon SageMaker can access and that can be passed as part of InputDataConfig in create_training_job.

**Note**

For all Amazon SageMaker algorithms, the ChannelName in InputDataConfig must be set to train. Some algorithms also support a validation or test input channels. These are typically used to evaluate the model's performance by using a hold-out dataset. Hold-out datasets are not used in the initial training but can be used to further tune the model.

**Trained Model Deserialization**

Amazon SageMaker models are stored as model.tar.gz in the S3 bucket specified in OutputDataConfig S3OutputPath parameter of the create_training_job call. The S3 bucket must be in the same AWS Region as the notebook instance. You can specify most of these model artifacts when creating a hosting model. You can also open and review them in your notebook instance. When model.tar.gz is untarred, it contains model_algo-1, which is a serialized Apache MXNet object. For example, you use the following to load the k-means model into memory and view it:

```python
import mxnet as mx
print(mx.ndarray.load('model_algo-1'))
```

**Common Data Formats for Inference**

Amazon SageMaker algorithms accept and produce several different MIME types for the HTTP payloads used in retrieving online and mini-batch predictions. You can use various AWS services to
transform or preprocess records prior to running inference. At a minimum, you need to convert the data for the following:

- Inference request serialization (handled by you)
- Inference request deserialization (handled by the algorithm)
- Inference response serialization (handled by the algorithm)
- Inference response deserialization (handled by you)

**Topics**

- Convert Data for Inference Request Serialization
- Convert Data for Inference Response Deserialization
- Common Request Formats for All Algorithms
- Use Batch Transform with Built-in Algorithms

**Convert Data for Inference Request Serialization**

Content type options for Amazon SageMaker algorithm inference requests include: `text/csv`, `application/json`, and `application/x-recordio-protobuf`. Algorithms that don't support all of these types can support other types. XGBoost, for example, only supports `text/csv` from this list, but also supports `text/libsvm`.

For `text/csv`, the value for the Body argument to `invoke_endpoint` should be a string with commas separating the values for each feature. For example, a record for a model with four features might look like `1.5,16.0,14,23.0`. Any transformations performed on the training data should also be performed on the data before obtaining inference. The order of the features matters and must remain unchanged.

`application/json` is significantly more flexible and provides multiple possible formats for developers to use in their applications. At a high level, in JavaScript, the payload might look like the following:

```javascript
let request = {
  // Instances might contain multiple rows that predictions are sought for.
  "instances": [
    {
      // Request and algorithm specific inference parameters.
      "configuration": {},
    }
  ],
```
// Data in the specific format required by the algorithm.
"data": {
    "<field name>": dataElement
}
}
]
}

You have the following options for specifying the `dataElement`:

**Protocol buffers equivalent**

// Has the same format as the protocol buffers implementation described for training.
let dataElement = {
    "keys": [],
    "values": [],
    "shape": []
}

**Simple numeric vector**

// An array containing numeric values is treated as an instance containing a
// single dense vector.
let dataElement = [1.5, 16.0, 14.0, 23.0]

// It will be converted to the following representation by the SDK.
let converted = {
    "features": {
        "values": dataElement
    }
}

**For multiple records**

let request = {
    "instances": [
        // First instance.
        {
            "features": [ 1.5, 16.0, 14.0, 23.0 ]
        },
        // Second instance.
        {
            "features": [ -2.0, 100.2, 15.2, 9.2 ]
        }
    ]
}
Convert Data for Inference Response Deserialization

Amazon SageMaker algorithms return JSON in several layouts. At a high level, the structure is:

```javascript
let response = {
  "predictions": [
    // Fields in the response object are defined on a per algorithm-basis.
  ]
}
```

The fields that are included in predictions differ across algorithms. The following are examples of output for the k-means algorithm.

**Single-record inference**

```javascript
let response = {
  "predictions": [
    // First instance prediction.
    {
      "closest_cluster": 5,
      "distance_to_cluster": 36.5
    }
  ]
}
```

**Multi-record inference**

```javascript
let response = {
  "predictions": [
    // First instance prediction.
    {
      "closest_cluster": 5,
      "distance_to_cluster": 36.5
    },
    // Second instance prediction.
    {
      "closest_cluster": 2,
      "distance_to_cluster": 90.3
    }
  ]
}
```
Multi-record inference with protobuf input

```json
{
    "features": [],
    "label": {
        "closest_cluster": {
            "values": [ 5.0 ] // e.g. the closest centroid/cluster was 1.0
        },
        "distance_to_cluster": {
            "values": [ 36.5 ]
        },
    },
    "uid": "abc123",
    "metadata": "{ "created_at": '2017-06-03' }"
}
```

SageMaker algorithms also support the JSONLINES format, where the per-record response content is same as that in JSON format. The multi-record structure is a concatenation of per-record response objects separated by newline characters. The response content for the built-in KMeans algorithm for 2 input data points is:

```
{"distance_to_cluster": 23.40593910217285, "closest_cluster": 0.0}
{"distance_to_cluster": 27.250282287597656, "closest_cluster": 0.0}
```

While running batch transform, we recommended using the jsonlines response type by setting the Accept field in the CreateTransformJobRequest to application/jsonlines.

**Common Request Formats for All Algorithms**

Most algorithms use several of the following inference request formats.

**JSON Request Format**

*Content type: application/JSON*

**Dense format**

```javascript
let request = {
    "instances": [
        {
            "features": [1.5, 16.0, 14.0, 23.0]
        }
    ]
}
```
let request = {
    "instances": [
        {
            "data": {
                "features": {
                    "values": [1.5, 16.0, 14.0, 23.0]
                }
            }
        }
    ]
}

Sparse format

{
    "instances": [
        {
            "data": {
                "features": {
                    "keys": [26, 182, 232, 243, 431],
                    "shape": [2000],
                    "values": [1, 1, 1, 4, 1]
                }
            }
        },
        {
            "data": {
                "features": {
                    "keys": [0, 182, 232, 243, 431],
                    "shape": [2000],
                    "values": [13, 1, 1, 4, 1]
                }
            }
        }
    ]
}

JSONLINES Request Format

**Content type:** application/JSONLINES

**Dense format**

A single record in dense format can be represented as either:
Sparse Format

A single record in sparse format is represented as:

```json
{ "data": { "features": { "keys": [26, 182, 232, 243, 431], "shape": [2000], "values": [1, 1, 1, 4, 1] } } }
```

Multiple records are represented as a concatenation of the above single-record representations, separated by newline characters:

```json
{ "data": { "features": { "keys": [0, 1, 3], "shape": [4], "values": [1, 4, 1] } } }
{ "data": { "features": { "values": [1.5, 16.0, 14.0, 23.0] } } }
{ "features": [1.5, 16.0, 14.0, 23.0] }

CSV Request Format

**Content type:** text/CSV; label_size=0

ℹ️ **Note**

CSV support is not available for factorization machines.

RECORDIO Request Format

**Content type:** application/x-recordio-protobuf

Use Batch Transform with Built-in Algorithms

While running batch transform, we recommend using the JSONLINES response type instead of JSON, if supported by the algorithm. This is accomplished by setting the `Accept` field in the `CreateTransformJobRequest` to `application/jsonlines`.

When you create a transform job, the `SplitType` must be set according to the `ContentType` of the input data. Similarly, depending on the `Accept` field in the `CreateTransformJobRequest`,
AssembleWith must be set accordingly. Please use the following table to help appropriately set these fields:

<table>
<thead>
<tr>
<th>ContentType</th>
<th>Recommended SplitType</th>
</tr>
</thead>
<tbody>
<tr>
<td>application/x-recordio-protobuf</td>
<td>RecordIO</td>
</tr>
<tr>
<td>text/csv</td>
<td>Line</td>
</tr>
<tr>
<td>application/jsonlines</td>
<td>Line</td>
</tr>
<tr>
<td>application/json</td>
<td>None</td>
</tr>
<tr>
<td>application/x-image</td>
<td>None</td>
</tr>
<tr>
<td>image/*</td>
<td>None</td>
</tr>
</tbody>
</table>

For more information on response formats for specific algorithms, see the following:

- DeepAR Inference Formats
- Factorization Machines Response Formats
- IP Insights Inference Data Formats
- K-Means Response Formats
- k-NN Request and Response Formats
- Linear learner response formats
- NTM Response Formats
- Data Formats for Object2Vec Inference
- Encoder Embeddings for Object2Vec
- PCA Response Formats
Instance Types for Built-in Algorithms

For training and hosting Amazon SageMaker algorithms, we recommend using the following Amazon EC2 instance types:

- ml.m5.xlarge, ml.m5.4xlarge, and ml.m5.12xlarge
- ml.c5.xlarge, ml.c5.2xlarge, and ml.c5.8xlarge
- ml.p3.xlarge, ml.p3.8xlarge, and ml.p3.16xlarge

Most Amazon SageMaker algorithms have been engineered to take advantage of GPU computing for training. For most algorithm training, we support P2, P3, G4dn, and G5 GPU instances. Despite higher per-instance costs, GPUs train more quickly, making them more cost effective. Exceptions are noted in this guide.

The size and type of data can have a great effect on which hardware configuration is most effective. When the same model is trained on a recurring basis, initial testing across a spectrum of instance types can discover configurations that are more cost-effective in the long run. Additionally, algorithms that train most efficiently on GPUs might not require GPUs for efficient inference. Experiment to determine the most cost effectiveness solution. To get an automatic instance recommendation or conduct custom load tests, use Amazon SageMaker Inference Recommender.

For more information on SageMaker hardware specifications, see Amazon SageMaker ML Instance Types.

Logs for Built-in Algorithms

Amazon SageMaker algorithms produce Amazon CloudWatch logs, which provide detailed information on the training process. To see the logs, in the AWS management console, choose CloudWatch, choose Logs, and then choose the /aws/sagemaker/TrainingJobs log group. Each training job has one log stream per node on which it was trained. The log stream's name begins with the value specified in the TrainingJobName parameter when the job was created.
The contents of logs vary by algorithms. However, you can typically find the following information:

- Confirmation of arguments provided at the beginning of the log
- Errors that occurred during training
- Measurement of an algorithm's accuracy or numerical performance
- Timings for the algorithm and any major stages within the algorithm

**Common Errors**

If a training job fails, some details about the failure are provided by the `FailureReason` return value in the training job description, as follows:

```python
sage = boto3.client('sagemaker')
sage.describe_training_job(TrainingJobName=job_name)['FailureReason']
```

Others are reported only in the CloudWatch logs. Common errors include the following:

1. Misspecifying a hyperparameter or specifying a hyperparameter that is invalid for the algorithm.

   **From the CloudWatch Log**

   ```
   [10/16/2017 23:45:17 ERROR 139623806805824 train.py:48] Additional properties are not allowed (u'mini_batch_siz' was unexpected)
   ```

2. Specifying an invalid value for a hyperparameter.

   **FailureReason**

   ```
   AlgorithmError: u'abc' is not valid under any of the given schemas\nFailed validating u'oneOf' in schema[u'properties'][u'feature_dim']:\n{u'oneOf': [[u'pattern': u'^([1-9][0-9]*)$', u'type': u'string']},
   ```
### FailureReason

```javascript
{u'minimum': 1, u'type': u'integer'}
```

3. Inaccurate protobuf file format.

#### From the CloudWatch log

- **[10/16/2017 23:57:17 ERROR 140373086025536 train.py:48] u'abc'**
  is not valid under any of the given schemas
- **[10/17/2017 18:01:04 ERROR 140234860816192 train.py:48] cannot**
  copy sequence with size 785 to array axis with dimension 784

### Built-in SageMaker Algorithms for Tabular Data

Amazon SageMaker provides built-in algorithms that are tailored to the analysis of tabular data. Tabular data refers to any datasets that are organized in tables consisting of rows (observations) and columns (features). The built-in SageMaker algorithms for tabular data can be used for either classification or regression problems.

- **AutoGluon-Tabular**—an open-source AutoML framework that succeeds by ensembling models and stacking them in multiple layers.
- **CatBoost**—an implementation of the gradient-boosted trees algorithm that introduces ordered boosting and an innovative algorithm for processing categorical features.
- **Factorization Machines Algorithm**—an extension of a linear model that is designed to economically capture interactions between features within high-dimensional sparse datasets.
- **K-Nearest Neighbors (k-NN) Algorithm**—a non-parametric method that uses the k nearest labeled points to assign a label to a new data point for classification or a predicted target value from the average of the k nearest points for regression.
- **LightGBM**—an implementation of the gradient-boosted trees algorithm that adds two novel techniques for improved efficiency and scalability: Gradient-based One-Side Sampling (GOSS) and Exclusive Feature Bundling (EFB).
- **Linear Learner Algorithm**—learns a linear function for regression or a linear threshold function for classification.
**TabTransformer**—a novel deep tabular data modeling architecture built on self-attention-based Transformers.

**XGBoost Algorithm**—an implementation of the gradient-boosted trees algorithm that combines an ensemble of estimates from a set of simpler and weaker models.

<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>AutoGluon-Tabular</td>
<td>training and (optional) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>CatBoost</td>
<td>training and (optional) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Factorization Machines</td>
<td>train and (optional) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf</td>
<td>CPU (GPU for dense data)</td>
<td>Yes</td>
</tr>
<tr>
<td>K-Nearest-Neighbors (k-NN)</td>
<td>train and (optional) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU (single GPU device on one or more instances)</td>
<td>Yes</td>
</tr>
<tr>
<td>LightGBM</td>
<td>training and (optional)</td>
<td>File</td>
<td>CSV</td>
<td>CPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
<td>Instance class</td>
<td>Parallelizable</td>
</tr>
<tr>
<td>----------------</td>
<td>--------------</td>
<td>---------------------</td>
<td>-----------</td>
<td>----------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Linear Learner</td>
<td>train and (optional) validation, test, or both</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>TabTransformer</td>
<td>training and (optional) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>XGBoost (0.90-1, 0.90-2, 1.0-1, 1.2-1, 1.2-21)</td>
<td>train and (optional) validation</td>
<td>File or Pipe</td>
<td>CSV, LibSVM, or Parquet</td>
<td>CPU (or GPU for 1.2-1)</td>
<td>Yes</td>
</tr>
</tbody>
</table>

**AutoGluon-Tabular**

*AutoGluon-Tabular* is a popular open-source AutoML framework that trains highly accurate machine learning models on an unprocessed tabular dataset. Unlike existing AutoML frameworks that primarily focus on model and hyperparameter selection, AutoGluon-Tabular succeeds by ensembling multiple models and stacking them in multiple layers.

**How to use SageMaker AutoGluon-Tabular**

You can use AutoGluon-Tabular as an Amazon SageMaker built-in algorithm. The following section describes how to use AutoGluon-Tabular with the SageMaker Python SDK. For information on
how to use AutoGluon-Tabular from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

- **Use AutoGluon-Tabular as a built-in algorithm**

  Use the AutoGluon-Tabular built-in algorithm to build an AutoGluon-Tabular training container as shown in the following code example. You can automatically spot the AutoGluon-Tabular built-in algorithm image URI using the SageMaker image_uris.retrieve API (or the get_image_uri API if using Amazon SageMaker Python SDK version 2).

  After specifying the AutoGluon-Tabular image URI, you can use the AutoGluon-Tabular container to construct an estimator using the SageMaker Estimator API and initiate a training job. The AutoGluon-Tabular built-in algorithm runs in script mode, but the training script is provided for you and there is no need to replace it. If you have extensive experience using script mode to create a SageMaker training job, then you can incorporate your own AutoGluon-Tabular training scripts.

```python
from sagemaker import image_uris, model_uris, script_uris

train_model_id, train_model_version, train_scope = "autogluon-classification-ensemble", "*", "training"
training_instance_type = "ml.p3.2xlarge"

# Retrieve the docker image
train_image_uri = image_uris.retrieve(
    region=None,
    framework=None,
    model_id=train_model_id,
    model_version=train_model_version,
    image_scope=train_scope,
    instance_type=training_instance_type
)

# Retrieve the training script
train_source_uri = script_uris.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    script_scope=train_scope
)

train_model_uri = model_uris.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    model_scope=train_scope
)
```
Use Built-in Algorithms
hyperparameters=hyperparameters,
output_path=s3_output_location
)

# Launch a SageMaker Training job by passing the S3 path of the training data
tabular_estimator.fit(
    {
        "training": training_dataset_s3_path,
        "validation": validation_dataset_s3_path,
    }, logs=True, job_name=training_job_name
)

For more information about how to set up the AutoGluon-Tabular as a built-in algorithm, see the following notebook examples. Any S3 bucket used in these examples must be in the same AWS Region as the notebook instance used to run them.

- Tabular classification with Amazon SageMaker AutoGluon-Tabular algorithm
- Tabular regression with Amazon SageMaker AutoGluon-Tabular algorithm

Input and Output interface for the AutoGluon-Tabular algorithm

Gradient boosting operates on tabular data, with the rows representing observations, one column representing the target variable or label, and the remaining columns representing features.

The SageMaker implementation of AutoGluon-Tabular supports CSV for training and inference:

- For **Training ContentType**, valid inputs must be *text/csv*.
- For **Inference ContentType**, valid inputs must be *text/csv*.

**Note**

For CSV training, the algorithm assumes that the target variable is in the first column and that the CSV does not have a header record.

For CSV inference, the algorithm assumes that CSV input does not have the label column.

Input format for training data, validation data, and categorical features

Be mindful of how to format your training data for input to the AutoGluon-Tabular model. You must provide the path to an Amazon S3 bucket that contains your training and validation data. You
can also include a list of categorical features. Use both the training and validation channels to provide your input data. Alternatively, you can use only the training channel.

**Use both the training and validation channels**

You can provide your input data by way of two S3 paths, one for the training channel and one for the validation channel. Each S3 path can either be an S3 prefix or a full S3 path pointing to one specific CSV file. The target variables should be in the first column of your CSV file. The predictor variables (features) should be in the remaining columns. The validation data is used to compute a validation score at the end of each boosting iteration. Early stopping is applied when the validation score stops improving.

If your predictors include categorical features, you can provide a JSON file named `categorical_index.json` in the same location as your training data file. If you provide a JSON file for categorical features, your training channel must point to an S3 prefix and not a specific CSV file. This file should contain a Python dictionary where the key is the string "cat_index_list" and the value is a list of unique integers. Each integer in the value list should indicate the column index of the corresponding categorical features in your training data CSV file. Each value should be a positive integer (greater than zero because zero represents the target value), less than the `Int32.MaxValue` (2,147,483,647), and less than the total number of columns. There should only be one categorical index JSON file.

**Use only the training channel:**

You can alternatively provide your input data by way of a single S3 path for the training channel. This S3 path should point to a directory with a subdirectory named `training/` that contains a CSV file. You can optionally include another subdirectory in the same location called `validation/` that also has a CSV file. If the validation data is not provided, then 20% of your training data is randomly sampled to serve as the validation data. If your predictors include categorical features, you can provide a JSON file named `categorical_index.json` in the same location as your data subdirectories.

**Note**

For CSV training input mode, the total memory available to the algorithm (instance count multiplied by the memory available in the InstanceType) must be able to hold the training dataset.
Amazon SageMaker AutoGluon-Tabular uses the autogluon.tabular.TabularPredictor module to serialize or deserialize the model, which can be used for saving or loading the model.

**To use a model trained with SageMaker AutoGluon-Tabular with the AutoGluon framework**

- Use the following Python code:

```python
import tarfile
from autogluon.tabular import TabularPredictor

t = tarfile.open('model.tar.gz', 'r:gz')
t.extractall()

model = TabularPredictor.load(model_file_path)

# prediction with test data
# dtest should be a pandas DataFrame with column names feature_0, feature_1, ..., feature_d
pred = model.predict(dtest)
```

**Amazon EC2 instance recommendation for the AutoGluon-Tabular algorithm**

SageMaker AutoGluon-Tabular supports single-instance CPU and single-instance GPU training. Despite higher per-instance costs, GPUs train more quickly, making them more cost effective. To take advantage of GPU training, specify the instance type as one of the GPU instances (for example, P3). SageMaker AutoGluon-Tabular currently does not support multi-GPU training.

**AutoGluon-Tabular sample notebooks**

The following table outlines a variety of sample notebooks that address different use cases of Amazon SageMaker AutoGluon-Tabular algorithm.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tabular classification with Amazon SageMaker AutoGluon-Tabular algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker AutoGluon-Tabular algorithm to train and host a tabular classification model.</td>
</tr>
</tbody>
</table>
Tabular regression with Amazon SageMaker AutoGluon-Tabular algorithm

This notebook demonstrates the use of the Amazon SageMaker AutoGluon-Tabular algorithm to train and host a tabular regression model.

For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all of the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

How AutoGluon-Tabular works

AutoGluon-Tabular performs advanced data processing, deep learning, and multi-layer model ensemble methods. It automatically recognizes the data type in each column for robust data preprocessing, including special handling of text fields.

AutoGluon fits various models ranging from off-the-shelf boosted trees to customized neural networks. These models are ensembled in a novel way: models are stacked in multiple layers and trained in a layer-wise manner that guarantees raw data can be translated into high-quality predictions within a given time constraint. This process mitigates overfitting by splitting the data in various ways with careful tracking of out-of-fold examples.

The AutoGluon-Tabular algorithm performs well in machine learning competitions because of its robust handling of a variety of data types, relationships, and distributions. You can use AutoGluon-Tabular for regression, classification (binary and multiclass), and ranking problems.

Refer to the following diagram illustrating how the multi-layer stacking strategy works.
For more information, see *AutoGluon-Tabular: Robust and Accurate AutoML for Structured Data*.

**AutoGluon-Tabular hyperparameters**

The following table contains the subset of hyperparameters that are required or most commonly used for the Amazon SageMaker AutoGluon-Tabular algorithm. Users set these parameters to facilitate the estimation of model parameters from data. The SageMaker AutoGluon-Tabular algorithm is an implementation of the open-source *AutoGluon-Tabular* package.

---

**Note**

The default hyperparameters are based on example datasets in the *AutoGluon-Tabular sample notebooks*.

---

By default, the SageMaker AutoGluon-Tabular algorithm automatically chooses an evaluation metric based on the type of classification problem. The algorithm detects the type of classification problem based on the number of labels in your data. For regression problems, the evaluation metric is root mean squared error. For binary classification problems, the evaluation metric is area under the receiver operating characteristic curve (AUC). For multiclass classification problems, the evaluation metric is accuracy. You can use the `eval_metric` hyperparameter to change the default evaluation metric. Refer to the following table for more information on AutoGluon-Tabular hyperparameters, including descriptions, valid values, and default values.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>eval_metric</td>
<td>The evaluation metric for validation data. If <code>eval_metric</code> is set to the default &quot;auto&quot; value, then the algorithm automatically chooses an evaluation metric based on the type of classification problem:</td>
</tr>
<tr>
<td></td>
<td>• &quot;root_mean_squared_error&quot; for regression</td>
</tr>
<tr>
<td></td>
<td>• &quot;roc_auc&quot; for binary classification</td>
</tr>
<tr>
<td></td>
<td>• &quot;accuracy&quot; for multi-class classification</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, refer to the <a href="#">AutoGluon documentation</a> for valid values.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;auto&quot;.</td>
</tr>
<tr>
<td>presets</td>
<td>List of preset configurations for various arguments in fit().</td>
</tr>
<tr>
<td></td>
<td>• &quot;best_quality&quot; : high predictive accuracy, slower inference times and higher disk usage</td>
</tr>
<tr>
<td></td>
<td>• &quot;high_quality&quot; : high predictive accuracy and fast inference</td>
</tr>
<tr>
<td></td>
<td>• &quot;good_quality&quot; : good predictive accuracy and very fast inference</td>
</tr>
<tr>
<td></td>
<td>• &quot;medium_quality&quot; : medium predictive accuracy, very fast inference and training time</td>
</tr>
<tr>
<td></td>
<td>• &quot;optimize_for_deployment&quot; : delete unused models and remove training artifacts</td>
</tr>
<tr>
<td></td>
<td>• &quot;interpretable&quot; : fits only interpretable rule-based models from the imodels package</td>
</tr>
<tr>
<td></td>
<td>For more details, see <a href="#">AutoGluon Predictors</a>.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: (&quot;best_quality&quot;, &quot;high_quality&quot;, &quot;good_quality&quot;, &quot;medium_quality&quot;)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>quality</td>
<td>&quot;quality&quot;, &quot;optimize_for_deployment&quot;, or &quot;interpretable&quot;). Default value: &quot;medium_quality&quot;.</td>
</tr>
<tr>
<td>auto_stack</td>
<td>Whether AutoGluon should automatically utilize bagging and multi-layer stack ensembling to boost predictive accuracy. Set auto_stack to &quot;True&quot; if you are willing to tolerate longer training times in order to maximize predictive accuracy. This automatically sets the num_bag_folds and num_stack_levels arguments based on dataset properties. Valid values: string, &quot;True&quot; or &quot;False&quot;. Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>num_bag_folds</td>
<td>Number of folds used for bagging of models. When num_bag_folds is equal to k, training time is roughly increased by a factor of k. Set num_bag_folds to 0 to deactivate bagging. This is disabled by default, but we recommend using values between 5 and 10 to maximize predictive performance. Increasing num_bag_folds results in models with lower bias, but that are more prone to overfitting. One is an invalid value for this parameter, and will raise a ValueError. Values greater than 10 may produce diminishing returns and can even harm overall results due to overfitting. To further improve predictions, avoid increasing num_bag_folds and instead increase num_bag_sets. Valid values: string, any integer between (and including) &quot;0&quot; and &quot;10&quot;. Default value: &quot;0&quot;.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>num_bag_sets</td>
<td>Number of repeats of kfold bagging to perform (values must be greater than or equal to 1). The total number of models trained during bagging is equal to num_bag_folds * num_bag_sets. This parameter defaults to one if time_limit is not specified. This parameter is disabled if num_bag_folds is not specified. Values greater than one result in superior predictive performance, especially on smaller problems and with stacking enabled. Valid values: integer, range: [1, 20]. Default value: 1.</td>
</tr>
<tr>
<td>num_stack_levels</td>
<td>Number of stacking levels to use in stack ensemble. Roughly increases model training time by factor of num_stack_levels + 1. Set this parameter to 0 to deactivate stack ensembling. This parameter is deactivated by default, but we recommend using values between 1 and 3 to maximize predictive performance. To prevent overfitting and a ValueError, num_bag_folds must be greater than or equal to 2. Valid values: float, range: [0, 3]. Default value: 0.</td>
</tr>
<tr>
<td>refit_full</td>
<td>Whether or not to retrain all models on all of the data (training and validation) after the normal training procedure. For more details, see AutoGluon Predictors. Valid values: string, &quot;True&quot; or &quot;False&quot;. Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>set_best_to_refit_full</td>
<td>Whether or not to change the default model that the predictor uses for prediction. If <code>set_best_to_refit_full</code> is set to &quot;True&quot;, the default model changes to the model that exhibited the highest validation score as a result of refitting (activated by <code>refit_full</code>). Only valid if <code>refit_full</code> is set. Valid values: string, &quot;True&quot; or &quot;False&quot;. Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>save_space</td>
<td>Whether or not to reduce the memory and disk size of predictor by deleting auxiliary model files that aren't needed for prediction on new data. This has no impact on inference accuracy. We recommend setting <code>save_space</code> to &quot;True&quot; if the only goal is to use the trained model for prediction. Certain advanced functionality may no longer be available if <code>save_space</code> is set to &quot;True&quot;. Refer to the <code>predictor.save_space()</code> documentation for more details. Valid values: string, &quot;True&quot; or &quot;False&quot;. Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>verbosity</td>
<td>The verbosity of print messages. <code>verbosity</code> levels range from 0 to 4, with higher levels corresponding to more detailed print statements. A <code>verbosity</code> of 0 suppresses warnings. Valid values: integer, any of the following: (0, 1, 2, 3, or 4). Default value: 2.</td>
</tr>
</tbody>
</table>

**Tuning an AutoGluon-Tabular model**

Although AutoGluon-Tabular can be used with model tuning, its design can deliver good performance using stacking and ensemble methods, meaning hyperparameter optimization is not necessary. Rather than focusing on model tuning, AutoGluon-Tabular succeeds by stacking models in multiple layers and training in a layer-wise manner.
For more information about AutoGluon-Tabular hyperparameters, see AutoGluon-Tabular hyperparameters.

CatBoost

CatBoost is a popular and high-performance open-source implementation of the Gradient Boosting Decision Tree (GBDT) algorithm. GBDT is a supervised learning algorithm that attempts to accurately predict a target variable by combining an ensemble of estimates from a set of simpler and weaker models.

CatBoost introduces two critical algorithmic advances to GBDT:

1. The implementation of ordered boosting, a permutation-driven alternative to the classic algorithm
2. An innovative algorithm for processing categorical features

Both techniques were created to fight a prediction shift caused by a special kind of target leakage present in all currently existing implementations of gradient boosting algorithms.

How to use SageMaker CatBoost

You can use CatBoost as an Amazon SageMaker built-in algorithm. The following section describes how to use CatBoost with the SageMaker Python SDK. For information on how to use CatBoost from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

• Use CatBoost as a built-in algorithm

Use the CatBoost built-in algorithm to build a CatBoost training container as shown in the following code example. You can automatically spot the CatBoost built-in algorithm image URI using the SageMaker image_uris.retrieve API (or the get_image_uri API if using Amazon SageMaker Python SDK version 2).

After specifying the CatBoost image URI, you can use the CatBoost container to construct an estimator using the SageMaker Estimator API and initiate a training job. The CatBoost built-in algorithm runs in script mode, but the training script is provided for you and there is no need to replace it. If you have extensive experience using script mode to create a SageMaker training job, then you can incorporate your own CatBoost training scripts.

```python
from sagemaker import image_uris, model_uris, script_uris
```
train_model_id, train_model_version, train_scope = "catboost-classification-model", "*", "training"
training_instance_type = "ml.m5.xlarge"

# Retrieve the docker image
train_image_uri = image_uri.retrieve(
    region=None,
    framework=None,
    model_id=train_model_id,
    model_version=train_model_version,
    image_scope=train_scope,
    instance_type=training_instance_type
)

# Retrieve the training script
train_source_uri = script_uri.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    script_scope=train_scope
)

train_model_uri = model_uri.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    model_scope=train_scope
)

# Sample training data is available in this bucket
training_data_bucket = f"jumpstart-cache-prod-{aws_region}"
training_data_prefix = "training-datasets/tabular_multiclass/"

training_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}/train"
validation_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}/validation"

output_bucket = sess.default_bucket()
output_prefix = "jumpstart-example-tabular-training"

s3_output_location = f"s3://{output_bucket}/{output_prefix}/output"

from sagemaker import hyperparameters

# Retrieve the default hyperparameters for training the model
hyperparameters = hyperparameters.retrieve_default(
    model_id=train_model_id, model_version=train_model_version)
# [Optional] Override default hyperparameters with custom values

```python
hyperparameters[  
    "iterations"  
] = "500"
print(hyperparameters)

from sagemaker.estimator import Estimator
from sagemaker.utils import name_from_base

training_job_name = name_from_base(f"built-in-algo-{train_model_id}-training")

# Create SageMaker Estimator instance

# Launch a SageMaker Training job by passing the S3 path of the training data
```

For more information about how to set up CatBoost as a built-in algorithm, see the following notebook examples.

- [Tabular classification with Amazon SageMaker LightGBM and CatBoost algorithm](#)
- [Tabular regression with Amazon SageMaker LightGBM and CatBoost algorithm](#)
Input and Output interface for the CatBoost algorithm

Gradient boosting operates on tabular data, with the rows representing observations, one column representing the target variable or label, and the remaining columns representing features.

The SageMaker implementation of CatBoost supports CSV for training and inference:

- For **Training ContentType**, valid inputs must be text/csv.
- For **Inference ContentType**, valid inputs must be text/csv.

**Note**

For CSV training, the algorithm assumes that the target variable is in the first column and that the CSV does not have a header record.

For CSV inference, the algorithm assumes that CSV input does not have the label column.

Input format for training data, validation data, and categorical features

Be mindful of how to format your training data for input to the CatBoost model. You must provide the path to an Amazon S3 bucket that contains your training and validation data. You can also include a list of categorical features. Use both the training and validation channels to provide your input data. Alternatively, you can use only the training channel.

Use both the training and validation channels

You can provide your input data by way of two S3 paths, one for the training channel and one for the validation channel. Each S3 path can either be an S3 prefix that points to one or more CSV files or a full S3 path pointing to one specific CSV file. The target variables should be in the first column of your CSV file. The predictor variables (features) should be in the remaining columns. If multiple CSV files are provided for the training or validation channels, the CatBoost algorithm concatenates the files. The validation data is used to compute a validation score at the end of each boosting iteration. Early stopping is applied when the validation score stops improving.

If your predictors include categorical features, you can provide a JSON file named categorical_index.json in the same location as your training data file or files. If you provide a JSON file for categorical features, your training channel must point to an S3 prefix and not a specific CSV file. This file should contain a Python dictionary where the key is the string "cat_index_list" and the value is a list of unique integers. Each integer in the value list should
indicate the column index of the corresponding categorical features in your training data CSV file. Each value should be a positive integer (greater than zero because zero represents the target value), less than the `Int32.MaxValue` (2147483647), and less than the total number of columns. There should only be one categorical index JSON file.

**Use only the training channel:**

You can alternatively provide your input data by way of a single S3 path for the training channel. This S3 path should point to a directory with a subdirectory named `training/` that contains one or more CSV files. You can optionally include another subdirectory in the same location called `validation/` that also has one or more CSV files. If the validation data is not provided, then 20% of your training data is randomly sampled to serve as the validation data. If your predictors include categorical features, you can provide a JSON file named `categorical_index.json` in the same location as your data subdirectories.

⚠️ **Note**

For CSV training input mode, the total memory available to the algorithm (instance count multiplied by the memory available in the `InstanceType`) must be able to hold the training dataset.

SageMaker CatBoost uses the `catboost.CatBoostClassifier` and `catboost.CatBoostRegressor` modules to serialize or deserialize the model, which can be used for saving or loading the model.

**To use a model trained with SageMaker CatBoost with catboost**

- Use the following Python code:

```python
import tarfile
from catboost import CatBoostClassifier

t = tarfile.open('model.tar.gz', 'r:gz')
t.extractall()

file_path = os.path.join(model_file_path, "model")
model = CatBoostClassifier()
model.load_model(file_path)
```
# prediction with test data
# dtest should be a pandas DataFrame with column names feature_0, feature_1, ..., feature_d
pred = model.predict(dtest)

Amazon EC2 instance recommendation for the CatBoost algorithm

SageMaker CatBoost currently only trains using CPUs. CatBoost is a memory-bound (as opposed to compute-bound) algorithm. So, a general-purpose compute instance (for example, M5) is a better choice than a compute-optimized instance (for example, C5). Further, we recommend that you have enough total memory in selected instances to hold the training data.

CatBoost sample notebooks

The following table outlines a variety of sample notebooks that address different use cases of Amazon SageMaker CatBoost algorithm.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tabular classification with Amazon SageMaker LightGBM and CatBoost algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker CatBoost algorithm to train and host a tabular classification model.</td>
</tr>
<tr>
<td>Tabular regression with Amazon SageMaker LightGBM and CatBoost algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker CatBoost algorithm to train and host a tabular regression model.</td>
</tr>
</tbody>
</table>

For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all of the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

How CatBoost Works

CatBoost implements a conventional Gradient Boosting Decision Tree (GBDT) algorithm with the addition of two critical algorithmic advances:

1. The implementation of ordered boosting, a permutation-driven alternative to the classic algorithm
2. An innovative algorithm for processing categorical features

Both techniques were created to fight a prediction shift caused by a special kind of target leakage present in all currently existing implementations of gradient boosting algorithms.

The CatBoost algorithm performs well in machine learning competitions because of its robust handling of a variety of data types, relationships, distributions, and the diversity of hyperparameters that you can fine-tune. You can use CatBoost for regression, classification (binary and multiclass), and ranking problems.

For more information on gradient boosting, see How XGBoost Works. For in-depth details about the additional GOSS and EFB techniques used in the CatBoost method, see CatBoost: unbiased boosting with categorical features.

CatBoost hyperparameters

The following table contains the subset of hyperparameters that are required or most commonly used for the Amazon SageMaker CatBoost algorithm. Users set these parameters to facilitate the estimation of model parameters from data. The SageMaker CatBoost algorithm is an implementation of the open-source CatBoost package.

Note

The default hyperparameters are based on example datasets in the CatBoost sample notebooks.

By default, the SageMaker CatBoost algorithm automatically chooses an evaluation metric and loss function based on the type of classification problem. The CatBoost algorithm detects the type of classification problem based on the number of labels in your data. For regression problems, the evaluation metric and loss functions are both root mean squared error. For binary classification problems, the evaluation metric is Area Under the Curve (AUC) and the loss function is log loss. For multiclass classification problems, the evaluation metric and loss functions are multiclass cross entropy. You can use the eval_metric hyperparameter to change the default evaluation metric. Refer to the following table for more information on LightGBM hyperparameters, including descriptions, valid values, and default values.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>iterations</td>
<td>The maximum number of trees that can be built.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: Positive integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 500.</td>
</tr>
<tr>
<td>early_stopping_rounds</td>
<td>The training will stop if one metric of one validation data point does not improve in the last early_stopping_rounds round. If early_stopping_rounds is less than or equal to zero, this hyperparameter is ignored.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 5.</td>
</tr>
<tr>
<td>eval_metric</td>
<td>The evaluation metric for validation data. If eval_metric is set to the default &quot;auto&quot; value, then the algorithm automatically chooses an evaluation metric based on the type of classification problem:</td>
</tr>
<tr>
<td></td>
<td>• &quot;RMSE&quot; for regression</td>
</tr>
<tr>
<td></td>
<td>• &quot;AUC&quot; for binary classification</td>
</tr>
<tr>
<td></td>
<td>• &quot;MultiClass&quot; for multi-class classification</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, refer to the <a href="https://catboost.ai/docs/#/en/algorithm-overview">CatBoost documentation</a> for valid values.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;auto&quot;.</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The rate at which the model weights are updated after working through each batch of training examples.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: (0.0, 1.0).</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.009.</td>
</tr>
<tr>
<td>depth</td>
<td>Depth of the tree.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: (1, 16).</td>
</tr>
<tr>
<td></td>
<td>Default value: 6.</td>
</tr>
<tr>
<td>l2_leaf_reg</td>
<td>Coefficient for the L2 regularization term of the cost function.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: Positive integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 3.</td>
</tr>
<tr>
<td>random_strength</td>
<td>The amount of randomness to use for scoring splits when the tree structure is selected. Use this parameter to avoid overfitting the model.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: Positive floating point number.</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0</td>
</tr>
<tr>
<td>max_leaves</td>
<td>The maximum number of leaves in the resulting tree. Can only be used with the &quot;Lossguide&quot; growing policy.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: [2, 64].</td>
</tr>
<tr>
<td></td>
<td>Default value: 31.</td>
</tr>
<tr>
<td>rsm</td>
<td>Random subspace method. The percentage of features to use at each split selection, when features are selected over again at random.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: (0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0</td>
</tr>
<tr>
<td>sampling_frequency</td>
<td>Frequency to sample weights and objects when building trees.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, either: (&quot;PerTreeLevel&quot; or &quot;PerTree&quot; ).</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;PerTreeLevel&quot; .</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>min_data_in_leaf</td>
<td>The minimum number of training samples in a leaf. CatBoost does not search for new splits in leaves with a sample count less than the specified value. Can only be used with the &quot;Lossguide&quot; and &quot;Depthwise&quot; growing policies. Valid values: integer, range: (1 or ∞). Default value: 1.</td>
</tr>
<tr>
<td>bagging_temperature</td>
<td>Defines the settings of the Bayesian bootstrap. Use the Bayesian bootstrap to assign random weights to objects. If bagging_temperature is set to 1.0, then the weights are sampled from an exponential distribution. If bagging_temperature is set to 0.0, then all weights are 1.0. Valid values: float, range: Non-negative float. Default value: 1.0.</td>
</tr>
<tr>
<td>boosting_type</td>
<td>The boosting scheme. &quot;Auto&quot; means that the boosting_type is selected based on processing unit type, the number of objects in the training dataset, and the selected learning mode. Valid values: string, any of the following: (&quot;Auto&quot;, &quot;Ordered&quot;, &quot;Plain&quot;). Default value: &quot;Auto&quot;.</td>
</tr>
<tr>
<td>scale_pos_weight</td>
<td>The weight for positive class in binary classification. The value is used as a multiplier for the weights of objects from positive class. Valid values: float, range: Positive float. Default value: 1.0.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>max_bin</td>
<td>The number of splits for numerical features. &quot;Auto&quot; means that max_bin is selected based on the processing unit type and other parameters. For details, see the CatBoost documentation. Valid values: string, either: (&quot;Auto&quot; or string of integer from &quot;1&quot; to &quot;65535&quot; inclusively). Default value: &quot;Auto&quot;.</td>
</tr>
<tr>
<td>grow_policy</td>
<td>The tree growing policy. Defines how to perform greedy tree construction. Valid values: string, any of the following: (&quot;SymmetricTree&quot;, &quot;Depthwise&quot;, or &quot;Lossguide&quot;). Default value: &quot;SymmetricTree&quot;.</td>
</tr>
<tr>
<td>random_seed</td>
<td>The random seed used for training. Valid values: integer, range: Non-negative integer. Default value: 1.0.</td>
</tr>
<tr>
<td>thread_count</td>
<td>The number of threads to use during the training. If thread_count is -1, then the number of threads is equal to the number of processor cores. thread_count cannot be 0. Valid values: integer, either: (-1 or positive integer). Default value: -1.</td>
</tr>
<tr>
<td>verbose</td>
<td>The verbosity of print messages, with higher levels corresponding to more detailed print statements. Valid values: integer, range: Positive integer. Default value: 1.</td>
</tr>
</tbody>
</table>
Tune a CatBoost model

Automatic model tuning, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your training and validation datasets. Model tuning focuses on the following hyperparameters:

- A learning loss function to optimize during model training
- An evaluation metric that is used to evaluate model performance during validation
- A set of hyperparameters and a range of values for each to use when tuning the model automatically

Automatic model tuning searches your chosen hyperparameters to find the combination of values that results in a model that optimizes the chosen evaluation metric.

Note
The learning loss function is automatically assigned based on the type of classification task, which is determined by the number of unique integers in the label column. For more information, see CatBoost hyperparameters.

Note
Automatic model tuning for CatBoost is only available from the Amazon SageMaker SDKs, not from the SageMaker console.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Evaluation metrics computed by the CatBoost algorithm

The SageMaker CatBoost algorithm computes the following metrics to use for model validation. The evaluation metric is automatically assigned based on the type of classification task, which is determined by the number of unique integers in the label column.
<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
<th>Regex Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>RMSE</td>
<td>root mean square error</td>
<td>minimize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>MAE</td>
<td>mean absolute error</td>
<td>minimize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>MedianAbsoluteError</td>
<td>median absolute error</td>
<td>minimize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>R2</td>
<td>r2 score</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>Logloss</td>
<td>binary cross entropy</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>Precision</td>
<td>precision</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>Recall</td>
<td>recall</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>F1</td>
<td>f1 score</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>AUC</td>
<td>auc score</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
</tbody>
</table>

**Use Built-in Algorithms**

2695
<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
<th>Regex Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>MultiClass</td>
<td>multiclass cross entropy</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>Accuracy</td>
<td>accuracy</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
<tr>
<td>BalancedAccuracy</td>
<td>balanced accuracy</td>
<td>maximize</td>
<td>&quot;bestTest = ([0-9.]+)&quot;</td>
</tr>
</tbody>
</table>

**Tunable CatBoost hyperparameters**

Tune the CatBoost model with the following hyperparameters. The hyperparameters that have the greatest effect on optimizing the CatBoost evaluation metrics are: `learning_rate`, `depth`, `l2_leaf_reg`, and `random_strength`. For a list of all the CatBoost hyperparameters, see the [CatBoost hyperparameters](#).  

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.001, MaxValue: 0.01</td>
</tr>
<tr>
<td>depth</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 4, MaxValue: 10</td>
</tr>
<tr>
<td>l2_leaf_reg</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 2, MaxValue: 10</td>
</tr>
<tr>
<td>random_strength</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 10</td>
</tr>
</tbody>
</table>
Factorization Machines Algorithm

The Factorization Machines algorithm is a general-purpose supervised learning algorithm that you can use for both classification and regression tasks. It is an extension of a linear model that is designed to capture interactions between features within high dimensional sparse datasets economically. For example, in a click prediction system, the Factorization Machines model can capture click rate patterns observed when ads from a certain ad-category are placed on pages from a certain page-category. Factorization machines are a good choice for tasks dealing with high dimensional sparse datasets, such as click prediction and item recommendation.

 {}: **Note**

The Amazon SageMaker implementation of the Factorization Machines algorithm considers only pair-wise (2nd order) interactions between features.

Topics

- Input/Output Interface for the Factorization Machines Algorithm
- EC2 Instance Recommendation for the Factorization Machines Algorithm
- Factorization Machines Sample Notebooks
- How Factorization Machines Work
- Factorization Machines Hyperparameters
- Tune a Factorization Machines Model
- Factorization Machines Response Formats

Input/Output Interface for the Factorization Machines Algorithm

The Factorization Machines algorithm can be run in either in binary classification mode or regression mode. In each mode, a dataset can be provided to the **test** channel along with the **train** channel dataset. The scoring depends on the mode used. In regression mode, the testing dataset is scored using Root Mean Square Error (RMSE). In binary classification mode, the test dataset is scored using Binary Cross Entropy (Log Loss), Accuracy (at threshold=0.5) and F1 Score (at threshold =0.5).

For **training**, the Factorization Machines algorithm currently supports only the **recordIO-protobuf** format with **Float32** tensors. Because their use case is predominantly on sparse data,
CSV is not a good candidate. Both File and Pipe mode training are supported for recordIO-wrapped protobuf.

For **inference**, the Factorization Machines algorithm supports the application/json and x-recordio-protobuf formats.

- For the **binary classification** problem, the algorithm predicts a score and a label. The label is a number and can be either 0 or 1. The score is a number that indicates how strongly the algorithm believes that the label should be 1. The algorithm computes score first and then derives the label from the score value. If the score is greater than or equal to 0.5, the label is 1.

- For the **regression** problem, just a score is returned and it is the predicted value. For example, if Factorization Machines is used to predict a movie rating, score is the predicted rating value.

Please see [Factorization Machines Sample Notebooks](#) for more details on training and inference file formats.

**EC2 Instance Recommendation for the Factorization Machines Algorithm**

The Amazon SageMaker Factorization Machines algorithm is highly scalable and can train across distributed instances. We recommend training and inference with CPU instances for both sparse and dense datasets. In some circumstances, training with one or more GPUs on dense data might provide some benefit. Training with GPUs is available only on dense data. Use CPU instances for sparse data. The Factorization Machines algorithm supports P2, P3, G4dn, and G5 instances for training and inference.

**Factorization Machines Sample Notebooks**

For a sample notebook that uses the SageMaker Factorization Machines algorithm to analyze the images of handwritten digits from zero to nine in the MNIST dataset, see [An Introduction to Factorization Machines with MNIST](#). For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](#). Once you have created a notebook instance and opened it, select the **SageMaker Examples** tab to see a list of all the SageMaker samples. Example notebooks that use Factorization Machines algorithm are located in the **Introduction to Amazon algorithms** section. To open a notebook, click on its **Use** tab and select **Create copy**.
How Factorization Machines Work

The prediction task for a Factorization Machines model is to estimate a function $\hat{y}$ from a feature set $x_i$ to a target domain. This domain is real-valued for regression and binary for classification. The Factorization Machines model is supervised and so has a training dataset $(x_i, y_j)$ available. The advantages this model presents lie in the way it uses a factorized parametrization to capture the pairwise feature interactions. It can be represented mathematically as follows:

$$\hat{y} = w_0 + \sum_i w_i x_i + \sum_i \sum_{j > i} <v_i, v_j> x_i x_j$$

The three terms in this equation correspond respectively to the three components of the model:

- The $w_0$ term represents the global bias.
- The $w_i$ linear terms model the strength of the $i^{th}$ variable.
- The $<v_i, v_j>$ factorization terms model the pairwise interaction between the $i^{th}$ and $j^{th}$ variable.

The global bias and linear terms are the same as in a linear model. The pairwise feature interactions are modeled in the third term as the inner product of the corresponding factors learned for each feature. Learned factors can also be considered as embedding vectors for each feature. For example, in a classification task, if a pair of features tends to co-occur more often in positive labeled samples, then the inner product of their factors would be large. In other words, their embedding vectors would be close to each other in cosine similarity. For more information about the Factorization Machines model, see Factorization Machines.

For regression tasks, the model is trained by minimizing the squared error between the model prediction $\hat{y}_n$ and the target value $y_n$. This is known as the square loss:

$$L = \frac{1}{N} \sum_n (y_n - \hat{y}_n)^2$$

For a classification task, the model is trained by minimizing the cross entropy loss, also known as the log loss:

$$L = \frac{1}{N} \sum_n [y_n \log \hat{p}_n + (1 - y_n) \log (1 - \hat{p}_n)]$$

where:

$$\hat{p}_n = \frac{1}{1 + e^{-\hat{y}_n}}$$
For more information about loss functions for classification, see Loss functions for classification.

Factorization Machines Hyperparameters

The following table contains the hyperparameters for the Factorization Machines algorithm. These are parameters that are set by users to facilitate the estimation of model parameters from data. The required hyperparameters that must be set are listed first, in alphabetical order. The optional hyperparameters that can be set are listed next, also in alphabetical order.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>feature_dim</td>
<td>The dimension of the input feature space. This could be very high with sparse input.</td>
</tr>
<tr>
<td></td>
<td>Required</td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer. Suggested value range: [10000,10000000]</td>
</tr>
<tr>
<td>num_factors</td>
<td>The dimensionality of factorization.</td>
</tr>
<tr>
<td></td>
<td>Required</td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer. Suggested value range: [2,1000], 64 typically generates good outcomes and is a good starting point.</td>
</tr>
<tr>
<td>predictor_type</td>
<td>The type of predictor.</td>
</tr>
<tr>
<td></td>
<td>Required</td>
</tr>
<tr>
<td></td>
<td>Valid values: String: binary_classifier or regressor</td>
</tr>
<tr>
<td>bias_init_method</td>
<td>The initialization method for the bias term:</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
|                      | • normal: Initializes weights with random values sampled from a normal distribution with a mean of zero and standard deviation specified by bias_init_sigma.  
|                      | • uniform: Initializes weights with random values uniformly sampled from a range specified by [-bias_init_scale, +bias_init_scale].  
|                      | • constant: Initializes the weights to a scalar value specified by bias_init_value.  
|                      | **Optional**                                                                                                                                                                                                                                                                                                                                 |
|                      | Valid values: uniform, normal, or constant                                                                                                                                                                                                                                                                                                   |
|                      | Default value: normal                                                                                                                                                                                                                                                                                                                          |
| bias_init_scale      | Range for initialization of the bias term. Takes effect if bias_init_method is set to uniform.                                                                                                                                                                                                                                             |
|                      | **Optional**                                                                                                                                                                                                                                                                                                                                 |
|                      | Valid values: Non-negative float. Suggested value range: [1e-8, 512].                                                                                                                                                                                                                                                                     |
|                      | Default value: None                                                                                                                                                                                                                                                                                                                            |
| bias_init_sigma      | The standard deviation for initialization of the bias term. Takes effect if bias_init_method is set to normal.                                                                                                                                                                     |
|                      | **Optional**                                                                                                                                                                                                                                                                                                                                 |
|                      | Valid values: Non-negative float. Suggested value range: [1e-8, 512].                                                                                                                                                                                                                                                                     |
|                      | Default value: 0.01
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bias_init_value</td>
<td>The initial value of the bias term. Takes effect if bias_init_method is set to constant. Optional Valid values: Float. Suggested value range: [1e-8, 512]. Default value: None</td>
</tr>
<tr>
<td>bias_lr</td>
<td>The learning rate for the bias term. Optional Valid values: Non-negative float. Suggested value range: [1e-8, 512]. Default value: 0.1</td>
</tr>
<tr>
<td>bias_wd</td>
<td>The weight decay for the bias term. Optional Valid values: Non-negative float. Suggested value range: [1e-8, 512]. Default value: 0.01</td>
</tr>
<tr>
<td>clip_gradient</td>
<td>Gradient clipping optimizer parameter. Clips the gradient by projecting onto the interval [-clip_gradient, +clip_gradient]. Optional Valid values: Float Default value: None</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| epochs         | The number of training epochs to run.  
   **Optional**  
   Valid values: Positive integer  
   Default value: 1 |
| eps            | Epsilon parameter to avoid division by 0.  
   **Optional**  
   Valid values: Float. Suggested value: small.  
   Default value: None |
| factors_init_method | The initialization method for factorization terms:  
   - **normal**: Initializes weights with random values sampled from a normal distribution with a mean of zero and standard deviation specified by `factors_init_sigma`.  
   - **uniform**: Initializes weights with random values uniformly sampled from a range specified by `[-factors_init_scale, +factors_init_scale]`.  
   - **constant**: Initializes the weights to a scalar value specified by `factors_init_value`.  
   **Optional**  
   Valid values: uniform, normal, or constant.  
   Default value: normal |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>factors_init_scale</td>
<td>The range for initialization of factorization terms. Takes effect if <code>factors_init_method</code> is set to <code>uniform</code>.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td>factors_init_sigma</td>
<td>The standard deviation for initialization of factorization terms. Takes effect if <code>factors_init_method</code> is set to <code>normal</code>.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td>factors_init_value</td>
<td>The initial value of factorization terms. Takes effect if <code>factors_init_method</code> is set to <code>constant</code>.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td>factors_lr</td>
<td>The learning rate for factorization terms.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>factors_wd</td>
<td>The weight decay for factorization terms.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.00001</td>
</tr>
<tr>
<td>linear_lr</td>
<td>The learning rate for linear terms.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.001</td>
</tr>
<tr>
<td>linear_init_method</td>
<td>The initialization method for linear terms:</td>
</tr>
<tr>
<td></td>
<td>• normal Initializes weights with random values sampled from a normal distribution with a mean of zero and standard deviation specified by linear_init_sigma.</td>
</tr>
<tr>
<td></td>
<td>• uniform Initializes weights with random values uniformly sampled from a range specified by [−linear_init_scale, +linear_init_scale].</td>
</tr>
<tr>
<td></td>
<td>• constant Initializes the weights to a scalar value specified by linear_init_value.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: uniform, normal, or constant.</td>
</tr>
<tr>
<td></td>
<td>Default value: normal</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>linear_init_scale</td>
<td>Range for initialization of linear terms. Takes effect if <code>linear_init_method</code> is set to <code>uniform</code>.</td>
</tr>
<tr>
<td>Optional</td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>linear_init_sigma</td>
<td>The standard deviation for initialization of linear terms. Takes effect if <code>linear_init_method</code> is set to <code>normal</code>.</td>
</tr>
<tr>
<td>Optional</td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.01</td>
</tr>
<tr>
<td>linear_init_value</td>
<td>The initial value of linear terms. Takes effect if <code>linear_init_method</code> is set to <code>constant</code>.</td>
</tr>
<tr>
<td>Optional</td>
<td>Valid values: Float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>linear_wd</td>
<td>The weight decay for linear terms.</td>
</tr>
<tr>
<td>Optional</td>
<td>Valid values: Non-negative float. Suggested value range: [1e-8, 512].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.001</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| mini_batch_size | The size of mini-batch used for training.  
**Optional**  
Valid values: Positive integer  
Default value: 1000 |
| rescale_grad | Gradient rescaling optimizer parameter. If set, multiplies the gradient with $\text{rescale\_grad}$ before updating. Often choose to be $1.0 / \text{batch\_size}$.  
**Optional**  
Valid values: Float  
Default value: None |

**Tune a Factorization Machines Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics Computed by the Factorization Machines Algorithm**

The Factorization Machines algorithm has both binary classification and regression predictor types. The predictor type determines which metric you can use for automatic model tuning. The algorithm reports a $\text{test:rmse}$ regressor metric, which is computed during training. When tuning the model for regression tasks, choose this metric as the objective.
The Factorization Machines algorithm reports three binary classification metrics, which are computed during training. When tuning the model for binary classification tasks, choose one of these as the objective.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:rmse</td>
<td>Root Mean Square Error</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:binary_classification_accuracy</td>
<td>Accuracy</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:binary_classification_cross_entropy</td>
<td>Cross Entropy</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:binary_f_beta</td>
<td>Beta</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

**Tunable Factorization Machines Hyperparameters**

You can tune the following hyperparameters for the Factorization Machines algorithm. The initialization parameters that contain the terms bias, linear, and factorization depend on their initialization method. There are three initialization methods: uniform, normal, and constant. These initialization methods are not themselves tunable. The parameters that are tunable are dependent on this choice of the initialization method. For example, if the initialization method is uniform, then only the scale parameters are tunable. Specifically, if bias_init_method==uniform, then bias_init_scale, linear_init_scale, and factors_init_scale are tunable. Similarly, if the initialization method is normal, then
only sigma parameters are tunable. If the initialization method is constant, then only value parameters are tunable. These dependencies are listed in the following table.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
<th>Dependency</th>
</tr>
</thead>
<tbody>
<tr>
<td>bias_init_scale</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>bias_init_method==uniform</td>
</tr>
<tr>
<td>bias_init_sigma</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>bias_init_method==normal</td>
</tr>
<tr>
<td>bias_init_value</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>bias_init_method==constant</td>
</tr>
<tr>
<td>bias_lr</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>None</td>
</tr>
<tr>
<td>bias_wd</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>None</td>
</tr>
<tr>
<td>epoch</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 1000</td>
<td>None</td>
</tr>
<tr>
<td>factors_init_scale</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>bias_init_method==uniform</td>
</tr>
<tr>
<td>factors_init_sigma</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>bias_init_method==normal</td>
</tr>
<tr>
<td>factors_init_value</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8, MaxValue: 512</td>
<td>bias_init_method==constant</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
<td>Dependency</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------------------</td>
<td>---------------------------</td>
<td>------------</td>
</tr>
<tr>
<td>factors_lr</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>MaxValue: 512</td>
<td></td>
</tr>
<tr>
<td>factors_wd</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>MaxValue: 512</td>
<td></td>
</tr>
<tr>
<td>linear_in</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td>bias_init _method==</td>
</tr>
<tr>
<td>it_scale</td>
<td></td>
<td>MaxValue: 512</td>
<td>uniform</td>
</tr>
<tr>
<td>linear_in</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td>bias_init _method==</td>
</tr>
<tr>
<td>it_sigma</td>
<td></td>
<td>MaxValue: 512</td>
<td>normal</td>
</tr>
<tr>
<td>linear_in</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td>bias_init _method==</td>
</tr>
<tr>
<td>it_value</td>
<td></td>
<td>MaxValue: 512</td>
<td>constant</td>
</tr>
<tr>
<td>linear_lr</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>MaxValue: 512</td>
<td></td>
</tr>
<tr>
<td>linear_wd</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-8,</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>MaxValue: 512</td>
<td></td>
</tr>
<tr>
<td>mini_batch</td>
<td>IntegerParameterRange</td>
<td>MinValue: 100,</td>
<td></td>
</tr>
<tr>
<td>h_size</td>
<td></td>
<td>MaxValue: 10000</td>
<td></td>
</tr>
</tbody>
</table>

**Factorization Machines Response Formats**

**JSON Response Format**

Binary classification

```javascript
let response = {
    "predictions": [
        {
            "score": 0.4,
```
Regression

```javascript
let response = {
    "predictions": [
        {
            "score": 0.4
        }
    ]
}
```

**JSONLINES Response Format**

**Binary classification**

```json
{"score": 0.4, "predicted_label": 0}
```

**Regression**

```json
{"score": 0.4}
```

**RECORDIO Response Format**

**Binary classification**

```json
[
    Record = {
        features = {},
        label = {
            'score': {
                keys: [],
                values: [0.4]  # float32
            },
            'predicted_label': {
                keys: [],
                values: [0.0]  # float32
            }
        }
    }
]```
Regression

[ ]

K-Nearest Neighbors (k-NN) Algorithm

Amazon SageMaker k-nearest neighbors (k-NN) algorithm is an index-based algorithm. It uses a non-parametric method for classification or regression. For classification problems, the algorithm queries the $k$ points that are closest to the sample point and returns the most frequently used label of their class as the predicted label. For regression problems, the algorithm queries the $k$ closest points to the sample point and returns the average of their feature values as the predicted value.

Training with the k-NN algorithm has three steps: sampling, dimension reduction, and index building. Sampling reduces the size of the initial dataset so that it fits into memory. For dimension reduction, the algorithm decreases the feature dimension of the data to reduce the footprint of the k-NN model in memory and inference latency. We provide two methods of dimension reduction methods: random projection and the fast Johnson-Lindenstrauss transform. Typically, you use dimension reduction for high-dimensional ($d >1000$) datasets to avoid the “curse of dimensionality” that troubles the statistical analysis of data that becomes sparse as dimensionality increases. The main objective of k-NN's training is to construct the index. The index enables efficient lookups of distances between points whose values or class labels have not yet been determined and the k nearest points to use for inference.

Topics

- Input/Output Interface for the k-NN Algorithm
Input/Output Interface for the k-NN Algorithm

SageMaker k-NN supports train and test data channels.

- Use a *train channel* for data that you want to sample and construct into the k-NN index.
- Use a *test channel* to emit scores in log files. Scores are listed as one line per mini-batch: accuracy for classifier, mean-squared error (mse) for regressor for score.

For training inputs, k-NN supports text/csv and application/x-recordio-protobuf data formats. For input type text/csv, the first label_size columns are interpreted as the label vector for that row. You can use either File mode or Pipe mode to train models on data that is formatted as recordIO-wrapped-protobuf or as CSV.

For inference inputs, k-NN supports the application/json, application/x-recordio-protobuf, and text/csv data formats. The text/csv format accepts a label_size and encoding parameter. It assumes a label_size of 0 and a UTF-8 encoding.

For inference outputs, k-NN supports the application/json and application/x-recordio-protobuf data formats. These two data formats also support a verbose output mode. In verbose output mode, the API provides the search results with the distances vector sorted from smallest to largest, and corresponding elements in the labels vector.

For batch transform, k-NN supports the application/jsonlines data format for both input and output. An example input is as follows:

```
content-type: application/jsonlines

{"features": [1.5, 16.0, 14.0, 23.0]}
```
An example output is as follows:

```json
accept: application/json

{"predicted_label": 0.0}
{"predicted_label": 2.0}
```

For more information on input and output file formats, see Data Formats for k-NN Training Input for training, k-NN Request and Response Formats for inference, and the k-NN Sample Notebooks.

**k-NN Sample Notebooks**

For a sample notebook that uses the SageMaker k-nearest neighbor algorithm to predict wilderness cover types from geological and forest service data, see the K-Nearest Neighbor Covertype.

Use a Jupyter notebook instance to run the example in SageMaker. To learn how to create and open a Jupyter notebook instance in SageMaker, see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker example notebooks. Find K-Nearest Neighbor notebooks in the Introduction to Amazon algorithms section. To open a notebook, click on its Use tab and select Create copy.

**How the k-NN Algorithm Works**

**Step 1: Sample**

To specify the total number of data points to be sampled from the training dataset, use the sample_size parameter. For example, if the initial dataset has 1,000 data points and the sample_size is set to 100, where the total number of instances is 2, each worker would sample 50 points. A total set of 100 data points would be collected. Sampling runs in linear time with respect to the number of data points.

**Step 2: Perform Dimension Reduction**

The current implementation of the k-NN algorithm has two methods of dimension reduction. You specify the method in the dimension_reduction_type hyperparameter. The sign method specifies a random projection, which uses a linear projection using a matrix of random signs,
and the fjlt method specifies a fast Johnson-Lindenstrauss transform, a method based on the Fourier transform. Both methods preserve the L2 and inner product distances. The fjlt method should be used when the target dimension is large and has better performance with CPU inference. The methods differ in their computational complexity. The sign method requires $O(ndk)$ time to reduce the dimension of a batch of $n$ points of dimension $d$ into a target dimension $k$. The fjlt method requires $O(nd \log(d))$ time, but the constants involved are larger. Using dimension reduction introduces noise into the data and this noise can reduce prediction accuracy.

**Step 3: Build an Index**

During inference, the algorithm queries the index for the k-nearest-neighbors of a sample point. Based on the references to the points, the algorithm makes the classification or regression prediction. It makes the prediction based on the class labels or values provided. k-NN provides three different types of indexes: a flat index, an inverted index, and an inverted index with product quantization. You specify the type with the `index_type` parameter.

**Serialize the Model**

When the k-NN algorithm finishes training, it serializes three files to prepare for inference.

- `model_algo-1`: Contains the serialized index for computing the nearest neighbors.
- `model_algo-1.labels`: Contains serialized labels (`np.float32` binary format) for computing the predicted label based on the query result from the index.
- `model_algo-1.json`: Contains the JSON-formatted model metadata which stores the k and `predictor_type` hyper-parameters from training for inference along with other relevant state.

With the current implementation of k-NN, you can modify the metadata file to change the way predictions are computed. For example, you can change $k$ to 10 or change `predictor_type` to `regressor`.

```json
{
    "k": 5,
    "predictor_type": "classifier",
    "dimension_reduction": {
        "type": "sign", 
        "seed": 3,
        "target_dim": 10,
        "input_dim": 20
    },
    "normalize": false,
    "version": "1.0"
}
```
EC2 Instance Recommendation for the k-NN Algorithm

We recommend training on a CPU instance (such as ml.m5.2xlarge) or on a GPU instance. The k-NN algorithm supports P2, P3, G4dn, and G5 GPU instance families for training and inference.

Inference requests from CPUs generally have a lower average latency than requests from GPUs because there is a tax on CPU-to-GPU communication when you use GPU hardware. However, GPUs generally have higher throughput for larger batches.

**k-NN Hyperparameters**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>feature_dim</td>
<td>The number of features in the input data.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer.</td>
</tr>
<tr>
<td>k</td>
<td>The number of nearest neighbors.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer.</td>
</tr>
<tr>
<td>predictor_type</td>
<td>The type of inference to use on the data labels.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: <code>classifier</code> for classification or <code>regressor</code> for regression.</td>
</tr>
<tr>
<td>sample_size</td>
<td>The number of data points to be sampled from the training data set.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer.</td>
</tr>
<tr>
<td>dimension_reduction_target</td>
<td>The target dimension to reduce to.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong> when you specify the <code>dimension_reduction_type</code> parameter.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>** Valid values: positive integer greater than 0 and less than feature_dim.</td>
<td></td>
</tr>
<tr>
<td>** dimension_reduction_type</td>
<td>The type of dimension reduction method.</td>
</tr>
<tr>
<td>** Optional</td>
<td></td>
</tr>
<tr>
<td>Valid values: sign for random projection or fjlt for the fast Johnson-Lindenstrauss transform.</td>
<td></td>
</tr>
<tr>
<td>Default value: No dimension reduction</td>
<td></td>
</tr>
<tr>
<td>** faiss_index_ivf_nlists</td>
<td>The number of centroids to construct in the index when index_type is faiss.IVFFlat or faiss.IVFPQ.</td>
</tr>
<tr>
<td>** Optional</td>
<td></td>
</tr>
<tr>
<td>Valid values: positive integer</td>
<td></td>
</tr>
<tr>
<td>Default value: auto, which resolves to sqrt(sample_size)</td>
<td></td>
</tr>
<tr>
<td>** faiss_index_pq_m</td>
<td>The number of vector sub-components to construct in the index when index_type is set to faiss.IVFPQ.</td>
</tr>
<tr>
<td>** The FaceBook AI Similarity Search (FAISS) library requires that the value of faiss_index_pq_m is a divisor of the data dimension. If faiss_index_pq_m is not a divisor of the data dimension, we increase the data dimension to smallest integer divisible by faiss_index_pq_m. If no dimension reduction is applied, the algorithm adds a padding of zeros. If dimension reduction is applied, the algorithm increase the value of the dimension_reduction_target hyper-parameter.</td>
<td>** Optional</td>
</tr>
<tr>
<td>Valid values: One of the following positive integers: 1, 2, 3, 4, 8, 12, 16, 20, 24, 28, 32, 40, 48, 56, 64, 96</td>
<td></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td><strong>index_metric</strong></td>
<td>The metric to measure the distance between points when finding nearest neighbors. When training with <code>index_type</code> set to <code>faiss.IVFPQ</code>, the <code>INNER_PRODUCT</code> distance and <code>COSINE</code> similarity are not supported.</td>
</tr>
<tr>
<td><strong>index_type</strong></td>
<td>The type of index.</td>
</tr>
<tr>
<td><strong>mini_batch_size</strong></td>
<td>The number of observations per mini-batch for the data iterator.</td>
</tr>
</tbody>
</table>

**Optional**

Valid values: `L2` for Euclidean-distance, `INNER_PRODUCT` for inner-product distance, `COSINE` for cosine similarity.

Default value: `L2`

Valid values: `faiss.Flat`, `faiss.IVFFlat`, `faiss.IVFPQ`.

Default values: `faiss.Flat`

Valid values: positive integer

Default value: 5000

**Tune a k-NN Model**

The Amazon SageMaker k-nearest neighbors algorithm is a supervised algorithm. The algorithm consumes a test data set and emits a metric about the accuracy for a classification task or about the mean squared error for a regression task. These accuracy metrics compare the model predictions for their respective task to the ground truth provided by the empirical test data. To find the best model that reports the highest accuracy or lowest error on the test dataset, run a hyperparameter tuning job for k-NN.
Automatic model tuning, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric appropriate for the prediction task of the algorithm. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric. The hyperparameters are used only to help estimate model parameters and are not used by the trained model to make predictions.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Metrics Computed by the k-NN Algorithm

The k-nearest neighbors algorithm computes one of two metrics in the following table during training depending on the type of task specified by the predictor_type hyper-parameter.

- **classifier** specifies a classification task and computes `test:accuracy`
- **regressor** specifies a regression task and computes `test:mse`.

Choose the predictor_type value appropriate for the type of task undertaken to calculate the relevant objective metric when tuning a model.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>test:accuracy</code></td>
<td>When predictor_type is set to classifier, k-NN compares the predicted label, based on the average of the k-nearest neighbors' labels, to the ground truth label provided in the test channel data. The accuracy reported ranges from 0.0 (0%) to 1.0 (100%).</td>
<td>Maximize</td>
</tr>
<tr>
<td><code>test:mse</code></td>
<td>When predictor_type is set to regressor, k-NN compares the predicted label, based on the average of the k-nearest neighbors' labels, to the ground truth label provided in the test channel data. The mean squared error is computed by comparing the two labels.</td>
<td>Minimize</td>
</tr>
</tbody>
</table>
Tunable k-NN Hyperparameters

Tune the Amazon SageMaker k-nearest neighbor model with the following hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>k</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 1024</td>
</tr>
<tr>
<td>sample_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 256, MaxValue: 2000000</td>
</tr>
</tbody>
</table>

Data Formats for k-NN Training Input

All Amazon SageMaker built-in algorithms adhere to the common input training formats described in Common Data Formats - Training. This topic contains a list of the available input formats for the SageMaker k-nearest-neighbor algorithm.

**CSV Data Format**

```plaintext
columns: text/csv; label_size=1
```

```
4,1.2,1.3,9.6,20.3
```

The first `label_size` columns are interpreted as the label vector for that row.

**RECORDIO Data Format**

```plaintext
text-type: application/x-recordio-protobuf
```

```
[       
    Record = {      
        features = {      
            'values': {      
                values: [1.2, 1.3, 9.6, 20.3]  # float32
            },
        },
    },
]`
```python
label = {
    'values': {
        'values': [4]  # float32
    }
}
```

**k-NN Request and Response Formats**

All Amazon SageMaker built-in algorithms adhere to the common input inference format described in [Common Data Formats - Inference](#). This topic contains a list of the available output formats for the SageMaker k-nearest-neighbor algorithm.

**INPUT: CSV Request Format**

```text
content-type: text/csv

1.2,1.3,9.6,20.3
```

This accepts a `label_size` or encoding parameter. It assumes a `label_size` of 0 and a utf-8 encoding.

**INPUT: JSON Request Format**

```json
content-type: application/json

{
    "instances": [
        {
            "data": {
                "features": {
                    "values": [-3, -1, -4, 2]
                }
            }
        },
        {
            "features": [3.0, 0.1, 0.04, 0.002]
        }
    ]
}
```

**INPUT: JSONLINES Request Format**

```json
content-type: application/jsonlines

{"features": [1.5, 16.0, 14.0, 23.0]}
```
Amazon SageMaker

Developer Guide

{"data": {"features": {"values": [1.5, 16.0, 14.0, 23.0]}}

INPUT: RECORDIO Request Format
content-type: application/x-recordio-protobuf
[
Record = {
features = {
'values': {
values: [-3, -1, -4, 2] # float32
}
},
label = {}
},
Record = {
features = {
'values': {
values: [3.0, 0.1, 0.04, 0.002] # float32
}
},
label = {}
},
]

OUTPUT: JSON Response Format
accept: application/json
{
"predictions": [
{"predicted_label": 0.0},
{"predicted_label": 2.0}
]
}

OUTPUT: JSONLINES Response Format
accept: application/jsonlines
{"predicted_label": 0.0}
{"predicted_label": 2.0}

Use Built-in Algorithms

2722


OUTPUT: VERBOSE JSON Response Format

In verbose mode, the API provides the search results with the distances vector sorted from smallest to largest, with corresponding elements in the labels vector. In this example, k is set to 3.

accept: application/json; verbose=true

```
{
    "predictions": [
        {
            "predicted_label": 0.0,
            "distances": [3.11792408, 3.89746071, 6.32548437],
            "labels": [0.0, 1.0, 0.0]
        },
        {
            "predicted_label": 2.0,
            "distances": [1.08470316, 3.04917915, 5.25393973],
            "labels": [2.0, 2.0, 0.0]
        }
    ]
}
```

OUTPUT: RECORDIO-PROTobuf Response Format

content-type: application/x-recordio-protobuf

```
[
    Record = {
        features = {},
        label = {
            'predicted_label': {
                values: [0.0]  # float32
            }
        }
    },
    Record = {
        features = {},
        label = {
            'predicted_label': {
                values: [2.0]  # float32
            }
        }
    }
]
```
OUTPUT: VERBOSE RECORDIO-PROTobuf Response Format

In verbose mode, the API provides the search results with the distances vector sorted from smallest to largest, with corresponding elements in the labels vector. In this example, k is set to 3.

```
accept: application/x-recordio-protobuf; verbose=true
```

```
[  
  Record = {  
    features = {},  
    label = {  
      'predicted_label': {  
        values: [0.0]  # float32  
      },  
      'distances': {  
        values: [3.11792408, 3.89746071, 6.32548437]  # float32  
      },  
      'labels': {  
        values: [0.0, 1.0, 0.0]  # float32  
      }  
    }  
  },  
  Record = {  
    features = {},  
    label = {  
      'predicted_label': {  
        values: [0.0]  # float32  
      },  
      'distances': {  
        values: [1.08470316, 3.04917915, 5.25393973]  # float32  
      },  
      'labels': {  
        values: [2.0, 2.0, 0.0]  # float32  
      }  
    }  
  }
]
```

SAMPLE OUTPUT for the k-NN Algorithm

For regressor tasks:
LightGBM

LightGBM is a popular and efficient open-source implementation of the Gradient Boosting Decision Tree (GBDT) algorithm. GBDT is a supervised learning algorithm that attempts to accurately predict a target variable by combining an ensemble of estimates from a set of simpler and weaker models. LightGBM uses additional techniques to significantly improve the efficiency and scalability of conventional GBDT.

How to use SageMaker LightGBM

You can use LightGBM as an Amazon SageMaker built-in algorithm. The following section describes how to use LightGBM with the SageMaker Python SDK. For information on how to use LightGBM from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

- Use LightGBM as a built-in algorithm

Use the LightGBM built-in algorithm to build a LightGBM training container as shown in the following code example. You can automatically spot the LightGBM built-in algorithm image URI using the SageMaker image_uris.retrieve API (or the get_image_uri API if using Amazon SageMaker Python SDK version 2).

After specifying the LightGBM image URI, you can use the LightGBM container to construct an estimator using the SageMaker Estimator API and initiate a training job. The LightGBM built-in algorithm runs in script mode, but the training script is provided for you and there is no need to replace it. If you have extensive experience using script mode to create a SageMaker training job, then you can incorporate your own LightGBM training scripts.

```python
from sagemaker import image_uris, model_uris, script_uris

train_model_id, train_model_version, train_scope = "lightgbm-classification-model", "*", "training"
training_instance_type = "ml.m5.xlarge"
```
# Retrieve the docker image
train_image_uri = image_uris.retrieve(
    region=None,
    framework=None,
    model_id=train_model_id,
    model_version=train_model_version,
    image_scope=train_scope,
    instance_type=training_instance_type
)

# Retrieve the training script
train_source_uri = script_uris.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    script_scope=train_scope
)

train_model_uri = model_uris.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    model_scope=train_scope
)

# Sample training data is available in this bucket
training_data_bucket = f"jumpstart-cache-prod-{aws_region}"
training_data_prefix = "training-datasets/tabular_multiclass/"

training_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}/train"
validation_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}/validation"

output_bucket = sess.default_bucket()
output_prefix = "jumpstart-example-tabular-training"
s3_output_location = f"s3://{output_bucket}/{output_prefix}/output"

from sagemaker import hyperparameters

# Retrieve the default hyperparameters for training the model
hyperparameters = hyperparameters.retrieve_default(
    model_id=train_model_id, model_version=train_model_version
)

# [Optional] Override default hyperparameters with custom values
```python
hyperparameters[
    "num_boost_round"
] = "500"
print(hyperparameters)

from sagemaker.estimator import Estimator
from sagemaker.utils import name_from_base

training_job_name = name_from_base(f"built-in-algo-{train_model_id}-training")

# Create SageMaker Estimator instance
tabular_estimator = Estimator(
    role=aws_role,
    image_uri=train_image_uri,
    source_dir=train_source_uri,
    model_uri=train_model_uri,
    entry_point="transfer_learning.py",
    instance_count=1, # for distributed training, specify an instance_count greater than 1
    instance_type=training_instance_type,
    max_run=360000,
    hyperparameters=hyperparameters,
    output_path=s3_output_location
)

# Launch a SageMaker Training job by passing the S3 path of the training data
tabular_estimator.fit(
    {
        "train": training_dataset_s3_path,
        "validation": validation_dataset_s3_path,
    }, logs=True, job_name=training_job_name
)
```

For more information about how to set up the LightGBM as a built-in algorithm, see the following notebook examples.

- [Tabular classification with Amazon SageMaker LightGBM and CatBoost algorithm](#)
- [Tabular regression with Amazon SageMaker LightGBM and CatBoost algorithm](#)
Input and Output interface for the LightGBM algorithm

Gradient boosting operates on tabular data, with the rows representing observations, one column representing the target variable or label, and the remaining columns representing features.

The SageMaker implementation of LightGBM supports CSV for training and inference:

- For **Training ContentType**, valid inputs must be `text/csv`.
- For **Inference ContentType**, valid inputs must be `text/csv`.

### Note

For CSV training, the algorithm assumes that the target variable is in the first column and that the CSV does not have a header record.

For CSV inference, the algorithm assumes that CSV input does not have the label column.

Input format for training data, validation data, and categorical features

Be mindful of how to format your training data for input to the LightGBM model. You must provide the path to an Amazon S3 bucket that contains your training and validation data. You can also include a list of categorical features. Use both the `train` and `validation` channels to provide your input data. Alternatively, you can use only the `train` channel.

### Note

Both `train` and `training` are valid channel names for LightGBM training.

Use both the `train` and `validation` channels

You can provide your input data by way of two S3 paths, one for the `train` channel and one for the `validation` channel. Each S3 path can either be an S3 prefix that points to one or more CSV files or a full S3 path pointing to one specific CSV file. The target variables should be in the first column of your CSV file. The predictor variables (features) should be in the remaining columns. If multiple CSV files are provided for the `train` or `validation` channels, the LightGBM algorithm concatenates the files. The validation data is used to compute a validation score at the end of each boosting iteration. Early stopping is applied when the validation score stops improving.
If your predictors include categorical features, you can provide a JSON file named `categorical_index.json` in the same location as your training data file or files. If you provide a JSON file for categorical features, your `train` channel must point to an S3 prefix and not a specific CSV file. This file should contain a Python dictionary where the key is the string "cat_index_list" and the value is a list of unique integers. Each integer in the value list should indicate the column index of the corresponding categorical features in your training data CSV file. Each value should be a positive integer (greater than zero because zero represents the target value), less than the `Int32.MaxValue` (2147483647), and less than the total number of columns. There should only be one categorical index JSON file.

**Use only the train channel:**

You can alternatively provide your input data by way of a single S3 path for the `train` channel. This S3 path should point to a directory with a subdirectory named `train/` that contains one or more CSV files. You can optionally include another subdirectory in the same location called `validation/` that also has one or more CSV files. If the validation data is not provided, then 20% of your training data is randomly sampled to serve as the validation data. If your predictors include categorical features, you can provide a JSON file named `categorical_index.json` in the same location as your data subdirectories.

**Note**

For CSV training input mode, the total memory available to the algorithm (instance count multiplied by the memory available in the `InstanceType`) must be able to hold the training dataset.

SageMaker LightGBM uses the Python Joblib module to serialize or deserialize the model, which can be used for saving or loading the model.

**To use a model trained with SageMaker LightGBM with the JobLib module**

- Use the following Python code:

```python
import joblib
import tarfile

t = tarfile.open('model.tar.gz', 'r:gz')
t.extractall()
```
model = joblib.load(model_file_path)

# prediction with test data
# dtest should be a pandas DataFrame with column names feature_0, feature_1, ..., feature_d
pred = model.predict(dtest)

**Amazon EC2 instance recommendation for the LightGBM algorithm**

SageMaker LightGBM currently supports single-instance and multi-instance CPU training. For multi-instance CPU training (distributed training), specify an instance_count greater than 1 when you define your Estimator. For more information on distributed training with LightGBM, see Amazon SageMaker LightGBM Distributed training using Dask.

LightGBM is a memory-bound (as opposed to compute-bound) algorithm. So, a general-purpose compute instance (for example, M5) is a better choice than a compute-optimized instance (for example, C5). Further, we recommend that you have enough total memory in selected instances to hold the training data.

**LightGBM sample notebooks**

The following table outlines a variety of sample notebooks that address different use cases of Amazon SageMaker LightGBM algorithm.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tabular classification with Amazon SageMaker LightGBM and CatBoost algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker LightGBM algorithm to train and host a tabular classification model.</td>
</tr>
<tr>
<td>Tabular regression with Amazon SageMaker LightGBM and CatBoost algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker LightGBM algorithm to train and host a tabular regression model.</td>
</tr>
<tr>
<td>Amazon SageMaker LightGBM Distributed training using Dask</td>
<td>This notebook demonstrates distributed training with the Amazon SageMaker LightGBM algorithm using the Dask framework.</td>
</tr>
</tbody>
</table>
For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all of the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

**How LightGBM works**

LightGBM implements a conventional Gradient Boosting Decision Tree (GBDT) algorithm with the addition of two novel techniques: Gradient-based One-Side Sampling (GOSS) and Exclusive Feature Bundling (EFB). These techniques are designed to significantly improve the efficiency and scalability of GBDT.

The LightGBM algorithm performs well in machine learning competitions because of its robust handling of a variety of data types, relationships, distributions, and the diversity of hyperparameters that you can fine-tune. You can use LightGBM for regression, classification (binary and multiclass), and ranking problems.

For more information on gradient boosting, see How XGBoost Works. For in-depth details about the additional GOSS and EFB techniques used in the LightGBM method, see *LightGBM: A Highly Efficient Gradient Boosting Decision Tree*.

**LightGBM hyperparameters**

The following table contains the subset of hyperparameters that are required or most commonly used for the Amazon SageMaker LightGBM algorithm. Users set these parameters to facilitate the estimation of model parameters from data. The SageMaker LightGBM algorithm is an implementation of the open-source LightGBM package.

<table>
<thead>
<tr>
<th>Hyperparameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>

**Note**

The default hyperparameters are based on example datasets in the LightGBM sample notebooks.

By default, the SageMaker LightGBM algorithm automatically chooses an evaluation metric and objective function based on the type of classification problem. The LightGBM algorithm detects the type of classification problem based on the number of labels in your data. For regression problems, the evaluation metric is root mean squared error and the objective function is L2 loss. For binary classification problems, the evaluation metric and objective function are both binary...
cross entropy. For multiclass classification problems, the evaluation metric is multiclass cross entropy and the objective function is softmax. You can use the `metric` hyperparameter to change the default evaluation metric. Refer to the following table for more information on LightGBM hyperparameters, including descriptions, valid values, and default values.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>num_boost_round</code></td>
<td>The maximum number of boosting iterations. <strong>Note:</strong> Internally, LightGBM constructs <code>num_class * num_boost_round</code> trees for multi-class classification problems.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: Positive integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 100.</td>
</tr>
<tr>
<td><code>early_stopping_rounds</code></td>
<td>The training will stop if one metric of one validation data point does not improve in the last <code>early_stopping_rounds</code> round. If <code>early_stopping_rounds</code> is less than or equal to zero, this hyperparameter is ignored.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 10.</td>
</tr>
<tr>
<td><code>metric</code></td>
<td>The evaluation metric for validation data. If <code>metric</code> is set to the default &quot;auto&quot; value, then the algorithm automatically chooses an evaluation metric based on the type of classification problem:</td>
</tr>
<tr>
<td></td>
<td>• <code>rmse</code> for regression</td>
</tr>
<tr>
<td></td>
<td>• <code>binary_logloss</code> for binary classification</td>
</tr>
<tr>
<td></td>
<td>• <code>multi_logloss</code> for multi-class classification</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: (&quot;auto&quot;, &quot;rmse&quot;, &quot;l1&quot;, &quot;l2&quot;, &quot;huber&quot;, &quot;fair&quot;, &quot;binary_logloss&quot;, &quot;binary_error&quot;, &quot;auc&quot;, &quot;average_precision&quot;, &quot;multi_logloss&quot;, &quot;multi_error&quot;, &quot;auc_mu&quot;, or &quot;cross_entropy&quot;).</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The rate at which the model weights are updated after working through each batch of training examples.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: ((0.0, 1.0)).</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.1.</td>
</tr>
<tr>
<td>num_leaves</td>
<td>The maximum number of leaves in one tree.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: ((1, 131072)).</td>
</tr>
<tr>
<td></td>
<td>Default value: 64.</td>
</tr>
<tr>
<td>feature_fraction</td>
<td>A subset of features to be selected on each iteration (tree). Must be less than 1.0.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: ((0.0, 1.0)).</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.9.</td>
</tr>
<tr>
<td>bagging_fraction</td>
<td>A subset of features similar to feature_fraction, but bagging_fraction randomly selects part of the data without resampling.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: ((0.0, 1.0)].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.9.</td>
</tr>
<tr>
<td>bagging_freq</td>
<td>The frequency to perform bagging. At every bagging_freq iteration, LightGBM randomly selects a percentage of the data to use for the next bagging_freq iteration. This percentage is determined by the bagging_fraction hyperparameter. If bagging_freq is zero, then bagging is deactivated.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: Non-negative integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>max_depth</td>
<td>The maximum depth for a tree model. This is used to deal with overfitting when the amount of data is small. If max_depth is less than or equal to zero, this means there is no limit for maximum depth. Valid values: integer. Default value: 6.</td>
</tr>
<tr>
<td>min_data_in_leaf</td>
<td>The minimum amount of data in one leaf. Can be used to deal with overfitting. Valid values: integer, range: Non-negative integer. Default value: 3.</td>
</tr>
<tr>
<td>max_delta_step</td>
<td>Used to limit the max output of tree leaves. If max_delta_step is less than or equal to 0, then there is no constraint. The final max output of leaves is learning_rate * max_delta_step . Valid values: float. Default value: 0.0.</td>
</tr>
<tr>
<td>lambda_l1</td>
<td>L1 regularization. Valid values: float, range: Non-negative float. Default value: 0.0.</td>
</tr>
<tr>
<td>lambda_l2</td>
<td>L2 regularization. Valid values: float, range: Non-negative float. Default value: 0.0.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>boosting</td>
<td>Boosting type</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: (&quot;gbdt&quot;, &quot;rf&quot;, &quot;dart&quot;, or &quot;goss&quot;).</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;gbdt&quot;.</td>
</tr>
<tr>
<td>min_gain_to_split</td>
<td>The minimum gain to perform a split. Can be used to speed up training.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, float: Non-negative float.</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0.</td>
</tr>
<tr>
<td>scale_pos_weight</td>
<td>The weight of the labels with positive class. Used only for binary classification tasks. scale_pos_weight cannot be used if is_unbalance is set to &quot;True&quot;.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: Positive float.</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0.</td>
</tr>
<tr>
<td>tree_learner</td>
<td>Tree learner type.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: (&quot;serial&quot;, &quot;feature&quot;, &quot;data&quot;, or &quot;voting&quot;).</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;serial&quot;.</td>
</tr>
<tr>
<td>feature_fraction_bynode</td>
<td>Selects a subset of random features on each tree node. For example, if feature_fraction_bynode is 0.8, then 80% of features are selected. Can be used to deal with overfitting.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: (0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>is_unbalance</td>
<td>Set to &quot;True&quot; if training data is unbalanced. Used only for binary classification tasks. is_unbalance cannot be used with scale_pos_weight. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>max_bin</td>
<td>The maximum number of bins used to bucket feature values. A small number of bins may reduce training accuracy, but may increase general performance. Can be used to deal with overfitting. Valid values: integer, range: (1, ∞). Default value: 255.</td>
</tr>
<tr>
<td>tweedie_variance_power</td>
<td>Controls the variance of the Tweedie distribution. Set this closer to 2.0 to shift toward a gamma distribution. Set this closer to 1.0 to shift toward a Poisson distribution. Used only for regression tasks. Valid values: float, range: [1.0, 2.0). Default value: 1.5.</td>
</tr>
<tr>
<td>num_threads</td>
<td>Number of parallel threads used to run LightGBM. Value 0 means default number of threads in OpenMP. Valid values: integer, range: Non-negative integer. Default value: 0.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>verbosity</td>
<td>The verbosity of print messages. If the verbosity is less than 0, then print messages only show fatal errors. If verbosity is set to 0, then print messages include errors and warnings. If verbosity is 1, then print messages show more information. A verbosity greater than 1 shows the most information in print messages and can be used for debugging.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.</td>
</tr>
</tbody>
</table>

**Tune a LightGBM model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your training and validation datasets. Model tuning focuses on the following hyperparameters:

- A learning objective function to optimize during model training
- An evaluation metric that is used to evaluate model performance during validation
- A set of hyperparameters and a range of values for each to use when tuning the model automatically

Automatic model tuning searches your specified hyperparameters to find the combination of values that results in a model that optimizes the chosen evaluation metric.

---

### Note

The learning objective function is automatically assigned based on the type of classification task, which is determined by the number of unique integers in the label column. For more information, see [LightGBM hyperparameters](#).
Note

Automatic model tuning for LightGBM is only available from the Amazon SageMaker SDKs, not from the SageMaker console.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Evaluation metrics computed by the LightGBM algorithm

The SageMaker LightGBM algorithm computes the following metrics to use for model validation. The evaluation metric is automatically assigned based on the type of classification task, which is determined by the number of unique integers in the label column.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
<th>Regex Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>rmse</td>
<td>root mean square error</td>
<td>minimize</td>
<td>&quot;rmse: ([0-9..]+)&quot;</td>
</tr>
<tr>
<td>l1</td>
<td>mean absolute error</td>
<td>minimize</td>
<td>&quot;l1: ([0-9..]+)&quot;</td>
</tr>
<tr>
<td>l2</td>
<td>mean squared error</td>
<td>minimize</td>
<td>&quot;l2: ([0-9..]+)&quot;</td>
</tr>
<tr>
<td>huber</td>
<td>huber loss</td>
<td>minimize</td>
<td>&quot;huber: ([0-9..]+)&quot;</td>
</tr>
<tr>
<td>fair</td>
<td>fair loss</td>
<td>minimize</td>
<td>&quot;fair: ([0-9..]+)&quot;</td>
</tr>
<tr>
<td>binary_logloss</td>
<td>binary cross entropy</td>
<td>maximize</td>
<td>&quot;binary_logloss: ([0-9..]+)&quot;</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Optimization Direction</td>
<td>Regex Pattern</td>
</tr>
<tr>
<td>------------------</td>
<td>------------------------------</td>
<td>------------------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>binary_error</td>
<td>binary error</td>
<td>minimize</td>
<td>&quot;binary_error: ([0-9.]+)+&quot;</td>
</tr>
<tr>
<td>auc</td>
<td>AUC</td>
<td>maximize</td>
<td>&quot;auc: ([0-9].[]+)+&quot;</td>
</tr>
<tr>
<td>average_precision</td>
<td>average precision score</td>
<td>maximize</td>
<td>&quot;average_precision : ([0-9].[]+)+&quot;</td>
</tr>
<tr>
<td>multi_log_loss</td>
<td>multiclass cross entropy</td>
<td>maximize</td>
<td>&quot;multi_log_loss: ([0-9].[]+)+&quot;</td>
</tr>
<tr>
<td>multi_error</td>
<td>multiclass error score</td>
<td>minimize</td>
<td>&quot;multi_error: ([0-9].[]+)+&quot;</td>
</tr>
<tr>
<td>auc_mu</td>
<td>AUC-mu</td>
<td>maximize</td>
<td>&quot;auc_mu: ([0-9].[]+)+&quot;</td>
</tr>
<tr>
<td>cross_entropy</td>
<td>cross entropy</td>
<td>minimize</td>
<td>&quot;cross_entropy: ([0-9].[]+)+&quot;</td>
</tr>
</tbody>
</table>

**Tunable LightGBM hyperparameters**

Tune the LightGBM model with the following hyperparameters. The hyperparameters that have the greatest effect on optimizing the LightGBM evaluation metrics are: learning_rate,
num_leaves, feature_fraction, bagging_fraction, bagging_freq, max_depth and min_data_in_leaf. For a list of all the LightGBM hyperparameters, see LightGBM hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.001, MaxValue: 0.01</td>
</tr>
<tr>
<td>num_leaves</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 10, MaxValue: 100</td>
</tr>
<tr>
<td>feature_fraction</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 1.0</td>
</tr>
<tr>
<td>bagging_fraction</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 1.0</td>
</tr>
<tr>
<td>bagging_freq</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 0, MaxValue: 10</td>
</tr>
<tr>
<td>max_depth</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 15, MaxValue: 100</td>
</tr>
<tr>
<td>min_data_in_leaf</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 10, MaxValue: 200</td>
</tr>
</tbody>
</table>

**Linear Learner Algorithm**

*Linear models* are supervised learning algorithms used for solving either classification or regression problems. For input, you give the model labeled examples \((x, y)\). \(x\) is a high-dimensional vector and \(y\) is a numeric label. For binary classification problems, the label must be either 0 or 1. For multiclass classification problems, the labels must be from 0 to num_classes - 1. For regression problems, \(y\) is a real number. The algorithm learns a linear function, or, for classification problems, a linear threshold function, and maps a vector \(x\) to an approximation of the label \(y\).

The Amazon SageMaker linear learner algorithm provides a solution for both classification and regression problems. With the SageMaker algorithm, you can simultaneously explore different
training objectives and choose the best solution from a validation set. You can also explore a large number of models and choose the best. The best model optimizes either of the following:

- Continuous objectives, such as mean square error, cross entropy loss, absolute error.
- Discrete objectives suited for classification, such as F1 measure, precision, recall, or accuracy.

Compared with methods that provide a solution for only continuous objectives, the SageMaker linear learner algorithm provides a significant increase in speed over naive hyperparameter optimization techniques. It is also more convenient.

The linear learner algorithm requires a data matrix, with rows representing the observations, and columns representing the dimensions of the features. It also requires an additional column that contains the labels that match the data points. At a minimum, Amazon SageMaker linear learner requires you to specify input and output data locations, and objective type (classification or regression) as arguments. The feature dimension is also required. For more information, see CreateTrainingJob. You can specify additional parameters in the HyperParameters string map of the request body. These parameters control the optimization procedure, or specifics of the objective function that you train on. For example, the number of epochs, regularization, and loss type.

If you're using Managed Spot Training, the linear learner algorithm supports using checkpoints to take a snapshot of the state of the model.

**Topics**

- Input/Output interface for the linear learner algorithm
- EC2 instance recommendation for the linear learner algorithm
- Linear learner sample notebooks
- How linear learner works
- Linear learner hyperparameters
- Tune a linear learner model
- Linear learner response formats

**Input/Output interface for the linear learner algorithm**

The Amazon SageMaker linear learner algorithm supports three data channels: train, validation (optional), and test (optional). If you provide validation data, the S3DataDistributionType
should be FullyReplicated. The algorithm logs validation loss at every epoch, and uses a sample of the validation data to calibrate and select the best model. If you don’t provide validation data, the algorithm uses a sample of the training data to calibrate and select the model. If you provide test data, the algorithm logs include the test score for the final model.

**For training**, the linear learner algorithm supports both recordIO-wrapped protobuf and CSV formats. For the application/x-recordio-protobuf input type, only Float32 tensors are supported. For the text/csv input type, the first column is assumed to be the label, which is the target variable for prediction. You can use either File mode or Pipe mode to train linear learner models on data that is formatted as recordIO-wrapped-protobuf or as CSV.

**For inference**, the linear learner algorithm supports the application/json, application/x-recordio-protobuf, and text/csv formats. When you make predictions on new data, the format of the response depends on the type of model. **For regression** (predictor_type='regressor'), the score is the prediction produced by the model. **For classification** (predictor_type='binary_classifier' or predictor_type='multiclass_classifier'), the model returns a score and also a predicted_label. The predicted_label is the class predicted by the model and the score measures the strength of that prediction.

- **For binary classification**, predicted_label is 0 or 1, and score is a single floating point number that indicates how strongly the algorithm believes that the label should be 1.
- **For multiclass classification**, the predicted_class will be an integer from 0 to num_classes-1, and score will be a list of one floating point number per class.

To interpret the score in classification problems, you have to consider the loss function used. If the loss hyperparameter value is logistic for binary classification or softmax_loss for multiclass classification, then the score can be interpreted as the probability of the corresponding class. These are the loss values used by the linear learner when the loss value is auto default value. But if the loss is set to hinge_loss, then the score cannot be interpreted as a probability. This is because hinge loss corresponds to a Support Vector Classifier, which does not produce probability estimates.

For more information on input and output file formats, see [Linear learner response formats](#). For more information on inference formats, and the [Linear learner sample notebooks](#).
EC2 instance recommendation for the linear learner algorithm

The linear learner algorithm supports both CPU and GPU instances for training and inference. For GPU, the linear learner algorithm supports P2, P3, G4dn, and G5 GPU families.

During testing, we have not found substantial evidence that multi-GPU instances are faster than single-GPU instances. Results can vary, depending on your specific use case.

Linear learner sample notebooks

The following table outlines a variety of sample notebooks that address different use cases of Amazon SageMaker linear learner algorithm.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>An Introduction with the MNIST dataset</td>
<td>Using the MNIST dataset, we train a binary classifier to predict a single digit.</td>
</tr>
<tr>
<td>How to Build a Multiclass Classifier?</td>
<td>Using UCI's Covertype dataset, we demonstrate how to train a multiclass classifier.</td>
</tr>
<tr>
<td>How to Build a Machine Learning (ML) Pipeline for Inference?</td>
<td>Using a Scikit-learn container, we demonstrate how to build an end-to-end ML pipeline.</td>
</tr>
</tbody>
</table>

For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all of the SageMaker samples. The topic modeling example notebooks using the linear learning algorithm are located in the Introduction to Amazon algorithms section. To open a notebook, choose its Use tab and choose Create copy.

How linear learner works

There are three steps involved in the implementation of the linear learner algorithm: preprocess, train, and validate.

Step 1: Preprocess

Normalization, or feature scaling, is an important preprocessing step for certain loss functions that ensures the model being trained on a dataset does not become dominated by the weight of
a single feature. The Amazon SageMaker Linear Learner algorithm has a normalization option to assist with this preprocessing step. If normalization is turned on, the algorithm first goes over a small sample of the data to learn the mean value and standard deviation for each feature and for the label. Each of the features in the full dataset is then shifted to have mean of zero and scaled to have a unit standard deviation.

**Note**
For best results, ensure your data is shuffled before training. Training with unshuffled data may cause training to fail.

You can configure whether the linear learner algorithm normalizes the feature data and the labels using the `normalize_data` and `normalize_label` hyperparameters, respectively. Normalization is enabled by default for both features and labels for regression. Only the features can be normalized for binary classification and this is the default behavior.

**Step 2: Train**

With the linear learner algorithm, you train with a distributed implementation of stochastic gradient descent (SGD). You can control the optimization process by choosing the optimization algorithm. For example, you can choose to use Adam, AdaGrad, stochastic gradient descent, or other optimization algorithms. You also specify their hyperparameters, such as momentum, learning rate, and the learning rate schedule. If you aren't sure which algorithm or hyperparameter value to use, choose a default that works for the majority of datasets.

During training, you simultaneously optimize multiple models, each with slightly different objectives. For example, you vary L1 or L2 regularization and try out different optimizer settings.

**Step 3: Validate and set the threshold**

When training multiple models in parallel, the models are evaluated against a validation set to select the most optimal model once training is complete. For regression, the most optimal model is the one that achieves the best loss on the validation set. For classification, a sample of the validation set is used to calibrate the classification threshold. The most optimal model selected is the one that achieves the best binary classification selection criteria on the validation set. Examples of such criteria include the F1 measure, accuracy, and cross-entropy loss.
Note

If the algorithm is not provided a validation set, then evaluating and selecting the most optimal model is not possible. To take advantage of parallel training and model selection ensure you provide a validation set to the algorithm.

Linear learner hyperparameters

The following table contains the hyperparameters for the linear learner algorithm. These are parameters that are set by users to facilitate the estimation of model parameters from data. The required hyperparameters that must be set are listed first, in alphabetical order. The optional hyperparameters that can be set are listed next, also in alphabetical order. When a hyperparameter is set to auto, Amazon SageMaker will automatically calculate and set the value of that hyperparameter.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_classes</td>
<td>The number of classes for the response variable. The algorithm assumes that classes are labeled 0, ..., num_classes - 1. Required when predictor_type is multiclass_classifier. Otherwise, the algorithm ignores it. Valid values: Integers from 3 to 1,000,000</td>
</tr>
<tr>
<td>predictor_type</td>
<td>Specifies the type of target variable as a binary classification, multiclass classification, or regression. Required Valid values: binary_classifier, multiclass_classifier, or regressor</td>
</tr>
<tr>
<td>accuracy_top_k</td>
<td>When computing the top-k accuracy metric for multiclass classification, the value of k. If the model assigns one of the top-k scores to the true label, an example is scored as correct. Optional</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>balance_multiclass_weights</td>
<td>Specifies whether to use class weights, which give each class equal importance in the loss function. Used only when the predictor_type is multiclass_classifier. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: true, false</td>
</tr>
<tr>
<td></td>
<td>Default value: false</td>
</tr>
<tr>
<td>beta_1</td>
<td>The exponential decay rate for first-moment estimates. Applies only when the optimizer value is adam. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or floating-point value between 0 and 1.0</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>beta_2</td>
<td>The exponential decay rate for second-moment estimates. Applies only when the optimizer value is adam. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or floating-point integer between 0 and 1.0</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>bias_lr_mult</td>
<td>Allows a different learning rate for the bias term. The actual learning rate for the bias is learning_rate * bias_lr_mult. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>bias_wd_mult</td>
<td>Allows different regularization for the bias term. The actual L2 regularization weight for the bias is $wd \times bias_wd_mult$. By default, there is no regularization on the bias term.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or non-negative floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>binary_classifier_</td>
<td>When predictor_type is set to binary_classifier, the model evaluation criteria for the validation dataset (or for the training dataset if you don’t provide a validation dataset). Criteria include:</td>
</tr>
<tr>
<td>model_selection_criteria</td>
<td>• accuracy—The model with the highest accuracy.</td>
</tr>
<tr>
<td></td>
<td>• f_beta—The model with the highest F1 score. The default is F1.</td>
</tr>
<tr>
<td></td>
<td>• precision_at_target_recall —The model with the highest precision at a given recall target.</td>
</tr>
<tr>
<td></td>
<td>• recall_at_target_precision —The model with the highest recall at a given precision target.</td>
</tr>
<tr>
<td></td>
<td>• loss_function —The model with the lowest value of the loss function used in training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: accuracy, f_beta, precision_at_target_recall, recall_at_target_precision, or loss_function</td>
</tr>
<tr>
<td></td>
<td>Default value: accuracy</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>If no improvement is made in the relevant metric, the number of epochs to wait before ending training. If you have provided a value for <code>binary_classifier_model_selection_criteria</code> the metric is that value. Otherwise, the metric is the same as the value specified for the <code>loss</code> hyperparameter. The metric is evaluated on the validation data. If you haven't provided validation data, the metric is always the same as the value specified for the <code>loss</code> hyperparameter and is evaluated on the training data. To disable early stopping, set <code>early_stopping_patience</code> to a value greater than the value specified for <code>epochs</code>. <strong>Optional</strong> Valid values: Positive integer Default value: 3</td>
</tr>
<tr>
<td>early_stopping_tolerance</td>
<td>The relative tolerance to measure an improvement in loss. If the ratio of the improvement in loss divided by the previous best loss is smaller than this value, early stopping considers the improvement to be zero. <strong>Optional</strong> Valid values: Positive floating-point integer Default value: 0.001</td>
</tr>
<tr>
<td>epochs</td>
<td>The maximum number of passes over the training data. <strong>Optional</strong> Valid values: Positive integer Default value: 15</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| *f_beta*      | The value of beta to use when calculating F score metrics for binary or multiclass classification. Also used if the value specified for `binary_classifier_model_selection_criteria` is *f_beta*.  
Optional  
Valid values: Positive floating-point integers  
Default value: 1.0 |
| *feature_dim* | The number of features in the input data.  
Optional  
Valid values: auto or positive integer  
Default values: auto |
| *huber_delta* | The parameter for Huber loss. During training and metric evaluation, compute L2 loss for errors smaller than delta and L1 loss for errors larger than delta.  
Optional  
Valid values: Positive floating-point integer  
Default value: 1.0 |
| *init_bias*   | Initial weight for the bias term.  
Optional  
Valid values: Floating-point integer  
Default value: 0 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| **init_method** | Sets the initial distribution function used for model weights. Functions include:  
- *uniform*—Uniformly distributed between (-scale, +scale)  
- *normal*—Normal distribution, with mean 0 and sigma  

**Optional**  
Valid values: *uniform* or *normal*  
Default value: *uniform* |
| **init_scale** | Scales an initial uniform distribution for model weights. Applies only when the **init_method** hyperparameter is set to *uniform*.  

**Optional**  
Valid values: Positive floating-point integer  
Default value: 0.07 |
| **init_sigma** | The initial standard deviation for the normal distribution. Applies only when the **init_method** hyperparameter is set to *normal*.  

**Optional**  
Valid values: Positive floating-point integer  
Default value: 0.01 |
| **l1** | The L1 regularization parameter. If you don't want to use L1 regularization, set the value to 0.  

**Optional**  
Valid values: auto or non-negative float  
Default value: auto |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>The step size used by the optimizer for parameter updates.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: auto, whose value depends on the optimizer chosen.</td>
</tr>
<tr>
<td>loss</td>
<td>Specifies the loss function.</td>
</tr>
<tr>
<td></td>
<td>The available loss functions and their default values depend on the value of predictor_type:</td>
</tr>
<tr>
<td></td>
<td>• If the predictor_type is set to regressor, the available options are auto, squared_loss, absolute_loss, eps_insensitive_squared_loss, eps_insensitive_absolute_loss, quantile_loss, and huber_loss. The default value for auto is squared_loss.</td>
</tr>
<tr>
<td></td>
<td>• If the predictor_type is set to binary_classifier, the available options are auto, logistic, and hinge_loss. The default value for auto is logistic.</td>
</tr>
<tr>
<td></td>
<td>• If the predictor_type is set to multiclass_classifier, the available options are auto and softmax_loss. The default value for auto is softmax_loss.</td>
</tr>
<tr>
<td></td>
<td>Valid values: auto, logistic, squared_loss, absolute_loss, hinge_loss, eps_insensitive_squared_loss, eps_insensitive_absolute_loss, quantile_loss, or huber_loss</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>loss_inse</td>
<td>The parameter for the epsilon-insensitive loss type. During training and metric evaluation, any error smaller than this value is considered to be zero.</td>
</tr>
<tr>
<td>nsitivity</td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.01</td>
</tr>
<tr>
<td>lr_scheduler_factor</td>
<td>For every lr_scheduler_step hyperparameter, the learning rate decreases by this quantity. Applies only when the use_lr_scheduler hyperparameter is set to true.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive floating-point integer between 0 and 1</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>lr_scheduler_minimum_lr</td>
<td>The learning rate never decreases to a value lower than the value set for lr_scheduler_minimum_lr. Applies only when the use_lr_scheduler hyperparameter is set to true.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default values: auto</td>
</tr>
<tr>
<td>lr_scheduler_step</td>
<td>The number of steps between decreases of the learning rate. Applies only when the use_lr_scheduler hyperparameter is set to true.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>margin</td>
<td>The margin for the hinge_loss function.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>The number of observations per mini-batch for the data iterator.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1000</td>
</tr>
<tr>
<td>momentum</td>
<td>The momentum of the sgd optimizer.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or a floating-point integer between 0 and 1.0</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>normalize_data</td>
<td>Normalizes the feature data before training. Data normalization shifts the</td>
</tr>
<tr>
<td></td>
<td>data for each feature to have a mean of zero and scales it to have</td>
</tr>
<tr>
<td></td>
<td>unit standard deviation.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto, true, or false</td>
</tr>
<tr>
<td></td>
<td>Default value: true</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>normalize_label</td>
<td>Normalizes the label. Label normalization shifts the label to have a mean of zero and scales it to have unit standard deviation. The auto default value normalizes the label for regression problems but does not for classification problems. If you set the normalize_label hyperparameter to true for classification problems, the algorithm ignores it.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto, true, or false</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>num_calibration_samples</td>
<td>The number of observations from the validation dataset to use for model calibration (when finding the best threshold).</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>num_models</td>
<td>The number of models to train in parallel. For the default, auto, the algorithm decides the number of parallel models to train. One model is trained according to the given training parameter (regularization, optimizer, loss), and the rest by close parameters.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or positive integer</td>
</tr>
<tr>
<td></td>
<td>Default values: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>num_point_for_scaler</td>
<td>The number of data points to use for calculating normalization or unbiasing of terms.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10,000</td>
</tr>
<tr>
<td>optimizer</td>
<td>The optimization algorithm to use.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values:</td>
</tr>
<tr>
<td></td>
<td>• auto—The default value.</td>
</tr>
<tr>
<td></td>
<td>• sgd—Stochastic gradient descent.</td>
</tr>
<tr>
<td></td>
<td>• adam—Adaptive momentum estimation.</td>
</tr>
<tr>
<td></td>
<td>• rmsprop—A gradient-based optimization technique that uses a moving average of squared gradients to normalize the gradient.</td>
</tr>
<tr>
<td></td>
<td>Default value: auto. The default setting for auto is adam.</td>
</tr>
<tr>
<td>positive_example_weight_mult</td>
<td>The weight assigned to positive examples when training a binary classifier. The weight of negative examples is fixed at 1. If you want the algorithm to choose a weight so that errors in classifying negative vs. positive examples have equal impact on training loss, specify balanced. If you want the algorithm to choose the weight that optimizes performance, specify auto.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: balanced, auto, or a positive floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>quantile</td>
<td>The quantile for quantile loss. For quantile q, the model attempts to produce predictions so that the value of true_label is greater than the prediction with probability q.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Floating-point integer between 0 and 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.5</td>
</tr>
<tr>
<td>target_precision</td>
<td>The target precision. If binary_classifier_model_selection_criteria is recall_at_target_precision, then precision is held at this value while recall is maximized.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Floating-point integer between 0 and 1.0</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.8</td>
</tr>
<tr>
<td>target_recall</td>
<td>The target recall. If binary_classifier_model_selection_criteria is precision_at_target_recall, then recall is held at this value while precision is maximized.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Floating-point integer between 0 and 1.0</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.8</td>
</tr>
<tr>
<td>unbias_data</td>
<td>Unbiases the features before training so that the mean is 0. By default data is unbiased as the use_bias hyperparameter is set to true.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto, true, or false</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>unbias_label</td>
<td>Unbiases labels before training so that the mean is 0. Applies to regression only if the use_bias hyperparameter is set to true.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto, true, or false</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>use_bias</td>
<td>Specifies whether the model should include a bias term, which is the intercept term in the linear equation.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: true or false</td>
</tr>
<tr>
<td></td>
<td>Default value: true</td>
</tr>
<tr>
<td>use_lr_scheduler</td>
<td>Whether to use a scheduler for the learning rate. If you want to use a scheduler, specify true.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: true or false</td>
</tr>
<tr>
<td></td>
<td>Default value: true</td>
</tr>
<tr>
<td>wd</td>
<td>The weight decay parameter, also known as the L2 regularization parameter. If you don't want to use L2 regularization, set the value to 0.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or non-negative floating-point integer</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
</tbody>
</table>
Tune a linear learner model

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

The linear learner algorithm also has an internal mechanism for tuning hyperparameters separate from the automatic model tuning feature described here. By default, the linear learner algorithm tunes hyperparameters by training multiple models in parallel. When you use automatic model tuning, the linear learner internal tuning mechanism is turned off automatically. This sets the number of parallel models, `num_models`, to 1. The algorithm ignores any value that you set for `num_models`.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics computed by the linear learner algorithm**

The linear learner algorithm reports the metrics in the following table, which are computed during training. Choose one of them as the objective metric. To avoid overfitting, we recommend tuning the model against a validation metric instead of a training metric.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:absoluteloss</td>
<td>The absolute loss of the final model on the test dataset. This objective metric is only valid for regression.</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:binaryclassification_accuracy</td>
<td>The accuracy of the final model on the test dataset. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:binaryf_beta</td>
<td>The F-beta score of the final model on the test dataset. By default, it is the F1 score, which is the harmonic mean of precision and recall.</td>
<td>Maximize</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Optimization Direction</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td></td>
<td>This objective metric is only valid for binary classification.</td>
<td></td>
</tr>
<tr>
<td>test:dcg</td>
<td>The discounted cumulative gain of the final model on the test dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:macro_f_beta</td>
<td>The F-beta score of the final model on the test dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:macro_precision</td>
<td>The precision score of the final model on the test dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:macro_recall</td>
<td>The recall score of the final model on the test dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:mse</td>
<td>The mean square error of the final model on the test dataset. This objective metric is only valid for regression.</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:multiclass_accuracy</td>
<td>The accuracy of the final model on the test dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:multiclass_top_k_accuracy</td>
<td>The accuracy among the top k labels predicted on the test dataset. If you choose this metric as the objective, we recommend setting the value of k using the accuracy_top_k hyperparameter. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Optimization Direction</td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>test:objective_loss</td>
<td>The mean value of the objective loss function on the test dataset after the model is trained. By default, the loss is logistic loss for binary classification and squared loss for regression. To set the loss to other types, use the loss hyperparameter.</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:precision</td>
<td>The precision of the final model on the test dataset. If you choose this metric as the objective, we recommend setting a target recall by setting the binary_classifier_model_selection hyperparameter to precision_at_target_recall and setting the value for the target_recall hyperparameter. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:recall</td>
<td>The recall of the final model on the test dataset. If you choose this metric as the objective, we recommend setting a target precision by setting the binary_classifier_model_selection hyperparameter to recall_at_target_precision and setting the value for the target_precision hyperparameter. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:roc_auc_score</td>
<td>The area under receiving operating characteristic curve (ROC curve) of the final model on the test dataset. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Optimization Direction</td>
</tr>
<tr>
<td>-------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>validation:absolute_loss</td>
<td>The absolute loss of the final model on the validation dataset. This objective metric is only valid for regression.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:binary_classification_accuracy</td>
<td>The accuracy of the final model on the validation dataset. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:binary_f_beta</td>
<td>The F-beta score of the final model on the validation dataset. By default, the F-beta score is the F1 score, which is the harmonic mean of the validation:precision and validation:recall metrics. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:dcg</td>
<td>The discounted cumulative gain of the final model on the validation dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:macro_f_beta</td>
<td>The F-beta score of the final model on the validation dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:macro_precision</td>
<td>The precision score of the final model on the validation dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:macro_recall</td>
<td>The recall score of the final model on the validation dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:mse</td>
<td>The mean square error of the final model on the validation dataset. This objective metric is only valid for regression.</td>
<td>Minimize</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Optimization Direction</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------------------</td>
</tr>
<tr>
<td>validaton:multiclass_accuracy</td>
<td>The accuracy of the final model on the validation dataset. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validaton:multiclass_top_k_accuracy</td>
<td>The accuracy among the top k labels predicted on the validation dataset. If you choose this metric as the objective, we recommend setting the value of k using the accuracy_top_k hyperparameter. This objective metric is only valid for multiclass classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validaton:objective_loss</td>
<td>The mean value of the objective loss function on the validation dataset every epoch. By default, the loss is logistic loss for binary classification and squared loss for regression. To set loss to other types, use the loss hyperparameter.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validaton:precision</td>
<td>The precision of the final model on the validation dataset. If you choose this metric as the objective, we recommend setting a target recall by setting the binary_classifier_model_selection hyperparameter to precision_at_target_recall and setting the value for the target_recall hyperparameter. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Optimization Direction</td>
</tr>
<tr>
<td>-----------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>validation:recall</td>
<td>The recall of the final model on the validation dataset. If you choose this metric as the objective, we recommend setting a target precision by setting the <code>binary_classifier_model_selection</code> hyperparameter to <code>recall_at_target_precision</code> and setting the value for the <code>target_precision</code> hyperparameter. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:rmse</td>
<td>The root mean square error of the final model on the validation dataset. This objective metric is only valid for regression.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:roc_auc_score</td>
<td>The area under receiving operating characteristic curve (ROC curve) of the final model on the validation dataset. This objective metric is only valid for binary classification.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

**Tuning linear learner hyperparameters**

You can tune a linear learner model with the following hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>wd</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-7, MaxValue: 1</td>
</tr>
<tr>
<td>l1</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-7, MaxValue: 1</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-5, MaxValue: 1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
</tr>
<tr>
<td>----------------------</td>
<td>----------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 100, MaxValue: 5000</td>
</tr>
<tr>
<td>use_bias</td>
<td>CategoricalParameterRanges</td>
<td>[True, False]</td>
</tr>
<tr>
<td>positive_example_weight_mult</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-5, MaxValue: 1e5</td>
</tr>
</tbody>
</table>

**Linear learner response formats**

**JSON response formats**

All Amazon SageMaker built-in algorithms adhere to the common input inference format described in [Common Data Formats - Inference](#). The following are the available output formats for the SageMaker linear learner algorithm.

**Binary Classification**

```javascript
let response = {
    "predictions": [
        {
            "score": 0.4,
            "predicted_label": 0
        }
    ]
}
```

**Multiclass Classification**

```javascript
let response = {
    "predictions": [
        {
            "score": [0.1, 0.2, 0.4, 0.3],
            "predicted_label": 2
        }
    ]
}
```
Regression

```javascript
let response = {
    "predictions": [
        {
            "score": 0.4
        }
    ]
}
```

JSONLINES response formats

**Binary Classification**

```json
{"score": 0.4, "predicted_label": 0}
```

**Multiclass Classification**

```json
{"score": [0.1, 0.2, 0.4, 0.3], "predicted_label": 2}
```

**Regression**

```json
{"score": 0.4}
```

RECORDIO response formats

**Binary Classification**

```json
[
    Record = {
        features = {},
        label = {
            'score': {
                keys: [],
                values: [0.4]  # float32
            },
            'predicted_label': {
                keys: [],
                values: [0.0]  # float32
            }
        }
    }
]
```
Multiclass Classification

```json
[
    Record = {
        "features": [],
        "label": {
            "score": {
                "values": [0.1, 0.2, 0.3, 0.4]
            },
            "predicted_label": {
                "values": [3]
            }
        },
        "uid": "abc123",
        "metadata": "{created_at: '2017-06-03'}"
    }
]
```

Regression

```json
[
    Record = {
        "features": {},
        "label": {
            'score': {
                'keys': [],
                'values': [0.4]  # float32
            }
        }
    }
]
```

TabTransformer

**TabTransformer** is a novel deep tabular data modeling architecture for supervised learning. The TabTransformer architecture is built on self-attention-based Transformers. The Transformer layers transform the embeddings of categorical features into robust contextual embeddings to achieve...
higher prediction accuracy. Furthermore, the contextual embeddings learned from TabTransformer are highly robust against both missing and noisy data features, and provide better interpretability.

**How to use SageMaker TabTransformer**

You can use TabTransformer as an Amazon SageMaker built-in algorithm. The following section describes how to use TabTransformer with the SageMaker Python SDK. For information on how to use TabTransformer from the Amazon SageMaker Studio Classic UI, see [SageMaker JumpStart](https://docs.aws.amazon.com/sagemaker/latest/dg/jumpstart-overview.html).

- **Use TabTransformer as a built-in algorithm**

  Use the TabTransformer built-in algorithm to build a TabTransformer training container as shown in the following code example. You can automatically spot the TabTransformer built-in algorithm image URI using the SageMaker `image_uris.retrieve` API (or the `get_image_uri` API if using Amazon SageMaker Python SDK version 2).

  After specifying the TabTransformer image URI, you can use the TabTransformer container to construct an estimator using the SageMaker Estimator API and initiate a training job. The TabTransformer built-in algorithm runs in script mode, but the training script is provided for you and there is no need to replace it. If you have extensive experience using script mode to create a SageMaker training job, then you can incorporate your own TabTransformer training scripts.

  ```python
  from sagemaker import image_uris, model_uris, script_uris

  train_model_id, train_model_version, train_scope = "pytorch-tabtransformerclassification-model", "*", "training"
  training_instance_type = "ml.p3.2xlarge"

  # Retrieve the docker image
  train_image_uri = image_uris.retrieve(
      region=None,
      framework=None,
      model_id=train_model_id,
      model_version=train_model_version,
      image_scope=train_scope,
      instance_type=training_instance_type
  )

  # Retrieve the training script
  train_source_uri = script_uris.retrieve(
      model_id=train_model_id, model_version=train_model_version,
      script_scope=train_scope
  )
  ```
train_model_uri = model_uris.retrieve(
    model_id=train_model_id, model_version=train_model_version,
    model_scope=train_scope
)

# Sample training data is available in this bucket
training_data_bucket = f"jumpstart-cache-prod-{aws_region}"
training_data_prefix = "training-datasets/tabular_binary/

training_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}/train"
validation_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}/validation"

output_bucket = sess.default_bucket()
output_prefix = "jumpstart-example-tabular-training"

s3_output_location = f"s3://{output_bucket}/{output_prefix}/output"

from sagemaker import hyperparameters

# Retrieve the default hyperparameters for training the model
hyperparameters = hyperparameters.retrieve_default(
    model_id=train_model_id, model_version=train_model_version
)

# [Optional] Override default hyperparameters with custom values
hyperparameters["n_epochs"] = "50"
print(hyperparameters)

from sagemaker.estimator import Estimator
from sagemaker.utils import name_from_base

training_job_name = name_from_base(f"built-in-algo-{train_model_id}-training")

# Create SageMaker Estimator instance
tabular_estimator = Estimator(
    role=aws_role,
    image_uri=train_image_uri,
    source_dir=train_source_uri,
model_uri=train_model_uri,
entry_point="transfer_learning.py",
instance_count=1,
instance_type=training_instance_type,
max_run=360000,
hyperparameters=hyperparameters,
output_path=s3_output_location
)

# Launch a SageMaker Training job by passing the S3 path of the training data
tabular_estimator.fit(
    {
        "training": training_dataset_s3_path,
        "validation": validation_dataset_s3_path,
    }, logs=True, job_name=training_job_name
)

For more information about how to set up the TabTransformer as a built-in algorithm, see the following notebook examples.

- Tabular classification with Amazon SageMaker TabTransformer algorithm
- Tabular regression with Amazon SageMaker TabTransformer algorithm

**Input and Output interface for the TabTransformer algorithm**

TabTransformer operates on tabular data, with the rows representing observations, one column representing the target variable or label, and the remaining columns representing features.

The SageMaker implementation of TabTransformer supports CSV for training and inference:

- For **Training ContentType**, valid inputs must be *text/csv*.
- For **Inference ContentType**, valid inputs must be *text/csv*.

> **Note**

For CSV training, the algorithm assumes that the target variable is in the first column and that the CSV does not have a header record.

For CSV inference, the algorithm assumes that CSV input does not have the label column.
Input format for training data, validation data, and categorical features

Be mindful of how to format your training data for input to the TabTransformer model. You must provide the path to an Amazon S3 bucket that contains your training and validation data. You can also include a list of categorical features. Use both the training and validation channels to provide your input data. Alternatively, you can use only the training channel.

Use both the training and validation channels

You can provide your input data by way of two S3 paths, one for the training channel and one for the validation channel. Each S3 path can either be an S3 prefix that points to one or more CSV files or a full S3 path pointing to one specific CSV file. The target variables should be in the first column of your CSV file. The predictor variables (features) should be in the remaining columns. If multiple CSV files are provided for the training or validation channels, the TabTransformer algorithm concatenates the files. The validation data is used to compute a validation score at the end of each boosting iteration. Early stopping is applied when the validation score stops improving.

If your predictors include categorical features, you can provide a JSON file named categorical_index.json in the same location as your training data file or files. If you provide a JSON file for categorical features, your training channel must point to an S3 prefix and not a specific CSV file. This file should contain a Python dictionary where the key is the string "cat_index_list" and the value is a list of unique integers. Each integer in the value list should indicate the column index of the corresponding categorical features in your training data CSV file. Each value should be a positive integer (greater than zero because zero represents the target value), less than the Int32.MaxValue (2147483647), and less than the total number of columns. There should only be one categorical index JSON file.

Use only the training channel:

You can alternatively provide your input data by way of a single S3 path for the training channel. This S3 path should point to a directory with a subdirectory named training/ that contains one or more CSV files. You can optionally include another subdirectory in the same location called validation/ that also has one or more CSV files. If the validation data is not provided, then 20% of your training data is randomly sampled to serve as the validation data. If your predictors include categorical features, you can provide a JSON file named categorical_index.json in the same location as your data subdirectories.
Note

For CSV training input mode, the total memory available to the algorithm (instance count multiplied by the memory available in the InstanceType) must be able to hold the training dataset.

Amazon EC2 instance recommendation for the TabTransformer algorithm

SageMaker TabTransformer supports single-instance CPU and single-instance GPU training. Despite higher per-instance costs, GPUs train more quickly, making them more cost effective. To take advantage of GPU training, specify the instance type as one of the GPU instances (for example, P3). SageMaker TabTransformer currently does not support multi-GPU training.

TabTransformer sample notebooks

The following table outlines a variety of sample notebooks that address different use cases of Amazon SageMaker TabTransformer algorithm.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tabular classification with Amazon SageMaker TabTransformer algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker TabTransformer algorithm to train and host a tabular classification model.</td>
</tr>
<tr>
<td>Tabular regression with Amazon SageMaker TabTransformer algorithm</td>
<td>This notebook demonstrates the use of the Amazon SageMaker TabTransformer algorithm to train and host a tabular regression model.</td>
</tr>
</tbody>
</table>

For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all of the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

How TabTransformer works

TabTransformer is a novel deep tabular data modeling architecture for supervised learning. The TabTransformer is built upon self-attention based Transformers. The Transformer layers transform
the embeddings of categorical features into robust contextual embeddings to achieve higher prediction accuracy. Furthermore, the contextual embeddings learned from TabTransformer are highly robust against both missing and noisy data features, and provide better interpretability.

TabTransformer performs well in machine learning competitions because of its robust handling of a variety of data types, relationships, distributions, and the diversity of hyperparameters that you can fine-tune. You can use TabTransformer for regression, classification (binary and multiclass), and ranking problems.

The following diagram illustrates the TabTransformer architecture.

![TabTransformer Architecture Diagram](image)

**Figure 1: The architecture of TabTransformer.**

For more information, see *TabTransformer: Tabular Data Modeling Using Contextual Embeddings.*
TabTransformer hyperparameters

The following table contains the subset of hyperparameters that are required or most commonly used for the Amazon SageMaker TabTransformer algorithm. Users set these parameters to facilitate the estimation of model parameters from data. The SageMaker TabTransformer algorithm is an implementation of the open-source TabTransformer package.

Note

The default hyperparameters are based on example datasets in the TabTransformer sample notebooks.

The SageMaker TabTransformer algorithm automatically chooses an evaluation metric and objective function based on the type of classification problem. The TabTransformer algorithm detects the type of classification problem based on the number of labels in your data. For regression problems, the evaluation metric is r square and the objective function is mean square error. For binary classification problems, the evaluation metric and objective function are both binary cross entropy. For multiclass classification problems, the evaluation metric and objective function are both multiclass cross entropy.

Note

The TabTransformer evaluation metric and objective functions are not currently available as hyperparameters. Instead, the SageMaker TabTransformer built-in algorithm automatically detects the type of classification task (regression, binary, or multiclass) based on the number of unique integers in the label column and assigns an evaluation metric and objective function.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>n_epochs</td>
<td>Number of epochs to train the deep neural network.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: Positive integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 5.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>patience</td>
<td>The training will stop if one metric of one validation data point does not improve in the last patience round.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: (2, 60).</td>
</tr>
<tr>
<td></td>
<td>Default value: 10.</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The rate at which the model weights are updated after working through each batch of training examples.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: Positive floating point number.</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.001.</td>
</tr>
<tr>
<td>batch_size</td>
<td>The number of examples propagated through the network.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: (1, 2048).</td>
</tr>
<tr>
<td></td>
<td>Default value: 256.</td>
</tr>
<tr>
<td>input_dim</td>
<td>The dimension of embeddings to encode the categorical and/or continuous columns.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: &quot;16&quot;, &quot;32&quot;, &quot;64&quot;, &quot;128&quot;, &quot;256&quot;, or &quot;512&quot;.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;32&quot;.</td>
</tr>
<tr>
<td>n_blocks</td>
<td>The number of Transformer encoder blocks.</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer, range: (1, 12).</td>
</tr>
<tr>
<td></td>
<td>Default value: 4.</td>
</tr>
<tr>
<td>attn_dropout</td>
<td>Dropout rate applied to the Multi-Head Attention layers.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: (0, 1).</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.2.</td>
</tr>
</tbody>
</table>
### Tune a TabTransformer model

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your training and validation datasets. Model tuning focuses on the following hyperparameters:

- A learning objective function to optimize during model training
- An evaluation metric that is used to evaluate model performance during validation
- A set of hyperparameters and a range of values for each to use when tuning the model automatically

Automatic model tuning searches your chosen hyperparameters to find the combination of values that results in a model that optimizes the chosen evaluation metric.

### Parameter Name

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>mlp_dropout</td>
<td>Dropout rate applied to the FeedForward network within the encoder layers and the final MLP layers on top of Transformer encoders. Valid values: float, range: (0, 1). Default value: 0.1.</td>
</tr>
<tr>
<td>frac_shared_embed</td>
<td>The fraction of embeddings shared by all the different categories for one particular column. Valid values: float, range: (0, 1). Default value: 0.25.</td>
</tr>
</tbody>
</table>

*Note*

The learning objective function and evaluation metric are both automatically assigned based on the type of classification task, which is determined by the number of unique integers in the label column. For more information, see [TabTransformer hyperparameters](#).
Note

Automatic model tuning for TabTransformer is only available from the Amazon SageMaker SDKs, not from the SageMaker console.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Evaluation metrics computed by the TabTransformer algorithm

The SageMaker TabTransformer algorithm computes the following metrics to use for model validation. The evaluation metric is automatically assigned based on the type of classification task, which is determined by the number of unique integers in the label column.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
<th>Regex Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>r2</td>
<td>r square</td>
<td>maximize</td>
<td>&quot;metrics= {'r2': (\S +)}&quot;</td>
</tr>
<tr>
<td>f1_score</td>
<td>binary cross entropy</td>
<td>maximize</td>
<td>&quot;metrics= {'f1': (\S +)}&quot;</td>
</tr>
<tr>
<td>accuracy_score</td>
<td>multiclass cross entropy</td>
<td>maximize</td>
<td>&quot;metrics= {'accuracy': (\S +)}&quot;</td>
</tr>
</tbody>
</table>

Tunable TabTransformer hyperparameters

Tune the TabTransformer model with the following hyperparameters. The hyperparameters that have the greatest effect on optimizing the TabTransformer evaluation metrics are: learning_rate, input_dim, n_blocks, attn_dropout, mlp_dropout, and frac_shared_embed. For a list of all the TabTransformer hyperparameters, see TabTransformer hyperparameters.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.001, MaxValue: 0.01</td>
</tr>
<tr>
<td>input_dim</td>
<td>CategoricalParameterRanges</td>
<td>[16, 32, 64, 128, 256, 512]</td>
</tr>
<tr>
<td>n_blocks</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 12</td>
</tr>
<tr>
<td>attn_dropout</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.8</td>
</tr>
<tr>
<td>mlp_dropout</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.8</td>
</tr>
<tr>
<td>frac_shared_embed</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.5</td>
</tr>
</tbody>
</table>

**XGBoost Algorithm**

The XGBoost (eXtreme Gradient Boosting) is a popular and efficient open-source implementation of the gradient boosted trees algorithm. Gradient boosting is a supervised learning algorithm that attempts to accurately predict a target variable by combining an ensemble of estimates from a set of simpler and weaker models. The XGBoost algorithm performs well in machine learning competitions because of its robust handling of a variety of data types, relationships, distributions, and the variety of hyperparameters that you can fine-tune. You can use XGBoost for regression, classification (binary and multiclass), and ranking problems.

You can use the new release of the XGBoost algorithm either as a Amazon SageMaker built-in algorithm or as a framework to run training scripts in your local environments. This implementation has a smaller memory footprint, better logging, improved hyperparameter validation, and an expanded set of metrics than the original versions. It provides an XGBoost estimator that executes a training script in a managed XGBoost environment. The current release of SageMaker XGBoost is based on the original XGBoost versions 1.0, 1.2, 1.3, 1.5, and 1.7.
Supported versions

- Framework (open source) mode: 1.0-1, 1.2-1, 1.2-2, 1.3-1, 1.5-1, 1.7-1
- Algorithm mode: 1.0-1, 1.2-1, 1.2-2, 1.3-1, 1.5-1, 1.7-1

⚠️ Warning
Due to required compute capacity, version 1.7-1 of SageMaker XGBoost is not compatible with GPU instances from the P2 instance family for training or inference.

⚠️ Important
When you retrieve the SageMaker XGBoost image URI, do not use :latest or :1 for the image URI tag. You must specify one of the Supported versions to choose the SageMaker-managed XGBoost container with the native XGBoost package version that you want to use. To find the package version migrated into the SageMaker XGBoost containers, see Docker Registry Paths and Example Code, choose your AWS Region, and navigate to the XGBoost (algorithm) section.

⚠️ Warning
The XGBoost 0.90 versions are deprecated. Supports for security updates or bug fixes for XGBoost 0.90 is discontinued. It is highly recommended to upgrade the XGBoost version to one of the newer versions.

_erase_ Note
XGBoost v1.1 is not supported on SageMaker because XGBoost 1.1 has a broken capability to run prediction when the test input has fewer features than the training data in LIBSVM inputs. This capability has been restored in XGBoost v1.2. Consider using SageMaker XGBoost 1.2-2 or later.
How to Use SageMaker XGBoost

With SageMaker, you can use XGBoost as a built-in algorithm or framework. By using XGBoost as a framework, you have more flexibility and access to more advanced scenarios, such as k-fold cross-validation, because you can customize your own training scripts. The following sections describe how to use XGBoost with the SageMaker Python SDK. For information on how to use XGBoost from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

- Use XGBoost as a framework

Use XGBoost as a framework to run your customized training scripts that can incorporate additional data processing into your training jobs. In the following code example, you can find how SageMaker Python SDK provides the XGBoost API as a framework in the same way it provides other framework APIs, such as TensorFlow, MXNet, and PyTorch.

```python
import boto3
import sagemaker
from sagemaker.xgboost.estimator import XGBoost
from sagemaker.session import Session
from sagemaker.inputs import TrainingInput

# initialize hyperparameters
hyperparameters = {
    "max_depth":"5",
    "eta":"0.2",
    "gamma":"4",
    "min_child_weight":"6",
    "subsample":"0.7",
    "verbosity":"1",
    "objective":"reg:squarederror",
    "num_round":"50"
}

# set an output path where the trained model will be saved
bucket = sagemaker.Session().default_bucket()
prefix = 'DEMO-xgboost-as-a-framework'
output_path = 's3://{}/{}/{}/output'.format(bucket, prefix, 'abalone-xgb-framework')

# construct a SageMaker XGBoost estimator
# specify the entry_point to your xgboost training script
estimator = XGBoost(entry_point = "your_xgboost_abalone_script.py",
                    framework_version='1.7-1',
                    hyperparameters=hyperparameters,
                    sagemaker_session=Session(),
                    role='SAGEMAKER_ROLE',
                    output_path=output_path)
```

Use Built-in Algorithms
```
role=sagemaker.get_execution_role(),
instance_count=1,
instance_type='ml.m5.2xlarge',
output_path=output_path)

# define the data type and paths to the training and validation datasets
content_type = "libsvm"
train_input = TrainingInput("s3://{}/{}/{}".format(bucket, prefix, 'train'),
    content_type=content_type)
validation_input = TrainingInput("s3://{}/{}/{}".format(bucket, prefix, 'validation'), content_type=content_type)

# execute the XGBoost training job
estimator.fit({'train': train_input, 'validation': validation_input})
```

For an end-to-end example of using SageMaker XGBoost as a framework, see Regression with Amazon SageMaker XGBoost

- **Use XGBoost as a built-in algorithm**

Use the XGBoost built-in algorithm to build an XGBoost training container as shown in the following code example. You can automatically spot the XGBoost built-in algorithm image URI using the SageMaker image_uris.retrieve API (or the get_image_uri API if using Amazon SageMaker Python SDK version 1). If you want to ensure if the image_uris.retrieve API finds the correct URI, see Common parameters for built-in algorithms and look up xgboost from the full list of built-in algorithm image URIs and available regions.

After specifying the XGBoost image URI, you can use the XGBoost container to construct an estimator using the SageMaker Estimator API and initiate a training job. This XGBoost built-in algorithm mode does not incorporate your own XGBoost training script and runs directly on the input datasets.

⚠️ **Important**

When you retrieve the SageMaker XGBoost image URI, do not use :latest or :1 for the image URI tag. You must specify one of the Supported versions to choose the SageMaker-managed XGBoost container with the native XGBoost package version that you want to use. To find the package version migrated into the SageMaker XGBoost containers, see Docker Registry Paths and Example Code, choose your AWS Region, and navigate to the XGBoost (algorithm) section.
import sagemaker
import boto3
from sagemaker import image_uris
from sagemaker.session import Session
from sagemaker.inputs import TrainingInput

# initialize hyperparameters
hyperparameters = {
    "max_depth": "5",
    "eta": "0.2",
    "gamma": "4",
    "min_child_weight": "6",
    "subsample": "0.7",
    "objective": "reg:squarederror",
    "num_round": "50"}

# set an output path where the trained model will be saved
bucket = sagemaker.Session().default_bucket()
prefix = 'DEMO-xgboost-as-a-built-in-algo'
output_path = 's3://{}/{}/{}/output'.format(bucket, prefix, 'abalone-xgb-built-in-algo')

# this line automatically looks for the XGBoost image URI and builds an XGBoost container.
# specify the repo_version depending on your preference.
xgboost_container = sagemaker.image_uris.retrieve("xgboost", region, "1.7-1")

# construct a SageMaker estimator that calls the xgboost-container
estimator = sagemaker.estimator.Estimator(image_uri=xgboost_container,
                                      hyperparameters=hyperparameters,
                                      role=sagemaker.get_execution_role(),
                                      instance_count=1,
                                      instance_type='ml.m5.2xlarge',
                                      volume_size=5, # 5 GB
                                      output_path=output_path)

# define the data type and paths to the training and validation datasets
content_type = "libsvm"
train_input = TrainingInput("s3://{}/{}/{}/".format(bucket, prefix, 'train'),
                           content_type=content_type)
validation_input = TrainingInput("s3://{}/{}/{}/".format(bucket, prefix, 'validation'),
                               content_type=content_type)
# execute the XGBoost training job
estimator.fit({'train': train_input, 'validation': validation_input})

For more information about how to set up the XGBoost as a built-in algorithm, see the following notebook examples.

- Managed Spot Training for XGBoost
- Regression with Amazon SageMaker XGBoost (Parquet input)

Input/Output Interface for the XGBoost Algorithm

Gradient boosting operates on tabular data, with the rows representing observations, one column representing the target variable or label, and the remaining columns representing features.

The SageMaker implementation of XGBoost supports the following data formats for training and inference:

- text/libsvm (default)
- text/csv
- application/x-parquet
- application/x-recordio-protobuf

Note
There are a few considerations to be aware of regarding training and inference input:

- For increased performance, we recommend using XGBoost with File mode, in which your data from Amazon S3 is stored on the training instance volumes.
- For training with columnar input, the algorithm assumes that the target variable (label) is the first column. For inference, the algorithm assumes that the input has no label column.
- For CSV data, the input should not have a header record.
- For LIBSVM training, the algorithm assumes that subsequent columns after the label column contain the zero-based index value pairs for features. So each row has the format: : <label> <index0>:<value0> <index1>:<value1>. 

Use Built-in Algorithms
For information on instance types and distributed training, see [EC2 Instance Recommendation for the XGBoost Algorithm](#).

For CSV training input mode, the total memory available to the algorithm (Instance Count * the memory available in the InstanceType) must be able to hold the training dataset. For libsvm training input mode, it's not required, but we recommend it.

For v1.3-1 and later, SageMaker XGBoost saves the model in the XGBoost internal binary format, using `Booster.save_model`. Previous versions use the Python pickle module to serialize/deserialize the model.

**Note**

Be mindful of versions when using an SageMaker XGBoost model in open source XGBoost. Versions 1.3-1 and later use the XGBoost internal binary format while previous versions use the Python pickle module.

---

**To use a model trained with SageMaker XGBoost v1.3-1 or later in open source XGBoost**

- Use the following Python code:

  ```python
  import xgboost as xgb
  xgb_model = xgb.Booster()
  xgb_model.load_model(model_file_path)
  xgb_model.predict(dtest)
  ```

**To use a model trained with previous versions of SageMaker XGBoost in open source XGBoost**

- Use the following Python code:

  ```python
  import pickle as pkl
  import tarfile

  t = tarfile.open('model.tar.gz', 'r:gz')
  t.extractall()
  ```
To differentiate the importance of labelled data points use Instance Weight Supports

- SageMaker XGBoost allows customers to differentiate the importance of labelled data points by assigning each instance a weight value. For text/libsvm input, customers can assign weight values to data instances by attaching them after the labels. For example, label:weight idx_0:val_0 idx_1:val_1.... For text/csv input, customers need to turn on the csv_weights flag in the parameters and attach weight values in the column after labels. For example: label,weight,val_0,val_1,...).

EC2 Instance Recommendation for the XGBoost Algorithm

SageMaker XGBoost supports CPU and GPU training and inference. Instance recommendations depend on training and inference needs, as well as the version of the XGBoost algorithm. Choose one of the following options for more information:

- CPU training
- GPU training
- Distributed CPU training
- Distributed GPU training
- Inference

Training

The SageMaker XGBoost algorithm supports CPU and GPU training.

CPU training

SageMaker XGBoost 1.0-1 or earlier only trains using CPUs. It is a memory-bound (as opposed to compute-bound) algorithm. So, a general-purpose compute instance (for example, M5) is a better choice than a compute-optimized instance (for example, C4). Further, we recommend that you have enough total memory in selected instances to hold the training data. Although it supports the use
of disk space to handle data that does not fit into main memory (the out-of-core feature available with the libsvm input mode), writing cache files onto disk slows the algorithm processing time.

**GPU training**

SageMaker XGBoost version 1.2-2 or later supports GPU training. Despite higher per-instance costs, GPUs train more quickly, making them more cost effective.

SageMaker XGBoost version 1.2-2 or later supports P2, P3, G4dn, and G5 GPU instance families.

SageMaker XGBoost version 1.7-1 or later supports P3, G4dn, and G5 GPU instance families. Note that due to compute capacity requirements, version 1.7-1 or later does not support the P2 instance family.

To take advantage of GPU training, specify the instance type as one of the GPU instances (for example, P3) and set the `tree_method` hyperparameter to `gpu_hist` in your existing XGBoost script.

**Distributed training**

SageMaker XGBoost supports CPU and GPU instances for distributed training.

**Distributed CPU training**

To run CPU training on multiple instances, set the `instance_count` parameter for the estimator to a value greater than one. The input data must be divided between the total number of instances.

**Divide input data across instances**

Divide the input data using the following steps:

1. Break the input data down into smaller files. The number of files should be at least equal to the number of instances used for distributed training. Using multiple smaller files as opposed to one large file also decreases the data download time for the training job.

2. When creating your `TrainingInput`, set the distribution parameter to `ShardedByS3Key`. This parameter ensures that each instance gets approximately $1/n$ of the number of files in S3 if there are $n$ instances specified in the training job.

**Distributed GPU training**

You can use distributed training with either single-GPU or multi-GPU instances.
Distributed training with single-GPU instances

SageMaker XGBoost versions 1.2-2 through 1.3-1 only support single-GPU instance training. This means that even if you select a multi-GPU instance, only one GPU is used per instance.

If you use XGBoost versions 1.2-2 through 1.3-1, or if you do not need to use multi-GPU instances, then you must divide your input data between the total number of instances. For more information, see Divide input data across instances.

⚠️ Note

Versions 1.2-2 through 1.3-1 of SageMaker XGBoost only use one GPU per instance even if you choose a multi-GPU instance.

Distributed training with multi-GPU instances

Starting with version 1.5-1, SageMaker XGBoost offers distributed GPU training with Dask. With Dask you can utilize all GPUs when using one or more multi-GPU instances. Dask also works when using single-GPU instances.

Train with Dask using the following steps:

1. Either omit the distribution parameter in your TrainingInput or set it to FullyReplicated.
2. When defining your hyperparameters, set use_dask_gpu_training to "true".

⚠️ Important

Distributed training with Dask only supports CSV and Parquet input formats. If you use other data formats such as LIBSVM or PROTOBUF, the training job fails. For Parquet data, ensure that the column names are saved as strings. Columns that have names of other data types will fail to load.
Important

Distributed training with Dask does not support pipe mode. If pipe mode is specified, the training job fails.

There are a few considerations to be aware of when training SageMaker XGBoost with Dask. Be sure to split your data into smaller files. Dask reads each Parquet file as a partition. There is a Dask worker for every GPU, so the number of files should be greater than the total number of GPUs (instance count * number of GPUs per instance). Having a very large number of files can also degrade performance. For more information, see Dask Best Practices.

Variations in output

The specified `tree_method` hyperparameter determines the algorithm that is used for XGBoost training. The tree methods `approx`, `hist` and `gpu_hist` are all approximate methods and use sketching for quantile calculation. For more information, see Tree Methods in the XGBoost documentation. Sketching is an approximate algorithm. Therefore, you can expect variations in the model depending on factors such as the number of workers chosen for distributed training. The significance of the variation is data-dependent.

Inference

SageMaker XGBoost supports CPU and GPU instances for inference. For information about the instance types for inference, see Amazon SageMaker ML Instance Types.

XGBoost Sample Notebooks

The following table outlines a variety of sample notebooks that address different use cases of Amazon SageMaker XGBoost algorithm.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><a href="#">How to Create a Custom XGBoost container?</a></td>
<td>This notebook shows you how to build a custom XGBoost Container with Amazon SageMaker Batch Transform.</td>
</tr>
<tr>
<td>Notebook Title</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Regression with XGBoost using Parquet</td>
<td>This notebook shows you how to use the Abalone dataset in Parquet to train a XGBoost model.</td>
</tr>
<tr>
<td>How to Train and Host a Multiclass Classification Model?</td>
<td>This notebook shows how to use the MNIST dataset to train and host a multiclass classification model.</td>
</tr>
<tr>
<td>How to train a Model for Customer Churn Prediction?</td>
<td>This notebook shows you how to train a model to Predict Mobile Customer Departure in an effort to identify unhappy customers.</td>
</tr>
<tr>
<td>An Introduction to Amazon SageMaker Managed Spot infrastructure for XGBoost Training</td>
<td>This notebook shows you how to use Spot Instances for training with a XGBoost Container.</td>
</tr>
<tr>
<td>How to use Amazon SageMaker Debugger to debug XGBoost Training Jobs?</td>
<td>This notebook shows you how to use Amazon SageMaker Debugger to monitor training jobs to detect inconsistencies using built-in debugging rules.</td>
</tr>
<tr>
<td>How to use Amazon SageMaker Debugger to debug XGBoost Training Jobs in Real-Time?</td>
<td>This notebook shows you how to use the MNIST dataset and Amazon SageMaker Debugger to perform real-time analysis of XGBoost training jobs while training jobs are running.</td>
</tr>
</tbody>
</table>

For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all of the SageMaker samples. The topic modeling example notebooks using the linear learning algorithm are located in the Introduction to Amazon algorithms section. To open a notebook, choose its Use tab and choose Create copy.
How XGBoost Works

XGBoost is a popular and efficient open-source implementation of the gradient boosted trees algorithm. Gradient boosting is a supervised learning algorithm, which attempts to accurately predict a target variable by combining the estimates of a set of simpler, weaker models.

When using gradient boosting for regression, the weak learners are regression trees, and each regression tree maps an input data point to one of its leafs that contains a continuous score. XGBoost minimizes a regularized (L1 and L2) objective function that combines a convex loss function (based on the difference between the predicted and target outputs) and a penalty term for model complexity (in other words, the regression tree functions). The training proceeds iteratively, adding new trees that predict the residuals or errors of prior trees that are then combined with previous trees to make the final prediction. It’s called gradient boosting because it uses a gradient descent algorithm to minimize the loss when adding new models.

Below is a brief illustration on how gradient tree boosting works.

\[
F_m(X) = F_{m-1}(X) + \alpha_m h_m(X, r_{m-1}),
\]

where \(\alpha_i\) and \(r_i\) are the regularization parameters and residuals computed with the \(i^{th}\) tree respectively, and \(h_i\) is a function that is trained to predict residuals, \(r_i\) using \(X\) for the \(i^{th}\) tree. To compute \(\alpha_i\) we use the residuals computed, \(r_i\) and compute the following: 

\[
\arg \min \alpha \sum_{i=1}^{m} L(Y_i, F_{i-1}(X_i) + \alpha h_i(X_i, r_{i-1}))
\]

where \(L(Y, F(X))\) is a differentiable loss function.
For more detail on XGBoost, see:

- XGBoost: A Scalable Tree Boosting System
- Gradient Tree Boosting
- Introduction to Boosted Trees

**XGBoost Hyperparameters**

The following table contains the subset of hyperparameters that are required or most commonly used for the Amazon SageMaker XGBoost algorithm. These are parameters that are set by users to facilitate the estimation of model parameters from data. The required hyperparameters that must be set are listed first, in alphabetical order. The optional hyperparameters that can be set are listed next, also in alphabetical order. The SageMaker XGBoost algorithm is an implementation of the open-source DMLC XGBoost package. For details about full set of hyperparameter that can be configured for this version of XGBoost, see [XGBoost Parameters](#).

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>num_class</code></td>
<td>The number of classes. Required if <code>objective</code> is set to <code>multi:softmax</code> or <code>multi:softprob</code>. Valid values: Integer.</td>
</tr>
<tr>
<td><code>num_round</code></td>
<td>The number of rounds to run the training. Required. Valid values: Integer.</td>
</tr>
<tr>
<td><code>alpha</code></td>
<td>L1 regularization term on weights. Increasing this value makes models more conservative. Optional. Valid values: Float. Default value: 0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>base_score</td>
<td>The initial prediction score of all instances, global bias.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float.</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.5</td>
</tr>
<tr>
<td>booster</td>
<td>Which booster to use. The <code>gbtree</code> and <code>dart</code> values use a tree-based model, while <code>gblinear</code> uses a linear function.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of &quot;gbtree&quot;, &quot;gblinear&quot;, or &quot;dart&quot;.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;gbtree&quot;</td>
</tr>
<tr>
<td>colsample_bylevel</td>
<td>Subsample ratio of columns for each split, in each level.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>colsample_bynode</td>
<td>Subsample ratio of columns from each node.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: (0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>colsample_bytree</td>
<td>Subsample ratio of columns when constructing each tree.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>csv_weights</td>
<td>When this flag is enabled, XGBoost differentiates the importance of instances for csv input by taking the second column (the column after labels) in training data as the instance weights.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 or 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>deterministic_histogram</td>
<td>When this flag is enabled, XGBoost builds histogram on GPU deterministically. Used only if tree_method is set to gpu_hist.</td>
</tr>
<tr>
<td></td>
<td>For a full list of valid inputs, please refer to XGBoost Parameters.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Range: &quot;true&quot; or &quot;false&quot;.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;true&quot;</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping_rounds</td>
<td>The model trains until the validation score stops improving. Validation error needs to decrease at least every <code>early_stopping_rounds</code> to continue training. SageMaker hosting uses the best model for inference.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: -</td>
</tr>
<tr>
<td>eta</td>
<td>Step size shrinkage used in updates to prevent overfitting. After each boosting step, you can directly get the weights of new features. The <code>eta</code> parameter actually shrinks the feature weights to make the boosting process more conservative.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.3</td>
</tr>
<tr>
<td>eval_metric</td>
<td>Evaluation metrics for validation data. A default metric is assigned according to the objective:</td>
</tr>
<tr>
<td></td>
<td>• <code>rmse</code>: for regression</td>
</tr>
<tr>
<td></td>
<td>• <code>error</code>: for classification</td>
</tr>
<tr>
<td></td>
<td>• <code>map</code>: for ranking</td>
</tr>
<tr>
<td></td>
<td>For a list of valid inputs, see <a href="https://docs.aws.amazon.com/sagemaker/latest/dg/algorithm-parameters-list.html#xgboost">XGBoost Learning Task Parameters</a>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String.</td>
</tr>
<tr>
<td></td>
<td>Default value: Default according to objective.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>gamma</td>
<td>Minimum loss reduction required to make a further partition on a leaf node of the tree. The larger, the more conservative the algorithm is.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: $[0, \infty)$.</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>grow_policy</td>
<td>Controls the way that new nodes are added to the tree. Currently supported only if <code>tree_method</code> is set to <code>hist</code>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either &quot;depthwise&quot; or &quot;lossguide&quot;.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;depthwise&quot;</td>
</tr>
<tr>
<td>interaction_constraints</td>
<td>Specify groups of variables that are allowed to interact.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Nested list of integers. Each integer represents a feature, and each nested list contains features that are allowed to interact e.g., [[1,2], [3,4,5]].</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>lambda</td>
<td>L2 regularization term on weights. Increasing this value makes models more conservative.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float.</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>lambda_bias</td>
<td>L2 regularization term on bias. Optional Valid values: Float. Range: [0.0, 1.0]. Default value: 0</td>
</tr>
<tr>
<td>max_bin</td>
<td>Maximum number of discrete bins to bucket continuous features. Used only if tree_method is set to hist. Optional Valid values: Integer. Default value: 256</td>
</tr>
<tr>
<td>max_delta_step</td>
<td>Maximum delta step allowed for each tree's weight estimation. When a positive integer is used, it helps make the update more conservative. The preferred option is to use it in logistic regression. Set it to 1-10 to help control the update. Optional Valid values: Integer. Range: [0,∞). Default value: 0</td>
</tr>
<tr>
<td>max_depth</td>
<td>Maximum depth of a tree. Increasing this value makes the model more complex and likely to be overfit. 0 indicates no limit. A limit is required when grow_policy =depth-wise. Optional Valid values: Integer. Range: [0,∞) Default value: 6</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>max_leaves</td>
<td>Maximum number of nodes to be added. Relevant only if grow_policy is set to lossguide.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>min_child_weight</td>
<td>Minimum sum of instance weight (hessian) needed in a child. If the tree partition step results in a leaf node with the sum of instance weight less than min_child_weight, the building process gives up further partitioning. In linear regression models, this simply corresponds to a minimum number of instances needed in each node. The larger the algorithm, the more conservative it is.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,∞).</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>monotone_constraints</td>
<td>Specifies monotonicity constraints on any feature.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Tuple of Integers. Valid integers: -1 (decreasing constraint), 0 (no constraint), 1 (increasing constraint).</td>
</tr>
<tr>
<td></td>
<td>E.g., (0, 1): No constraint on first predictor, and an increasing constraint on the second. (-1, 1): Decreasing constraint on first predictor, and an increasing constraint on the second.</td>
</tr>
<tr>
<td></td>
<td>Default value: (0, 0)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>normalize_type</td>
<td>Type of normalization algorithm.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either <em>tree</em> or <em>forest</em>.</td>
</tr>
<tr>
<td></td>
<td>Default value: <em>tree</em></td>
</tr>
<tr>
<td>nthread</td>
<td>Number of parallel threads used to run <em>xgboost</em>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer.</td>
</tr>
<tr>
<td></td>
<td>Default value: Maximum number of threads.</td>
</tr>
<tr>
<td>objective</td>
<td>Specifies the learning task and the corresponding learning objective. Examples: <em>reg:logistic</em>, <em>multi:softmax</em>, <em>reg:squarederror</em>. For a full list of valid inputs, refer to <a href="#">XGBoost Learning Task Parameters</a>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;*reg:squarederror&quot;</td>
</tr>
<tr>
<td>one_drop</td>
<td>When this flag is enabled, at least one tree is always dropped during the dropout.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 or 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>process_type</td>
<td>The type of boosting process to run.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either &quot;default&quot; or &quot;update&quot;.</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;default&quot;</td>
</tr>
<tr>
<td>rate_drop</td>
<td>The dropout rate that specifies the fraction of previous trees to drop during the dropout.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0</td>
</tr>
<tr>
<td>refresh_leaf</td>
<td>This is a parameter of the 'refresh' updater plug-in. When set to true (1), tree leaves and tree node stats are updated. When set to false (0), only tree node stats are updated.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0/1</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>sample_type</td>
<td>Type of sampling algorithm.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either uniform or weighted.</td>
</tr>
<tr>
<td></td>
<td>Default value: uniform</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>scale_pos_weight</td>
<td>Controls the balance of positive and negative weights. It's useful for unbalanced classes. A typical value to consider: ( \frac{\text{sum(negative cases)}}{\text{sum(positive cases)}} ). Optional Valid values: float Default value: 1</td>
</tr>
<tr>
<td>seed</td>
<td>Random number seed.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>single_precision_histogram</td>
<td>When this flag is enabled, XGBoost uses single precision to build histograms instead of double precision. Used only if tree_method is set to hist or gpu_hist. For a full list of valid inputs, please refer to XGBoost Parameters. Optional Valid values: String. Range: &quot;true&quot; or &quot;false&quot; Default value: &quot;false&quot;</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>sketch_eps</td>
<td>Used only for approximate greedy algorithm. This translates into $O\left(\frac{1}{\text{sketch}_\text{eps}}\right)$ number of bins. Compared to directly select number of bins, this comes with theoretical guarantee with sketch accuracy.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float, Range: [0, 1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.03</td>
</tr>
<tr>
<td>skip_drop</td>
<td>Probability of skipping the dropout procedure during a boosting iteration.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0</td>
</tr>
<tr>
<td>subsample</td>
<td>Subsample ratio of the training instance. Setting it to 0.5 means that XGBoost randomly collects half of the data instances to grow trees. This prevents overfitting.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>tree_method</td>
<td>The tree construction algorithm used in XGBoost.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: One of auto, exact, approx, hist, or gpu_hist.</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>tweedie_variance_power</strong></td>
<td>Parameter that controls the variance of the Tweedie distribution.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong>  <strong>Valid values:</strong> Float. Range: (1, 2). <strong>Default value:</strong> 1.5</td>
</tr>
<tr>
<td><strong>updater</strong></td>
<td>A comma-separated string that defines the sequence of tree updaters to run. This provides a modular way to construct and to modify the trees.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong>  <strong>Valid values:</strong> comma-separated string. <strong>Default value:</strong> grow_colmaker, prune</td>
</tr>
<tr>
<td><strong>use_dask_gpu_training</strong></td>
<td>Set use_dask_gpu_training to &quot;true&quot; if you want to run distributed GPU training with Dask. Dask GPU training is only supported for versions 1.5-1 and later. Do not set this value to &quot;true&quot; for versions preceding 1.5-1. For more information, see <a href="#">Distributed GPU training</a>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong>  <strong>Valid values:</strong> String. Range: &quot;true&quot; or &quot;false&quot; <strong>Default value:</strong> &quot;false&quot;</td>
</tr>
</tbody>
</table>
### Parameter Name | Description
--- | ---
verbosity | Verbosity of printing messages.

Valid values: 0 (silent), 1 (warning), 2 (info), 3 (debug).

**Optional**

Default value: 1

---

**Tune an XGBoost Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your training and validation datasets. You choose three types of hyperparameters:

- a learning objective function to optimize during model training
- an `eval_metric` to use to evaluate model performance during validation
- a set of hyperparameters and a range of values for each to use when tuning the model automatically

You choose the evaluation metric from set of evaluation metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the evaluation metric.

**Note**

Automatic model tuning for XGBoost 0.90 is only available from the Amazon SageMaker SDKs, not from the SageMaker console.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Evaluation Metrics Computed by the XGBoost Algorithm**

The XGBoost algorithm computes the following metrics to use for model validation. When tuning the model, choose one of these metrics to evaluate the model. For full list of valid `eval_metric` values, refer to [XGBoost Learning Task Parameters](#).
<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:accuracy</td>
<td>Classification rate, calculated as #(right)/#(all cases).</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:auc</td>
<td>Area under the curve.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:error</td>
<td>Binary classification error rate, calculated as #(wrong cases)/#(all cases).</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:f1</td>
<td>Indicator of classification accuracy, calculated as the harmonic mean of precision and recall.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:logloss</td>
<td>Negative log-likelihood.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:mae</td>
<td>Mean absolute error.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:map</td>
<td>Mean average precision.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:meanerror</td>
<td>Multiclass classification error rate, calculated as #(wrong cases)/#(all cases).</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:mlogloss</td>
<td>Negative log-likelihood for multiclass classification.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:mse</td>
<td>Mean squared error.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:ndcg</td>
<td>Normalized Discounted Cumulative Gain.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:rmse</td>
<td>Root mean square error.</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

**Tunable XGBoost Hyperparameters**

Tune the XGBoost model with the following hyperparameters. The hyperparameters that have the greatest effect on optimizing the XGBoost evaluation metrics are: alpha, min_child_weight, subsample, eta, and num_round.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>alpha</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 1000</td>
</tr>
<tr>
<td>colsample _bylevel</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 1</td>
</tr>
<tr>
<td>colsample _bynode</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 1</td>
</tr>
<tr>
<td>colsample _bytree</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.5, MaxValue: 1</td>
</tr>
<tr>
<td>eta</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 0.5</td>
</tr>
<tr>
<td>gamma</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 5</td>
</tr>
<tr>
<td>lambda</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 1000</td>
</tr>
<tr>
<td>max_delta_step</td>
<td>IntegerParameterRanges</td>
<td>[0, 10]</td>
</tr>
<tr>
<td>max_depth</td>
<td>IntegerParameterRanges</td>
<td>[0, 10]</td>
</tr>
<tr>
<td>min_child _weight</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 120</td>
</tr>
<tr>
<td>num_round</td>
<td>IntegerParameterRanges</td>
<td>[1, 4000]</td>
</tr>
<tr>
<td>subsample</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.5, MaxValue: 1</td>
</tr>
</tbody>
</table>
Deprecated Versions of XGBoost and their Upgrades

This topic contains documentation for previous versions of Amazon SageMaker XGBoost that are still available but deprecated. It also provides instructions on how to upgrade deprecated versions of XGBoost, when possible, to more current versions.

Topics

- Upgrade XGBoost Version 0.90 to Version 1.5
- XGBoost Version 0.72

Upgrade XGBoost Version 0.90 to Version 1.5

If you are using the SageMaker Python SDK, to upgrade existing XGBoost 0.90 jobs to version 1.5, you must have version 2.x of the SDK installed and change the XGBoost version and framework_version parameters to 1.5-1. If you are using Boto3, you need to update the Docker image, and a few hyperparameters and learning objectives.

Topics

- Upgrade SageMaker Python SDK Version 1.x to Version 2.x
- Change the image tag to 1.5-1
- Change Docker Image for Boto3
- Update Hyperparameters and Learning Objectives

Upgrade SageMaker Python SDK Version 1.x to Version 2.x

If you are still using Version 1.x of the SageMaker Python SDK, you must to upgrade version 2.x of the SageMaker Python SDK. For information on the latest version of the SageMaker Python SDK, see Use Version 2.x of the SageMaker Python SDK. To install the latest version, run:

```
python -m pip install --upgrade sagemaker
```

Change the image tag to 1.5-1

If you are using the SageMaker Python SDK and using the XGBoost build-in algorithm, change the version parameter in image_uris.retrieve.

```
from sagemaker import image_uris
```
image_uris.retrieve(framework="xgboost", region="us-west-2", version="1.5-1")

estimator = sagemaker.estimator.Estimator(image_uri=xgboost_container,
    hyperparameters=hyperparameters,
    role=sagemaker.get_execution_role(),
    instance_count=1,
    instance_type='ml.m5.2xlarge',
    volume_size=5, # 5 GB
    output_path=output_path)

If you are using the SageMaker Python SDK and using XGBoost as a framework to run your customized training scripts, change the framework_version parameter in the XGBoost API.

estimator = XGBoost(entry_point = "your_xgboost_abalone_script.py",
    framework_version='1.5-1',
    hyperparameters=hyperparameters,
    role=sagemaker.get_execution_role(),
    instance_count=1,
    instance_type='ml.m5.2xlarge',
    output_path=output_path)

sagemaker.session.s3_input in SageMaker Python SDK version 1.x has been renamed to sagemaker.inputs.TrainingInput. You must use sagemaker.inputs.TrainingInput as in the following example.

content_type = "libsvm"
train_input = TrainingInput("s3://{}/{}/{}/".format(bucket, prefix, 'train'),
    content_type=content_type)
validation_input = TrainingInput("s3://{}/{}/{}/".format(bucket, prefix, 'validation'),
    content_type=content_type)

For the full list of SageMaker Python SDK version 2.x changes, see Use Version 2.x of the SageMaker Python SDK.

**Change Docker Image for Boto3**

If you are using Boto3 to train or deploy your model, change the docker image tag (1, 0.72, 0.90-1 or 0.90-2) to 1.5-1.

```json
{
    "AlgorithmSpecification": {
```
"TrainingImage": "746614075791.dkr.ecr.us-west-1.amazonaws.com/sagemaker-xgboost:1.5-1"
}
...
}

If you using the SageMaker Python SDK to retrieve registry path, change the version parameter in image_uris.retrieve.

def image_uris.retrieve(framework="xgboost", region="us-west-2", version="1.5-1")

Update Hyperparameters and Learning Objectives

The silent parameter has been deprecated and is no longer available in XGBoost 1.5 and later versions. Use verbosity instead. If you were using the reg:linear learning objective, it has been deprecated as well in favor of reg:squarederror. Use reg:squarederror instead.

```python
hyperparameters = {
    "verbosity": "2",
    "objective": "reg:squarederror",
    "num_round": "50",
    ...
}

estimator = sagemaker.estimator.Estimator(image_uri=xgboost_container,
                                            hyperparameters=hyperparameters,
                                            ...
)
```

XGBoost Version 0.72

⚠️ Important

The XGBoost 0.72 is deprecated by Amazon SageMaker. You can still use this old version of XGBoost (as a built-in algorithm) by pulling its image URI as shown in the following code sample. For XGBoost, the image URI ending with :1 is for the old version.

SageMaker Python SDK v1

```python
import boto3
from sagemaker.amazon.amazon_estimator import get_image_uri
```
xgb_image_uri = get_image_uri(boto3.Session().region_name, "xgboost", repo_version="1")

SageMaker Python SDK v2

import boto3
from sagemaker import image_uris

xgb_image_uri = image_uris.retrieve("xgboost", boto3.Session().region_name, "1")

If you want to use newer versions, you have to explicitly specify the image URI tags (see Supported versions).

This previous release of the Amazon SageMaker XGBoost algorithm is based on the 0.72 release. **XGBoost** (eXtreme Gradient Boosting) is a popular and efficient open-source implementation of the gradient boosted trees algorithm. Gradient boosting is a supervised learning algorithm that attempts to accurately predict a target variable by combining the estimates of a set of simpler, weaker models. XGBoost has done remarkably well in machine learning competitions because it robustly handles a variety of data types, relationships, and distributions, and because of the large number of hyperparameters that can be tweaked and tuned for improved fits. This flexibility makes XGBoost a solid choice for problems in regression, classification (binary and multiclass), and ranking.

Customers should consider using the new release of XGBoost Algorithm. They can use it as a SageMaker built-in algorithm or as a framework to run scripts in their local environments as they would typically, for example, do with a Tensorflow deep learning framework. The new implementation has a smaller memory footprint, better logging, improved hyperparameter validation, and an expanded set of metrics. The earlier implementation of XGBoost remains available to customers if they need to postpone migrating to the new version. But this previous implementation will remain tied to the 0.72 release of XGBoost.

**Input/Output Interface for the XGBoost Release 0.72**

Gradient boosting operates on tabular data, with the rows representing observations, one column representing the target variable or label, and the remaining columns representing features.
The SageMaker implementation of XGBoost supports CSV and libsvm formats for training and inference:

- For Training ContentType, valid inputs are text/libsvm (default) or text/csv.
- For Inference ContentType, valid inputs are text/libsvm or (the default) text/csv.

**Note**

For CSV training, the algorithm assumes that the target variable is in the first column and that the CSV does not have a header record. For CSV inference, the algorithm assumes that CSV input does not have the label column.

For libsvm training, the algorithm assumes that the label is in the first column. Subsequent columns contain the zero-based index value pairs for features. So each row has the format: `<label> <index0>:<value0> <index1>:<value1> ...` Inference requests for libsvm may or may not have labels in the libsvm format.

This differs from other SageMaker algorithms, which use the protobuf training input format to maintain greater consistency with standard XGBoost data formats.

For CSV training input mode, the total memory available to the algorithm (Instance Count * the memory available in the InstanceType) must be able to hold the training dataset. For libsvm training input mode, it's not required, but we recommend it.

SageMaker XGBoost uses the Python pickle module to serialize/deserialize the model, which can be used for saving/loading the model.

**To use a model trained with SageMaker XGBoost in open source XGBoost**

- Use the following Python code:

```python
import pickle as pkl
import tarfile
import xgboost

t = tarfile.open('model.tar.gz', 'r:gz')
t.extractall()
model = pkl.load(open(model_file_path, 'rb'))
```
# prediction with test data

```python
pred = model.predict(dtest)
```

To differentiate the importance of labelled data points use Instance Weight Supports

- SageMaker XGBoost allows customers to differentiate the importance of labelled data points by assigning each instance a weight value. For `text/libsvm` input, customers can assign weight values to data instances by attaching them after the labels. For example, `label:weight idx_0:val_0 idx_1:val_1...`. For `text/csv` input, customers need to turn on the `csv_weights` flag in the parameters and attach weight values in the column after labels. For example: `label,weight,val_0,val_1,...`.

## EC2 Instance Recommendation for the XGBoost Release 0.72

SageMaker XGBoost currently only trains using CPUs. It is a memory-bound (as opposed to compute-bound) algorithm. So, a general-purpose compute instance (for example, M4) is a better choice than a compute-optimized instance (for example, C4). Further, we recommend that you have enough total memory in selected instances to hold the training data. Although it supports the use of disk space to handle data that does not fit into main memory (the out-of-core feature available with the `libsvm` input mode), writing cache files onto disk slows the algorithm processing time.

## XGBoost Release 0.72 Sample Notebooks

For a sample notebook that shows how to use the latest version of SageMaker XGBoost as a built-in algorithm to train and host a regression model, see [Regression with Amazon SageMaker XGBoost algorithm](https://docs.aws.amazon.com/sagemaker/latest/dg/xgboost regression.html). To use the 0.72 version of XGBoost, you need to change the version in the sample code to 0.72. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](https://docs.aws.amazon.com/sagemaker/latest/dg/notebook-instances.html). Once you have created a notebook instance and opened it, select the [SageMaker Examples tab](https://docs.aws.amazon.com/sagemaker/latest/dg/sg-examples-quickstart.html) to see a list of all the SageMaker samples. The topic modeling example notebooks using the XGBoost algorithms are located in the [Introduction to Amazon algorithms section](https://docs.aws.amazon.com/sagemaker/latest/dg/sg-examples-topic-modeling.html). To open a notebook, click on its `Use` tab and select `Create copy`.

## XGBoost Release 0.72 Hyperparameters

The following table contains the hyperparameters for the XGBoost algorithm. These are parameters that are set by users to facilitate the estimation of model parameters from data. The
required hyperparameters that must be set are listed first, in alphabetical order. The optional hyperparameters that can be set are listed next, also in alphabetical order. The SageMaker XGBoost algorithm is an implementation of the open-source XGBoost package. Currently SageMaker supports version 0.72. For more detail about hyperparameter configuration for this version of XGBoost, see [XGBoost Parameters](#).

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>num_class</strong></td>
<td>The number of classes. <strong>Required</strong> if <code>objective</code> is set to <code>multi:softmax</code> or <code>multi:softprob</code>. Valid values: integer</td>
</tr>
<tr>
<td><strong>num_round</strong></td>
<td>The number of rounds to run the training. <strong>Required</strong> Valid values: integer</td>
</tr>
<tr>
<td><strong>alpha</strong></td>
<td>L1 regularization term on weights. Increasing this value makes models more conservative. <strong>Optional</strong> Valid values: float Default value: 0</td>
</tr>
<tr>
<td><strong>base_score</strong></td>
<td>The initial prediction score of all instances, global bias. <strong>Optional</strong> Valid values: float Default value: 0.5</td>
</tr>
<tr>
<td><strong>booster</strong></td>
<td>Which booster to use. The <code>gbtree</code> and <code>dart</code> values use a tree-based model, while <code>gblinear</code> uses a linear function.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Valid values:</strong> String. One of <strong>gbtree</strong>, <strong>gblinear</strong>, or <strong>dart</strong>.</td>
<td></td>
</tr>
<tr>
<td><strong>Default value:</strong> <strong>gbtree</strong></td>
<td></td>
</tr>
<tr>
<td><strong>colsample_bylevel</strong></td>
<td>Subsample ratio of columns for each split, in each level.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Valid values:</strong> Float. Range: [0,1].</td>
<td></td>
</tr>
<tr>
<td><strong>Default value:</strong> 1</td>
<td></td>
</tr>
<tr>
<td><strong>colsample_bytree</strong></td>
<td>Subsample ratio of columns when constructing each tree.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Valid values:</strong> Float. Range: [0,1].</td>
<td></td>
</tr>
<tr>
<td><strong>Default value:</strong> 1</td>
<td></td>
</tr>
<tr>
<td><strong>csv_weights</strong></td>
<td>When this flag is enabled, XGBoost differentiates the importance of instances for csv input by taking the second column (the column after labels) in training data as the instance weights.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Valid values:</strong> 0 or 1</td>
<td></td>
</tr>
<tr>
<td><strong>Default value:</strong> 0</td>
<td></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping_rounds</td>
<td>The model trains until the validation score stops improving. Validation error needs to decrease at least every <code>early_stopping_rounds</code> to continue training. SageMaker hosting uses the best model for inference.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: -</td>
</tr>
<tr>
<td>eta</td>
<td>Step size shrinkage used in updates to prevent overfitting. After each boosting step, you can directly get the weights of new features. The <code>eta</code> parameter actually shrinks the feature weights to make the boosting process more conservative.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.3</td>
</tr>
<tr>
<td>eval_metric</td>
<td>Evaluation metrics for validation data. A default metric is assigned according to the objective:</td>
</tr>
<tr>
<td></td>
<td>• <code>rmse</code>: for regression</td>
</tr>
<tr>
<td></td>
<td>• <code>error</code>: for classification</td>
</tr>
<tr>
<td></td>
<td>• <code>map</code>: for ranking</td>
</tr>
<tr>
<td></td>
<td>For a list of valid inputs, see <a href="https://github.com/dmlc/xgboost/blob/master/doc/parameter.rst">XGBoost Parameters</a>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: string</td>
</tr>
<tr>
<td></td>
<td>Default value: Default according to objective.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>gamma</td>
<td>Minimum loss reduction required to make a further partition on a leaf node of the tree. The larger, the more conservative the algorithm is.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0,∞).</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>grow_policy</td>
<td>Controls the way that new nodes are added to the tree. Currently supported only if tree_method is set to hist.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either depthwise or lossguide.</td>
</tr>
<tr>
<td></td>
<td>Default value: depthwise</td>
</tr>
<tr>
<td>lambda</td>
<td>L2 regularization term on weights. Increasing this value makes models more conservative.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>lambda_bias</td>
<td>L2 regularization term on bias.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>max_bin</td>
<td>Maximum number of discrete bins to bucket continuous features. Used only if <code>tree_method</code> is set to <code>hist</code>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 256</td>
</tr>
<tr>
<td>max_delta_step</td>
<td>Maximum delta step allowed for each tree's weight estimation. When a positive integer is used, it helps make the update more conservative. The preferred option is to use it in logistic regression. Set it to 1-10 to help control the update.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer. Range: [0,∞).</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>max_depth</td>
<td>Maximum depth of a tree. Increasing this value makes the model more complex and likely to be overfit. 0 indicates no limit. A limit is required when <code>grow_policy</code> = <code>depth-wise</code> .</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer. Range: [0,∞)</td>
</tr>
<tr>
<td></td>
<td>Default value: 6</td>
</tr>
<tr>
<td>max_leaves</td>
<td>Maximum number of nodes to be added. Relevant only if <code>grow_policy</code> is set to <code>lossguide</code> .</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>min_child_weight</td>
<td>Minimum sum of instance weight (hessian) needed in a child. If the tree partition step results in a leaf node with the sum of instance weight less than min_child_weight, the building process gives up further partitioning. In linear regression models, this simply corresponds to a minimum number of instances needed in each node. The larger the algorithm, the more conservative it is.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: ([0, \infty)).</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>normalize_type</td>
<td>Type of normalization algorithm.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either tree or forest.</td>
</tr>
<tr>
<td></td>
<td>Default value: tree</td>
</tr>
<tr>
<td>nthread</td>
<td>Number of parallel threads used to run xgboost.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: Maximum number of threads.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
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</tr>
<tr>
<td>objective</td>
<td>Specifies the learning task and the corresponding learning objective. Examples: <code>reg:logistic</code>, <code>reg:softmax</code>, <code>multi:squarederror</code>. For a full list of valid inputs, refer to <a href="#">XGBoost Parameters</a>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: string</td>
</tr>
<tr>
<td></td>
<td>Default value: <code>reg:squarederror</code></td>
</tr>
<tr>
<td>one_drop</td>
<td>When this flag is enabled, at least one tree is always dropped during the dropout.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 or 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>process_type</td>
<td>The type of boosting process to run.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either <code>default</code> or <code>update</code>.</td>
</tr>
<tr>
<td></td>
<td>Default value: <code>default</code></td>
</tr>
<tr>
<td>rate_drop</td>
<td>The dropout rate that specifies the fraction of previous trees to drop during the dropout.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: <code>[0.0, 1.0]</code>.</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>refresh_leaf</td>
<td>This is a parameter of the 'refresh' updater plug-in. When set to true (1), tree leaves and tree node stats are updated. When set to false(0), only tree node stats are updated.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0/1</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>sample_type</td>
<td>Type of sampling algorithm.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either uniform or weighted.</td>
</tr>
<tr>
<td></td>
<td>Default value: uniform</td>
</tr>
<tr>
<td>scale_pos_weight</td>
<td>Controls the balance of positive and negative weights. It's useful for unbalanced classes. A typical value to consider: sum(negative cases) / sum(positive cases).</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>seed</td>
<td>Random number seed.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td><strong>Parameter Name</strong></td>
<td><strong>Description</strong></td>
</tr>
<tr>
<td>--------------------</td>
<td>-----------------</td>
</tr>
</tbody>
</table>
| silent             | 0 means print running messages, 1 means silent mode.  
Valid values: 0 or 1  
**Optional**  
Default value: 0 |
| sketch_eps         | Used only for approximate greedy algorithm. This translates into \( O(1 / \text{sketch\_eps}) \) number of bins. Compared to directly select number of bins, this comes with theoretical guarantee with sketch accuracy.  
**Optional**  
Valid values: Float, Range: \([0, 1]\).  
Default value: 0.03 |
| skip_drop          | Probability of skipping the dropout procedure during a boosting iteration.  
**Optional**  
Valid values: Float. Range: \([0.0, 1.0]\).  
Default value: 0.0 |
| subsample          | Subsample ratio of the training instance. Setting it to 0.5 means that XGBoost randomly collects half of the data instances to grow trees. This prevents overfitting.  
**Optional**  
Valid values: Float. Range: \([0,1]\).  
Default value: 1 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>tree_method</td>
<td>The tree construction algorithm used in XGBoost.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: One of auto, exact, approx, or hist.</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>tweedie_variance_power</td>
<td>Parameter that controls the variance of the Tweedie distribution.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range: (1, 2).</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.5</td>
</tr>
<tr>
<td>updater</td>
<td>A comma-separated string that defines the sequence of tree updaters to run.</td>
</tr>
<tr>
<td></td>
<td>This provides a modular way to construct and to modify the trees.</td>
</tr>
<tr>
<td></td>
<td>For a full list of valid inputs, please refer to <a href="#">XGBoost Parameter</a>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: comma-separated string.</td>
</tr>
<tr>
<td></td>
<td>Default value: grow_colmaker, prune</td>
</tr>
</tbody>
</table>

**Tune an XGBoost Release 0.72 Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your training and validation datasets. You choose three types of hyperparameters:

- a learning objective function to optimize during model training
- an `eval_metric` to use to evaluate model performance during validation
• a set of hyperparameters and a range of values for each to use when tuning the model automatically

You choose the evaluation metric from set of evaluation metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the evaluation metric.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

**Metrics Computed by the XGBoost Release 0.72 Algorithm**

The XGBoost algorithm based on version 0.72 computes the following nine metrics to use for model validation. When tuning the model, choose one of these metrics to evaluate the model. For full list of valid eval_metric values, refer to XGBoost Learning Task Parameters

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:auc</td>
<td>Area under the curve.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:error</td>
<td>Binary classification error rate, calculated as #(wrong cases)/#(all cases).</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:logloss</td>
<td>Negative log-likelihood.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:mae</td>
<td>Mean absolute error.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:map</td>
<td>Mean average precision.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:merror</td>
<td>Multiclass classification error rate, calculated as #(wrong cases)/#(all cases).</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:mlogloss</td>
<td>Negative log-likelihood for multiclass classification.</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:ndcg</td>
<td>Normalized Discounted Cumulative Gain.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:rmse</td>
<td>Root mean square error.</td>
<td>Minimize</td>
</tr>
</tbody>
</table>
Tunable XGBoost Release 0.72 Hyperparameters

Tune the XGBoost model with the following hyperparameters. The hyperparameters that have the greatest effect on optimizing the XGBoost evaluation metrics are: alpha, min_child_weight, subsample, eta, and num_round.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>alpha</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 1000</td>
</tr>
<tr>
<td>colsample_bylevel</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 1</td>
</tr>
<tr>
<td>colsample_bytree</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.5, MaxValue: 1</td>
</tr>
<tr>
<td>eta</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 0.5</td>
</tr>
<tr>
<td>gamma</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 5</td>
</tr>
<tr>
<td>lambda</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 1000</td>
</tr>
<tr>
<td>max_delta_step</td>
<td>IntegerParameterRanges</td>
<td>[0, 10]</td>
</tr>
<tr>
<td>max_depth</td>
<td>IntegerParameterRanges</td>
<td>[0, 10]</td>
</tr>
<tr>
<td>min_child_weight</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0, MaxValue: 120</td>
</tr>
<tr>
<td>num_round</td>
<td>IntegerParameterRanges</td>
<td>[1, 4000]</td>
</tr>
<tr>
<td>subsample</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.5, MaxValue: 1</td>
</tr>
</tbody>
</table>
**Built-in SageMaker Algorithms for Text Data**

SageMaker provides algorithms that are tailored to the analysis of textual documents used in natural language processing, document classification or summarization, topic modeling or classification, and language transcription or translation.

- **BlazingText algorithm**—a highly optimized implementation of the Word2vec and text classification algorithms that scale to large datasets easily. It is useful for many downstream natural language processing (NLP) tasks.

- **Latent Dirichlet Allocation (LDA) Algorithm**—an algorithm suitable for determining topics in a set of documents. It is an *unsupervised algorithm*, which means that it doesn't use example data with answers during training.

- **Neural Topic Model (NTM) Algorithm**—another unsupervised technique for determining topics in a set of documents, using a neural network approach.

- **Object2Vec Algorithm**—a general-purpose neural embedding algorithm that can be used for recommendation systems, document classification, and sentence embeddings.

- **Sequence-to-Sequence Algorithm**—a supervised algorithm commonly used for neural machine translation.

- **Text Classification - TensorFlow**—a supervised algorithm that supports transfer learning with available pretrained models for text classification.

<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>BlazingText</td>
<td>train</td>
<td>File or Pipe</td>
<td>Text file (one sentence per line with space-separated tokens)</td>
<td>GPU (single instance only) or CPU</td>
<td>No</td>
</tr>
</tbody>
</table>

Use Built-in Algorithms
<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDA</td>
<td>train and (optional ly) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Neural Topic Model</td>
<td>train and (optional ly) validation, test, or both</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>GPU or CPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Object2Vec</td>
<td>train and (optional ly) validation, test, or both</td>
<td>File</td>
<td>JSON Lines</td>
<td>GPU or CPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Seq2Seq Modeling</td>
<td>train, validation, and vocab</td>
<td>File</td>
<td>recordIO-protobuf</td>
<td>GPU (single instance only)</td>
<td>No</td>
</tr>
<tr>
<td>Text Classification - TensorFlow</td>
<td>training and validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU</td>
<td>Yes (only across multiple GPUs on a single instance)</td>
</tr>
</tbody>
</table>
BlazingText algorithm

The Amazon SageMaker BlazingText algorithm provides highly optimized implementations of the Word2vec and text classification algorithms. The Word2vec algorithm is useful for many downstream natural language processing (NLP) tasks, such as sentiment analysis, named entity recognition, machine translation, etc. Text classification is an important task for applications that perform web searches, information retrieval, ranking, and document classification.

The Word2vec algorithm maps words to high-quality distributed vectors. The resulting vector representation of a word is called a word embedding. Words that are semantically similar correspond to vectors that are close together. That way, word embeddings capture the semantic relationships between words.

Many natural language processing (NLP) applications learn word embeddings by training on large collections of documents. These pretrained vector representations provide information about semantics and word distributions that typically improves the generalizability of other models that are later trained on a more limited amount of data. Most implementations of the Word2vec algorithm are not optimized for multi-core CPU architectures. This makes it difficult to scale to large datasets.

With the BlazingText algorithm, you can scale to large datasets easily. Similar to Word2vec, it provides the Skip-gram and continuous bag-of-words (CBOW) training architectures. BlazingText's implementation of the supervised multi-class, multi-label text classification algorithm extends the fastText text classifier to use GPU acceleration with custom CUDA kernels. You can train a model on more than a billion words in a couple of minutes using a multi-core CPU or a GPU. And, you achieve performance on par with the state-of-the-art deep learning text classification algorithms.

The BlazingText algorithm is not parallelizable. For more information on parameters related to training, see Docker Registry Paths for SageMaker Built-in Algorithms.

The SageMaker BlazingText algorithms provides the following features:

- Accelerated training of the fastText text classifier on multi-core CPUs or a GPU and Word2Vec on GPUs using highly optimized CUDA kernels. For more information, see BlazingText: Scaling and Accelerating Word2Vec using Multiple GPUs.
- Enriched Word Vectors with Subword Information by learning vector representations for character n-grams. This approach enables BlazingText to generate meaningful vectors for out-of-vocabulary (OOV) words by representing their vectors as the sum of the character n-gram (subword) vectors.
A batch_skipgram mode for the Word2Vec algorithm that allows faster training and distributed computation across multiple CPU nodes. The batch_skipgram mode does mini-batching using the Negative Sample Sharing strategy to convert level-1 BLAS operations into level-3 BLAS operations. This efficiently leverages the multiply-add instructions of modern architectures. For more information, see Parallelizing Word2Vec in Shared and Distributed Memory.

To summarize, the following modes are supported by BlazingText on different types instances:

<table>
<thead>
<tr>
<th>Modes</th>
<th>Word2Vec (Unsupervised Learning)</th>
<th>Text Classification (Supervised Learning)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single CPU instance</td>
<td>cbow, Skip-gram, Batch Skip-gram</td>
<td>supervised</td>
</tr>
<tr>
<td>Single GPU instance (with 1 or more GPUs)</td>
<td>cbow, Skip-gram</td>
<td>supervised with one GPU</td>
</tr>
<tr>
<td>Multiple CPU instances</td>
<td>Batch Skip-gram</td>
<td>None</td>
</tr>
</tbody>
</table>

For more information about the mathematics behind BlazingText, see BlazingText: Scaling and Accelerating Word2Vec using Multiple GPUs.

Topics

- Input/Output Interface for the BlazingText Algorithm
- EC2 Instance Recommendation for the BlazingText Algorithm
- BlazingText Sample Notebooks
- BlazingText Hyperparameters
- Tune a BlazingText Model
Input/Output Interface for the BlazingText Algorithm

The BlazingText algorithm expects a single preprocessed text file with space-separated tokens. Each line in the file should contain a single sentence. If you need to train on multiple text files, concatenate them into one file and upload the file in the respective channel.

Training and Validation Data Format

Training and Validation Data Format for the Word2Vec Algorithm

For Word2Vec training, upload the file under the train channel. No other channels are supported. The file should contain a training sentence per line.

Training and Validation Data Format for the Text Classification Algorithm

For supervised mode, you can train with file mode or with the augmented manifest text format.

Train with File Mode

For supervised mode, the training/validation file should contain a training sentence per line along with the labels. Labels are words that are prefixed by the string __label__. Here is an example of a training/validation file:

```
__label__4  linux ready for prime time , intel says , despite all the linux hype , the open-source movement has yet to make a huge splash in the desktop market . that may be about to change , thanks to chipmaking giant intel corp .
__label__2  bowled by the slower one again , kolkata , november 14 the past caught up with sourav ganguly as the indian skippers return to international cricket was short lived .
```

Note

The order of labels within the sentence doesn't matter.

Upload the training file under the train channel, and optionally upload the validation file under the validation channel.
Train with Augmented Manifest Text Format

Supervised mode for CPU instances also supports the augmented manifest format, which enables you to do training in pipe mode without needing to create RecordIO files. While using the format, an S3 manifest file needs to be generated that contains the list of sentences and their corresponding labels. The manifest file format should be in JSON Lines format in which each line represents one sample. The sentences are specified using the source tag and the label can be specified using the label tag. Both source and label tags should be provided under the AttributeNames parameter value as specified in the request.

```
{"source":"linux ready for prime time, intel says, despite all the linux hype", "label":1}
{"source":"bowled by the slower one again, kolkata, november 14 the past caught up with sourav ganguly", "label":2}
```

Multi-label training is also supported by specifying a JSON array of labels.

```
{"source":"linux ready for prime time, intel says, despite all the linux hype", "label": [1, 3]}
{"source":"bowled by the slower one again, kolkata, november 14 the past caught up with sourav ganguly", "label": [2, 4, 5]}
```

For more information on augmented manifest files, see [Provide Dataset Metadata to Training Jobs with an Augmented Manifest File](#).

Model Artifacts and Inference

Model Artifacts for the Word2Vec Algorithm

For Word2Vec training, the model artifacts consist of vectors.txt, which contains words-to-vectors mapping, and vectors.bin, a binary used by BlazingText for hosting, inference, or both. vectors.txt stores the vectors in a format that is compatible with other tools like Gensim and Spacy. For example, a Gensim user can run the following commands to load the vectors.txt file:

```
from gensim.models import KeyedVectors
word_vectors = KeyedVectors.load_word2vec_format('vectors.txt', binary=False)
word_vectors.most_similar(positive=['woman', 'king'], negative=['man'])
word_vectors.doesnt_match("breakfast cereal dinner lunch".split())
```

If the evaluation parameter is set to True, an additional file, eval.json, is created. This file contains the similarity evaluation results (using Spearman's rank correlation coefficients) on WS-353
dataset. The number of words from the WS-353 dataset that aren't there in the training corpus are reported.

For inference requests, the model accepts a JSON file containing a list of strings and returns a list of vectors. If the word is not found in vocabulary, inference returns a vector of zeros. If subwords is set to True during training, the model is able to generate vectors for out-of-vocabulary (OOV) words.

**Sample JSON Request**

Mime-type: application/json

```json
{
  "instances": ["word1", "word2", "word3"]
}
```

**Model Artifacts for the Text Classification Algorithm**

Training with supervised outputs creates a `model.bin` file that can be consumed by BlazingText hosting. For inference, the BlazingText model accepts a JSON file containing a list of sentences and returns a list of corresponding predicted labels and probability scores. Each sentence is expected to be a string with space-separated tokens, words, or both.

**Sample JSON Request**

Mime-type: application/json

```json
{
  "instances": ["the movie was excellent", "i did not like the plot ."]
}
```

By default, the server returns only one prediction, the one with the highest probability. For retrieving the top $k$ predictions, you can set $k$ in the configuration, as follows:

```json
{
  "instances": ["the movie was excellent", "i did not like the plot ."],
  "configuration": {"k": 2}
}
```
For BlazingText, the content-type and accept parameters must be equal. For batch transform, they both need to be application/jsonlines. If they differ, the Accept field is ignored. The format for input follows:

```json
content-type: application/jsonlines

{"source": "source_0"}
{"source": "source_1"}

if you need to pass the value of k for top-k, then you can do it in the following way:

{"source": "source_0", "k": 2}
{"source": "source_1", "k": 3}
```

The format for output follows:

```json
accept: application/jsonlines

{"prob": [prob_1], "label": ["__label__1"]}
{"prob": [prob_1], "label": ["__label__1"]}

If you have passed the value of k to be more than 1, then response will be in this format:

```json
{"prob": [prob_1, prob_2], "label": ["__label__1", "__label__2"]}
{"prob": [prob_1, prob_2], "label": ["__label__1", "__label__2"]}
```

For both supervised (text classification) and unsupervised (Word2Vec) modes, the binaries (*.bin) produced by BlazingText can be cross-consumed by fastText and vice versa. You can use binaries produced by BlazingText by fastText. Likewise, you can host the model binaries created with fastText using BlazingText.

Here is an example of how to use a model generated with BlazingText with fastText:

```bash
#Download the model artifact from S3
aws s3 cp s3://<YOUR_S3_BUCKET>/<PREFIX>/model.tar.gz model.tar.gz

#Unzip the model archive
tar -xzf model.tar.gz
```
# Use the model archive with fastText

```bash
fasttext predict ./model.bin test.txt
```

However, the binaries are only supported when training on CPU and single GPU; training on multi-GPU will not produce binaries.

**EC2 Instance Recommendation for the BlazingText Algorithm**

For **cbow** and **skipgram** modes, BlazingText supports single CPU and single GPU instances. Both of these modes support learning of subwords embeddings. To achieve the highest speed without compromising accuracy, we recommend that you use an ml.p3.2xlarge instance.

For **batch_skipgram** mode, BlazingText supports single or multiple CPU instances. When training on multiple instances, set the value of the `S3DataDistributionType` field of the `S3DataSource` object that you pass to `CreateTrainingJob` to `FullyReplicated`. BlazingText takes care of distributing data across machines.

For the supervised text classification mode, a C5 instance is recommended if the training dataset is less than 2 GB. For larger datasets, use an instance with a single GPU. BlazingText supports P2, P3, G4dn, and G5 instances for training and inference.

**BlazingText Sample Notebooks**

For a sample notebook that trains and deploys the SageMaker BlazingText algorithm to generate word vectors, see [Learning Word2Vec Word Representations using BlazingText](#). For instructions for creating and accessing Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](#). After creating and opening a notebook instance, choose the **SageMaker Examples** tab to see a list of all the SageMaker examples. The topic modeling example notebooks that use the Blazing Text are located in the **Introduction to Amazon algorithms** section. To open a notebook, choose its **Use** tab, then choose **Create copy**.

**BlazingText Hyperparameters**

When you start a training job with a `CreateTrainingJob` request, you specify a training algorithm. You can also specify algorithm-specific hyperparameters as string-to-string maps. The hyperparameters for the BlazingText algorithm depend on which mode you use: Word2Vec (unsupervised) and Text Classification (supervised).
## Word2Vec Hyperparameters

The following table lists the hyperparameters for the BlazingText Word2Vec training algorithm provided by Amazon SageMaker.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>mode</strong></td>
<td>The Word2vec architecture used for training.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: batch_skipgram, skipgram, or cbow</td>
</tr>
<tr>
<td><strong>batch_size</strong></td>
<td>The size of each batch when mode is set to batch_skipgram. Set to a number between 10 and 20.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 11</td>
</tr>
<tr>
<td><strong>buckets</strong></td>
<td>The number of hash buckets to use for subwords.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 2000000</td>
</tr>
<tr>
<td><strong>epochs</strong></td>
<td>The number of complete passes through the training data.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td><strong>evaluation</strong></td>
<td>Whether the trained model is evaluated using the WordSimilarity-353 Test.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
|                | Valid values: (Boolean) True or False  
|                | Default value: True |
| learning_rate  | The step size used for parameter updates.  
|                | Optional  
|                | Valid values: Positive float  
|                | Default value: 0.05 |
| min_char       | The minimum number of characters to use for subwords/character n-grams.  
|                | Optional  
|                | Valid values: positive integer  
|                | Default value: 3 |
| min_count      | Words that appear less than min_count times are discarded.  
|                | Optional  
|                | Valid values: Non-negative integer  
|                | Default value: 5 |
| max_char       | The maximum number of characters to use for subwords/character n-grams  
|                | Optional  
|                | Valid values: positive integer  
<p>|                | Default value: 6 |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>negative_samples</td>
<td>The number of negative samples for the negative sample sharing strategy.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td>sampling_threshold</td>
<td>The threshold for the occurrence of words. Words that appear with higher</td>
</tr>
<tr>
<td></td>
<td>frequency in the training data are randomly down-sampled.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive fraction. The recommended range is (0, 1e-3]</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0001</td>
</tr>
<tr>
<td>subwords</td>
<td>Whether to learn subword embeddings on not.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: (Boolean) True or False</td>
</tr>
<tr>
<td></td>
<td>Default value: False</td>
</tr>
<tr>
<td>vector_dim</td>
<td>The dimension of the word vectors that the algorithm learns.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 100</td>
</tr>
</tbody>
</table>
Amazon SageMaker
Developer Guide

Parameter Name | Description
--- | ---
window_size | The size of the context window. The context window is the number of words surrounding the target word used for training.

*Optional*
Valid values: Positive integer
Default value: 5

Text Classification Hyperparameters

The following table lists the hyperparameters for the Text Classification training algorithm provided by Amazon SageMaker.

### Note
Although some of the parameters are common between the Text Classification and Word2Vec modes, they might have different meanings depending on the context.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>mode</td>
<td>The training mode.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: supervised</td>
</tr>
<tr>
<td>buckets</td>
<td>The number of hash buckets to use for word n-grams.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 2000000</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| early_stopping | Whether to stop training if validation accuracy doesn't improve after a patience number of epochs. Note that a validation channel is required if early stopping is used.  
**Optional**  
Valid values: (Boolean) True or False  
Default value: False |
| epochs | The maximum number of complete passes through the training data.  
**Optional**  
Valid values: Positive integer  
Default value: 5 |
| learning_rate | The step size used for parameter updates.  
**Optional**  
Valid values: Positive float  
Default value: 0.05 |
| min_count | Words that appear less than min_count times are discarded.  
**Optional**  
Valid values: Non-negative integer  
Default value: 5 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| min_epochs     | The minimum number of epochs to train before early stopping logic is invoked.  
Optional  
Valid values: Positive integer  
Default value: 5 |
| patience       | The number of epochs to wait before applying early stopping when no progress is made on the validation set. Used only when `early_stopping` is True.  
Optional  
Valid values: Positive integer  
Default value: 4 |
| vector_dim     | The dimension of the embedding layer.  
Optional  
Valid values: Positive integer  
Default value: 100 |
| word_ngrams    | The number of word n-gram features to use.  
Optional  
Valid values: Positive integer  
Default value: 2 |

**Tune a BlazingText Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the
tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

**Metrics Computed by the BlazingText Algorithm**

The BlazingText Word2Vec algorithm (skipgram, cbow, and batch_skipgram modes) reports on a single metric during training: `train:mean_rho`. This metric is computed on WS-353 word similarity datasets. When tuning the hyperparameter values for the Word2Vec algorithm, use this metric as the objective.

The BlazingText Text Classification algorithm (supervised mode), also reports on a single metric during training: the `validation:accuracy`. When tuning the hyperparameter values for the text classification algorithm, use these metrics as the objective.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>train:mean_rho</code></td>
<td>The mean rho (Spearman's rank correlation coefficient) on WS-353 word similarity datasets</td>
<td>Maximize</td>
</tr>
<tr>
<td><code>validation:accuracy</code></td>
<td>The classification accuracy on the user-specified validation dataset</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

**Tunable BlazingText Hyperparameters**

**Tunable Hyperparameters for the Word2Vec Algorithm**

Tune an Amazon SageMaker BlazingText Word2Vec model with the following hyperparameters. The hyperparameters that have the greatest impact on Word2Vec objective metrics are: `mode`, `learning_rate`, `window_size`, `vector_dim`, and `negative_samples`. 
### Tunable Hyperparameters for the Text Classification Algorithm

Tune an Amazon SageMaker BlazingText text classification model with the following hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges or Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>buckets</td>
<td>IntegerParameterRange</td>
<td>[10000000-100000000]</td>
</tr>
<tr>
<td>epochs</td>
<td>IntegerParameterRange</td>
<td>[5-15]</td>
</tr>
</tbody>
</table>
### Latent Dirichlet Allocation (LDA) Algorithm

The Amazon SageMaker Latent Dirichlet Allocation (LDA) algorithm is an unsupervised learning algorithm that attempts to describe a set of observations as a mixture of distinct categories. LDA is most commonly used to discover a user-specified number of topics shared by documents within a text corpus. Here each observation is a document, the features are the presence (or occurrence count) of each word, and the categories are the topics. Since the method is unsupervised, the topics are not specified up front, and are not guaranteed to align with how a human may naturally categorize documents. The topics are learned as a probability distribution over the words that occur in each document. Each document, in turn, is described as a mixture of topics.

The exact content of two documents with similar topic mixtures will not be the same. But overall, you would expect these documents to more frequently use a shared subset of words, than when compared with a document from a different topic mixture. This allows LDA to discover these word groups and use them to form topics. As an extremely simple example, given a set of documents where the only words that occur within them are: *eat, sleep, play, meow,* and *bark*, LDA might produce topics like the following:

<table>
<thead>
<tr>
<th>Topic</th>
<th><em>eat</em></th>
<th><em>sleep</em></th>
<th><em>play</em></th>
<th><em>meow</em></th>
<th><em>bark</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Topic 1</td>
<td>0.1</td>
<td>0.3</td>
<td>0.2</td>
<td>0.4</td>
<td>0.0</td>
</tr>
<tr>
<td>Topic 2</td>
<td>0.2</td>
<td>0.1</td>
<td>0.4</td>
<td>0.0</td>
<td>0.3</td>
</tr>
</tbody>
</table>

---

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges or Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.005, MaxValue: 0.01</td>
</tr>
<tr>
<td>min_count</td>
<td>IntegerParameterRange</td>
<td>[0-100]</td>
</tr>
<tr>
<td>vector_dim</td>
<td>IntegerParameterRange</td>
<td>[32-300]</td>
</tr>
<tr>
<td>word_ngrams</td>
<td>IntegerParameterRange</td>
<td>[1-3]</td>
</tr>
</tbody>
</table>
You can infer that documents that are more likely to fall into Topic 1 are about cats (who are more likely to meow and sleep), and documents that fall into Topic 2 are about dogs (who prefer to play and bark). These topics can be found even though the words dog and cat never appear in any of the texts.

**Topics**

- Choosing between Latent Dirichlet Allocation (LDA) and Neural Topic Model (NTM)
- Input/Output Interface for the LDA Algorithm
- EC2 Instance Recommendation for the LDA Algorithm
- LDA Sample Notebooks
- How LDA Works
- LDA Hyperparameters
- Tune an LDA Model

**Choosing between Latent Dirichlet Allocation (LDA) and Neural Topic Model (NTM)**

Topic models are commonly used to produce topics from corpuses that (1) coherently encapsulate semantic meaning and (2) describe documents well. As such, topic models aim to minimize perplexity and maximize topic coherence.

Perplexity is an intrinsic language modeling evaluation metric that measures the inverse of the geometric mean per-word likelihood in your test data. A lower perplexity score indicates better generalization performance. Research has shown that the likelihood computed per word often does not align to human judgement, and can be entirely non-correlated, thus topic coherence has been introduced. Each inferred topic from your model consists of words, and topic coherence is computed to the top N words for that particular topic from your model. It is often defined as the average or median of the pairwise word-similarity scores of the words in that topic e.g., Pointwise Mutual Information (PMI). A promising model generates coherent topics or topics with high topic coherence scores.

While the objective is to train a topic model that minimizes perplexity and maximizes topic coherence, there is often a tradeoff with both LDA and NTM. Recent research by Amazon, Dinget et al., 2018 has shown that NTM is promising for achieving high topic coherence but LDA trained with collapsed Gibbs sampling achieves better perplexity. There is a tradeoff between perplexity and topic coherence. From a practicality standpoint regarding hardware and compute power, SageMaker NTM hardware is more flexible than LDA and can scale better because NTM can run on
CPU and GPU and can be parallelized across multiple GPU instances, whereas LDA only supports single-instance CPU training.

**Topics**
- Input/Output Interface for the LDA Algorithm
- EC2 Instance Recommendation for the LDA Algorithm
- LDA Sample Notebooks
- How LDA Works
- LDA Hyperparameters
- Tune an LDA Model

**Input/Output Interface for the LDA Algorithm**

LDA expects data to be provided on the train channel, and optionally supports a test channel, which is scored by the final model. LDA supports both recordIO-wrapped-protobuf (dense and sparse) and CSV file formats. For CSV, the data must be dense and have dimension equal to \( \text{number of records} \times \text{vocabulary size} \). LDA can be trained in File or Pipe mode when using recordIO-wrapped protobuf, but only in File mode for the CSV format.

For inference, text/csv, application/json, and application/x-recordio-protobuf content types are supported. Sparse data can also be passed for application/json and application/x-recordio-protobuf. LDA inference returns application/json or application/x-recordio-protobuf predictions, which include the topic_mixture vector for each observation.

Please see the [LDA Sample Notebooks](#) for more detail on training and inference formats.

**EC2 Instance Recommendation for the LDA Algorithm**

LDA currently only supports single-instance CPU training. CPU instances are recommended for hosting/inference.

**LDA Sample Notebooks**

For a sample notebook that shows how to train the SageMaker Latent Dirichlet Allocation algorithm on a dataset and then how to deploy the trained model to perform inferences about the topic mixtures in input documents, see the [An Introduction to SageMaker LDA](#). For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker,
see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. The topic modeling example notebooks using the NTM algorithms are located in the Introduction to Amazon algorithms section. To open a notebook, click on its Use tab and select Create copy.

How LDA Works

Amazon SageMaker LDA is an unsupervised learning algorithm that attempts to describe a set of observations as a mixture of different categories. These categories are themselves a probability distribution over the features. LDA is a generative probability model, which means it attempts to provide a model for the distribution of outputs and inputs based on latent variables. This is opposed to discriminative models, which attempt to learn how inputs map to outputs.

You can use LDA for a variety of tasks, from clustering customers based on product purchases to automatic harmonic analysis in music. However, it is most commonly associated with topic modeling in text corpuses. Observations are referred to as documents. The feature set is referred to as vocabulary. A feature is referred to as a word. And the resulting categories are referred to as topics.

Note

Lemmatization significantly increases algorithm performance and accuracy. Consider preprocessing any input text data. For more information, see Stemming and lemmatization.

An LDA model is defined by two parameters:

- $\alpha$—A prior estimate on topic probability (in other words, the average frequency that each topic within a given document occurs).
- $\beta$—a collection of $k$ topics where each topic is given a probability distribution over the vocabulary used in a document corpus, also called a "topic-word distribution."

LDA is a "bag-of-words" model, which means that the order of words does not matter. LDA is a generative model where each document is generated word-by-word by choosing a topic mixture $\theta \sim$ Dirichlet($\alpha$).

For each word in the document:

- Choose a topic $z \sim$ Multinomial($\theta$)
Choose the corresponding topic-word distribution $\beta_z$.

Draw a word $w \sim \text{Multinomial}(\beta_z)$.

When training the model, the goal is to find parameters $\alpha$ and $\beta$, which maximize the probability that the text corpus is generated by the model.

The most popular methods for estimating the LDA model use Gibbs sampling or Expectation Maximization (EM) techniques. The Amazon SageMaker LDA uses tensor spectral decomposition. This provides several advantages:

- **Theoretical guarantees on results.** The standard EM-method is guaranteed to converge only to local optima, which are often of poor quality.

- **Embarrassingly parallelizable.** The work can be trivially divided over input documents in both training and inference. The EM-method and Gibbs Sampling approaches can be parallelized, but not as easily.

- **Fast.** Although the EM-method has low iteration cost it is prone to slow convergence rates. Gibbs Sampling is also subject to slow convergence rates and also requires a large number of samples.

At a high-level, the tensor decomposition algorithm follows this process:

1. The goal is to calculate the spectral decomposition of a $V \times V \times V$ tensor, which summarizes the moments of the documents in our corpus. $V$ is vocabulary size (in other words, the number of distinct words in all of the documents). The spectral components of this tensor are the LDA parameters $\alpha$ and $\beta$, which maximize the overall likelihood of the document corpus. However, because vocabulary size tends to be large, this $V \times V \times V$ tensor is prohibitively large to store in memory.

2. Instead, it uses a $V \times V$ moment matrix, which is the two-dimensional analog of the tensor from step 1, to find a whitening matrix of dimension $V \times k$. This matrix can be used to convert the $V \times V$ moment matrix into a $k \times k$ identity matrix. $k$ is the number of topics in the model.

3. This same whitening matrix can then be used to find a smaller $k \times k \times k$ tensor. When spectrally decomposed, this tensor has components that have a simple relationship with the components of the $V \times V \times V$ tensor.

4. **Alternating Least Squares** is used to decompose the smaller $k \times k \times k$ tensor. This provides a substantial improvement in memory consumption and speed. The parameters $\alpha$ and $\beta$ can be found by “unwhitening” these outputs in the spectral decomposition.
After the LDA model's parameters have been found, you can find the topic mixtures for each document. You use stochastic gradient descent to maximize the likelihood function of observing a given topic mixture corresponding to these data.

Topic quality can be improved by increasing the number of topics to look for in training and then filtering out poor quality ones. This is in fact done automatically in SageMaker LDA: 25% more topics are computed and only the ones with largest associated Dirichlet priors are returned. To perform further topic filtering and analysis, you can increase the topic count and modify the resulting LDA model as follows:

```python
> import mxnet as mx
> alpha, beta = mx.ndarray.load('model.tar.gz')
> # modify alpha and beta
> mx.nd.save('new_model.tar.gz', [new_alpha, new_beta])
> # upload to S3 and create new SageMaker model using the console
```

For more information about algorithms for LDA and the SageMaker implementation, see the following:


**LDA Hyperparameters**

In the `CreateTrainingJob` request, you specify the training algorithm. You can also specify algorithm-specific hyperparameters as string-to-string maps. The following table lists the hyperparameters for the LDA training algorithm provided by Amazon SageMaker. For more information, see [How LDA Works](#).
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_topics</td>
<td>The number of topics for LDA to find within the data.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>feature_dim</td>
<td>The size of the vocabulary of the input document corpus.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>The total number of documents in the input document corpus.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>alpha0</td>
<td>Initial guess for the concentration parameter: the sum of the elements of the Dirichlet prior. Small values are more likely to generate sparse topic mixtures and large values (greater than 1.0) produce more uniform mixtures.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive float</td>
</tr>
<tr>
<td></td>
<td>Default value: 1.0</td>
</tr>
<tr>
<td>max_restarts</td>
<td>The number of restarts to perform during the Alternating Least Squares (ALS) spectral decomposition phase of the algorithm. Can be used to find better quality local minima at the expense of additional computation, but typically should not be adjusted.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
</tbody>
</table>
## Tune an LDA Model

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

LDA is an unsupervised topic modeling algorithm that attempts to describe a set of observations (documents) as a mixture of different categories (topics). The “per-word log-likelihood” (PWLL) metric measures the likelihood that a learned set of topics (an LDA model) accurately describes a test document dataset. Larger values of PWLL indicate that the test data is more likely to be described by the LDA model.
For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Metrics Computed by the LDA Algorithm

The LDA algorithm reports on a single metric during training: `test:pwll`. When tuning a model, choose this metric as the objective metric.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>test:pwll</code></td>
<td>Per-word log-likelihood on the test dataset. The likelihood that the test dataset is accurately described by the learned LDA model.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

Tunable LDA Hyperparameters

You can tune the following hyperparameters for the LDA algorithm. Both hyperparameters, `alpha0` and `num_topics`, can affect the LDA objective metric (`test:pwll`). If you don't already know the optimal values for these hyperparameters, which maximize per-word log-likelihood and produce an accurate LDA model, automatic model tuning can help find them.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>alpha0</code></td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.1, MaxValue: 10</td>
</tr>
<tr>
<td><code>num_topics</code></td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 150</td>
</tr>
</tbody>
</table>

Neural Topic Model (NTM) Algorithm

Amazon SageMaker NTM is an unsupervised learning algorithm that is used to organize a corpus of documents into topics that contain word groupings based on their statistical distribution. Documents that contain frequent occurrences of words such as "bike", "car", "train", "mileage", and "speed" are likely to share a topic on "transportation" for example. Topic modeling can be used to classify or summarize documents based on the topics detected or to retrieve information
or recommend content based on topic similarities. The topics from documents that NTM learns are characterized as a *latent representation* because the topics are inferred from the observed word distributions in the corpus. The semantics of topics are usually inferred by examining the top ranking words they contain. Because the method is unsupervised, only the number of topics, not the topics themselves, are prespecified. In addition, the topics are not guaranteed to align with how a human might naturally categorize documents.

Topic modeling provides a way to visualize the contents of a large document corpus in terms of the learned topics. Documents relevant to each topic might be indexed or searched for based on their soft topic labels. The latent representations of documents might also be used to find similar documents in the topic space. You can also use the latent representations of documents that the topic model learns for input to another supervised algorithm such as a document classifier. Because the latent representations of documents are expected to capture the semantics of the underlying documents, algorithms based in part on these representations are expected to perform better than those based on lexical features alone.

Although you can use both the Amazon SageMaker NTM and LDA algorithms for topic modeling, they are distinct algorithms and can be expected to produce different results on the same input data.

For more information on the mathematics behind NTM, see [Neural Variational Inference for Text Processing](#).

**Topics**

- [Input/Output Interface for the NTM Algorithm](#)
- [EC2 Instance Recommendation for the NTM Algorithm](#)
- [NTM Sample Notebooks](#)
- [NTM Hyperparameters](#)
- [Tune an NTM Model](#)
- [NTM Response Formats](#)

**Input/Output Interface for the NTM Algorithm**

Amazon SageMaker Neural Topic Model supports four data channels: train, validation, test, and auxiliary. The validation, test, and auxiliary data channels are optional. If you specify any of these optional channels, set the value of the S3DataDistributionType parameter for them to **FullyReplicated**. If you provide validation data, the loss on this data is logged at every epoch,
and the model stops training as soon as it detects that the validation loss is not improving. If you don't provide validation data, the algorithm stops early based on the training data, but this can be less efficient. If you provide test data, the algorithm reports the test loss from the final model.

The train, validation, and test data channels for NTM support both recordIO-wrapped-protobuf (dense and sparse) and CSV file formats. For CSV format, each row must be represented densely with zero counts for words not present in the corresponding document, and have dimension equal to: \((\text{number of records}) \times (\text{vocabulary size})\). You can use either File mode or Pipe mode to train models on data that is formatted as recordIO-wrapped-protobuf or as CSV. The auxiliary channel is used to supply a text file that contains vocabulary. By supplying the vocabulary file, users are able to see the top words for each of the topics printed in the log instead of their integer IDs. Having the vocabulary file also allows NTM to compute the Word Embedding Topic Coherence (WETC) scores, a new metric displayed in the log that captures similarity among the top words in each topic effectively. The ContentType for the auxiliary channel is text/plain, with each line containing a single word, in the order corresponding to the integer IDs provided in the data. The vocabulary file must be named vocab.txt and currently only UTF-8 encoding is supported.

For inference, text/csv, application/json, application/jsonlines, and application/x-recordio-protobuf content types are supported. Sparse data can also be passed for application/json and application/x-recordio-protobuf. NTM inference returns application/json or application/x-recordio-protobuf predictions, which include the topic_weights vector for each observation.

See the [blog post](#) and the companion [notebook](#) for more details on using the auxiliary channel and the WETC scores. For more information on how to compute the WETC score, see [Coherence-Aware Neural Topic Modeling](#). We used the pairwise WETC described in this paper for the Amazon SageMaker Neural Topic Model.

For more information on input and output file formats, see [NTM Response Formats](#) for inference and the [NTM Sample Notebooks](#).

### EC2 Instance Recommendation for the NTM Algorithm

NTM training supports both GPU and CPU instance types. We recommend GPU instances, but for certain workloads, CPU instances may result in lower training costs. CPU instances should be sufficient for inference. NTM training supports P2, P3, G4dn, and G5 GPU instance families for training and inference.
NTM Sample Notebooks

For a sample notebook that uses the SageMaker NTM algorithm to uncover topics in documents from a synthetic data source where the topic distributions are known, see the [Introduction to Basic Functionality of NTM](#). For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](#). Once you have created a notebook instance and opened it, select the [SageMaker Examples](#) tab to see a list of all the SageMaker samples. The topic modeling example notebooks using the NTM algorithms are located in the [Introduction to Amazon algorithms](#) section. To open a notebook, click on its Use tab and select Create copy.

### NTM Hyperparameters

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>feature_dim</td>
<td>The vocabulary size of the dataset.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer (min: 1, max: 1,000,000)</td>
</tr>
<tr>
<td>num_topics</td>
<td>The number of required topics.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer (min: 2, max: 1000)</td>
</tr>
<tr>
<td>batch_norm</td>
<td>Whether to use batch normalization during training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: true or false</td>
</tr>
<tr>
<td></td>
<td>Default value: false</td>
</tr>
<tr>
<td>clip_gradient</td>
<td>The maximum magnitude for each gradient component.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float (min: 1e-3)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| encoder_layers | The number of layers in the encoder and the output size of each layer. When set to *auto*, the algorithm uses two layers of sizes $3 \times \text{num_topics}$ and $2 \times \text{num_topics}$ respectively.  

**Optional**  
Valid values: Comma-separated list of positive integers or *auto*  
Default value: *auto* |
| encoder_layers_activation | The activation function to use in the encoder layers.  

**Optional**  
Valid values:  
- *sigmoid*: Sigmoid function  
- *tanh*: Hyperbolic tangent  
- *relu*: Rectified linear unit  
Default value: *sigmoid* |
| epochs | The maximum number of passes over the training data.  

**Optional**  
Valid values: Positive integer (min: 1)  
Default value: 50 |
| learning_rate | The learning rate for the optimizer.  

**Optional**  
Valid values: Float (min: 1e-6, max: 1.0)  
Default value: 0.001 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>mini_batch_size</td>
<td>The number of examples in each mini batch.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer (min: 1, max: 10000)</td>
</tr>
<tr>
<td></td>
<td>Default value: 256</td>
</tr>
<tr>
<td>num_patience_epochs</td>
<td>The number of successive epochs over which early stopping criterion is evaluated. Early stopping is triggered when the change in the loss function drops below the specified tolerance within the last number of epochs. To disable early stopping, set num_patience_epochs to a value larger than epochs.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer (min: 1)</td>
</tr>
<tr>
<td></td>
<td>Default value: 3</td>
</tr>
<tr>
<td>optimizer</td>
<td>The optimizer to use for training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values:</td>
</tr>
<tr>
<td></td>
<td>• sgd: <a href="#">Stochastic gradient descent</a></td>
</tr>
<tr>
<td></td>
<td>• adam: <a href="#">Adaptive momentum estimation</a></td>
</tr>
<tr>
<td></td>
<td>• adagrad: <a href="#">Adaptive gradient algorithm</a></td>
</tr>
<tr>
<td></td>
<td>• adadelta: <a href="#">An adaptive learning rate algorithm</a></td>
</tr>
<tr>
<td></td>
<td>• rmsprop: <a href="#">Root mean square propagation</a></td>
</tr>
<tr>
<td></td>
<td>Default value: adadelta</td>
</tr>
</tbody>
</table>
### Parameter Name | Description
--- | ---
rescale_gradient | The rescale factor for gradient.  
**Optional**  
Valid values: float (min: $1e^{-3}$, max: 1.0)  
Default value: 1.0

sub_sample | The fraction of the training data to sample for training per epoch.  
**Optional**  
Valid values: Float (min: 0.0, max: 1.0)  
Default value: 1.0

tolerance | The maximum relative change in the loss function. Early stopping is triggered when change in the loss function drops below this value within the last `num_patience_epochs` number of epochs.  
**Optional**  
Valid values: Float (min: $1e^{-6}$, max: 0.1)  
Default value: 0.001

weight_decay | The weight decay coefficient. Adds L2 regularization.  
**Optional**  
Valid values: Float (min: 0.0, max: 1.0)  
Default value: 0.0
**Tune an NTM Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

Amazon SageMaker NTM is an unsupervised learning algorithm that learns latent representations of large collections of discrete data, such as a corpus of documents. Latent representations use inferred variables that are not directly measured to model the observations in a dataset. Automatic model tuning on NTM helps you find the model that minimizes loss over the training or validation data. *Training loss* measures how well the model fits the training data. *Validation loss* measures how well the model can generalize to data that it is not trained on. Low training loss indicates that a model is a good fit to the training data. Low validation loss indicates that a model has not overfit the training data and so should be able to model documents successfully on which is has not been trained. Usually, it's preferable to have both losses be small. However, minimizing training loss too much might result in overfitting and increase validation loss, which would reduce the generality of the model.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics Computed by the NTM Algorithm**

The NTM algorithm reports a single metric that is computed during training: 
*validation:total_loss*. The total loss is the sum of the reconstruction loss and Kullback-Leibler divergence. When tuning hyperparameter values, choose this metric as the objective.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:total_loss</td>
<td>Total Loss on validation set</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

**Tunable NTM Hyperparameters**

You can tune the following hyperparameters for the NTM algorithm. Usually setting low *mini_batch_size* and small *learning_rate* values results in lower validation losses, although
it might take longer to train. Low validation losses don't necessarily produce more coherent topics as interpreted by humans. The effect of other hyperparameters on training and validation loss can vary from dataset to dataset. To see which values are compatible, see NTM Hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>encoder_layers_activation</td>
<td>CategoricalParameterRanges</td>
<td>['sigmoid', 'tanh', 'relu']</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-4, MaxValue: 0.1</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 16, MaxValue:2048</td>
</tr>
<tr>
<td>optimizer</td>
<td>CategoricalParameterRanges</td>
<td>['sgd', 'adam', 'adadelta']</td>
</tr>
<tr>
<td>rescale_gradient</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.1, MaxValue: 1.0</td>
</tr>
<tr>
<td>weight_decay</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 1.0</td>
</tr>
</tbody>
</table>

**NTM Response Formats**

All Amazon SageMaker built-in algorithms adhere to the common input inference format described in Common Data Formats - Inference. This topic contains a list of the available output formats for the SageMaker NTM algorithm.

**JSON Response Format**

```json
{
    "predictions": [
        {
            "topic_weights": [0.02, 0.1, 0,...],
            "topic_weights": [0.25, 0.067, 0,...]
        }
    ]
}"
```
Object2Vec Algorithm

The Amazon SageMaker Object2Vec algorithm is a general-purpose neural embedding algorithm that is highly customizable. It can learn low-dimensional dense embeddings of high-dimensional objects. The embeddings are learned in a way that preserves the semantics of the relationship between pairs of objects in the original space in the embedding space. You can use the learned embeddings to efficiently compute nearest neighbors of objects and to visualize natural clusters of related objects in low-dimensional space, for example. You can also use the embeddings as features of the corresponding objects in downstream supervised tasks, such as classification or regression.
Object2Vec generalizes the well-known Word2Vec embedding technique for words that is optimized in the SageMaker BlazingText algorithm. For a blog post that discusses how to apply Object2Vec to some practical use cases, see Introduction to Amazon SageMaker Object2Vec.

Topics

- I/O Interface for the Object2Vec Algorithm
- EC2 Instance Recommendation for the Object2Vec Algorithm
- Object2Vec Sample Notebooks
- How Object2Vec Works
- Object2Vec Hyperparameters
- Tune an Object2Vec Model
- Data Formats for Object2Vec Training
- Data Formats for Object2Vec Inference
- Encoder Embeddings for Object2Vec

I/O Interface for the Object2Vec Algorithm

You can use Object2Vec on many input data types, including the following examples.

<table>
<thead>
<tr>
<th>Input Data Type</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sentence-sentence pairs</td>
<td>&quot;A soccer game with multiple males playing.&quot; and &quot;Some men are playing a sport.&quot;</td>
</tr>
<tr>
<td>Labels-sequence pairs</td>
<td>The genre tags of the movie &quot;Titanic&quot;, such as &quot;Romance&quot; and &quot;Drama&quot;, and its short description: &quot;James Cameron's Titanic is an epic, action-packed romance set against the ill-fated maiden voyage of the R.M.S. Titanic. She was the most luxurious liner of her era, a ship of dreams, which ultimately carried over 1,500 people to their death in the ice cold waters of the North Atlantic in the early hours of April 15, 1912.&quot;</td>
</tr>
<tr>
<td>Customer-customer pairs</td>
<td>The customer ID of Jane and customer ID of Jackie.</td>
</tr>
<tr>
<td>Product-product pairs</td>
<td>The product ID of football and product ID of basketball.</td>
</tr>
</tbody>
</table>
The type of Amazon Elastic Compute Cloud (Amazon EC2) instance that you use depends on whether you are training or running inference.
When training a model using the Object2Vec algorithm on a CPU, start with an ml.m5.2xlarge instance. For training on a GPU, start with an ml.p2.xlarge instance. If the training takes too long on this instance, you can use a larger instance. Currently, the Object2Vec algorithm can train only on a single machine. However, it does offer support for multiple GPUs. Object2Vec supports P2, P3, G4dn, and G5 GPU instance families for training and inference.

For inference with a trained Object2Vec model that has a deep neural network, we recommend using ml.p3.2xlarge GPU instance. Due to GPU memory scarcity, the INFRINGEMENT_PREFERRED_MODE environment variable can be specified to optimize on whether the section called “GPU optimization: Classification or Regression” or the section called “GPU optimization: Encoder Embeddings” inference network is loaded into GPU.

Object2Vec Sample Notebooks

- Using Object2Vec to Encode Sentences into Fixed Length Embeddings

Note

To run the notebooks on a notebook instance, see Example Notebooks. To run the notebooks on Studio, see Create or Open an Amazon SageMaker Studio Classic Notebook.

How Object2Vec Works

When using the Amazon SageMaker Object2Vec algorithm, you follow the standard workflow: process the data, train the model, and produce inferences.

Topics

- Step 1: Process Data
- Step 2: Train a Model
- Step 3: Produce Inferences

Step 1: Process Data

During preprocessing, convert the data to the JSON Lines text file format specified in Data Formats for Object2Vec Training. To get the highest accuracy during training, also randomly shuffle the data before feeding it into the model. How you generate random permutations depends on the language. For python, you could use np.random.shuffle; for Unix, shuf.
Step 2: Train a Model

The SageMaker Object2Vec algorithm has the following main components:

- **Two input channels** – The input channels take a pair of objects of the same or different types as inputs, and pass them to independent and customizable encoders.

- **Two encoders** – The two encoders, enc0 and enc1, convert each object into a fixed-length embedding vector. The encoded embeddings of the objects in the pair are then passed into a comparator.

- **A comparator** – The comparator compares the embeddings in different ways and outputs scores that indicate the strength of the relationship between the paired objects. In the output score for a sentence pair. For example, 1 indicates a strong relationship between a sentence pair, and 0 represents a weak relationship.

During training, the algorithm accepts pairs of objects and their relationship labels or scores as inputs. The objects in each pair can be of different types, as described earlier. If the inputs to both encoders are composed of the same token-level units, you can use a shared token embedding layer by setting the tied_token_embedding_weight hyperparameter to True when you create the training job. This is possible, for example, when comparing sentences that both have word token-level units. To generate negative samples at a specified rate, set the negative_sampling_rate hyperparameter to the desired ratio of negative to positive samples. This hyperparameter expedites learning how to discriminate between the positive samples observed in the training data and the negative samples that are not likely to be observed.

Pairs of objects are passed through independent, customizable encoders that are compatible with the input types of corresponding objects. The encoders convert each object in a pair into a fixed-length embedding vector of equal length. The pair of vectors are passed to a comparator operator, which assembles the vectors into a single vector using the value specified in the he comparator_list hyperparameter. The assembled vector then passes through a multilayer perceptron (MLP) layer, which produces an output that the loss function compares with the labels that you provided. This comparison evaluates the strength of the relationship between the objects in the pair as predicted by the model. The following figure shows this workflow.
Step 3: Produce Inferences

After the model is trained, you can use the trained encoder to preprocess input objects or to perform two types of inference:

- To convert singleton input objects into fixed-length embeddings using the corresponding encoder
- To predict the relationship label or score between a pair of input objects

The inference server automatically figures out which of the types is requested based on the input data. To get the embeddings as output, provide only one input. To predict the relationship label or score, provide both inputs in the pair.
Object2Vec Hyperparameters

In the CreateTrainingJob request, you specify the training algorithm. You can also specify algorithm-specific hyperparameters as string-to-string maps. The following table lists the hyperparameters for the Object2Vec training algorithm.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>enc0_max_seq_len</td>
<td>The maximum sequence length for the enc0 encoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $1 \leq \text{integer} \leq 5000$</td>
</tr>
<tr>
<td>enc0_vocab_size</td>
<td>The vocabulary size of enc0 tokens.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $2 \leq \text{integer} \leq 3000000$</td>
</tr>
<tr>
<td>bucket_width</td>
<td>The allowed difference between data sequence length when bucketing is enabled. To enable bucketing, specify a non-zero value for this parameter.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $0 \leq \text{integer} \leq 100$</td>
</tr>
<tr>
<td></td>
<td>Default value: 0 (no bucketing)</td>
</tr>
<tr>
<td>comparator_list</td>
<td>A list used to customize the way in which two embeddings are compared. The Object2Vec comparator operator layer takes the encodings from both encoders as inputs and outputs a single vector. This vector is a concatenation of subvectors. The string values passed to the comparator_list and the order in which they are passed determine how these subvectors are assembled. For example, if comparator_list=&quot;hadamard, concat&quot; , then the comparator operator constructs the vector by concatenating the Hadamard product of two encodings and the concatenation of two...</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| encodings      | If, on the other hand, `comparator_list="hadamard"`, then the comparator operator constructs the vector as the hadamard product of only two encodings. **Optional** Valid values: A string that contains any combination of the names of the three binary operators: hadamard, concat, or abs_diff. The Object2Vec algorithm currently requires that the two vector encodings have the same dimension. These operators produce the subvectors as follows:  
  - `hadamard`: Constructs a vector as the Hadamard (element-wise) product of two encodings.  
  - `concat`: Constructs a vector as the concatenation of two encodings.  
  - `abs_diff`: Constructs a vector as the absolute difference between two encodings. Default value: "hadamard, concat, abs_diff" |
<p>| dropout        | The dropout probability for network layers. Dropout is a form of regularization used in neural networks that reduces overfitting by trimming codependent neurons. <strong>Optional</strong> Valid values: $0.0 \leq \text{float} \leq 1.0$ Default value: 0.0 |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>early_stopping_patience</td>
<td>The number of consecutive epochs without improvement allowed before early stopping is applied. Improvement is defined by with the early_stopping_tolerance hyperparameter.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: (1 \leq \text{integer} \leq 5)</td>
</tr>
<tr>
<td></td>
<td>Default value: 3</td>
</tr>
<tr>
<td>early_stopping_tolerance</td>
<td>The reduction in the loss function that an algorithm must achieve between consecutive epochs to avoid early stopping after the number of consecutive epochs specified in the early_stopping_patience hyperparameter concludes.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: (0.000001 \leq \text{float} \leq 0.1)</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.01</td>
</tr>
<tr>
<td>enc_dim</td>
<td>The dimension of the output of the embedding layer.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: (4 \leq \text{integer} \leq 10000)</td>
</tr>
<tr>
<td></td>
<td>Default value: 4096</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>enc0_network</td>
<td>The network model for the enc0 encoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: hcnn, bilstm, or pooled_embedding</td>
</tr>
<tr>
<td></td>
<td>• hcnn: A hierarchical convolutional neural network.</td>
</tr>
<tr>
<td></td>
<td>• bilstm: A bidirectional long short-term memory network (biLSTM), in which</td>
</tr>
<tr>
<td></td>
<td>the signal propagates backward and forward in time. This is an appropriate</td>
</tr>
<tr>
<td></td>
<td>recurrent neural network (RNN) architecture for sequential learning tasks.</td>
</tr>
<tr>
<td></td>
<td>• pooled_embedding: Averages the embeddings of all of the tokens in the</td>
</tr>
<tr>
<td></td>
<td>input.</td>
</tr>
<tr>
<td></td>
<td>Default value: hcnn</td>
</tr>
<tr>
<td>enc0_cnn_filter_width</td>
<td>The filter width of the convolutional neural network (CNN) enc0 encoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 1 ≤ integer ≤ 9</td>
</tr>
<tr>
<td></td>
<td>Default value: 3</td>
</tr>
<tr>
<td>enc0_freeze_pretrained_embedding</td>
<td>Whether to freeze enc0 pretrained embedding weights.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: True or False</td>
</tr>
<tr>
<td></td>
<td>Default value: True</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>enc0_layers</td>
<td>The number of layers in the enc0 encoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto or $1 \leq \text{integer} \leq 4$</td>
</tr>
<tr>
<td></td>
<td>• For hcnn, auto means 4.</td>
</tr>
<tr>
<td></td>
<td>• For bilstm, auto means 1.</td>
</tr>
<tr>
<td></td>
<td>• For pooled_embedding, auto ignores the number of layers.</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>enc0_pretrained_token_embedding_file</td>
<td>The filename of the pretrained enc0 token embedding file in the auxiliary data channel.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String with alphanumeric characters, underscore, or period.</td>
</tr>
<tr>
<td></td>
<td>[A-Za-z0-9._]</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;&quot; (empty string)</td>
</tr>
<tr>
<td>enc0_token_embedding_dim</td>
<td>The output dimension of the enc0 token embedding layer.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $2 \leq \text{integer} \leq 1000$</td>
</tr>
<tr>
<td></td>
<td>Default value: 300</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>enc0_vocab_file</td>
<td>The vocabulary file for mapping pretrained enc0 token embedding vectors to numerical vocabulary IDs.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String with alphanumeric characters, underscore, or period. [A-Za-z0-9._]</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;&quot; (empty string)</td>
</tr>
<tr>
<td>enc1_network</td>
<td>The network model for the enc1 encoder. If you want the enc1 encoder to use the same network model as enc0, including the hyperparameter values, set the value to enc0.</td>
</tr>
<tr>
<td></td>
<td><strong>Note</strong></td>
</tr>
<tr>
<td></td>
<td>Even when the enc0 and enc1 encoder networks have symmetric architectures, you can’t shared parameter values for these networks.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: enc0, hcnn, bilstm, or pooled_embedding</td>
</tr>
<tr>
<td></td>
<td>• enc0: The network model for the enc0 encoder.</td>
</tr>
<tr>
<td></td>
<td>• hcnn: A hierarchical convolutional neural network.</td>
</tr>
<tr>
<td></td>
<td>• bilstm: A bidirectional LSTM, in which the signal propagates backward and forward in time. This is an appropriate recurrent neural network (RNN) architecture for sequential learning tasks.</td>
</tr>
<tr>
<td></td>
<td>• pooled_embedding: The averages of the embeddings of all of the tokens in the input.</td>
</tr>
<tr>
<td></td>
<td>Default value: enc0</td>
</tr>
</tbody>
</table>

Use Built-in Algorithms
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>enc1_cnn_filter_width</td>
<td>The filter width of the CNN enc1 encoder.</td>
</tr>
<tr>
<td><strong>Conditional</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: 1 ≤ integer ≤ 9</td>
<td></td>
</tr>
<tr>
<td>Default value: 3</td>
<td></td>
</tr>
<tr>
<td>enc1_freeze_pretrained_embedding</td>
<td>Whether to freeze enc1 pretrained embedding weights.</td>
</tr>
<tr>
<td><strong>Conditional</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: True or False</td>
<td></td>
</tr>
<tr>
<td>Default value: True</td>
<td></td>
</tr>
<tr>
<td>enc1_layers</td>
<td>The number of layers in the enc1 encoder.</td>
</tr>
<tr>
<td><strong>Conditional</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: auto or 1 ≤ integer ≤ 4</td>
<td></td>
</tr>
<tr>
<td>• For hcnn, auto means 4.</td>
<td></td>
</tr>
<tr>
<td>• For bilstm, auto means 1.</td>
<td></td>
</tr>
<tr>
<td>• For pooled_embedding, auto ignores the number of layers.</td>
<td></td>
</tr>
<tr>
<td>Default value: auto</td>
<td></td>
</tr>
<tr>
<td>enc1_max_seq_len</td>
<td>The maximum sequence length for the enc1 encoder.</td>
</tr>
<tr>
<td><strong>Conditional</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: 1 ≤ integer ≤ 5000</td>
<td></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| enc1_pretrained_embedding_file | The name of the enc1 pretrained token embedding file in the auxiliary data channel.  

**Conditional**  
Valid values: String with alphanumeric characters, underscore, or period. [A-Za-z0-9\_\.]  
Default value: "" (empty string) |
| enc1_token_embedding_dim | The output dimension of the enc1 token embedding layer.  

**Conditional**  
Valid values: $2 \leq \text{integer} \leq 1000$  
Default value: 300 |
| enc1_vocab_file | The vocabulary file for mapping pretrained enc1 token embeddings to vocabulary IDs.  

**Conditional**  
Valid values: String with alphanumeric characters, underscore, or period. [A-Za-z0-9\_\.]  
Default value: "" (empty string) |
| enc1_vocab_size | The vocabulary size of enc0 tokens.  

**Conditional**  
Valid values: $2 \leq \text{integer} \leq 3000000$ |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>epochs</td>
<td>The number of epochs to run for training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $1 \leq \text{integer} \leq 100$</td>
</tr>
<tr>
<td></td>
<td>Default value: 30</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The learning rate for training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $1.0E-6 \leq \text{float} \leq 1.0$</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0004</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>The batch size that the dataset is split into for an optimizer during training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $1 \leq \text{integer} \leq 10000$</td>
</tr>
<tr>
<td></td>
<td>Default value: 32</td>
</tr>
<tr>
<td>mlp_activation</td>
<td>The type of activation function for the multilayer perceptron (MLP) layer.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: tanh, relu, or linear</td>
</tr>
<tr>
<td></td>
<td>• tanh: Hyperbolic tangent</td>
</tr>
<tr>
<td></td>
<td>• relu: Rectified linear unit (ReLU)</td>
</tr>
<tr>
<td></td>
<td>• linear: Linear function</td>
</tr>
<tr>
<td></td>
<td>Default value: linear</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>mlp_dim</td>
<td>The dimension of the output from MLP layers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 2 ≤ integer ≤ 10000</td>
</tr>
<tr>
<td></td>
<td>Default value: 512</td>
</tr>
<tr>
<td>mlp_layers</td>
<td>The number of MLP layers in the network.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 ≤ integer ≤ 10</td>
</tr>
<tr>
<td></td>
<td>Default value: 2</td>
</tr>
<tr>
<td>negative_sampling_rate</td>
<td>The ratio of negative samples, generated to assist in training the algorithm,</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 ≤ integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0 (off)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>num_classes</td>
<td>The number of classes for classification training. Amazon SageMaker ignores this hyperparameter for regression problems.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: $2 \leq \text{integer} \leq 30$</td>
</tr>
<tr>
<td></td>
<td>Default value: 2</td>
</tr>
<tr>
<td>optimizer</td>
<td>The optimizer type.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: adadelta, adagrad, adam, sgd, or rmsprop.</td>
</tr>
<tr>
<td></td>
<td>- adadelta: A per-dimension learning rate method for gradient descent</td>
</tr>
<tr>
<td></td>
<td>- adagrad: The adaptive gradient algorithm</td>
</tr>
<tr>
<td></td>
<td>- adam: The adaptive moment estimation algorithm</td>
</tr>
<tr>
<td></td>
<td>- sgd: Stochastic gradient descent</td>
</tr>
<tr>
<td></td>
<td>- rmsprop: Root mean square propagation</td>
</tr>
<tr>
<td></td>
<td>Default value: adam</td>
</tr>
<tr>
<td>output_layer</td>
<td>The type of output layer where you specify that the task is regression or classification.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: softmax or mean_squared_error</td>
</tr>
<tr>
<td></td>
<td>- softmax: The Softmax function used for classification.</td>
</tr>
<tr>
<td></td>
<td>- mean_squared_error: The MSE used for regression.</td>
</tr>
<tr>
<td></td>
<td>Default value: softmax</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>tied_token_embedding_weight</td>
<td>Whether to use a shared embedding layer for both encoders. If the inputs to both encoders use the same token-level units, use a shared token embedding layer. For example, for a collection of documents, if one encoder encodes sentences and another encodes whole documents, you can use a shared token embedding layer. That's because both sentences and documents are composed of word tokens from the same vocabulary.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: True or False</td>
</tr>
<tr>
<td></td>
<td>Default value: False</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| token_embedding_storage_type          | The mode of gradient update used during training: when the dense mode is used, the optimizer calculates the full gradient matrix for the token embedding layer even if most rows of the gradient are zero-valued. When sparse mode is used, the optimizer only stores rows of the gradient that are actually being used in the mini-batch. If you want the algorithm to perform lazy gradient updates, which calculate the gradients only in the non-zero rows and which speed up training, specify row_sparse. Setting the value to row_sparse constrains the values available for other hyperparameters, as follows:   

  • The optimizer hyperparameter must be set to adam, adagrad, or sgd. Otherwise, the algorithm throws a CustomerValueError.

  • The algorithm automatically disables bucketing, setting the bucket_width hyperparameter to 0.                                                                                                      |
| Optional                              | Valid values: dense or row_sparse                                                                                                                                                                                                                                                                                                           |
| Default value: dense                  |                                                                                                                                                                                                                                                                                                                                              |
| weight_decay                         | The weight decay parameter used for optimization.                                                                                                                                                                                                                                                                                            |
| Optional                              | Valid values: 0 ≤ float ≤ 10000                                                                                                                                                                                                                                                                                                             |
| Default value: 0 (no decay)           |                                                                                                                                                                                                                                                                                                                                              |

**Tune an Object2Vec Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. For the objective
metric, you use one of the metrics that the algorithm computes. Automatic model tuning searches
the chosen hyperparameters to find the combination of values that result in the model that
optimizes the objective metric.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

**Metrics Computed by the Object2Vec Algorithm**

The Object2Vec algorithm has both classification and regression metrics. The output_layer type
determines which metric you can use for automatic model tuning.

**Regressor Metrics Computed by the Object2Vec Algorithm**

The algorithm reports a mean squared error regressor metric, which is computed during testing
and validation. When tuning the model for regression tasks, choose this metric as the objective.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:mean_squared_error</td>
<td>The Mean Square Error</td>
<td>Minimize</td>
</tr>
<tr>
<td>validation:mean_squared_error</td>
<td>The Mean Square Error</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

**Classification Metrics Computed by the Object2Vec Algorithm**

The Object2Vec algorithm reports accuracy and cross-entropy classification metrics, which are
computed during test and validation. When tuning the model for classification tasks, choose one of
these as the objective.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:accuracy</td>
<td>Accuracy</td>
<td>Maximize</td>
</tr>
<tr>
<td>test:cross_entropy</td>
<td>Cross-entropy</td>
<td>Minimize</td>
</tr>
</tbody>
</table>
## Metric Name

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:accuracy</td>
<td>Accuracy</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:cross_entropy</td>
<td>Cross-entropy</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

### Tunable Object2Vec Hyperparameters

You can tune the following hyperparameters for the Object2Vec algorithm.

<table>
<thead>
<tr>
<th>Hyperparameter Name</th>
<th>Hyperparameter Type</th>
<th>Recommended Ranges and Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>dropout</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 1.0</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 5</td>
</tr>
<tr>
<td>early_stopping_tolerance</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.001, MaxValue: 0.1</td>
</tr>
<tr>
<td>enc_dim</td>
<td>IntegerParameterRange</td>
<td>MinValue: 4, MaxValue: 4096</td>
</tr>
<tr>
<td>enc0_cnn_filter_width</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 5</td>
</tr>
<tr>
<td>enc0_layers</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 4</td>
</tr>
<tr>
<td>Hyperparameter Name</td>
<td>Hyperparameter Type</td>
<td>Recommended Ranges and Values</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-----------------------------</td>
<td>----------------------------------------</td>
</tr>
<tr>
<td>enc0_token_embedding_dim</td>
<td>IntegerParameterRange</td>
<td>MinValue: 5, MaxValue: 300</td>
</tr>
<tr>
<td>enc1_cnn_filter_width</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 5</td>
</tr>
<tr>
<td>enc1_layers</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 4</td>
</tr>
<tr>
<td>enc1_token_embedding_dim</td>
<td>IntegerParameterRange</td>
<td>MinValue: 5, MaxValue: 300</td>
</tr>
<tr>
<td>epochs</td>
<td>IntegerParameterRange</td>
<td>MinValue: 4, MaxValue: 20</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-6, MaxValue: 1.0</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 8192</td>
</tr>
<tr>
<td>mlp_activation</td>
<td>CategoricalParameterRanges</td>
<td>[tanh, relu, linear]</td>
</tr>
<tr>
<td>mlp_dim</td>
<td>IntegerParameterRange</td>
<td>MinValue: 16, MaxValue: 1024</td>
</tr>
<tr>
<td>mlp_layers</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 4</td>
</tr>
</tbody>
</table>
## Hyperparameter Name | Hyperparameter Type | Recommended Ranges and Values
--- | --- | ---
optimizer | CategoricalParameterRanges | [adagrad, adam, rmsprop, sgd, adadelta]
weight_decay | ContinuousParameterRange | MinValue: 0.0, MaxValue: 1.0

### Data Formats for Object2Vec Training

#### Input: JSON Lines Request Format

Content-type: application/jsonlines

```json
[
  {
    "label": 0,
    "in0": [6, 17, 606, 19, 53, 67, 52, 12, 5, 10, 15, 10178, 7, 33, 652, 80, 15, 69, 821, 4],
    "in1": [16, 21, 13, 45, 14, 9, 80, 59, 164, 4]
  },
  {
    "label": 1,
    "in0": [22, 1016, 32, 13, 25, 11, 5, 64, 573, 45, 5, 80, 15, 67, 21, 7, 9, 107, 4],
    "in1": [22, 32, 13, 25, 1016, 573, 3252, 4]
  },
  {
    "label": 1,
    "in0": [774, 14, 21, 206],
    "in1": [21, 366, 125]
  }
]
```

The “in0” and “in1” are the inputs for encoder0 and encoder1, respectively. The same format is valid for both classification and regression problems. For regression, the field "label" can accept real valued inputs.

### Data Formats for Object2Vec Inference

#### GPU optimization: Classification or Regression

Due to GPU memory scarcity, the INFEERENCE_PREFERRED_MODE environment variable can be specified to optimize on whether the classification/regression or the the section called “Output: Encoder Embeddings” inference network is loaded into GPU. If the majority of your inference is for classification or regression, specify INFEERENCE_PREFERRED_MODE=classification. The following is a Batch Transform example of using 4 instances of p3.2xlarge that optimizes for classification/regression inference:

```python
transformer = o2v.transformer(instance_count=4,
```
instance_type="ml.p2.xlarge",
max_concurrent_transforms=2,
max_payload=1,  # 1MB
strategy='MultiRecord',
env={'INFERENCE_PREFERRED_MODE': 'classification'},  # only useful with GPU
output_path=output_s3_path)

Input: Classification or Regression Request Format

Content-type: application/json

{
    "instances": [
        {
            "in0": [6, 17, 606, 19, 53, 67, 52, 12, 5, 10, 15, 10178, 7, 33, 652, 80, 15, 69, 821, 4],
            "in1": [16, 21, 13, 45, 14, 9, 80, 59, 164, 4]},
        {
            "in0": [22, 1016, 32, 13, 25, 11, 5, 64, 573, 45, 5, 80, 15, 67, 21, 7, 9, 107, 4],
            "in1": [22, 32, 13, 25, 1016, 573, 3252, 4]},
        {
            "in0": [774, 14, 21, 206],
            "in1": [21, 366, 125]}
    ]
}

Content-type: application/jsonlines

["in0": [6, 17, 606, 19, 53, 67, 52, 12, 5, 10, 15, 10178, 7, 33, 652, 80, 15, 69, 821, 4],
        "in1": [16, 21, 13, 45, 14, 9, 80, 59, 164, 4}}
["in0": [22, 1016, 32, 13, 25, 11, 5, 64, 573, 45, 5, 80, 15, 67, 21, 7, 9, 107, 4],
        "in1": [22, 32, 13, 25, 1016, 573, 3252, 4]}
["in0": [774, 14, 21, 206],
        "in1": [21, 366, 125}]

For classification problems, the length of the scores vector corresponds to num_classes. For regression problems, the length is 1.

Output: Classification or Regression Response Format

Accept: application/json

{
    "predictions": [
        {
            "scores": [
                0.6533935070037842,
            ]
        }]
}

Use Built-in Algorithms
In both the classification and regression formats, the scores apply to individual labels.

**Encoder Embeddings for Object2Vec**

**GPU optimization: Encoder Embeddings**

An embedding is a mapping from discrete objects, such as words, to vectors of real numbers.

Due to GPU memory scarcity, the INFERENCEPREFERRED_MODE environment variable can be specified to optimize on whether the the section called “Inference Formats: Scoring” or the encoder embedding inference network is loaded into GPU. If the majority of your inference is for encoder embeddings, specify INFERENCEPREFERRED_MODE=embedding. The following is a Batch Transform example of using 4 instances of p3.2xlarge that optimizes for encoder embedding inference:

```python
import o2v
transformer = o2v.transformer(instance_count=4,
                            instance_type="ml.p2.xlarge",
                            max_concurrent_transforms=2,
                            max_payload=1,  # 1MB
                            strategy='MultiRecord',
                            env={"INFERENCEPREFERRED_MODE": 'embedding'},  # only useful with GPU
```

Use Built-in Algorithms
Input: Encoder Embeddings

Content-type: application/json; infer_max_seqlens=<FWD-LENGTH>,<BCK-LENGTH>

Where <FWD-LENGTH> and <BCK-LENGTH> are integers in the range [1,5000] and define the maximum sequence lengths for the forward and backward encoder.

```
{
   "instances": [
      {"in0": [6, 17, 606, 19, 53, 67, 52, 12, 5, 10, 15, 10178, 7, 33, 652, 80, 15, 69, 821, 4]},
      {"in0": [22, 1016, 32, 13, 25, 11, 5, 64, 573, 45, 5, 80, 15, 67, 21, 7, 9, 107, 4]},
      {"in0": [774, 14, 21, 206]}
   ]
}
```

Content-type: application/jsonlines; infer_max_seqlens=<FWD-LENGTH>,<BCK-LENGTH>

Where <FWD-LENGTH> and <BCK-LENGTH> are integers in the range [1,5000] and define the maximum sequence lengths for the forward and backward encoder.

```
{"in0": [6, 17, 606, 19, 53, 67, 52, 12, 5, 10, 15, 10178, 7, 33, 652, 80, 15, 69, 821, 4]}
{"in0": [22, 1016, 32, 13, 25, 11, 5, 64, 573, 45, 5, 80, 15, 67, 21, 7, 9, 107, 4]}
{"in0": [774, 14, 21, 206]}
```

In both of these formats, you specify only one input type: “in0” or “in1.” The inference service then invokes the corresponding encoder and outputs the embeddings for each of the instances.

Output: Encoder Embeddings

Content-type: application/json

```
{
   "predictions": [
      {"embeddings": [0.057368703186511, 0.030703511089086, 0.099890425801277, 0.063688032329082, 0.026327300816774, 0.003637571120634, 0.021305780857801, 0.004316598642617, 0.0, 0.003397724591195, 0.0, 0.000378780066967, 0.0, 0.0, 0.0, 0.007419463712722]},
      {"embeddings": [0.150190666317939, 0.05145975202321, 0.098204270005226, 0.064249359071254, 0.026327300816774, 0.01513972133398, 0.047553978860378, 0.0, 0.0, 0.011533712036907, 0.011472506448626, 0.010696629062294, 0.0, 0.0, 0.0, 0.008508535102009]}
   ]
}
```
The vector length of the embeddings output by the inference service is equal to the value of one of the following hyperparameters that you specify at training time: `enc0_token_embedding_dim`, `enc1_token_embedding_dim`, or `enc_dim`.

**Sequence-to-Sequence Algorithm**

Amazon SageMaker Sequence to Sequence is a supervised learning algorithm where the input is a sequence of tokens (for example, text, audio) and the output generated is another sequence of tokens. Example applications include: machine translation (input a sentence from one language and predict what that sentence would be in another language), text summarization (input a longer string of words and predict a shorter string of words that is a summary), speech-to-text (audio clips converted into output sentences in tokens). Recently, problems in this domain have been successfully modeled with deep neural networks that show a significant performance boost over previous methodologies. Amazon SageMaker seq2seq uses Recurrent Neural Networks (RNNs) and Convolutional Neural Network (CNN) models with attention as encoder-decoder architectures.

**Topics**

- [Input/Output Interface for the Sequence-to-Sequence Algorithm](#)
- [EC2 Instance Recommendation for the Sequence-to-Sequence Algorithm](#)
- [Sequence-to-Sequence Sample Notebooks](#)
- [How Sequence-to-Sequence Works](#)
- [Sequence-to-Sequence Hyperparameters](#)
- [Tune a Sequence-to-Sequence Model](#)

**Input/Output Interface for the Sequence-to-Sequence Algorithm**

**Training**
Amazon SageMaker seq2seq expects data in RecordIO-Protobuf format. However, the tokens are expected as integers, not as floating points, as is usually the case.

A script to convert data from tokenized text files to the protobuf format is included in the seq2seq example notebook. In general, it packs the data into 32-bit integer tensors and generates the necessary vocabulary files, which are needed for metric calculation and inference.

After preprocessing is done, the algorithm can be invoked for training. The algorithm expects three channels:

- **train**: It should contain the training data (for example, the train.rec file generated by the preprocessing script).
- **validation**: It should contain the validation data (for example, the val.rec file generated by the preprocessing script).
- **vocab**: It should contain two vocabulary files (vocab.src.json and vocab.trg.json)

If the algorithm doesn't find data in any of these three channels, training results in an error.

**Inference**

For hosted endpoints, inference supports two data formats. To perform inference using space separated text tokens, use the application/json format. Otherwise, use the recordio-protobuf format to work with the integer encoded data. Both modes support batching of input data. application/json format also allows you to visualize the attention matrix.

- **application/json**: Expects the input in JSON format and returns the output in JSON format. Both content and accept types should be application/json. Each sequence is expected to be a string with whitespace separated tokens. This format is recommended when the number of source sequences in the batch is small. It also supports the following additional configuration options:

  configuration: {attention_matrix: true}: Returns the attention matrix for the particular input sequence.

- **application/x-recordio-protobuf**: Expects the input in recordio-protobuf format and returns the output in recordio-protobuf format. Both content and accept types should be applications/x-recordio-protobuf. For this format, the source sequences must be converted into a list of integers for subsequent protobuf encoding. This format is recommended for bulk inference.
For batch transform, inference supports JSON Lines format. Batch transform expects the input in JSON Lines format and returns the output in JSON Lines format. Both content and accept types should be application/jsonlines. The format for input is as follows:

```json
content-type: application/jsonlines

{"source": "source_sequence_0"}
{"source": "source_sequence_1"}
```

The format for response is as follows:

```json
accept: application/jsonlines

{"target": "predicted_sequence_0"}
{"target": "predicted_sequence_1"}
```

For additional details on how to serialize and deserialize the inputs and outputs to specific formats for inference, see the Sequence-to-Sequence Sample Notebooks.

**EC2 Instance Recommendation for the Sequence-to-Sequence Algorithm**

The Amazon SageMaker seq2seq algorithm only supports on GPU instance types and can only train on a single machine. However, you can use instances with multiple GPUs. The seq2seq algorithm supports P2, P3, G4dn, and G5 GPU instance families.

**Sequence-to-Sequence Sample Notebooks**

For a sample notebook that shows how to use the SageMaker Sequence to Sequence algorithm to train a English-German translation model, see Machine Translation English-German Example Using SageMaker Seq2Seq. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. The topic modeling example notebooks using the NTM algorithms are located in the Introduction to Amazon algorithms section. To open a notebook, click on its Use tab and select Create copy.

**How Sequence-to-Sequence Works**

Typically, a neural network for sequence-to-sequence modeling consists of a few layers, including:
• **An embedding layer.** In this layer, the input matrix, which is input tokens encoded in a sparse way (for example, one-hot encoded) are mapped to a dense feature layer. This is required because a high-dimensional feature vector is more capable of encoding information regarding a particular token (word for text corpora) than a simple one-hot-encoded vector. It is also a standard practice to initialize this embedding layer with a pre-trained word vector like FastText or Glove or to initialize it randomly and learn the parameters during training.

• **An encoder layer.** After the input tokens are mapped into a high-dimensional feature space, the sequence is passed through an encoder layer to compress all the information from the input embedding layer (of the entire sequence) into a fixed-length feature vector. Typically, an encoder is made of RNN-type networks like long short-term memory (LSTM) or gated recurrent units (GRU). ([Colah's blog](https://colah.github.io/posts/2015-08-Understanding-LSTMs/) explains LSTM in a great detail.)

• **A decoder layer.** The decoder layer takes this encoded feature vector and produces the output sequence of tokens. This layer is also usually built with RNN architectures (LSTM and GRU).

The whole model is trained jointly to maximize the probability of the target sequence given the source sequence. This model was first introduced by Sutskever et al. in 2014.

**Attention mechanism.** The disadvantage of an encoder-decoder framework is that model performance decreases as and when the length of the source sequence increases because of the limit of how much information the fixed-length encoded feature vector can contain. To tackle this problem, in 2015, Bahdanau et al. proposed the attention mechanism. In an attention mechanism, the decoder tries to find the location in the encoder sequence where the most important information could be located and uses that information and previously decoded words to predict the next token in the sequence.

For more in details, see the whitepaper [Effective Approaches to Attention-based Neural Machine Translation](https://arxiv.org/abs/1609.04907) by Luong, et al. that explains and simplifies calculations for various attention mechanisms. Additionally, the whitepaper [Google's Neural Machine Translation System: Bridging the Gap between Human and Machine Translation](https://arxiv.org/abs/1609.08144) by Wu, et al. describes Google's architecture for machine translation, which uses skip connections between encoder and decoder layers.

### Sequence-to-Sequence Hyperparameters

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>batch_size</td>
<td>Mini batch size for gradient descent.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 64</td>
</tr>
<tr>
<td>beam_size</td>
<td>Length of the beam for beam search. Used during training for computing bleu and used during inference.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td>bleu_sample_size</td>
<td>Number of instances to pick from validation dataset to decode and compute bleu score during training. Set to -1 to use full validation set (if bleu is chosen as optimized_metric).</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>bucket_width</td>
<td>Returns (source,target) buckets up to ((\text{max_seq_len_source }, \text{max_seq_len_target})). The longer side of the data uses steps of bucket_width while the shorter side uses steps scaled down by the average target/source length ratio. If one sided reaches its maximum length before the other, width of extra buckets on that side is fixed to that side of (\text{max_len}).</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>bucketing_enabled</td>
<td>Set to <code>false</code> to disable bucketing, unroll to maximum length.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: <code>true</code> or <code>false</code></td>
</tr>
<tr>
<td></td>
<td>Default value: <code>true</code></td>
</tr>
<tr>
<td>checkpoint_frequency_num_batches</td>
<td>Checkpoint and evaluate every x batches. This checkpointing hyperparameter is passed to the SageMaker's seq2seq algorithm for early stopping and retrieving the best model. The algorithm's checkpointing runs locally in the algorithm's training container and is not compatible with SageMaker checkpointing. The algorithm temporarily saves checkpoints to a local path and stores the best model artifact to the model output path in S3 after the training job has stopped.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1000</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| checkpoint_threshold        | Maximum number of checkpoints model is allowed to not improve in optimized_metric on validation dataset before training is stopped. This checkpointing hyperparameter is passed to the SageMaker's seq2seq algorithm for early stopping and retrieving the best model. The algorithm's checkpointing runs locally in the algorithm's training container and is not compatible with SageMaker checkpointing. The algorithm temporarily saves checkpoints to a local path and stores the best model artifact to the model output path in S3 after the training job has stopped.  

**Optional**

Valid values: positive integer

Default value: 3                                                                                                                                                                                                                                           |
<p>| clip_gradient               | Clip absolute gradient values greater than this. Set to negative to disable.                                                                                                                                                                                                                                               |
|                             | <strong>Optional</strong>                                                                                                                                                                                                                                                                                                                                                   |
|                             | Valid values: float                                                                                                                                                                                                                                                                                  |
|                             | Default value: 1                                                                                                                                                                                                                                                                                   |
| cnn_activation_type         | The cnn activation type to be used.                                                                                                                                                                                                                                                                 |
|                             | <strong>Optional</strong>                                                                                                                                                                                                                                                                                      |
|                             | Valid values: String. One of glu, relu, softrelu, sigmoid, or tanh.                                                                                                                                                                                                                                 |
|                             | Default value: glu                                                                                                                                                                                                                                                                                 |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>cnn_hidden_dropout</td>
<td>Dropout probability for dropout between convolutional layers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range in [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>cnn_kernel_width_decoder</td>
<td>Kernel width for the cnn decoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td>cnn_kernel_width_encoder</td>
<td>Kernel width for the cnn encoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 3</td>
</tr>
<tr>
<td>cnn_num_hidden</td>
<td>Number of cnn hidden units for encoder and decoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 512</td>
</tr>
<tr>
<td>decoder_type</td>
<td>Decoder type.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either rnn or cnn.</td>
</tr>
<tr>
<td></td>
<td>Default value: rnn</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>embed_dropout_source</td>
<td>Dropout probability for source side embeddings.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range in [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>embed_dropout_target</td>
<td>Dropout probability for target side embeddings.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range in [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>encoder_type</td>
<td>Encoder type. The <em>rnn</em> architecture is based on attention mechanism by Bahdanau et al. and <em>cnn</em> architecture is based on Gehring et al.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either <em>rnn</em> or <em>cnn</em>.</td>
</tr>
<tr>
<td></td>
<td>Default value: <em>rnn</em></td>
</tr>
<tr>
<td>fixed_rate_lr_half_life</td>
<td>Half life for learning rate in terms of number of checkpoints for <em>fixed_rate_</em> * schedulers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>learning_rate</td>
<td>Initial learning rate.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0003</td>
</tr>
<tr>
<td>loss_type</td>
<td>Loss function for training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. cross-entropy</td>
</tr>
<tr>
<td></td>
<td>Default value: cross-entropy</td>
</tr>
<tr>
<td>lr_scheduler_type</td>
<td>Learning rate scheduler type. plateau_reduce means reduce the learning rate whenever optimized _metric on validation_accuracy plateaus. inv_t is inverse time decay. learning_rate / (1+decay_rate *t)</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of plateau_reduce, fixed_rate_inv_t, or fixed_rate_inv_sqrt_t.</td>
</tr>
<tr>
<td></td>
<td>Default value: plateau_reduce</td>
</tr>
<tr>
<td>max_num_batches</td>
<td>Maximum number of updates/batches to process. -1 for infinite.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: -1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| max_num_epochs       | Maximum number of epochs to pass through training data before fitting is stopped. Training continues until this number of epochs even if validation accuracy is not improving if this parameter is passed. Ignored if not passed. **Optional**

Valid values: Positive integer and less than or equal to max_num_epochs.

Default value: none |
| max_seq_len_source   | Maximum length for the source sequence length. Sequences longer than this length are truncated to this length. **Optional**

Valid values: positive integer

Default value: 100 |
| max_seq_len_target   | Maximum length for the target sequence length. Sequences longer than this length are truncated to this length. **Optional**

Valid values: positive integer

Default value: 100 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>min_num_epochs</td>
<td>Minimum number of epochs the training must run before it is stopped via early_stopping conditions. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>momentum</td>
<td>Momentum constant used for sgd. Don't pass this parameter if you are using adam or rmsprop. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: none</td>
</tr>
<tr>
<td>num_embed_source</td>
<td>Embedding size for source tokens. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 512</td>
</tr>
<tr>
<td>num_embed_target</td>
<td>Embedding size for target tokens. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 512</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>num_layers_decoder</td>
<td>Number of layers for Decoder <em>rnn</em> or <em>cnn</em>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>num_layers_encoder</td>
<td>Number of layers for Encoder <em>rnn</em> or <em>cnn</em>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
</tr>
<tr>
<td>optimized_metric</td>
<td>Metrics to optimize with early stopping.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of <em>perplexity</em>, <em>accuracy</em>, or <em>bleu</em>.</td>
</tr>
<tr>
<td></td>
<td>Default value: <em>perplexity</em></td>
</tr>
<tr>
<td>optimizer_type</td>
<td>Optimizer to choose from.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of <em>adam</em>, <em>sgd</em>, or <em>rmsprop</em>.</td>
</tr>
<tr>
<td></td>
<td>Default value: <em>adam</em></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>plateau_reduce_lr_factor</td>
<td>Factor to multiply learning rate with (for plateau reduce ).</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.5</td>
</tr>
<tr>
<td>plateau_reduce_lr_threshold</td>
<td>For plateau_reduce scheduler, multiply learning rate with reduce factor if optimized_metric didn't improve for this many checkpoints.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 3</td>
</tr>
<tr>
<td>rnn_attention_in_upper_layers</td>
<td>Pass the attention to upper layers of rnn, like Google NMT paper. Only applicable if more than one layer is used.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: boolean (true or false)</td>
</tr>
<tr>
<td></td>
<td>Default value: true</td>
</tr>
<tr>
<td>rnn_attention_num_hidden</td>
<td>Number of hidden units for attention layers. defaults to rnn_num_hidden.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: rnn_num_hidden</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>rnn_attention_type</td>
<td>Attention model for encoders. mlp refers to concat and bilinear refers to general from the Luong et al. paper.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of dot, fixed, mlp, or bilinear.</td>
</tr>
<tr>
<td></td>
<td>Default value: mlp</td>
</tr>
<tr>
<td>rnn_cell_type</td>
<td>Specific type of rnn architecture.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either lstm or gru.</td>
</tr>
<tr>
<td></td>
<td>Default value: lstm</td>
</tr>
<tr>
<td>rnn_decoder_state_init</td>
<td>How to initialize rnn decoder states from encoders.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of last, avg, or zero.</td>
</tr>
<tr>
<td></td>
<td>Default value: last</td>
</tr>
<tr>
<td>rnn_first_residual_layer</td>
<td>First rnn layer to have a residual connection, only applicable if number of layers in encoder or decoder is more than 1.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 2</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><code>rnn_num_hidden</code></td>
<td>The number of <em>rnn</em> hidden units for encoder and decoder. This must be a multiple of 2 because the algorithm uses bi-directional Long Term Short Term Memory (LSTM) by default.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive even integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 1024</td>
</tr>
<tr>
<td><code>rnn_residual_connections</code></td>
<td>Add residual connection to stacked <em>rnn</em>. Number of layers should be more than 1.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: boolean (true or false)</td>
</tr>
<tr>
<td></td>
<td>Default value: false</td>
</tr>
<tr>
<td><code>rnn_decoder_hidden_dropout</code></td>
<td>Dropout probability for hidden state that combines the context with the <em>rnn</em> hidden state in the decoder.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range in [0,1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td><code>training_metric</code></td>
<td>Metrics to track on training on validation data.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either perplexity or accuracy.</td>
</tr>
<tr>
<td></td>
<td>Default value: perplexity</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>weight_decay</td>
<td>Weight decay constant.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>weight_init_scale</td>
<td>Weight initialization scale (for uniform and xavier initialization).</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 2.34</td>
</tr>
<tr>
<td>weight_init_type</td>
<td>Type of weight initialization.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: String. Either uniform or xavier.</td>
</tr>
<tr>
<td></td>
<td>Default value: xavier</td>
</tr>
<tr>
<td>xavier_factor_type</td>
<td>Xavier factor type.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: String. One of in, out, or avg.</td>
</tr>
<tr>
<td></td>
<td>Default value: in</td>
</tr>
</tbody>
</table>

**Tune a Sequence-to-Sequence Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches
the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

**Metrics Computed by the Sequence-to-Sequence Algorithm**

The sequence to sequence algorithm reports three metrics that are computed during training. Choose one of them as an objective to optimize when tuning the hyperparameter values.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:accuracy</td>
<td>Accuracy computed on the validation dataset.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:bleu</td>
<td><strong>Bleu</strong> score computed on the validation dataset. Because BLEU computation is expensive, you can choose to compute BLEU on a random subsample of the validation dataset to speed up the overall training process. Use the bleu_sample_size parameter to specify the subsample.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:perplexity</td>
<td><strong>Perplexity</strong>, is a loss function computed on the validation dataset. Perplexity measures the cross-entropy between an empirical sample and the distribution predicted by a model and so provides a measure of how well a model predicts the sample values, Models that are good at predicting a sample have a low perplexity.</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

**Tunable Sequence-to-Sequence Hyperparameters**

You can tune the following hyperparameters for the SageMaker Sequence to Sequence algorithm. The hyperparameters that have the greatest impact on sequence to sequence objective
The metrics are: batch_size, optimizer_type, learning_rate, num_layers_encoder, and num_layers_decoder.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_layers_encoder</td>
<td>IntegerParameterRange</td>
<td>[1-10]</td>
</tr>
<tr>
<td>num_layers_decoder</td>
<td>IntegerParameterRange</td>
<td>[1-10]</td>
</tr>
<tr>
<td>batch_size</td>
<td>CategoricalParameterRange</td>
<td>[16,32,64,128,256, 512,1024,2048]</td>
</tr>
<tr>
<td>optimizer_type</td>
<td>CategoricalParameterRange</td>
<td>['adam', 'sgd', 'rmsprop']</td>
</tr>
<tr>
<td>weight_init_type</td>
<td>CategoricalParameterRange</td>
<td>['xavier', 'uniform']</td>
</tr>
<tr>
<td>weight_init_scale</td>
<td>ContinuousParameterRange</td>
<td>For the xavier type: MinValue: 2.0, MaxValue: 3.0 For the uniform type: MinValue: -1.0, MaxValue: 1.0</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.00005, MaxValue: 0.2</td>
</tr>
<tr>
<td>weight_decay</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.1</td>
</tr>
<tr>
<td>momentum</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.5, MaxValue: 0.9</td>
</tr>
<tr>
<td>clip_gradient</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1.0, MaxValue: 5.0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
</tr>
<tr>
<td>------------------------------</td>
<td>-----------------------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><code>rnn_num_hidden</code></td>
<td>CategoricalParameterRange</td>
<td>Applicable only to recurrent neural networks (RNNs). [128, 256, 512, 1024, 2048]</td>
</tr>
<tr>
<td><code>cnn_num_hidden</code></td>
<td>CategoricalParameterRange</td>
<td>Applicable only to convolutional neural networks (CNNs). [128, 256, 512, 1024, 2048]</td>
</tr>
<tr>
<td><code>num_embed_source</code></td>
<td>IntegerParameterRange</td>
<td>[256-512]</td>
</tr>
<tr>
<td><code>num_embed_target</code></td>
<td>IntegerParameterRange</td>
<td>[256-512]</td>
</tr>
<tr>
<td><code>embed_dropout_source</code></td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.5</td>
</tr>
<tr>
<td><code>embed_dropout_target</code></td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.5</td>
</tr>
<tr>
<td><code>rnn_decoder_hidden_dropout</code></td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.5</td>
</tr>
<tr>
<td><code>cnn_hidden_dropout</code></td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.5</td>
</tr>
<tr>
<td><code>lr_scheduler_type</code></td>
<td>CategoricalParameterRange</td>
<td>['plateau_reduce', 'fixed_rate_inv_t', 'fixed_rate_inv_sqrt_t']</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-----------------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>plateau_reduce_lr_factor</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.1, MaxValue: 0.5</td>
</tr>
<tr>
<td>plateau_reduce_lr_threshold</td>
<td>IntegerParameterRange</td>
<td>[1-5]</td>
</tr>
<tr>
<td>fixed_rate_lr_half_life</td>
<td>IntegerParameterRange</td>
<td>[10-30]</td>
</tr>
</tbody>
</table>

**Text Classification - TensorFlow**

The Amazon SageMaker Text Classification - TensorFlow algorithm is a supervised learning algorithm that supports transfer learning with many pretrained models from the TensorFlow Hub. Use transfer learning to fine-tune one of the available pretrained models on your own dataset, even if a large amount of text data is not available. The text classification algorithm takes a text string as input and outputs a probability for each of the class labels. Training datasets must be in CSV format.

**Topics**

- [How to use the SageMaker Text Classification - TensorFlow algorithm](#)
- [Input and output interface for the Text Classification - TensorFlow algorithm](#)
- [Amazon EC2 instance recommendation for the Text Classification - TensorFlow algorithm](#)
- [Text Classification - TensorFlow sample notebooks](#)
- [How Text Classification - TensorFlow Works](#)
- [TensorFlow Hub Models](#)
- [Text Classification - TensorFlow Hyperparameters](#)
- [Tune a Text Classification - TensorFlow model](#)
How to use the SageMaker Text Classification - TensorFlow algorithm

You can use Text Classification - TensorFlow as an Amazon SageMaker built-in algorithm. The following section describes how to use Text Classification - TensorFlow with the SageMaker Python SDK. For information on how to use Text Classification - TensorFlow from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

The Text Classification - TensorFlow algorithm supports transfer learning using any of the compatible pretrained TensorFlow models. For a list of all available pretrained models, see TensorFlow Hub Models. Every pretrained model has a unique model_id. The following example uses BERT Base Uncased (model_id: tensorflow-tc-bert-en-uncased-L-12-H-768-A-12-2) to fine-tune on a custom dataset. The pretrained models are all pre-downloaded from the TensorFlow Hub and stored in Amazon S3 buckets so that training jobs can run in network isolation. Use these pre-generated model training artifacts to construct a SageMaker Estimator.

First, retrieve the Docker image URI, training script URI, and pretrained model URI. Then, change the hyperparameters as you see fit. You can see a Python dictionary of all available hyperparameters and their default values with hyperparameters.retrieve_default. For more information, see Text Classification - TensorFlow Hyperparameters. Use these values to construct a SageMaker Estimator.

Note

Default hyperparameter values are different for different models. For example, for larger models, the default batch size is smaller.

This example uses the SST2 dataset, which contains positive and negative movie reviews. We pre-downloaded the dataset and made it available with Amazon S3. To fine-tune your model, call .fit using the Amazon S3 location of your training dataset. Any S3 bucket used in a notebook must be in the same AWS Region as the notebook instance that accesses it.

```python
from sagemaker import image_uris, model_uris, script_uris, hyperparameters
from sagemaker.estimator import Estimator

model_id, model_version = "tensorflow-tc-bert-en-uncased-L-12-H-768-A-12-2", "*
training_instance_type = "ml.p3.2xlarge"

# Retrieve the Docker image
```
train_image_uri =
    image_uris.retrieve(model_id=model_id, model_version=model_version, image_scope="training", instance_type=training_instance_type, region=None, framework=None)

# Retrieve the training script
train_source_uri = script_uris.retrieve(model_id=model_id, model_version=model_version, script_scope="training")

# Retrieve the pretrained model tarball for transfer learning
train_model_uri = model_uris.retrieve(model_id=model_id, model_version=model_version, model_scope="training")

# Retrieve the default hyperparameters for fine-tuning the model
hyperparameters = hyperparameters.retrieve_default(model_id=model_id, model_version=model_version)

# [Optional] Override default hyperparameters with custom values
hyperparameters["epochs"] = "5"

# Sample training data is available in this bucket
training_data_bucket = f"jumpstart-cache-prod-{aws_region}"
training_data_prefix = "training-datasets/SST2/"

training_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}"

output_bucket = sess.default_bucket()
output_prefix = "jumpstart-example-tc-training"
s3_output_location = f"s3://{output_bucket}/{output_prefix}/output"

# Create an Estimator instance
tf_tc_estimator = Estimator(
    role=aws_role,
    image_uri=train_image_uri,
    source_dir=train_source_uri,
    model_uri=train_model_uri,
    entry_point="transfer_learning.py",
    instance_count=1,
    instance_type=training_instance_type,
    max_run=360000,
    hyperparameters=hyperparameters,
    output_path=s3_output_location,
)

# Launch a training job
For more information about how to use the SageMaker Text Classification - TensorFlow algorithm for transfer learning on a custom dataset, see the [Introduction to JumpStart - Text Classification](#) notebook.

### Input and output interface for the Text Classification - TensorFlow algorithm

Each of the pretrained models listed in TensorFlow Hub Models can be fine-tuned to any dataset made up of text sentences with any number of classes. The pretrained model attaches a classification layer to the Text Embedding model and initializes the layer parameters to random values. The output dimension of the classification layer is determined based on the number of classes detected in the input data.

Be mindful of how to format your training data for input to the Text Classification - TensorFlow model.

- **Training data input format:** A directory containing a data.csv file. Each row of the first column should have integer class labels between 0 and the number of classes. Each row of the second column should have the corresponding text data.

The following is an example of an input CSV file. Note that the file should not have any header. The file should be hosted in an Amazon S3 bucket with a path similar to the following: s3://bucket_name/input_directory/. Note that the trailing / is required.

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>hide new secretions from the parental units</td>
</tr>
<tr>
<td>0</td>
<td>contains no wit, only labored gags</td>
</tr>
<tr>
<td>1</td>
<td>that loves its characters and communicates something rather beautiful about human nature</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>

### Incremental training

You can seed the training of a new model with artifacts from a model that you trained previously with SageMaker. Incremental training saves training time when you want to train a new model with the same or similar data.
You can only seed a SageMaker Text Classification - TensorFlow model with another Text Classification - TensorFlow model trained in SageMaker.

You can use any dataset for incremental training, as long as the set of classes remains the same. The incremental training step is similar to the fine-tuning step, but instead of starting with a pretrained model, you start with an existing fine-tuned model.

For more information on using incremental training with the SageMaker Text Classification - TensorFlow algorithm, see the Introduction to JumpStart - Text Classification sample notebook.

**Inference with the Text Classification - TensorFlow algorithm**

You can host the fine-tuned model that results from your TensorFlow Text Classification training for inference. Any raw text formats for inference must be content type `application/x-text`.

Running inference results in probability values, class labels for all classes, and the predicted label corresponding to the class index with the highest probability encoded in JSON format. The Text Classification - TensorFlow model processes a single string per request and outputs only one line. The following is an example of a JSON format response:

```json
accept: application/json;verbose

"probabilities": [prob_0, prob_1, prob_2, ...],
"labels": [label_0, label_1, label_2, ...],
"predicted_label": predicted_label
```

If accept is set to `application/json`, then the model only outputs probabilities.

**Amazon EC2 instance recommendation for the Text Classification - TensorFlow algorithm**

The Text Classification - TensorFlow algorithm supports all CPU and GPU instances for training, including:

- ml.p2.xlarge
- ml.p2.16xlarge
- ml.p3.2xlarge
We recommend GPU instances with more memory for training with large batch sizes. Both CPU (such as M5) and GPU (P2, P3, G4dn, or G5) instances can be used for inference. For a comprehensive list of SageMaker training and inference instances across AWS Regions, see Amazon SageMaker Pricing.

Text Classification - TensorFlow sample notebooks

For more information about how to use the SageMaker Text Classification - TensorFlow algorithm for transfer learning on a custom dataset, see the Introduction to JumpStart - Text Classification notebook.

For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

How Text Classification - TensorFlow Works

The Text Classification - TensorFlow algorithm takes text as classifies it into one of the output class labels. Deep learning networks such as BERT are highly accurate for text classification. There are also deep learning networks that are trained on large text datasets, such as TextNet, which has more than 11 million texts with about 11,000 categories. After a network is trained with TextNet data, you can then fine-tune the network on a dataset with a particular focus to perform more specific text classification tasks. The Amazon SageMaker Text Classification - TensorFlow algorithm supports transfer learning on many pretrained models that are available in the TensorFlow Hub.

According to the number of class labels in your training data, a text classification layer is attached to the pretrained TensorFlow model of your choice. The classification layer consists of a dropout layer, a dense layer, and a fully connected layer with 2-norm regularization, and is initialized with random weights. You can change the hyperparameter values for the dropout rate of the dropout layer and the L2 regularization factor for the dense layer.
You can fine-tune either the entire network (including the pretrained model) or only the top classification layer on new training data. With this method of transfer learning, training with smaller datasets is possible.

**TensorFlow Hub Models**

The following pretrained models are available to use for transfer learning with the Text Classification - TensorFlow algorithm.

The following models vary significantly in size, number of model parameters, training time, and inference latency for any given dataset. The best model for your use case depends on the complexity of your fine-tuning dataset and any requirements that you have on training time, inference latency, or model accuracy.

<table>
<thead>
<tr>
<th>Model Name</th>
<th>model_id</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>BERT Base Uncased</td>
<td>tensorflow-tc-bert-en-uncased-L-12-H-768-A-12-2</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>BERT Base Cased</td>
<td>tensorflow-tc-bert-en-cased-L-12-H-768-A-12-2</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>BERT Base Multilingual Cased</td>
<td>tensorflow-tc-bert-multi-cased-L-12-H-768-A-12-2</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Model Name</td>
<td>model_id</td>
<td>Source</td>
</tr>
<tr>
<td>----------------------------</td>
<td>----------------------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>Small BERT L-4_H-256_A-4</td>
<td>tensorflow-tc-small-bert-bert-en-uncased-L-4-H-256-A-4</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-4_H-768_A-12</td>
<td>tensorflow-tc-small-bert-bert-en-uncased-L-4-H-768-A-12</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-6_H-256_A-4</td>
<td>tensorflow-tc-small-bert-bert-en-uncased-L-6-H-256-A-4</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-6_H-768_A-12</td>
<td>tensorflow-tc-small-bert-bert-en-uncased-L-6-H-768-A-12</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Model Name</td>
<td>model_id</td>
<td>Source</td>
</tr>
<tr>
<td>----------------------------</td>
<td>---------------------------------------------------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>Small BERT L-8_H-256_A-4</td>
<td>tensorflow-tc-smallbert-bert-en-uncased-L-8-H-256-A-4</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-8_H-768_A-12</td>
<td>tensorflow-tc-smallbert-bert-en-uncased-L-8-H-768-A-12</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-10_H-256_A-4</td>
<td>tensorflow-tc-smallbert-bert-en-uncased-L-10-H-256-A-4</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-10_H-768_A-12</td>
<td>tensorflow-tc-smallbert-bert-en-uncased-L-10-H-768-A-12</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Model Name</td>
<td>model_id</td>
<td>Source</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>-----------------------------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>Small BERT L-12_H-256_A-4</td>
<td>tensorflow-tc-small-bert-bert-en-uncased-L-12-H-256-A-4</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>Small BERT L-12_H-768_A-12</td>
<td>tensorflow-tc-small-bert-bert-en-uncased-L-12-H-768-A-12</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>BERT Large Cased</td>
<td>tensorflow-tc-bert-en-cased-L-24-H-1024-A-16-2</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>ALBERT Base</td>
<td>tensorflow-tc-albert-en-base</td>
<td>TensorFlow Hub link</td>
</tr>
<tr>
<td>ELECTRA Small++</td>
<td>tensorflow-tc-electra-small-1</td>
<td>TensorFlow Hub link</td>
</tr>
</tbody>
</table>
## Text Classification - TensorFlow Hyperparameters

Hyperparameters are parameters that are set before a machine learning model begins learning. The following hyperparameters are supported by the Amazon SageMaker built-in Object Detection - TensorFlow algorithm. See [Tune a Text Classification - TensorFlow model](#) for information on hyperparameter tuning.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>batch_size</td>
<td>The batch size for training. For training on instances with multiple GPUs, this batch size is used across the GPUs. Valid values: positive integer. Default value: 32.</td>
</tr>
<tr>
<td>beta_1</td>
<td>The beta1 for the &quot;adam&quot; and &quot;adamw&quot; optimizers. Represents the exponential decay rate for the first moment estimates. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>beta_2</td>
<td>The beta2 for the &quot;adam&quot; and &quot;adamw&quot; optimizers. Represents the exponential decay rate for the second moment estimates. Ignored for other optimizers.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.999.</td>
</tr>
<tr>
<td>dropout_rate</td>
<td>The dropout rate for the dropout layer in the top classification layer. Used only when <code>reinitialize_top_layer</code> is set to &quot;True&quot;.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.2</td>
</tr>
<tr>
<td>early_stopping</td>
<td>Set to &quot;True&quot; to use early stopping logic during training. If &quot;False&quot;, early stopping is not used.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, either: (&quot;True&quot; or &quot;False&quot;).</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>early_stopping_min_delta</td>
<td>The minimum change needed to qualify as an improvement. An absolute change less than the value of <code>early_stopping_min_delta</code> does not qualify as improvement. Used only when <code>early_stopping</code> is set to &quot;True&quot;.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>The number of epochs to continue training with no improvement. Used only when early_stopping is set to &quot;True&quot;. Valid values: positive integer. Default value: 5.</td>
</tr>
<tr>
<td>epochs</td>
<td>The number of training epochs. Valid values: positive integer. Default value: 10.</td>
</tr>
<tr>
<td>epsilon</td>
<td>The epsilon for &quot;adam&quot;, &quot;rmsprop&quot;, &quot;adadelta&quot;, and &quot;adagrad&quot; optimizers. Usually set to a small value to avoid division by 0. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 1e-7.</td>
</tr>
<tr>
<td>initial_accumulator_value</td>
<td>The starting value for the accumulators, or the per-parameter momentum values, for the &quot;adagrad&quot; optimizer. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.0001.</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The optimizer learning rate. Valid values: float, range: [0.0, 1.0]. Default value: 0.001.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| momentum           | The momentum for the "sgd" and "nesterov" optimizers. Ignored for other optimizers.  
                      | Valid values: float, range: [0.0, 1.0].                                                
                      | Default value: 0.9.                                                                    |
| optimizer          | The optimizer type. For more information, see Optimizers in the TensorFlow documentation.  
                      | Valid values: string, any of the following: ("adamw", "adam",  
                      | "sgd", "nesterov", "rmsprop", "adagrad", "adadelta").  
                      | Default value: "adam".                                                                  |
| regularizers_l2    | The L2 regularization factor for the dense layer in the classification layer. Used only when reinitialize_top_layer is set to "True".  
                      | Valid values: float, range: [0.0, 1.0].                                                  
                      | Default value: 0.0001.                                                                  |
| reinitialize_top_layer | If set to "Auto", the top classification layer parameters are re-initialized during fine-tuning. For incremental training, top classification layer parameters are not re-initialized unless set to "True".  
                      | Valid values: string, any of the following: ("Auto", "True" or "False").                  
<pre><code>                  | Default value: &quot;Auto&quot;.                                                                 |
</code></pre>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\rho$</td>
<td>The discounting factor for the gradient of the &quot;adadelta&quot; and &quot;rmsprop&quot; optimizers. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.95.</td>
</tr>
<tr>
<td>train_only_on_top_layer</td>
<td>If &quot;True&quot;, only the top classification layer parameters are fine-tuned. If &quot;False&quot;, all model parameters are fine-tuned. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>validation_split_ratio</td>
<td>The fraction of training data to randomly split to create validation data. Only used if validation data is not provided through the validation channel. Valid values: float, range: [0.0, 1.0]. Default value: 0.2.</td>
</tr>
<tr>
<td>warmup_steps_fraction</td>
<td>The fraction of the total number of gradient update steps, where the learning rate increases from 0 to the initial learning rate as a warm up. Only used with the adamw optimizer. Valid values: float, range: [0.0, 1.0]. Default value: 0.1.</td>
</tr>
</tbody>
</table>

**Tune a Text Classification - TensorFlow model**

Automatic model tuning, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.
For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics computed by the Text Classification - TensorFlow algorithm**

Refer to the following chart to find which metrics are computed by the Text Classification - TensorFlow algorithm.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
<th>Regex Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:accuracy</td>
<td>The ratio of the number of correct predictions to the total number of predictions made.</td>
<td>Maximize</td>
<td>val_accuracy=([0-9.]+)</td>
</tr>
</tbody>
</table>

**Tunable Text Classification - TensorFlow hyperparameters**

Tune a text classification model with the following hyperparameters. The hyperparameters that have the greatest impact on text classification objective metrics are: `batch_size`, `learning_rate`, and `optimizer`. Tune the optimizer-related hyperparameters, such as `momentum`, `regularizers_l2`, `beta_1`, `beta_2`, and `eps` based on the selected optimizer. For example, use `beta_1` and `beta_2` only when `adamw` or `adam` is the optimizer.

For more information about which hyperparameters are used for each optimizer, see [Text Classification - TensorFlow Hyperparameters](#).

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 4, MaxValue: 128</td>
</tr>
<tr>
<td>beta_1</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
<tr>
<td>beta_2</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
</tbody>
</table>
### Parameter Name | Parameter Type | Recommended Ranges
---|---|---
eps | ContinuousParameterRanges | MinValue: 1e-8, MaxValue: 1.0
learning_rate | ContinuousParameterRanges | MinValue: 1e-6, MaxValue: 0.5
momentum | ContinuousParameterRanges | MinValue: 0.0, MaxValue: 0.999
optimizer | CategoricalParameterRanges | ['adamw', 'adam', 'sgd', 'rmsprop', 'nesterov', 'adagrad', 'adadelta']
regularizers_l2 | ContinuousParameterRanges | MinValue: 0.0, MaxValue: 0.999
train_only_on_top_layer | CategoricalParameterRanges | ['True', 'False']

### Built-in SageMaker Algorithms for Time-Series Data

SageMaker provides algorithms that are tailored to the analysis of time-series data for forecasting product demand, server loads, webpage requests, and more.

- **DeepAR Forecasting Algorithm**—a supervised learning algorithm for forecasting scalar (one-dimensional) time series using recurrent neural networks (RNN).
DeepAR Forecasting Algorithm

The Amazon SageMaker DeepAR forecasting algorithm is a supervised learning algorithm for forecasting scalar (one-dimensional) time series using recurrent neural networks (RNN). Classical forecasting methods, such as autoregressive integrated moving average (ARIMA) or exponential smoothing (ETS), fit a single model to each individual time series. They then use that model to extrapolate the time series into the future.

In many applications, however, you have many similar time series across a set of cross-sectional units. For example, you might have time series groupings for demand for different products, server loads, and requests for webpages. For this type of application, you can benefit from training a single model jointly over all of the time series. DeepAR takes this approach. When your dataset contains hundreds of related time series, DeepAR outperforms the standard ARIMA and ETS methods. You can also use the trained model to generate forecasts for new time series that are similar to the ones it has been trained on.

The training input for the DeepAR algorithm is one or, preferably, more target time series that have been generated by the same process or similar processes. Based on this input dataset, the algorithm trains a model that learns an approximation of this process/processes and uses it to predict how the target time series evolves. Each target time series can be optionally associated with a vector of static (time-independent) categorical features provided by the cat field and a vector of dynamic (time-dependent) time series provided by the dynamic_feat field. SageMaker trains the DeepAR model by randomly sampling training examples from each target time series in the training dataset. Each training example consists of a pair of adjacent context and prediction windows with fixed predefined lengths. To control how far in the past the network can see, use the context_length hyperparameter. To control how far in the future predictions can be made, use the prediction_length hyperparameter. For more information, see How the DeepAR Algorithm Works.
Input/Output Interface for the DeepAR Algorithm

DeepAR supports two data channels. The required **train** channel describes the training dataset. The optional **test** channel describes a dataset that the algorithm uses to evaluate model accuracy after training. You can provide training and test datasets in **JSON Lines** format. Files can be in gzip or **Parquet** file format.

When specifying the paths for the training and test data, you can specify a single file or a directory that contains multiple files, which can be stored in subdirectories. If you specify a directory, DeepAR uses all files in the directory as inputs for the corresponding channel, except those that start with a period (.) and those named _SUCCESS. This ensures that you can directly use output folders produced by Spark jobs as input channels for your DeepAR training jobs.

By default, the DeepAR model determines the input format from the file extension (.json, .json.gz, or .parquet) in the specified input path. If the path does not end in one of these extensions, you must explicitly specify the format in the SDK for Python. Use the content_type parameter of the **s3_input** class.

The records in your input files should contain the following fields:

- **start**—A string with the format **YYYY-MM-DD HH:MM:SS**. The start timestamp can't contain time zone information.
- **target**—An array of floating-point values or integers that represent the time series. You can encode missing values as null literals, or as "NaN" strings in JSON, or as nan floating-point values in Parquet.
• dynamic_feat (optional)—An array of arrays of floating-point values or integers that represents the vector of custom feature time series (dynamic features). If you set this field, all records must have the same number of inner arrays (the same number of feature time series). In addition, each inner array must be the same length as the associated target value plus prediction_length. Missing values are not supported in the features. For example, if target time series represents the demand of different products, an associated dynamic_feat might be a boolean time-series which indicates whether a promotion was applied (1) to the particular product or not (0):

```json
{"start": ..., "target": [1, 5, 10, 2], "dynamic_feat": [[0, 1, 1, 0]]}
```

• cat (optional)—An array of categorical features that can be used to encode the groups that the record belongs to. Categorical features must be encoded as a 0-based sequence of positive integers. For example, the categorical domain \{R, G, B\} can be encoded as \{0, 1, 2\}. All values from each categorical domain must be represented in the training dataset. That's because the DeepAR algorithm can forecast only for categories that have been observed during training. And, each categorical feature is embedded in a low-dimensional space whose dimensionality is controlled by the embedding_dimension hyperparameter. For more information, see DeepAR Hyperparameters.

If you use a JSON file, it must be in JSON Lines format. For example:

```json
{"start": "2009-11-01 00:00:00", "target": [4.3, "NaN", 5.1, ...], "cat": [0, 1], "dynamic_feat": [[1.1, 1.2, 0.5, ...]]}
{"start": "2012-01-30 00:00:00", "target": [1.0, -5.0, ...], "cat": [2, 3], "dynamic_feat": [[1.1, 2.05, ...]]}
{"start": "1999-01-30 00:00:00", "target": [2.0, 1.0], "cat": [1, 4], "dynamic_feat": [[1.3, 0.4]]}
```

In this example, each time series has two associated categorical features and one time series features.

For Parquet, you use the same three fields as columns. In addition, "start" can be the datetime type. You can compress Parquet files using gzip (gzip) or the Snappy compression library (snappy).

If the algorithm is trained without cat and dynamic_feat fields, it learns a "global" model, that is a model that is agnostic to the specific identity of the target time series at inference time and is conditioned only on its shape.
If the model is conditioned on the `cat` and `dynamic_feat` feature data provided for each time series, the prediction will probably be influenced by the character of time series with the corresponding `cat` features. For example, if the target time series represents the demand of clothing items, you can associate a two-dimensional `cat` vector that encodes the type of item (e.g. 0 = shoes, 1 = dress) in the first component and the color of an item (e.g. 0 = red, 1 = blue) in the second component. A sample input would look as follows:

```json
{ "start": ..., "target": ..., "cat": [0, 0], ... } # red shoes
{ "start": ..., "target": ..., "cat": [1, 1], ... } # blue dress
```

At inference time, you can request predictions for targets with `cat` values that are combinations of the `cat` values observed in the training data, for example:

```json
{ "start": ..., "target": ..., "cat": [0, 1], ... } # blue shoes
{ "start": ..., "target": ..., "cat": [1, 0], ... } # red dress
```

The following guidelines apply to training data:

- The start time and length of the time series can differ. For example, in marketing, products often enter a retail catalog at different dates, so their start dates naturally differ. But all series must have the same frequency, number of categorical features, and number of dynamic features.
- Shuffle the training file with respect to the position of the time series in the file. In other words, the time series should occur in random order in the file.
- Make sure to set the `start` field correctly. The algorithm uses the `start` timestamp to derive the internal features.
- If you use categorical features (`cat`), all time series must have the same number of categorical features. If the dataset contains the `cat` field, the algorithm uses it and extracts the cardinality of the groups from the dataset. By default, cardinality is "auto". If the dataset contains the `cat` field, but you don't want to use it, you can disable it by setting `cardinality` to "". If a model was trained using a `cat` feature, you must include it for inference.
- If your dataset contains the `dynamic_feat` field, the algorithm uses it automatically. All time series have to have the same number of feature time series. The time points in each of the feature time series correspond one-to-one to the time points in the target. In addition, the entry in the `dynamic_feat` field should have the same length as the target. If the dataset contains the `dynamic_feat` field, but you don't want to use it, disable it by setting `num_dynamic_feat` to "". If the model was trained with the `dynamic_feat` field, you must provide this field for
inference. In addition, each of the features has to have the length of the provided target plus the prediction_length. In other words, you must provide the feature value in the future.

If you specify optional test channel data, the DeepAR algorithm evaluates the trained model with different accuracy metrics. The algorithm calculates the root mean square error (RMSE) over the test data as follows:

$$\text{RMSE} = \sqrt{\frac{1}{nT} \sum_{i,t} (\hat{y}_{i,t} - y_{i,t})^2}$$

$y_{i,t}$ is the true value of time series $i$ at the time $t$. $\hat{y}_{i,t}$ is the mean prediction. The sum is over all $n$ time series in the test set and over the last $T$ time points for each time series, where $T$ corresponds to the forecast horizon. You specify the length of the forecast horizon by setting the prediction_length hyperparameter. For more information, see DeepAR Hyperparameters.

In addition, the algorithm evaluates the accuracy of the forecast distribution using weighted quantile loss. For a quantile in the range $[0, 1]$, the weighted quantile loss is defined as follows:

$$\text{wQuantileLoss}[\tau] = \frac{2 \sum_{i,t} Q_{i,t}^{(\tau)}}{\sum_{i,t} |y_{i,t}|}, \quad \text{with} \quad Q_{i,t}^{(\tau)} = \begin{cases} (1-\tau) |q_{i,t}^{(\tau)} - y_{i,t}| & \text{if } q_{i,t}^{(\tau)} > y_{i,t} \\ \tau |q_{i,t}^{(\tau)} - y_{i,t}| & \text{otherwise} \end{cases}$$

$q_{i,t}^{(\tau)}$ is the $\tau$-quantile of the distribution that the model predicts. To specify which quantiles to calculate loss for, set the test_quantiles hyperparameter. In addition to these, the average of the prescribed quantile losses is reported as part of the training logs. For information, see DeepAR Hyperparameters.

For inference, DeepAR accepts JSON format and the following fields:

- "instances", which includes one or more time series in JSON Lines format
- A name of "configuration", which includes parameters for generating the forecast

For more information, see DeepAR Inference Formats.

**Best Practices for Using the DeepAR Algorithm**

When preparing your time series data, follow these best practices to achieve the best results:

- Except for when splitting your dataset for training and testing, always provide the entire time series for training, testing, and when calling the model for inference. Regardless of how you set
context_length, don't break up the time series or provide only a part of it. The model uses data points further back than the value set in context_length for the lagged values feature.

• When tuning a DeepAR model, you can split the dataset to create a training dataset and a test dataset. In a typical evaluation, you would test the model on the same time series used for training, but on the future prediction_length time points that follow immediately after the last time point visible during training. You can create training and test datasets that satisfy this criteria by using the entire dataset (the full length of all time series that are available) as a test set and removing the last prediction_length points from each time series for training. During training, the model doesn't see the target values for time points on which it is evaluated during testing. During testing, the algorithm withholds the last prediction_length points of each time series in the test set and generates a prediction. Then it compares the forecast with the withheld values. You can create more complex evaluations by repeating time series multiple times in the test set, but cutting them at different endpoints. With this approach, accuracy metrics are averaged over multiple forecasts from different time points. For more information, see Tune a DeepAR Model.

• Avoid using very large values (>400) for the prediction_length because it makes the model slow and less accurate. If you want to forecast further into the future, consider aggregating your data at a lower frequency. For example, use 5min instead of 1min.

• Because lags are used, a model can look further back in the time series than the value specified for context_length. Therefore, you don't need to set this parameter to a large value. We recommend starting with the value that you used for prediction_length.

• We recommend training a DeepAR model on as many time series as are available. Although a DeepAR model trained on a single time series might work well, standard forecasting algorithms, such as ARIMA or ETS, might provide more accurate results. The DeepAR algorithm starts to outperform the standard methods when your dataset contains hundreds of related time series. Currently, DeepAR requires that the total number of observations available across all training time series is at least 300.

EC2 Instance Recommendations for the DeepAR Algorithm

You can train DeepAR on both GPU and CPU instances and in both single and multi-machine settings. We recommend starting with a single CPU instance (for example, ml.c4.2xlarge or ml.c4.4xlarge), and switching to GPU instances and multiple machines only when necessary. Using GPUs and multiple machines improves throughput only for larger models (with many cells per layer and many layers) and for large mini-batch sizes (for example, greater than 512).
For inference, DeepAR supports only CPU instances.

Specifying large values for `context_length`, `prediction_length`, `num_cells`, `num_layers`, or `mini_batch_size` can create models that are too large for small instances. In this case, use a larger instance type or reduce the values for these parameters. This problem also frequently occurs when running hyperparameter tuning jobs. In that case, use an instance type large enough for the model tuning job and consider limiting the upper values of the critical parameters to avoid job failures.

DeepAR Sample Notebooks

For a sample notebook that shows how to prepare a time series dataset for training the SageMaker DeepAR algorithm and how to deploy the trained model for performing inferences, see DeepAR demo on electricity dataset, which illustrates the advanced features of DeepAR on a real world dataset. For instructions on creating and accessing Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After creating and opening a notebook instance, choose the SageMaker Examples tab to see a list of all of the SageMaker examples. To open a notebook, choose its Use tab, and choose Create copy.

How the DeepAR Algorithm Works

During training, DeepAR accepts a training dataset and an optional test dataset. It uses the test dataset to evaluate the trained model. In general, the datasets don't have to contain the same set of time series. You can use a model trained on a given training set to generate forecasts for the future of the time series in the training set, and for other time series. Both the training and the test datasets consist of one or, preferably, more target time series. Each target time series can optionally be associated with a vector of feature time series and a vector of categorical features. For more information, see Input/Output Interface for the DeepAR Algorithm.

For example, the following is an element of a training set indexed by $i$ which consists of a target time series, $Z_{i,t}$, and two associated feature time series, $X_{i,1,t}$ and $X_{i,2,t}$:
The target time series might contain missing values, which are represented by line breaks in the time series. DeepAR supports only feature time series that are known in the future. This allows you to run "what if?" scenarios. What happens, for example, if I change the price of a product in some way?

Each target time series can also be associated with a number of categorical features. You can use these features to encode which groupings a time series belongs to. Categorical features allow the model to learn typical behavior for groups, which it can use to increase model accuracy. DeepAR implements this by learning an embedding vector for each group that captures the common properties of all time series in the group.

How Feature Time Series Work in the DeepAR Algorithm

To facilitate learning time-dependent patterns, such as spikes during weekends, DeepAR automatically creates feature time series based on the frequency of the target time series. It uses these derived feature time series with the custom feature time series that you provide during training and inference. The following figure shows two of these derived time series features: $u_{i,1,t}$ represents the hour of the day and $u_{i,2,t}$ the day of the week.
The DeepAR algorithm automatically generates these feature time series. The following table lists the derived features for the supported basic time frequencies.

<table>
<thead>
<tr>
<th>Frequency of the Time Series</th>
<th>Derived Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minute</td>
<td>minute-of-hour, hour-of-day, day-of-week, day-of-month, day-of-year</td>
</tr>
<tr>
<td>Hour</td>
<td>hour-of-day, day-of-week, day-of-month, day-of-year</td>
</tr>
<tr>
<td>Day</td>
<td>day-of-week, day-of-month, day-of-year</td>
</tr>
<tr>
<td>Week</td>
<td>day-of-month, week-of-year</td>
</tr>
<tr>
<td>Month</td>
<td>month-of-year</td>
</tr>
</tbody>
</table>

DeepAR trains a model by randomly sampling several training examples from each of the time series in the training dataset. Each training example consists of a pair of adjacent context and prediction windows with fixed predefined lengths. The `context_length` hyperparameter controls how far in the past the network can see, and the `prediction_length` hyperparameter controls how far in the future predictions can be made. During training, the algorithm ignores training set elements containing time series that are shorter than a specified prediction length. The following figure represents five samples with context lengths of 12 hours and prediction lengths of 6 hours drawn from element \( i \). For brevity, we've omitted the feature time series \( x_{i,1,t} \) and \( u_{i,2,t} \).
To capture seasonality patterns, DeepAR also automatically feeds lagged values from the target time series. In the example with hourly frequency, for each time index, $t = T$, the model exposes the $z_{i,t}$ values, which occurred approximately one, two, and three days in the past.

![Graph showing lagged values](image)

For inference, the trained model takes as input target time series, which might or might not have been used during training, and forecasts a probability distribution for the next `prediction_length` values. Because DeepAR is trained on the entire dataset, the forecast takes into account patterns learned from similar time series.

For information on the mathematics behind DeepAR, see [DeepAR: Probabilistic Forecasting with Autoregressive Recurrent Networks](#).

**DeepAR Hyperparameters**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>context_length</code></td>
<td>The number of time-points that the model gets to see before making the prediction. The value for this parameter should be about the same as the <code>prediction_length</code>. The model also receives lagged inputs from the target, so <code>context_length</code> can be much smaller than typical seasonalities. For example, a daily time series can have yearly seasonality. The model automatically includes a lag of one year, so the context length can be shorter than a year. The lag values that the model picks depend on the frequency of the time series. For example, lag values for daily frequency are previous week, 2 weeks, 3 weeks, 4 weeks, and year. Required</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>epochs</td>
<td>The maximum number of passes over the training data. The optimal value depends on your data size and learning rate. See also <code>early_stopping_patience</code>. Typical values range from 10 to 1000.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td>prediction_length</td>
<td>The number of time-steps that the model is trained to predict, also called the forecast horizon. The trained model always generates forecasts with this length. It can't generate longer forecasts. The <code>prediction_length</code> is fixed when a model is trained and it cannot be changed later.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
</tbody>
</table>

Valid values: Positive integer
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>time_freq</td>
<td>The granularity of the time series in the dataset. Use <code>time_freq</code> to select appropriate date features and lags. The model supports the following basic frequencies. It also supports multiples of these basic frequencies. For example, <code>5min</code> specifies a frequency of 5 minutes.</td>
</tr>
<tr>
<td></td>
<td>• <code>M</code>: monthly</td>
</tr>
<tr>
<td></td>
<td>• <code>W</code>: weekly</td>
</tr>
<tr>
<td></td>
<td>• <code>D</code>: daily</td>
</tr>
<tr>
<td></td>
<td>• <code>H</code>: hourly</td>
</tr>
<tr>
<td></td>
<td>• <code>min</code>: every minute</td>
</tr>
</tbody>
</table>

**Required**

Valid values: An integer followed by `M, W, D, H, or min`. For example, `5min`. 
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| **cardinality** | When using the categorical features (cat), cardinality is an array specifying the number of categories (groups) per categorical feature. Set this to `auto` to infer the cardinality from the data. The `auto` mode also works when no categorical features are used in the dataset. This is the recommended setting for the parameter.  

Set cardinality to `ignore` to force DeepAR to not use categorical features, even if they are present in the data.  

To perform additional data validation, it is possible to explicitly set this parameter to the actual value. For example, if two categorical features are provided where the first has 2 and the other has 3 possible values, set this to `[2, 3]`.  

For more information on how to use categorical feature, see the data-section on the main documentation page of DeepAR.  

Optional  

Valid values: `auto`, `ignore`, array of positive integers, empty string, or  

Default value: `auto` |
| **dropout_rate** | The dropout rate to use during training. The model uses zoneout regularization. For each iteration, a random subset of hidden neurons are not updated. Typical values are less than 0.2.  

Optional  

Valid values: float  

Default value: 0.1 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>early_stopping_patience</td>
<td>If this parameter is set, training stops when no progress is made within the specified number of epochs. The model that has the lowest loss is returned as the final model.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td>embedding_dimension</td>
<td>Size of embedding vector learned per categorical feature (same value is used for all categorical features).</td>
</tr>
<tr>
<td></td>
<td>The DeepAR model can learn group-level time series patterns when a categorical grouping feature is provided. To do this, the model learns an embedding vector of size <code>embedding_dimension</code> for each group, capturing the common properties of all time series in the group. A larger <code>embedding_dimension</code> allows the model to capture more complex patterns. However, because increasing the <code>embedding_dimension</code> increases the number of parameters in the model, more training data is required to accurately learn these parameters. Typical values for this parameter are between 10-100.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The learning rate used in training. Typical values range from 1e-4 to 1e-1.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default value: 1e-3</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| likelihood       | The model generates a probabilistic forecast, and can provide quantiles of the distribution and return samples. Depending on your data, select an appropriate likelihood (noise model) that is used for uncertainty estimates. The following likelihoods can be selected:  
  - *gaussian*: Use for real-valued data.  
  - *beta*: Use for real-valued targets between 0 and 1 inclusive.  
  - *negative-binomial*: Use for count data (non-negative integers).  
  - *student-T*: An alternative for real-valued data that works well for bursty data.  
  - *deterministic-L1*: A loss function that does not estimate uncertainty and only learns a point forecast. |
<p>|                  | <strong>Optional</strong>                                                                                                                                                                                                |
|                  | Valid values: One of <em>gaussian</em>, <em>beta</em>, <em>negative-binomial</em>, <em>student-T</em>, or <em>deterministic-L1</em>.                                                                                                             |
|                  | Default value: student-T                                                                                                                                                                                     |
| mini_batch_size  | The size of mini-batches used during training. Typical values range from 32 to 512.                                                                                                                           |
|                  | <strong>Optional</strong>                                                                                                                                                                                                |
|                  | Valid values: positive integer                                                                                                                                                                              |
|                  | Default value: 128                                                                                                                                                                                           |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_cells</td>
<td>The number of cells to use in each hidden layer of the RNN. Typical values range from 30 to 100.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 40</td>
</tr>
<tr>
<td>num_dynamic_feat</td>
<td>The number of dynamic features provided in the data. Set this to auto to infer the number of dynamic features from the data. The auto mode also works when no dynamic features are used in the dataset. This is the recommended setting for the parameter.</td>
</tr>
<tr>
<td></td>
<td>To force DeepAR to not use dynamic features, even if they are present in the data, set num_dynamic_feat to ignore.</td>
</tr>
<tr>
<td></td>
<td>To perform additional data validation, it is possible to explicitly set this parameter to the actual integer value. For example, if two dynamic features are provided, set this to 2.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: auto, ignore, positive integer, or empty string</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>num_eval_samples</td>
<td>The number of samples that are used per time-series when calculating test accuracy metrics. This parameter does not have any influence on the training or the final model. In particular, the model can be queried with a different number of samples. This parameter only affects the reported accuracy scores on the test channel after training. Smaller values result in faster evaluation, but then the evaluation scores are typically worse and more uncertain. When evaluating with higher quantiles, for example 0.95, it may be important to increase the number of evaluation samples. Optional Valid values: integer Default value: 100</td>
</tr>
<tr>
<td>num_layers</td>
<td>The number of hidden layers in the RNN. Typical values range from 1 to 4. Optional Valid values: positive integer Default value: 2</td>
</tr>
<tr>
<td>test_quantiles</td>
<td>Quantiles for which to calculate quantile loss on the test channel. Optional Valid values: array of floats Default value: [0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9]</td>
</tr>
</tbody>
</table>
Tune a DeepAR Model

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics Computed by the DeepAR Algorithm**

The DeepAR algorithm reports three metrics, which are computed during training. When tuning a model, choose one of these as the objective. For the objective, use either the forecast accuracy on a provided test channel (recommended) or the training loss. For recommendations for the training/test split for the DeepAR algorithm, see [Best Practices for Using the DeepAR Algorithm](#).

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:RMSE</td>
<td>The root mean square error between the forecast and the actual target computed on the test set.</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:mean_wQuantileLoss</td>
<td>The average overall quantile losses computed on the test set. To control which quantiles are used, set the test_quantiles hyperparameter.</td>
<td>Minimize</td>
</tr>
<tr>
<td>train:final_loss</td>
<td>The training negative log-likelihood loss averaged over the last training epoch for the model.</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

**Tunable Hyperparameters for the DeepAR Algorithm**

Tune a DeepAR model with the following hyperparameters. The hyperparameters that have the greatest impact, listed in order from the most to least impactful, on DeepAR objective metrics are: epochs, context_length, mini_batch_size, learning_rate, and num_cells.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>epochs</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 1000</td>
</tr>
<tr>
<td>context_length</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 200</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 32, MaxValue: 1028</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-5, MaxValue: 1e-1</td>
</tr>
<tr>
<td>num_cells</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 30, MaxValue: 200</td>
</tr>
<tr>
<td>num_layers</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 8</td>
</tr>
<tr>
<td>dropout_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.00, MaxValue: 0.2</td>
</tr>
<tr>
<td>embedding_dimension</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 50</td>
</tr>
</tbody>
</table>

**DeepAR Inference Formats**

**DeepAR JSON Request Formats**

Query a trained model by using the model's endpoint. The endpoint takes the following JSON request format.

In the request, the instances field corresponds to the time series that should be forecast by the model.

If the model was trained with categories, you must provide a cat for each instance. If the model was trained without the cat field, it should be omitted.
If the model was trained with a custom feature time series (dynamic_feat), you have to provide the same number of dynamic_feat values for each instance. Each of them should have a length given by \text{length(target)} + \text{prediction_length}, where the last \text{prediction_length} values correspond to the time points in the future that will be predicted. If the model was trained without custom feature time series, the field should not be included in the request.

```json
{
    "instances": [
        {
            "start": "2009-11-01 00:00:00",
            "target": [4.0, 10.0, "NaN", 100.0, 113.0],
            "cat": [0, 1],
            "dynamic_feat": [[1.0, 1.1, 2.1, 0.5, 3.1, 4.1, 1.2, 5.0, ...]]
        },
        {
            "start": "2012-01-30",
            "target": [1.0],
            "cat": [2, 1],
            "dynamic_feat": [[2.0, 3.1, 4.5, 1.5, 1.8, 3.2, 0.1, 3.0, ...]]
        },
        {
            "start": "1999-01-30",
            "target": [2.0, 1.0],
            "cat": [1, 3],
            "dynamic_feat": [[1.0, 0.1, -2.5, 0.3, 2.0, -1.2, -0.1, -3.0, ...]]
        }
    ],
    "configuration": {
        "num_samples": 50,
        "output_types": ["mean", "quantiles", "samples"],
        "quantiles": ["0.5", "0.9"]
    }
}
```

The configuration field is optional. configuration.num_samples sets the number of sample paths that the model generates to estimate the mean and quantiles. configuration.output_types describes the information that will be returned in the request. Valid values are "mean" "quantiles" and "samples". If you specify "quantiles", each of the quantile values in configuration.quantiles is returned as a time series. If you specify "samples", the model also returns the raw samples used to calculate the other outputs.
DeepAR JSON Response Formats

The following is the format of a response, where [... ] are arrays of numbers:

```json
{
   "predictions": [
   {
      "quantiles": {
         "0.9": [...],
         "0.5": [...]
      },
      "samples": [...],
      "mean": [...]
   },
   {
      "quantiles": {
         "0.9": [...],
         "0.5": [...]
      },
      "samples": [...],
      "mean": [...]
   },
   {
      "quantiles": {
         "0.9": [...],
         "0.5": [...]
      },
      "samples": [...],
      "mean": [...]
   }
   ]
}
```

DeepAR has a response timeout of 60 seconds. When passing multiple time series in a single request, the forecasts are generated sequentially. Because the forecast for each time series typically takes about 300 to 1000 milliseconds or longer, depending on the model size, passing too many time series in a single request can cause time outs. It's better to send fewer time series per request and send more requests. Because the DeepAR algorithm uses multiple workers per instance, you can achieve much higher throughput by sending multiple requests in parallel.

By default, DeepAR uses one worker per CPU for inference, if there is sufficient memory per CPU. If the model is large and there isn't enough memory to run a model on each
CPU, the number of workers is reduced. The number of workers used for inference can be overwritten using the environment variable MODEL_SERVER_WORKERS. For example, by setting MODEL_SERVER_WORKERS=1) when calling the SageMaker CreateModel API.

Batch Transform with the DeepAR Algorithm

DeepAR forecasting supports getting inferences by using batch transform from data using the JSON Lines format. In this format, each record is represented on a single line as a JSON object, and lines are separated by newline characters. The format is identical to the JSON Lines format used for model training. For information, see Input/Output Interface for the DeepAR Algorithm. For example:

```json
{"start": "2009-11-01 00:00:00", "target": [4.3, "NaN", 5.1, ...], "cat": [0, 1],
  "dynamic_feat": [[1.1, 1.2, 0.5, ..]]}
{"start": "2012-01-30 00:00:00", "target": [1.0, -5.0, ...], "cat": [2, 3],
  "dynamic_feat": [[1.1, 2.05, ...]]}
{"start": "1999-01-30 00:00:00", "target": [2.0, 1.0], "cat": [1, 4], "dynamic_feat":
  [[1.3, 0.4]]}
```

Note

When creating the transformation job with CreateTransformJob, set the BatchStrategy value to SingleRecord and set the SplitType value in the TransformInput configuration to Line, as the default values currently cause runtime failures.

Similar to the hosted endpoint inference request format, the cat and the dynamic_feat fields for each instance are required if both of the following are true:

- The model is trained on a dataset that contained both the cat and the dynamic_feat fields.
- The corresponding cardinality and num_dynamic_feat values used in the training job are not set to "".

Unlike hosted endpoint inference, the configuration field is set once for the entire batch inference job using an environment variable named DEEPAR_INDUCTION_CONFIG. The value of DEEPAR_INDUCTION_CONFIG can be passed when the model is created by calling
CreateTransformJob API. If DEEPAR_INFERENCE_CONFIG is missing in the container environment, the inference container uses the following default:

```json
{
    "num_samples": 100,
    "output_types": ["mean", "quantiles"],
    "quantiles": ["0.1", "0.2", "0.3", "0.4", "0.5", "0.6", "0.7", "0.8", "0.9"]
}
```

The output is also in JSON Lines format, with one line per prediction, in an order identical to the instance order in the corresponding input file. Predictions are encoded as objects identical to the ones returned by responses in online inference mode. For example:

```json
{ "quantiles": { "0.1": [...], "0.2": [...] }, "samples": [...], "mean": [...] }
```

Note that in the TransformInput configuration of the SageMaker CreateTransformJob request clients must explicitly set the AssembleWith value to Line, as the default value None concatenates all JSON objects on the same line.

For example, here is a SageMaker CreateTransformJob request for a DeepAR job with a custom DEEPAR_INFERENCE_CONFIG:

```json
{
    "BatchStrategy": "SingleRecord",
    "Environment": {
        "DEEPAR_INFERENCE_CONFIG": "{ "num_samples": 200, "output_types": ["mean"] }",
        ...
    },
    "TransformInput": {
        "SplitType": "Line",
        ...
    },
    "TransformOutput": {
        "AssembleWith": "Line",
        ...
    },
    ...
}
```
Unsupervised Built-in SageMaker Algorithms

Amazon SageMaker provides several built-in algorithms that can be used for a variety of unsupervised learning tasks such as clustering, dimension reduction, pattern recognition, and anomaly detection.

- **IP Insights**—learns the usage patterns for IPv4 addresses. It is designed to capture associations between IPv4 addresses and various entities, such as user IDs or account numbers.
- **K-Means Algorithm**—finds discrete groupings within data, where members of a group are as similar as possible to one another and as different as possible from members of other groups.
- **Principal Component Analysis (PCA) Algorithm**—reduces the dimensionality (number of features) within a dataset by projecting data points onto the first few principal components. The objective is to retain as much information or variation as possible. For mathematicians, principal components are eigenvectors of the data's covariance matrix.
- **Random Cut Forest (RCF) Algorithm**—detects anomalous data points within a data set that diverge from otherwise well-structured or patterned data.

<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>IP Insights</td>
<td>train and (optionally) validation</td>
<td>File</td>
<td>CSV</td>
<td>CPU or GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>K-Means</td>
<td>train and (optionally) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU or GPU (single GPU device on one or more instances)</td>
<td>No</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
<td>Instance class</td>
<td>Parallelizable</td>
</tr>
<tr>
<td>----------------</td>
<td>--------------</td>
<td>---------------------</td>
<td>-----------</td>
<td>----------------</td>
<td>----------------</td>
</tr>
<tr>
<td>PCA</td>
<td>train and (optional ly) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>GPU or CPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Random Cut Forest</td>
<td>train and (optional ly) test</td>
<td>File or Pipe</td>
<td>recordIO-protobuf or CSV</td>
<td>CPU</td>
<td>Yes</td>
</tr>
</tbody>
</table>

**IP Insights**

Amazon SageMaker IP Insights is an unsupervised learning algorithm that learns the usage patterns for IPv4 addresses. It is designed to capture associations between IPv4 addresses and various entities, such as user IDs or account numbers. You can use it to identify a user attempting to log into a web service from an anomalous IP address, for example. Or you can use it to identify an account that is attempting to create computing resources from an unusual IP address. Trained IP Insight models can be hosted at an endpoint for making real-time predictions or used for processing batch transforms.

SageMaker IP insights ingests historical data as (entity, IPv4 Address) pairs and learns the IP usage patterns of each entity. When queried with an (entity, IPv4 Address) event, a SageMaker IP Insights model returns a score that infers how anomalous the pattern of the event is. For example, when a user attempts to log in from an IP address, if the IP Insights score is high enough, a web login server might decide to trigger a multi-factor authentication system. In more advanced solutions, you can feed the IP Insights score into another machine learning model. For example, you can combine the IP Insight score with other features to rank the findings of another security system, such as those from Amazon GuardDuty.

The SageMaker IP Insights algorithm can also learn vector representations of IP addresses, known as embeddings. You can use vector-encoded embeddings as features in downstream machine learning tasks that use the information observed in the IP addresses. For example, you can use them in tasks such as measuring similarities between IP addresses in clustering and visualization tasks.

**Topics**
Input/Output Interface for the IP Insights Algorithm

Training and Validation

The SageMaker IP Insights algorithm supports training and validation data channels. It uses the optional validation channel to compute an area-under-curve (AUC) score on a predefined negative sampling strategy. The AUC metric validates how well the model discriminates between positive and negative samples. Training and validation data content types need to be in text/csv format. The first column of the CSV data is an opaque string that provides a unique identifier for the entity. The second column is an IPv4 address in decimal-dot notation. IP Insights currently supports only File mode. For more information and some examples, see IP Insights Training Data Formats.

Inference

For inference, IP Insights supports text/csv, application/json, and application/jsonlines data content types. For more information about the common data formats for inference provided by SageMaker, see Common Data Formats for Inference. IP Insights inference returns output formatted as either application/json or application/jsonlines. Each record in the output data contains the corresponding dot_product (or compatibility score) for each input data point. For more information and some examples, see IP Insights Inference Data Formats.

EC2 Instance Recommendation for the IP Insights Algorithm

The SageMaker IP Insights algorithm can run on both GPU and CPU instances. For training jobs, we recommend using GPU instances. However, for certain workloads with large training datasets, distributed CPU instances might reduce training costs. For inference, we recommend using CPU instances. IP Insights supports P2, P3, G4dn, and G5 GPU families.
GPU Instances for the IP Insights Algorithm

IP Insights supports all available GPUs. If you need to speed up training, we recommend starting with a single GPU instance, such as ml.p3.2xlarge, and then moving to a multi-GPU environment, such as ml.p3.8xlarge and ml.p3.16xlarge. Multi-GPUs automatically divide the mini batches of training data across themselves. If you switch from a single GPU to multiple GPUs, the mini_batch_size is divided equally into the number of GPUs used. You may want to increase the value of the mini_batch_size to compensate for this.

CPU Instances for the IP Insights Algorithm

The type of CPU instance that we recommend depends largely on the instance's available memory and the model size. The model size is determined by two hyperparameters: vector_dim and num_entity_vectors. The maximum supported model size is 8 GB. The following table lists typical EC2 instance types that you would deploy based on these input parameters for various model sizes. In Table 1, the value for vector_dim in the first column range from 32 to 2048 and the values for num_entity_vectors in the first row range from 10,000 to 50,000,000.

<table>
<thead>
<tr>
<th>vector_dim</th>
<th>10,000</th>
<th>50,000</th>
<th>100,000</th>
<th>500,000</th>
<th>1,000,000</th>
<th>5,000,000</th>
<th>10,000,000</th>
<th>50,000,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>32</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
</tr>
<tr>
<td>64</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
</tr>
<tr>
<td>128</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
</tr>
<tr>
<td>256</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
</tr>
<tr>
<td>512</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
<td>ml.m5.large</td>
</tr>
</tbody>
</table>

Use Built-in Algorithms
The values for the mini_batch_size, num_ip_encoder_layers, random_negative_sampling_rate, and shuffled_negative_sampling_rate hyperparameters also affect the amount of memory required. If these values are large, you might need to use a larger instance type than normal.

**IP Insights Sample Notebooks**

For a sample notebook that shows how to train the SageMaker IP Insights algorithm and perform inferences with it, see [An Introduction to the SageMakerIP Insights Algorithm](#). For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](#). After creating a notebook instance, choose the SageMaker Examples tab to see a list of all the SageMaker examples. To open a notebook, choose its Use tab and choose Create copy.

**How IP Insights Works**

Amazon SageMaker IP Insights is an unsupervised algorithm that consumes observed data in the form of (entity, IPv4 address) pairs that associates entities with IP addresses. IP Insights determines how likely it is that an entity would use a particular IP address by learning latent vector representations for both entities and IP addresses. The distance between these two representations can then serve as the proxy for how likely this association is.

The IP Insights algorithm uses a neural network to learn the latent vector representations for entities and IP addresses. Entities are first hashed to a large but fixed hash space and then encoded by a simple embedding layer. Character strings such as user names or account IDs can be fed directly into IP Insights as they appear in log files. You don't need to preprocess the data for entity
identifiers. You can provide entities as an arbitrary string value during both training and inference. The hash size should be configured with a value that is high enough to ensure that the number of collisions, which occur when distinct entities are mapped to the same latent vector, remain insignificant. For more information about how to select appropriate hash sizes, see Feature Hashing for Large Scale Multitask Learning. For representing IP addresses, on the other hand, IP Insights uses a specially designed encoder network to uniquely represent each possible IPv4 address by exploiting the prefix structure of IP addresses.

During training, IP Insights automatically generates negative samples by randomly pairing entities and IP addresses. These negative samples represent data that is less likely to occur in reality. The model is trained to discriminate between positive samples that are observed in the training data and these generated negative samples. More specifically, the model is trained to minimize the cross entropy, also known as the log loss, defined as follows:

$$\mathcal{L} = \frac{1}{N} \sum_n [y_n \log p_n + (1 - y_n) \log (1 - p_n)]$$

$y_n$ is the label that indicates whether the sample is from the real distribution governing observed data ($y_n=1$) or from the distribution generating negative samples ($y_n=0$). $p_n$ is the probability that the sample is from the real distribution, as predicted by the model.

Generating negative samples is an important process that is used to achieve an accurate model of the observed data. If negative samples are extremely unlikely, for example, if all of the IP addresses in negative samples are 10.0.0.0, then the model trivially learns to distinguish negative samples and fails to accurately characterize the actual observed dataset. To keep negative samples more realistic, IP Insights generates negative samples both by randomly generating IP addresses and randomly picking IP addresses from training data. You can configure the type of negative sampling and the rates at which negative samples are generated with the `random_negative_sampling_rate` and `shuffled_negative_sampling_rate` hyperparameters.

Given an nth (entity, IP address pair), the IP Insights model outputs a score, $S_n$, that indicates how compatible the entity is with the IP address. This score corresponds to the log odds ratio for a given (entity, IP address) of the pair coming from a real distribution as compared to coming from a negative distribution. It is defined as follows:
\[ S_n = \log \left( \frac{P_{real}(n)}{P_{neg}(n)} \right) \]

The score is essentially a measure of the similarity between the vector representations of the nth entity and IP address. It can be interpreted as how much more likely it would be to observe this event in reality than in a randomly generated dataset. During training, the algorithm uses this score to calculate an estimate of the probability of a sample coming from the real distribution, \( p_n \), to use in the cross entropy minimization, where:

\[ p_n = \frac{1}{1 + e^{-S_n}} \]

**IP Insights Hyperparameters**

In the CreateTransformJob request, you specify the training algorithm. You can also specify algorithm-specific hyperparameters as string-to-string maps. The following table lists the hyperparameters for the Amazon SageMaker IP Insights algorithm.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_entity_vectors</td>
<td>The number of entity vector representations (entity embedding vectors) to train. Each entity in the training set is randomly assigned to one of these vectors using a hash function. Because of hash collisions, it might be possible to have multiple entities assigned to the same vector. This would cause the same vector to represent multiple entities. This generally has a negligible effect on model performance, as long as the collision rate is not too severe. To keep the collision rate low, set this value as high as possible. However, the model size, and, therefore, the memory requirement, for both training and inference, scales linearly with this hyperparameter. We recommend that you set this value to twice the number of unique entity identifiers.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Required</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: 1 ≤ positive integer ≤ 250,000,000</td>
<td></td>
</tr>
<tr>
<td>vector_dim</td>
<td>The size of embedding vectors to represent entities and IP addresses. The larger the value, the more information that can be encoded using these representations. In practice, model size scales linearly with this parameter and limits how large the dimension can be. In addition, using vector representations that are too large can cause the model to overfit, especially for small training datasets. Overfitting occurs when a model doesn't learn any pattern in the data but effectively memorizes the training data and, therefore, cannot generalize well and performs poorly during inference. The recommended value is 128.</td>
</tr>
<tr>
<td><strong>Required</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: 4 ≤ positive integer ≤ 4096</td>
<td></td>
</tr>
<tr>
<td>batch_metrics_publish_interval</td>
<td>The interval (every X batches) at which the Apache MXNet Speedometer function prints the training speed of the network (samples/second).</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td>Valid values: positive integer ≥ 1</td>
<td></td>
</tr>
<tr>
<td>Default value: 1,000</td>
<td></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| epochs           | The number of passes over the training data. The optimal value depends on your data size and learning rate. Typical values range from 5 to 100. Optional
                   | Valid values: positive integer ≥ 1                                                                                                                                                                |
|                  | Default value: 10                                                                                                                                                                                           |
| learning_rate    | The learning rate for the optimizer. IP Insights use a gradient-descent-based Adam optimizer. The learning rate effectively controls the step size to update model parameters at each iteration. Too large a learning rate can cause the model to diverge because the training is likely to overshoot a minima. On the other hand, too small a learning rate slows down convergence. Typical values range from 1e-4 to 1e-1. Optional
<pre><code>               | Valid values: 1e-6 ≤ float ≤ 10.0                                                                                                                                                                      |
</code></pre>
<p>|                  | Default value: 0.001                                                                                                                                                                                       |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
<th>Optional</th>
<th>Valid values</th>
<th>Default value</th>
</tr>
</thead>
<tbody>
<tr>
<td>mini_batch_size</td>
<td>The number of examples in each mini batch. The training procedure processes data in mini batches. The optimal value depends on the number of unique account identifiers in the dataset. In general, the larger the mini_batch_size, the faster the training and the greater the number of possible shuffled-negative-sample combinations. However, with a large mini_batch_size, the training is more likely to converge to a poor local minimum and perform relatively worse for inference.</td>
<td>Yes</td>
<td>1 ≤ positive integer ≤ 500000</td>
<td>10,000</td>
</tr>
<tr>
<td>num_ip_encoder_layers</td>
<td>The number of fully connected layers used to encode the IP address embedding. The larger the number of layers, the greater the model's capacity to capture patterns among IP addresses. However, using a large number of layers increases the chance of overfitting.</td>
<td>Yes</td>
<td>0 ≤ positive integer ≤ 100</td>
<td>1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>random_negative_sampling_rate</td>
<td>The number of random negative samples, $R$, to generate per input example. The training procedure relies on negative samples to prevent the vector representations of the model collapsing to a single point. Random negative sampling generates $R$ random IP addresses for each input account in the mini batch. The sum of the <code>random_negative_sampling_rate</code> ($R$) and <code>shuffled_negative_sampling_rate</code> ($S$) must be in the interval: $1 \leq R + S \leq 500$.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Valid values: $0 \leq$ positive integer $\leq 500$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>shuffled_negative_sampling_rate</td>
<td>The number of shuffled negative samples, $S$, to generate per input example. In some cases, it helps to use more realistic negative samples that are randomly picked from the training data itself. This kind of negative sampling is achieved by shuffling the data within a mini batch. Shuffled negative sampling generates $S$ negative IP addresses by shuffling the IP address and account pairings within a mini batch. The sum of the <code>random_negative_sampling_rate</code> ($R$) and <code>shuffled_negative_sampling_rate</code> ($S$) must be in the interval: $1 \leq R + S \leq 500$.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Valid values: $0 \leq$ positive integer $\leq 500$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Default value: 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| weight_decay   | The weight decay coefficient. This parameter adds an L2 regularization factor that is required to prevent the model from overfitting the training data.  
Optional  
Valid values: $0.0 \leq \text{float} \leq 10.0$  
Default value: 0.00001 |

**Tune an IP Insights Model**

*Automatic model tuning*, also called hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics Computed by the IP Insights Algorithm**

The Amazon SageMaker IP Insights algorithm is an unsupervised learning algorithm that learns associations between IP addresses and entities. The algorithm trains a discriminator model, which learns to separate observed data points (positive samples) from randomly generated data points (negative samples). Automatic model tuning on IP Insights helps you find the model that can most accurately distinguish between unlabeled validation data and automatically generated negative samples. The model accuracy on the validation dataset is measured by the area under the receiver operating characteristic curve. This `validation:discriminator_auc` metric can take values between 0.0 and 1.0, where 1.0 indicates perfect accuracy.

The IP Insights algorithm computes a `validation:discriminator_auc` metric during validation, the value of which is used as the objective function to optimize for hyperparameter tuning.
### Metric Name

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:discriminator_auc</td>
<td>Area under the receiver operating characteristic curve on the validation dataset. The validation dataset is not labeled. Area Under the Curve (AUC) is a metric that describes the model's ability to discriminate validation data points from randomly generated data points.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

### Tunable IP Insights Hyperparameters

You can tune the following hyperparameters for the SageMaker IP Insights algorithm.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>epochs</td>
<td>IntegerParameterRange</td>
<td>MinValue: 1, MaxValue: 100</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-4, MaxValue: 0.1</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 100, MaxValue: 50000</td>
</tr>
<tr>
<td>num_entity_vectors</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 10000, MaxValue: 1000000</td>
</tr>
<tr>
<td>num_ip_encoder_layers</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 10</td>
</tr>
<tr>
<td>random_negative_sampling_rate</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 0, MaxValue: 10</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>-------------------------------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>shuffled_negative_sampling_rate</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 0, MaxValue: 10</td>
</tr>
<tr>
<td>vector_dim</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 8, MaxValue: 256</td>
</tr>
<tr>
<td>weight_decay</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 1.0</td>
</tr>
</tbody>
</table>

**IP Insights Data Formats**

This section provides examples of the available input and output data formats used by the IP Insights algorithm during training and inference.

**Topics**
- IP Insights Training Data Formats
- IP Insights Inference Data Formats

**IP Insights Training Data Formats**

The following are the available data input formats for the IP Insights algorithm. Amazon SageMaker built-in algorithms adhere to the common input training format described in Common Data Formats for Training. However, the SageMaker IP Insights algorithm currently supports only the CSV data input format.

**IP Insights Training Data Input Formats**

**INPUT: CSV**

The CSV file must have two columns. The first column is an opaque string that corresponds to an entity's unique identifier. The second column is the IPv4 address of the entity's access event in decimal-dot notation.

content-type: text/csv
IP Insights Inference Data Formats

The following are the available input and output formats for the IP Insights algorithm. Amazon SageMaker built-in algorithms adhere to the common input inference format described in Common Data Formats for Inference. However, the SageMaker IP Insights algorithm does not currently support RecordIO format.

IP Insights Input Request Formats

INPUT: CSV Format

The CSV file must have two columns. The first column is an opaque string that corresponds to an entity's unique identifier. The second column is the IPv4 address of the entity's access event in decimal-dot notation.

```
content-type: text/csv

entity_id_1, 192.168.1.2
entity_id_2, 10.10.1.2
```

INPUT: JSON Format

JSON data can be provided in different formats. IP Insights follows the common SageMaker formats. For more information about inference formats, see Common Data Formats for Inference.

```
content-type: application/json

{
   "instances": [
      {
         "data": {"features": {"values": ["entity_id_1", "192.168.1.2"]}}
      },
      {
         "data": {"features": ["entity_id_2", "10.10.1.2"]}
      }
   ]
}
```

INPUT: JSONLINES Format

The JSON Lines content type is useful for running batch transform jobs. For more information on SageMaker inference formats, see Common Data Formats for Inference. For more information on running batch transform jobs, see Use Batch Transform.
IP Insights Output Response Formats

OUTPUT: JSON Response Format

The default output of the SageMaker IP Insights algorithm is the dot_product between the input entity and IP address. The dot_product signifies how compatible the model considers the entity and IP address. The dot_product is unbounded. To make predictions about whether an event is anomalous, you need to set a threshold based on your defined distribution. For information about how to use the dot_product for anomaly detection, see the An Introduction to the SageMakerIP Insights Algorithm.

Advanced users can access the model's learned entity and IP embeddings by providing the additional content-type parameter verbose=True to the Accept heading. You can use the entity_embedding and ip_embedding for debugging, visualizing, and understanding the model. Additionally, you can use these embeddings in other machine learning techniques, such as classification or clustering.

Use Built-in Algorithms
K-Means Algorithm

K-means is an unsupervised learning algorithm. It attempts to find discrete groupings within data, where members of a group are as similar as possible to one another and as different as possible from members of other groups. You define the attributes that you want the algorithm to use to determine similarity.

Amazon SageMaker uses a modified version of the web-scale k-means clustering algorithm. Compared with the original version of the algorithm, the version used by Amazon SageMaker is more accurate. Like the original algorithm, it scales to massive datasets and delivers improvements in training time. To do this, the version used by Amazon SageMaker streams mini-batches (small, random subsets) of the training data. For more information about mini-batch k-means, see Web-scale k-means Clustering.

The k-means algorithm expects tabular data, where rows represent the observations that you want to cluster, and the columns represent attributes of the observations. The $n$ attributes in each row represent a point in $n$-dimensional space. The Euclidean distance between these points represents the similarity of the corresponding observations. The algorithm groups observations with similar
attribute values (the points corresponding to these observations are closer together). For more information about how k-means works in Amazon SageMaker, see How K-Means Clustering Works.

Topics
- Input/Output Interface for the K-Means Algorithm
- EC2 Instance Recommendation for the K-Means Algorithm
- K-Means Sample Notebooks
- How K-Means Clustering Works
- K-Means Hyperparameters
- Tune a K-Means Model
- K-Means Response Formats

Input/Output Interface for the K-Means Algorithm

For training, the k-means algorithm expects data to be provided in the train channel (recommended S3DataDistributionType=ShardedByS3Key), with an optional test channel (recommended S3DataDistributionType=FullyReplicated) to score the data on. Both recordIO-wrapped-protobuf and CSV formats are supported for training. You can use either File mode or Pipe mode to train models on data that is formatted as recordIO-wrapped-protobuf or as CSV.

For inference, text/csv, application/json, and application/x-recordio-protobuf are supported. k-means returns a closest_cluster label and the distance_to_cluster for each observation.

For more information on input and output file formats, see K-Means Response Formats for inference and the K-Means Sample Notebooks. The k-means algorithm does not support multiple instance learning, in which the training set consists of labeled “bags”, each of which is a collection of unlabeled instances.

EC2 Instance Recommendation for the K-Means Algorithm

We recommend training k-means on CPU instances. You can train on GPU instances, but should limit GPU training to single-GPU instances (such as ml.g4dn.xlarge) because only one GPU is used per instance. The k-means algorithm supports P2, P3, G4dn, and G5 instances for training and inference.
**K-Means Sample Notebooks**

For a sample notebook that uses the SageMaker K-means algorithm to segment the population of counties in the United States by attributes identified using principle component analysis, see [Analyze US census data for population segmentation using Amazon SageMaker](#). For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](#). Once you have created a notebook instance and opened it, select the **SageMaker Examples** tab to see a list of all the SageMaker samples. To open a notebook, click on its **Use** tab and select **Create copy**.

**How K-Means Clustering Works**

K-means is an algorithm that trains a model that groups similar objects together. The k-means algorithm accomplishes this by mapping each observation in the input dataset to a point in the \( n \)-dimensional space (where \( n \) is the number of attributes of the observation). For example, your dataset might contain observations of temperature and humidity in a particular location, which are mapped to points \((t, h)\) in 2-dimensional space.

__Note__

Clustering algorithms are unsupervised. In unsupervised learning, labels that might be associated with the objects in the training dataset aren't used. For more information, see [Unsupervised learning](#).

In k-means clustering, each cluster has a center. During model training, the k-means algorithm uses the distance of the point that corresponds to each observation in the dataset to the cluster centers as the basis for clustering. You choose the number of clusters \((k)\) to create.

For example, suppose that you want to create a model to recognize handwritten digits and you choose the MNIST dataset for training. The dataset provides thousands of images of handwritten digits (0 through 9). In this example, you might choose to create 10 clusters, one for each digit \((0, 1, \ldots, 9)\). As part of model training, the k-means algorithm groups the input images into 10 clusters.

Each image in the MNIST dataset is a 28x28-pixel image, with a total of 784 pixels. Each image corresponds to a point in a 784-dimensional space, similar to a point in a 2-dimensional space \((x, y)\). To find a cluster to which a point belongs, the k-means algorithm finds the distance of that point...
from all of the cluster centers. It then chooses the cluster with the closest center as the cluster to which the image belongs.

**Note**

Amazon SageMaker uses a customized version of the algorithm where, instead of specifying that the algorithm create $k$ clusters, you might choose to improve model accuracy by specifying extra cluster centers ($K = k \times x$). However, the algorithm ultimately reduces these to $k$ clusters.

In SageMaker, you specify the number of clusters when creating a training job. For more information, see [CreateTrainingJob](#). In the request body, you add the `HyperParameters` string map to specify the $k$ and `extra_center_factor` strings.

The following is a summary of how k-means works for model training in SageMaker:

1. It determines the initial $K$ cluster centers.

**Note**

In the following topics, $K$ clusters refer to $k \times x$, where you specify $k$ and $x$ when creating a model training job.

2. It iterates over input training data and recalculates cluster centers.

3. It reduces resulting clusters to $k$ (if the data scientist specified the creation of $k \times x$ clusters in the request).

The following sections also explain some of the parameters that a data scientist might specify to configure a model training job as part of the `HyperParameters` string map.

**Topics**

- [Step 1: Determine the Initial Cluster Centers](#)
- [Step 2: Iterate over the Training Dataset and Calculate Cluster Centers](#)
- [Step 3: Reduce the Clusters from $K$ to $k$](#)
Step 1: Determine the Initial Cluster Centers

When using k-means in SageMaker, the initial cluster centers are chosen from the observations in a small, randomly sampled batch. Choose one of the following strategies to determine how these initial cluster centers are selected:

- The random approach—Randomly choose $K$ observations in your input dataset as cluster centers. For example, you might choose a cluster center that points to the 784-dimensional space that corresponds to any 10 images in the MNIST training dataset.

- The k-means++ approach, which works as follows:
  1. Start with one cluster and determine its center. You randomly select an observation from your training dataset and use the point corresponding to the observation as the cluster center. For example, in the MNIST dataset, randomly choose a handwritten digit image. Then choose the point in the 784-dimensional space that corresponds to the image as your cluster center. This is cluster center 1.
  2. Determine the center for cluster 2. From the remaining observations in the training dataset, pick an observation at random. Choose one that is different than the one you previously selected. This observation corresponds to a point that is far away from cluster center 1. Using the MNIST dataset as an example, you do the following:
     - For each of the remaining images, find the distance of the corresponding point from cluster center 1. Square the distance and assign a probability that is proportional to the square of the distance. That way, an image that is different from the one that you previously selected has a higher probability of getting selected as cluster center 2.
     - Choose one of the images randomly, based on probabilities assigned in the previous step. The point that corresponds to the image is cluster center 2.
  3. Repeat Step 2 to find cluster center 3. This time, find the distances of the remaining images from cluster center 2.
  4. Repeat the process until you have the $K$ cluster centers.

To train a model in SageMaker, you create a training job. In the request, you provide configuration information by specifying the following HyperParameters string maps:

- To specify the number of clusters to create, add the k string.
- For greater accuracy, add the optional extra_center_factor string.
• To specify the strategy that you want to use to determine the initial cluster centers, add the `init_method` string and set its value to `random` or `k-means++`.

For more information about the SageMaker k-means estimator, see [K-means](https://docs.aws.amazon.com/sagemaker/latest/dg/kmeans.html) in the Amazon SageMaker Python SDK documentation.

You now have an initial set of cluster centers.

**Step 2: Iterate over the Training Dataset and Calculate Cluster Centers**

The cluster centers that you created in the preceding step are mostly random, with some consideration for the training dataset. In this step, you use the training dataset to move these centers toward the true cluster centers. The algorithm iterates over the training dataset, and recalculates the $K$ cluster centers.

1. Read a mini-batch of observations (a small, randomly chosen subset of all records) from the training dataset and do the following.

   **Note**
   
   When creating a model training job, you specify the batch size in the `mini_batch_size` string in the `HyperParameters` string map.

   a. Assign all of the observations in the mini-batch to one of the clusters with the closest cluster center.

   b. Calculate the number of observations assigned to each cluster. Then, calculate the proportion of new points assigned per cluster.

   For example, consider the following clusters:

   Cluster $c1 = 100$ previously assigned points. You added 25 points from the mini-batch in this step.

   Cluster $c2 = 150$ previously assigned points. You added 40 points from the mini-batch in this step.

   Cluster $c3 = 450$ previously assigned points. You added 5 points from the mini-batch in this step.
Calculate the proportion of new points assigned to each of clusters as follows:

\[
\begin{align*}
p_1 & = \text{proportion of points assigned to } c_1 = 25/(100+25) \\
p_2 & = \text{proportion of points assigned to } c_2 = 40/(150+40) \\
p_3 & = \text{proportion of points assigned to } c_3 = 5/(450+5)
\end{align*}
\]

c. Compute the center of the new points added to each cluster:

\[
\begin{align*}
d_1 & = \text{center of the new points added to cluster } 1 \\
d_2 & = \text{center of the new points added to cluster } 2 \\
d_3 & = \text{center of the new points added to cluster } 3
\end{align*}
\]

d. Compute the weighted average to find the updated cluster centers as follows:

\[
\begin{align*}
\text{Center of cluster 1} & = ((1 - p_1) \times \text{center of cluster 1}) + (p_1 \times d_1) \\
\text{Center of cluster 2} & = ((1 - p_2) \times \text{center of cluster 2}) + (p_2 \times d_2) \\
\text{Center of cluster 3} & = ((1 - p_3) \times \text{center of cluster 3}) + (p_3 \times d_3)
\end{align*}
\]

2. Read the next mini-batch, and repeat Step 1 to recalculate the cluster centers.

3. For more information about mini-batch k-means, see Web-scale k-means Clustering).

**Step 3: Reduce the Clusters from K to k**

If the algorithm created K clusters—(K = k\times x) where x is greater than 1—then it reduces the K clusters to k clusters. (For more information, see extra_center_factor in the preceding discussion.) It does this by applying Lloyd's method with kmeans++ initialization to the K cluster centers. For more information about Lloyd's method, see k-means clustering.

**K-Means Hyperparameters**

In the CreateTrainingJob request, you specify the training algorithm that you want to use. You can also specify algorithm-specific hyperparameters as string-to-string maps. The following table lists the hyperparameters for the k-means training algorithm provided by Amazon SageMaker. For more information about how k-means clustering works, see How K-Means Clustering Works.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>feature_dim</td>
<td>The number of features in the input data.</td>
</tr>
<tr>
<td></td>
<td>Required</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>k</strong></td>
<td>The number of required clusters.</td>
</tr>
<tr>
<td><strong>Required</strong></td>
<td></td>
</tr>
<tr>
<td><strong>epochs</strong></td>
<td>The number of passes done over the training data.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td><strong>eval_metrics</strong></td>
<td>A JSON list of metric types used to report a score for the model. Allowed</td>
</tr>
<tr>
<td></td>
<td>values are msd for Means Square Deviation and ssd for Sum of Square Distance.</td>
</tr>
<tr>
<td></td>
<td>If test data is provided, the score is reported for each of the metrics requested.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
<tr>
<td><strong>extra_center_factor</strong></td>
<td>The algorithm creates K centers = num_clusters * extra_center_factor as it runs and reduces the number of centers from K to k when finalizing the model.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
</tr>
</tbody>
</table>

**Valid values:**
- Positive integer
- Either `"msd"` or `"ssd"` or `"msd", "ssd"`.
- Either a positive integer or `auto`.

**Default value:**
- 1
- `auto`
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>half_life_time_size</td>
<td>Used to determine the weight given to an observation when computing a cluster mean. This weight decays exponentially as more points are observed. When a point is first observed, it is assigned a weight of 1 when computing the cluster mean. The decay constant for the exponential decay function is chosen so that after observing half_life_time_size points, its weight is 1/2. If set to 0, there is no decay.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>init_method</td>
<td>Method by which the algorithm chooses the initial cluster centers. The standard k-means approach chooses them at random. An alternative k-means++ method chooses the first cluster center at random. Then it spreads out the position of the remaining initial clusters by weighting the selection of centers with a probability distribution that is proportional to the square of the distance of the remaining data points from existing centers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either random or kmeans++.</td>
</tr>
<tr>
<td></td>
<td>Default value: random</td>
</tr>
<tr>
<td>local_lloyd_init_method</td>
<td>The initialization method for Lloyd's expectation-maximization (EM) procedure used to build the final model containing k centers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either random or kmeans++.</td>
</tr>
<tr>
<td></td>
<td>Default value: kmeans++</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>local_lloyd_max_iter</strong></td>
<td>The maximum number of iterations for Lloyd's expectation-maximization (EM) procedure used to build the final model containing ( k ) centers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 300</td>
</tr>
<tr>
<td><strong>local_lloyd_num_trials</strong></td>
<td>The number of times the Lloyd's expectation-maximization (EM) procedure with the least loss is run when building the final model containing ( k ) centers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Either a positive integer or auto.</td>
</tr>
<tr>
<td></td>
<td>Default value: auto</td>
</tr>
<tr>
<td><strong>local_lloyd_tol</strong></td>
<td>The tolerance for change in loss for early stopping of Lloyd's expectation-maximization (EM) procedure used to build the final model containing ( k ) centers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float. Range in [0, 1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0001</td>
</tr>
<tr>
<td><strong>mini_batch_size</strong></td>
<td>The number of observations per mini-batch for the data iterator.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5000</td>
</tr>
</tbody>
</table>
Tune a K-Means Model

Automatic model tuning, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

The Amazon SageMaker k-means algorithm is an unsupervised algorithm that groups data into clusters whose members are as similar as possible. Because it is unsupervised, it doesn't use a validation dataset that hyperparameters can optimize against. But it does take a test dataset and emits metrics that depend on the squared distance between the data points and the final cluster centroids at the end of each training run. To find the model that reports the tightest clusters on the test dataset, you can use a hyperparameter tuning job. The clusters optimize the similarity of their members.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Metrics Computed by the K-Means Algorithm

The k-means algorithm computes the following metrics during training. When tuning a model, choose one of these metrics as the objective metric.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:msd</td>
<td>Mean squared distances between each record in the test set and the closest center of the model.</td>
<td>Minimize</td>
</tr>
<tr>
<td>test:ssd</td>
<td>Sum of the squared distances between each record in the test set and the closest center of the model.</td>
<td>Minimize</td>
</tr>
</tbody>
</table>

Tunable K-Means Hyperparameters

Tune the Amazon SageMaker k-means model with the following hyperparameters. The hyperparameters that have the greatest impact on k-means objective metrics are:
mini_batch_size, extra_center_factor, and init_method. Tuning the hyperparameter epochs generally results in minor improvements.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>epochs</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 10</td>
</tr>
<tr>
<td>extra_center_factor</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 4, MaxValue: 10</td>
</tr>
<tr>
<td>init_method</td>
<td>CategoricalParameterRanges</td>
<td>['kmeans++', 'random']</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 3000, MaxValue: 15000</td>
</tr>
</tbody>
</table>

**K-Means Response Formats**

All SageMaker built-in algorithms adhere to the common input inference format described in Common Data Formats - Inference. This topic contains a list of the available output formats for the SageMaker k-means algorithm.

**JSON Response Format**

```json
{
    "predictions": [
        {
            "closest_cluster": 1.0,
            "distance_to_cluster": 3.0,
        },
        {
            "closest_cluster": 2.0,
            "distance_to_cluster": 5.0,
        },
        // More predictions...
    ]
}
```
JSONLINES Response Format

```json
{
  "closest_cluster": 1.0,
  "distance_to_cluster": 3.0
}
{
  "closest_cluster": 2.0,
  "distance_to_cluster": 5.0
}
```

RECORDIO Response Format

```json
[
  Record = {
    features = {},
    label = {
      'closest_cluster': {
        keys: [],
        values: [1.0, 2.0]  # float32
      },
      'distance_to_cluster': {
        keys: [],
        values: [3.0, 5.0]  # float32
      }
    }
  }
]
```

CSV Response Format

The first value in each line corresponds to `closest_cluster`.

The second value in each line corresponds to `distance_to_cluster`.

```
1.0,3.0
2.0,5.0
```

Principal Component Analysis (PCA) Algorithm

PCA is an unsupervised machine learning algorithm that attempts to reduce the dimensionality (number of features) within a dataset while still retaining as much information as possible. This is done by finding a new set of features called *components*, which are composites of the original features that are uncorrelated with one another. They are also constrained so that the first component accounts for the largest possible variability in the data, the second component the second most variability, and so on.
In Amazon SageMaker, PCA operates in two modes, depending on the scenario:

- **regular**: For datasets with sparse data and a moderate number of observations and features.
- **randomized**: For datasets with both a large number of observations and features. This mode uses an approximation algorithm.

PCA uses tabular data.

The rows represent observations you want to embed in a lower dimensional space. The columns represent features that you want to find a reduced approximation for. The algorithm calculates the covariance matrix (or an approximation thereof in a distributed manner), and then performs the singular value decomposition on this summary to produce the principal components.

**Topics**

- **Input/Output Interface for the PCA Algorithm**
- **EC2 Instance Recommendation for the PCA Algorithm**
- **PCA Sample Notebooks**
- **How PCA Works**
- **PCA Hyperparameters**
- **PCA Response Formats**

**Input/Output Interface for the PCA Algorithm**

For training, PCA expects data provided in the train channel, and optionally supports a dataset passed to the test dataset, which is scored by the final algorithm. Both `recordIO-wrapped-protobuf` and CSV formats are supported for training. You can use either File mode or Pipe mode to train models on data that is formatted as `recordIO-wrapped-protobuf` or as CSV.

For inference, PCA supports `text/csv`, `application/json`, and `application/x-recordio-protobuf`. Results are returned in either `application/json` or `application/x-recordio-protobuf` format with a vector of "projections."

For more information on input and output file formats, see [PCA Response Formats](#) for inference and the [PCA Sample Notebooks](#).
EC2 Instance Recommendation for the PCA Algorithm

PCA supports CPU and GPU instances for training and inference. Which instance type is most performant depends heavily on the specifics of the input data. For GPU instances, PCA supports P2, P3, G4dn, and G5.

PCA Sample Notebooks

For a sample notebook that shows how to use the SageMaker Principal Component Analysis algorithm to analyze the images of handwritten digits from zero to nine in the MNIST dataset, see An Introduction to PCA with MNIST. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. The topic modeling example notebooks using the NTM algorithms are located in the Introduction to Amazon algorithms section. To open a notebook, click on its Use tab and select Create copy.

How PCA Works

Principal Component Analysis (PCA) is a learning algorithm that reduces the dimensionality (number of features) within a dataset while still retaining as much information as possible.

PCA reduces dimensionality by finding a new set of features called components, which are composites of the original features, but are uncorrelated with one another. The first component accounts for the largest possible variability in the data, the second component the second most variability, and so on.

It is an unsupervised dimensionality reduction algorithm. In unsupervised learning, labels that might be associated with the objects in the training dataset aren't used.

Given the input of a matrix with rows

\[ \mathbf{x}_1, \ldots, \mathbf{x}_n \]

each of dimension 1 \* d, the data is partitioned into mini-batches of rows and distributed among the training nodes (workers). Each worker then computes a summary of its data. The summaries of the different workers are then unified into a single solution at the end of the computation.

Modes

The Amazon SageMaker PCA algorithm uses either of two modes to calculate these summaries, depending on the situation:
• **regular**: for datasets with sparse data and a moderate number of observations and features.

• **randomized**: for datasets with both a large number of observations and features. This mode uses an approximation algorithm.

As the algorithm's last step, it performs the singular value decomposition on the unified solution, from which the principal components are then derived.

### Mode 1: Regular

The workers jointly compute both
\[
\sum x_i^T x_i
\]
and
\[
\sum x_i
\].

#### Note
Because
\[x_i\]
are \(1 \times d\) row vectors,
\[
x_i^T x_i\]
is a matrix (not a scalar). Using row vectors within the code allows us to obtain efficient caching.

The covariance matrix is computed as
\[
\sum x_i^T x_i - \frac{1}{n} (\sum x_i)^T \sum x_i
\]
, and its top \(\text{num}_\text{components}\) singular vectors form the model.

#### Note
If \(\text{subtract}_\text{mean}\) is \(\text{False}\), we avoid computing and subtracting
\[
\sum x_i
\].
Use this algorithm when the dimension $d$ of the vectors is small enough so that $d^3$ can fit in memory.

**Mode 2: Randomized**

When the number of features in the input dataset is large, we use a method to approximate the covariance metric. For every mini-batch $x_i$ of dimension $b \times d$, we randomly initialize a $(\text{num\_components} + \text{extra\_components}) \times b$ matrix that we multiply by each mini-batch, to create a $(\text{num\_components} + \text{extra\_components}) \times d$ matrix. The sum of these matrices is computed by the workers, and the servers perform SVD on the final $(\text{num\_components} + \text{extra\_components}) \times d$ matrix. The top right $\text{num\_components}$ singular vectors of it are the approximation of the top singular vectors of the input matrix.

Let
\[
\ell = \text{num\_components} + \text{extra\_components}.
\]

Given a mini-batch $x_i$ of dimension $b \times d$, the worker draws a random matrix $H_i$ of dimension $\ell \times b$.

Depending on whether the environment uses a GPU or CPU and the dimension size, the matrix is either a random sign matrix where each entry is $\pm 1$ or a FJLT (fast Johnson Lindenstrauss transform; for information, see FJLT Transforms and the follow-up papers). The worker then computes $H_i x_i$ and maintains

\[
B = \sum H_i x_i.
\]

The worker also maintains

\[
h^T
\]

, the sum of columns of $H_1, \ldots, H_T$ (T being the total number of mini-batches), and $s$, the sum of all input rows. After processing the entire shard of data, the worker sends the server $B$, $h$, $s$, and $n$ (the number of input rows).
Denote the different inputs to the server as $B^1, h^1, s^1, n^1$.

The server computes $B, h, s, n$ the sums of the respective inputs. It then computes $C = B - (1/n)h^T s$

, and finds its singular value decomposition. The top-right singular vectors and singular values of $C$ are used as the approximate solution to the problem.

### PCA Hyperparameters

In the `CreateTrainingJob` request, you specify the training algorithm. You can also specify algorithm-specific HyperParameters as string-to-string maps. The following table lists the hyperparameters for the PCA training algorithm provided by Amazon SageMaker. For more information about how PCA works, see [How PCA Works](#).

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>feature_dim</td>
<td>Input dimension.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>Number of rows in a mini-batch.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>num_components</td>
<td>The number of principal components to compute.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>algorithm_mode</td>
<td>Mode for computing the principal components.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: <em>regular</em> or <em>randomized</em></td>
</tr>
<tr>
<td></td>
<td>Default value: <em>regular</em></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>extra_components</td>
<td>As the value increases, the solution becomes more accurate but the runtime and memory consumption increase linearly. The default, -1, means the maximum of 10 and num_components. Valid for randomized mode only.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Non-negative integer or -1</td>
</tr>
<tr>
<td></td>
<td>Default value: -1</td>
</tr>
<tr>
<td>subtract_mean</td>
<td>Indicates whether the data should be unbiased both during training and at inference.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: One of true or false</td>
</tr>
<tr>
<td></td>
<td>Default value: true</td>
</tr>
</tbody>
</table>

**PCA Response Formats**

All Amazon SageMaker built-in algorithms adhere to the common input inference format described in [Common Data Formats - Inference](#). This topic contains a list of the available output formats for the SageMaker PCA algorithm.

**JSON Response Format**

Accept—application/json

```json
{
    "projections": [
        {
            "projection": [1.0, 2.0, 3.0, 4.0, 5.0]
        },
        {
            "projection": [6.0, 7.0, 8.0, 9.0, 0.0]
        },
        ...
    ]
}
```
Random Cut Forest (RCF) Algorithm

Amazon SageMaker Random Cut Forest (RCF) is an unsupervised algorithm for detecting anomalous data points within a data set. These are observations which diverge from otherwise well-structured or patterned data. Anomalies can manifest as unexpected spikes in time series data, breaks in periodicity, or unclassifiable data points. They are easy to describe in that, when
viewed in a plot, they are often easily distinguishable from the "regular" data. Including these anomalies in a data set can drastically increase the complexity of a machine learning task since the "regular" data can often be described with a simple model.

With each data point, RCF associates an anomaly score. Low score values indicate that the data point is considered "normal." High values indicate the presence of an anomaly in the data. The definitions of "low" and "high" depend on the application but common practice suggests that scores beyond three standard deviations from the mean score are considered anomalous.

While there are many applications of anomaly detection algorithms to one-dimensional time series data such as traffic volume analysis or sound volume spike detection, RCF is designed to work with arbitrary-dimensional input. Amazon SageMaker RCF scales well with respect to number of features, data set size, and number of instances.

Topics

• Input/Output Interface for the RCF Algorithm
• Instance Recommendations for the RCF Algorithm
• RCF Sample Notebooks
• How RCF Works
• RCF Hyperparameters
• Tune an RCF Model
• RCF Response Formats

Input/Output Interface for the RCF Algorithm

Amazon SageMaker Random Cut Forest supports the train and test data channels. The optional test channel is used to compute accuracy, precision, recall, and F1-score metrics on labeled data. Train and test data content types can be either application/x-recordio-protobuf or text/csv formats. For the test data, when using text/csv format, the content must be specified as text/csv;label_size=1 where the first column of each row represents the anomaly label: "1" for an anomalous data point and "0" for a normal data point. You can use either File mode or Pipe mode to train RCF models on data that is formatted as recordIO-wrapped-protobuf or as CSV.

The train channel only supports S3DataDistributionType=ShardedByS3Key and the test channel only supports S3DataDistributionType=FullyReplicated. The following example specifies the S3 distribution type for the train channel using the Amazon SageMaker Python SDK.
**Note**

The `sagemaker.inputs.s3_input` method was renamed to `sagemaker.inputs.TrainingInput` in SageMaker Python SDK v2.

```python
import sagemaker

# specify Random Cut Forest training job information and hyperparameters
rcf = sagemaker.estimator.Estimator(...)

# explicitly specify "ShardedByS3Key" distribution type
train_data = sagemaker.inputs.TrainingInput(
    s3_data=s3_training_data_location,
    content_type='text/csv;label_size=0',
    distribution='ShardedByS3Key')

# run the training job on input data stored in S3
rcf.fit({'train': train_data})
```

To avoid common errors around execution roles, ensure that you have the execution roles required, `AmazonSageMakerFullAccess` and `AmazonEC2ContainerRegistryFullAccess`. To avoid common errors around your image not existing or its permissions being incorrect, ensure that your ECR image is not larger than the allocated disk space on the training instance. To avoid this, run your training job on an instance that has sufficient disk space. In addition, if your ECR image is from a different AWS account's Elastic Container Service (ECS) repository, and you do not set repository permissions to grant access, this will result in an error. See the [ECR repository permissions](#) for more information on setting a repository policy statement.

See the [S3DataSource](#) for more information on customizing the S3 data source attributes. Finally, in order to take advantage of multi-instance training the training data must be partitioned into at least as many files as instances.

For inference, RCF supports `application/x-recordio-protobuf`, `text/csv` and `application/json` input data content types. See the [Common Data Formats for Built-in Algorithms](#) documentation for more information. RCF inference returns `application/x-recordio-protobuf` or `application/json` formatted output. Each record in these output data contains the corresponding anomaly scores for each input data point. See [Common Data Formats--Inference](#) for more information.
For more information on input and output file formats, see RCF Response Formats for inference and the RCF Sample Notebooks.

Instance Recommendations for the RCF Algorithm

For training, we recommend the ml.m4, ml.c4, and ml.c5 instance families. For inference we recommend using a ml.c5.xl instance type in particular, for maximum performance as well as minimized cost per hour of usage. Although the algorithm could technically run on GPU instance types it does not take advantage of GPU hardware.

RCF Sample Notebooks

For an example of how to train an RCF model and perform inferences with it, see the An Introduction to SageMaker Random Cut Forests notebook. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. To open a notebook, click on its Use tab and select Create copy.

How RCF Works

Amazon SageMaker Random Cut Forest (RCF) is an unsupervised algorithm for detecting anomalous data points within a dataset. These are observations which diverge from otherwise well-structured or patterned data. Anomalies can manifest as unexpected spikes in time series data, breaks in periodicity, or unclassifiable data points. They are easy to describe in that, when viewed in a plot, they are often easily distinguishable from the "regular" data. Including these anomalies in a dataset can drastically increase the complexity of a machine learning task since the "regular" data can often be described with a simple model.

The main idea behind the RCF algorithm is to create a forest of trees where each tree is obtained using a partition of a sample of the training data. For example, a random sample of the input data is first determined. The random sample is then partitioned according to the number of trees in the forest. Each tree is given such a partition and organizes that subset of points into a k-d tree. The anomaly score assigned to a data point by the tree is defined as the expected change in complexity of the tree as a result adding that point to the tree; which, in approximation, is inversely proportional to the resulting depth of the point in the tree. The random cut forest assigns an anomaly score by computing the average score from each constituent tree and scaling the result with respect to the sample size. The RCF algorithm is based on the one described in reference [1].
Sample Data Randomly

The first step in the RCF algorithm is to obtain a random sample of the training data. In particular, suppose we want a sample of size $K$ from $N$ total data points. If the training data is small enough, the entire dataset can be used, and we could randomly draw $K$ elements from this set. However, frequently the training data is too large to fit all at once, and this approach isn't feasible. Instead, we use a technique called reservoir sampling.

Reservoir sampling is an algorithm for efficiently drawing random samples from a dataset $S = \{S_1, \ldots, S_N\}$ where the elements in the dataset can only be observed one at a time or in batches. In fact, reservoir sampling works even when $N$ is not known a priori. If only one sample is requested, such as when $K = 1$, the algorithm is like this:

**Algorithm: Reservoir Sampling**

- **Input:** dataset or data stream $S = \{S_1, \ldots, S_N\}$
- **Initialize the random sample** $X = S_1$
- **For each observed sample** $S_n, n = 2, \ldots, N$
  - Pick a uniform random number $\xi \in [0, 1]$
  - **If** $\xi < 1/n$
  - **Set** $X = S_n$
This algorithm selects a random sample such that
\[ P(X = S_n) = 1/N \]
for all
\[ n = 1, \ldots, N \]
When
\[ K > 1 \]
the algorithm is more complicated. Additionally, a distinction must be made between random sampling that is with and without replacement. RCF performs an augmented reservoir sampling without replacement on the training data based on the algorithms described in [2].

**Train a RCF Model and Produce Inferences**

The next step in RCF is to construct a random cut forest using the random sample of data. First, the sample is partitioned into a number of equal-sized partitions equal to the number of trees in the forest. Then, each partition is sent to an individual tree. The tree recursively organizes its partition into a binary tree by partitioning the data domain into bounding boxes.

This procedure is best illustrated with an example. Suppose a tree is given the following two-dimensional dataset. The corresponding tree is initialized to the root node:

A two-dimensional dataset where the majority of data lies in a cluster (blue) except for one anomalous data point (orange). The tree is initialized with a root node.
The RCF algorithm organizes these data in a tree by first computing a bounding box of the data, selecting a random dimension (giving more weight to dimensions with higher "variance"), and then randomly determining the position of a hyperplane "cut" through that dimension. The two resulting subspaces define their own sub tree. In this example, the cut happens to separate a lone point from the remainder of the sample. The first level of the resulting binary tree consists of two nodes, one which will consist of the subtree of points to the left of the initial cut and the other representing the single point on the right.

A random cut partitioning the two-dimensional dataset. An anomalous data point is more likely to lie isolated in a bounding box at a smaller tree depth than other points.

Bounding boxes are then computed for the left and right halves of the data and the process is repeated until every leaf of the tree represents a single data point from the sample. Note that if the lone point is sufficiently far away then it is more likely that a random cut would result in point isolation. This observation provides the intuition that tree depth is, loosely speaking, inversely proportional to the anomaly score.

When performing inference using a trained RCF model the final anomaly score is reported as the average across scores reported by each tree. Note that it is often the case that the new data point does not already reside in the tree. To determine the score associated with the new point the data point is inserted into the given tree and the tree is efficiently (and temporarily) reassembled in a manner equivalent to the training process described above. That is, the resulting tree is as if the input data point were a member of the sample used to construct the tree in the first place. The reported score is inversely proportional to the depth of the input point within the tree.
Choose Hyperparameters

The primary hyperparameters used to tune the RCF model are `num_trees` and `num_samples_per_tree`. Increasing `num_trees` has the effect of reducing the noise observed in anomaly scores since the final score is the average of the scores reported by each tree. While the optimal value is application-dependent we recommend using 100 trees to begin with as a balance between score noise and model complexity. Note that inference time is proportional to the number of trees. Although training time is also affected it is dominated by the reservoir sampling algorithm describe above.

The parameter `num_samples_per_tree` is related to the expected density of anomalies in the dataset. In particular, `num_samples_per_tree` should be chosen such that $1/num_samples_per_tree$ approximates the ratio of anomalous data to normal data. For example, if 256 samples are used in each tree then we expect our data to contain anomalies $1/256$ or approximately 0.4% of the time. Again, an optimal value for this hyperparameter is dependent on the application.

References


RCF Hyperparameters

In the `CreateTrainingJob` request, you specify the training algorithm. You can also specify algorithm-specific hyperparameters as string-to-string maps. The following table lists the hyperparameters for the Amazon SageMaker RCF algorithm. For more information, including recommendations on how to choose hyperparameters, see *How RCF Works*. 
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| feature_dim                | The number of features in the data set. (If you use the Random Cut Forest estimator, this value is calculated for you and need not be specified.)  
**Required**  
Valid values: Positive integer (min: 1, max: 10000)                                                                                                                                                                                                                       |
| eval_metrics               | A list of metrics used to score a labeled test data set. The following metrics can be selected for output:  
- accuracy - returns fraction of correct predictions.  
- precision_recall_fscore - returns the positive and negative precision, recall, and F1-scores.  
**Optional**  
Valid values: a list with possible values taken from accuracy or precision_recall_fscore.  
Default value: Both accuracy, precision_recall_fscore are calculated.                                                                                                                                                                                                 |
| num_samples_per_tree       | Number of random samples given to each tree from the training data set.  
**Optional**  
Valid values: Positive integer (min: 1, max: 2048)  
Default value: 256                                                                                                                                                                                                                                                       |
| num_trees                  | Number of trees in the forest.  
**Optional**  
Valid values: Positive integer (min: 50, max: 1000)                                                                                                                                                                                                                        |
Tune an RCF Model

*Automatic model tuning*, also known as hyperparameter tuning or hyperparameter optimization, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

The Amazon SageMaker RCF algorithm is an unsupervised anomaly-detection algorithm that requires a labeled test dataset for hyperparameter optimization. RCF calculates anomaly scores for test data points and then labels the data points as anomalous if their scores are beyond three standard deviations from the mean score. This is known as the three-sigma limit heuristic. The F1-score is based on the difference between calculated labels and actual labels. The hyperparameter tuning job finds the model that maximizes that score. The success of hyperparameter optimization depends on the applicability of the three-sigma limit heuristic to the test dataset.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

**Metrics Computed by the RCF Algorithm**

The RCF algorithm computes the following metric during training. When tuning the model, choose this metric as the objective metric.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>test:f1</td>
<td>F1-score on the test dataset, based on the difference between calculated labels and actual labels.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

**Tunable RCF Hyperparameters**

You can tune a RCF model with the following hyperparameters.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_samples_per_tree</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 2048</td>
</tr>
<tr>
<td>num_trees</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 50, MaxValue: 1000</td>
</tr>
</tbody>
</table>

**RCF Response Formats**

All Amazon SageMaker built-in algorithms adhere to the common input inference format described in [Common Data Formats - Inference](#). Note that SageMaker Random Cut Forest supports both dense and sparse JSON and RecordIO formats. This topic contains a list of the available output formats for the SageMaker RCF algorithm.

**JSON Response Format**

ACCEPT: application/json.

```json
{
    "scores": [
        {
            "score": 0.02
        },
        {
            "score": 0.25
        }
    ]
}
```
JSONLINES Response Format

ACCEPT: application/jsonlines.

{
"score": 0.02,
"score": 0.25
}

RECORDIO Response Format

ACCEPT: application/x-recordio-protobuf.

[ 
  Record = {
    features = {},
    label = {
      'score': {
        keys: [],
        values: [0.25]  # float32
      }
    }
  }
]
Record = {

features = {},

label = {

'score': {

    keys: [],

    values: [0.23]  # float32
}

}

}
Built-in SageMaker Algorithms for Computer Vision

SageMaker provides image processing algorithms that are used for image classification, object detection, and computer vision.

- **Image Classification - MXNet**—uses example data with answers (referred to as a *supervised algorithm*). Use this algorithm to classify images.

- **Image Classification - TensorFlow**—uses pretrained TensorFlow Hub models to fine-tune for specific tasks (referred to as a *supervised algorithm*). Use this algorithm to classify images.

- **Object Detection - MXNet**—detects and classifies objects in images using a single deep neural network. It is a supervised learning algorithm that takes images as input and identifies all instances of objects within the image scene.

- **Object Detection - TensorFlow**—detects bounding boxes and object labels in an image. It is a supervised learning algorithm that supports transfer learning with available pretrained TensorFlow models.

- **Semantic Segmentation Algorithm**—provides a fine-grained, pixel-level approach to developing computer vision applications.

<table>
<thead>
<tr>
<th>Algorithm name</th>
<th>Channel name</th>
<th>Training input mode</th>
<th>File type</th>
<th>Instance class</th>
<th>Parallelizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image Classification - MXNet</td>
<td>train and validation, (optionally) train_lst, validation_lst, and model</td>
<td>File or Pipe</td>
<td>recordIO or image files (.jpg or .png)</td>
<td>GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
<td>Instance class</td>
<td>Parallelizable</td>
</tr>
<tr>
<td>------------------------------</td>
<td>---------------------------------------</td>
<td>---------------------</td>
<td>--------------------------</td>
<td>----------------</td>
<td>-----------------------------------------------------</td>
</tr>
<tr>
<td>Image Classification - TensorFlow</td>
<td>training and validation</td>
<td>File</td>
<td>image files (.jpg, .jpeg, or .png)</td>
<td>CPU or GPU</td>
<td>Yes (only across multiple GPUs on a single instance)</td>
</tr>
<tr>
<td>Object Detection</td>
<td>train and validation, (optionally)</td>
<td>File or Pipe</td>
<td>recordIO or image files (.jpg or .png)</td>
<td>GPU</td>
<td>Yes</td>
</tr>
<tr>
<td>Object Detection - TensorFlow</td>
<td>training and validation</td>
<td>File</td>
<td>image files (.jpg, .jpeg, or .png)</td>
<td>GPU</td>
<td>Yes (only across multiple GPUs on a single instance)</td>
</tr>
<tr>
<td>Algorithm name</td>
<td>Channel name</td>
<td>Training input mode</td>
<td>File type</td>
<td>Instance class</td>
<td>Parallelizable</td>
</tr>
<tr>
<td>---------------</td>
<td>--------------</td>
<td>---------------------</td>
<td>-----------</td>
<td>----------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Semantic Segmentation</td>
<td>train and validation, train_annotation, validation, and optionally label_map and model</td>
<td>File or Pipe</td>
<td>Image files</td>
<td>GPU (single instance only)</td>
<td>No</td>
</tr>
</tbody>
</table>

**Image Classification - MXNet**

The Amazon SageMaker image classification algorithm is a supervised learning algorithm that supports multi-label classification. It takes an image as input and outputs one or more labels assigned to that image. It uses a convolutional neural network that can be trained from scratch or trained using transfer learning when a large number of training images are not available.

The recommended input format for the Amazon SageMaker image classification algorithms is Apache MXNet RecordIO. However, you can also use raw images in .jpg or .png format. Refer to this discussion for a broad overview of efficient data preparation and loading for machine learning systems.

**Note**

To maintain better interoperability with existing deep learning frameworks, this differs from the protobuf data formats commonly used by other Amazon SageMaker algorithms.

For more information on convolutional networks, see:

- [Deep residual learning for image recognition](#) Kaiming He, et al., 2016 IEEE Conference on Computer Vision and Pattern Recognition
Input/Output Interface for the Image Classification Algorithm

The SageMaker Image Classification algorithm supports both RecordIO (application/x-recordio) and image (image/png, image/jpeg, and application/x-image) content types for training in file mode, and supports the RecordIO (application/x-recordio) content type for training in pipe mode. However, you can also train in pipe mode using the image files (image/png, image/jpeg, and application/x-image), without creating RecordIO files, by using the augmented manifest format.

Distributed training is supported for file mode and pipe mode. When using the RecordIO content type in pipe mode, you must set the S3DataDistributionType of the S3DataSource to FullyReplicated. The algorithm supports a fully replicated model where your data is copied onto each machine.

The algorithm supports image/png, image/jpeg, and application/x-image for inference.

Train with RecordIO Format

If you use the RecordIO format for training, specify both train and validation channels as values for the InputDataConfig parameter of the CreateTrainingJob request. Specify one RecordIO (.rec) file in the train channel and one RecordIO file in the validation channel. Set the content type for both channels to application/x-recordio.
Train with Image Format

If you use the Image format for training, specify train, validation, train_lst, and validation_lst channels as values for the InputDataConfig parameter of the CreateTrainingJob request. Specify the individual image data (.jpg or .png files) for the train and validation channels. Specify one .lst file in each of the train_lst and validation_lst channels. Set the content type for all four channels to application/x-image.

Note
SageMaker reads the training and validation data separately from different channels, so you must store the training and validation data in different folders.

A .lst file is a tab-separated file with three columns that contains a list of image files. The first column specifies the image index, the second column specifies the class label index for the image, and the third column specifies the relative path of the image file. The image index in the first column must be unique across all of the images. The set of class label indices are numbered successively and the numbering should start with 0. For example, 0 for the cat class, 1 for the dog class, and so on for additional classes.

The following is an example of a .lst file:

```
5    1   your_image_directory/train_img_dog1.jpg
1000 0   your_image_directory/train_img_cat1.jpg
22   1   your_image_directory/train_img_dog2.jpg
```

For example, if your training images are stored in s3://<your_bucket>/train/class_dog, s3://<your_bucket>/train/class_cat, and so on, specify the path for your train channel as s3://<your_bucket>/train, which is the top-level directory for your data. In the .lst file, specify the relative path for an individual file named train_image_dog1.jpg in the class_dog class directory as class_dog/train_image_dog1.jpg. You can also store all your image files under one subdirectory inside the train directory. In that case, use that subdirectory for the relative path. For example, s3://<your_bucket>/train/your_image_directory.

Train with Augmented Manifest Image Format

The augmented manifest format enables you to do training in Pipe mode using image files without needing to create RecordIO files. You need to specify both train and validation channels as values
for the InputDataConfig parameter of the CreateTrainingJob request. While using the format, an S3 manifest file needs to be generated that contains the list of images and their corresponding annotations. The manifest file format should be in JSON Lines format in which each line represents one sample. The images are specified using the 'source-ref' tag that points to the S3 location of the image. The annotations are provided under the "AttributeNames" parameter value as specified in the CreateTrainingJob request. It can also contain additional metadata under the metadata tag, but these are ignored by the algorithm. In the following example, the "AttributeNames" are contained in the list of image and annotation references ["source-ref", "class"]. The corresponding label value is "0" for the first image and “1” for the second image:

```json
{"source-ref":"s3://image/filename1.jpg",  "class":"0"}
{"source-ref":"s3://image/filename2.jpg",  "class":"1",  "class-metadata": {"class-name":  "cat",  "type" : "groundtruth/image-classification"}}
```

The order of "AttributeNames" in the input files matters when training the ImageClassification algorithm. It accepts piped data in a specific order, with image first, followed by label. So the "AttributeNames" in this example are provided with "source-ref" first, followed by "class". When using the ImageClassification algorithm with Augmented Manifest, the value of the RecordWrapperType parameter must be "RecordIO".

Multi-label training is also supported by specifying a JSON array of values. The num_classes hyperparameter must be set to match the total number of classes. There are two valid label formats: multi-hot and class-id.

In the multi-hot format, each label is a multi-hot encoded vector of all classes, where each class takes the value of 0 or 1. In the following example, there are three classes. The first image is labeled with classes 0 and 2, while the second image is labeled with class 2 only:

```json
{"image-ref": "s3://mybucket/sample01/image1.jpg",  "class": "[1, 0, 1]"}
{"image-ref": "s3://mybucket/sample02/image2.jpg",  "class": "[0, 0, 1]"}
```

In the class-id format, each label is a list of the class ids, from [0, num_classes), which apply to the data point. The previous example would instead look like this:

```json
{"image-ref": "s3://mybucket/sample01/image1.jpg",  "class": "[0, 2]"}
{"image-ref": "s3://mybucket/sample02/image2.jpg",  "class": "[2]"}
```
The multi-hot format is the default, but can be explicitly set in the content type with the label-format parameter: "application/x-recordio; label-format=multi-hot". The class-id format, which is the format outputted by GroundTruth, must be set explicitly: "application/x-recordio; label-format=class-id".

For more information on augmented manifest files, see Provide Dataset Metadata to Training Jobs with an Augmented Manifest File.

Incremental Training

You can also seed the training of a new model with the artifacts from a model that you trained previously with SageMaker. Incremental training saves training time when you want to train a new model with the same or similar data. SageMaker image classification models can be seeded only with another built-in image classification model trained in SageMaker.

To use a pretrained model, in the CreateTrainingJob request, specify the ChannelName as "model" in the InputDataConfig parameter. Set the ContentType for the model channel to application/x-sagemaker-model. The input hyperparameters of both the new model and the pretrained model that you upload to the model channel must have the same settings for the num_layers, image_shape and num_classes input parameters. These parameters define the network architecture. For the pretrained model file, use the compressed model artifacts (in .tar.gz format) output by SageMaker. You can use either RecordIO or image formats for input data.

Inference with the Image Classification Algorithm

The generated models can be hosted for inference and support encoded .jpg and .png image formats as image/png, image/jpeg, and application/x-image content-type. The input image is resized automatically. The output is the probability values for all classes encoded in JSON format, or in JSON Lines text format for batch transform. The image classification model processes a single image per request and so outputs only one line in the JSON or JSON Lines format. The following is an example of a response in JSON Lines format:

```json
accept: application/jsonlines

{"prediction": [prob_0, prob_1, prob_2, prob_3, ...]}
```

For more details on training and inference, see the image classification sample notebook instances referenced in the introduction.
EC2 Instance Recommendation for the Image Classification Algorithm

For image classification, we support P2, P3, G4dn, and G5 instances. We recommend using GPU instances with more memory for training with large batch sizes. You can also run the algorithm on multi-GPU and multi-machine settings for distributed training. Both CPU (such as C4) and GPU (P2, P3, G4dn, or G5) instances can be used for inference.

Image Classification Sample Notebooks

For a sample notebook that uses the SageMaker image classification algorithm, see Build and Register an MXNet Image Classification Model via SageMaker Pipelines. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. The example image classification notebooks are located in the Introduction to Amazon algorithms section. To open a notebook, click on its Use tab and select Create copy.

How Image Classification Works

The image classification algorithm takes an image as input and classifies it into one of the output categories. Deep learning has revolutionized the image classification domain and has achieved great performance. Various deep learning networks such as ResNet, DenseNet, Inception, and so on, have been developed to be highly accurate for image classification. At the same time, there have been efforts to collect labeled image data that are essential for training these networks. ImageNet is one such large dataset that has more than 11 million images with about 11,000 categories. Once a network is trained with ImageNet data, it can then be used to generalize with other datasets as well, by simple re-adjustment or fine-tuning. In this transfer learning approach, a network is initialized with weights (in this example, trained on ImageNet), which can be later fine-tuned for an image classification task in a different dataset.

Image classification in Amazon SageMaker can be run in two modes: full training and transfer learning. In full training mode, the network is initialized with random weights and trained on user data from scratch. In transfer learning mode, the network is initialized with pre-trained weights and just the top fully connected layer is initialized with random weights. Then, the whole network is fine-tuned with new data. In this mode, training can be achieved even with a smaller dataset. This is because the network is already trained and therefore can be used in cases without sufficient training data.
# Image Classification Hyperparameters

Hyperparameters are parameters that are set before a machine learning model begins learning. The following hyperparameters are supported by the Amazon SageMaker built-in Image Classification algorithm. See [Tune an Image Classification Model](#) for information on image classification hyperparameter tuning.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| num_classes            | Number of output classes. This parameter defines the dimensions of the network output and is typically set to the number of classes in the dataset. Besides multi-class classification, multi-label classification is supported too. Please refer to [Input/Output Interface for the Image Classification Algorithm](#) for details on how to work with multi-label classification with augmented manifest files. Required
|                        | Valid values: positive integer                                                                                                                                                                               |
| num_training_samples   | Number of training examples in the input dataset. If there is a mismatch between this value and the number of samples in the training set, then the behavior of the lr_scheduler_step parameter is undefined and distributed training accuracy might be affected. Required
|                        | Valid values: positive integer                                                                                                                                                                               |
| augmentation_type      | Data augmentation type. The input images can be augmented in multiple ways as specified below.                                                                                                              |
|                        | • crop: Randomly crop the image and flip the image horizontally  
|                        | • crop_color: In addition to 'crop', three random values in the range [-36, 36], [-50, 50], and [-50, 50] are added to the image.  

---

Use Built-in Algorithms
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>corresponding Hue-Saturation-Lightness channels respectively</td>
</tr>
<tr>
<td></td>
<td>• <strong>crop_color_transform</strong>: In addition to <strong>crop_color</strong>, random transformations, including rotation, shear, and aspect ratio variations are applied to the image. The maximum angle of rotation is 10 degrees, the maximum shear ratio is 0.1, and the maximum aspect changing ratio is 0.25.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: <strong>crop</strong>, <strong>crop_color</strong>, or <strong>crop_color_transform</strong>.</td>
</tr>
<tr>
<td></td>
<td>Default value: no default value</td>
</tr>
<tr>
<td><strong>beta_1</strong></td>
<td>The beta1 for adam, that is the exponential decay rate for the first moment estimates.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float. Range in [0, 1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.9</td>
</tr>
<tr>
<td><strong>beta_2</strong></td>
<td>The beta2 for adam, that is the exponential decay rate for the second moment estimates.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float. Range in [0, 1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.999</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>checkpoint_frequency</td>
<td>Period to store model parameters (in number of epochs). Note that all checkpoint files are saved as part of the final model file &quot;model.tar.gz&quot; and uploaded to S3 to the specified model location. This increases the size of the model file proportionally to the number of checkpoints saved during training.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer no greater than epochs.</td>
</tr>
<tr>
<td></td>
<td>Default value: no default value (Save checkpoint at the epoch that has the best validation accuracy)</td>
</tr>
<tr>
<td>early_stopping</td>
<td>True to use early stopping logic during training. False not to use it.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: True or False</td>
</tr>
<tr>
<td></td>
<td>Default value: False</td>
</tr>
<tr>
<td>early_stopping_min_epochs</td>
<td>The minimum number of epochs that must be run before the early stopping logic can be invoked. It is used only when early_stopping = True.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>The number of epochs to wait before ending training if no improvement is made in the relevant metric. It is used only when early_stopping = True. Optional. Valid values: positive integer. Default value: 5.</td>
</tr>
<tr>
<td>early_stopping_tolerance</td>
<td>Relative tolerance to measure an improvement in accuracy validation metric. If the ratio of the improvement in accuracy divided by the previous best accuracy is smaller than the early_stopping_tolerance value set, early stopping considers there is no improvement. It is used only when early_stopping = True. Optional. Valid values: 0 ≤ float ≤ 1. Default value: 0.0.</td>
</tr>
<tr>
<td>epochs</td>
<td>Number of training epochs. Optional. Valid values: positive integer. Default value: 30.</td>
</tr>
<tr>
<td>eps</td>
<td>The epsilon for adam and rmsprop. It is usually set to a small value to avoid division by 0. Optional. Valid values: float. Range in [0, 1]. Default value: 1e-8.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| gamma          | The gamma for `rmsprop`, the decay factor for the moving average of the squared gradient.  
**Optional** 
Valid values: float. Range in [0, 1].  
Default value: 0.9 |
| image_shape    | The input image dimensions, which is the same size as the input layer of the network. The format is defined as `num_channels, height, width`. The image dimension can take on any value as the network can handle varied dimensions of the input. However, there may be memory constraints if a larger image dimension is used. Pretrained models can use only a fixed 224 x 224 image size. Typical image dimensions for image classification are `'3,224,224'`. This is similar to the ImageNet dataset.  
For training, if any input image is smaller than this parameter in any dimension, training fails. If an image is larger, a portion of the image is cropped, with the cropped area specified by this parameter. If hyperparameter `augmentation_type` is set, random crop is taken; otherwise, central crop is taken.  
At inference, input images are resized to the `image_shape` that was used during training. Aspect ratio is not preserved, and images are not cropped.  
**Optional** 
Valid values: string  
Default value: ‘3,224,224’ |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>kv_store</strong></td>
<td>Weight update synchronization mode during distributed training. The weight updates can be updated either synchronously or asynchronously across machines. Synchronous updates typically provide better accuracy than asynchronous updates but can be slower. See distributed training in MXNet for more details. This parameter is not applicable to single machine training.</td>
</tr>
<tr>
<td></td>
<td>• <strong>dist_sync</strong> : The gradients are synchronized after every batch with all the workers. With <code>dist_sync</code>, batch-size now means the batch size used on each machine. So if there are n machines and we use batch size b, then <code>dist_sync</code> behaves like local with batch size n*b</td>
</tr>
<tr>
<td></td>
<td>• <strong>dist_async</strong> : Performs asynchronous updates. The weights are updated whenever gradients are received from any machine and the weight updates are atomic. However, the order is not guaranteed.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: <code>dist_sync</code> or <code>dist_async</code></td>
</tr>
<tr>
<td></td>
<td>Default value: no default value</td>
</tr>
<tr>
<td><strong>learning_rate</strong></td>
<td>Initial learning rate.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float. Range in [0, 1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>lr_scheduler_factor</td>
<td>The ratio to reduce learning rate used in conjunction with the lr_scheduler_step parameter, defined as lr_new = lr_old * lr_scheduler_factor.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float. Range in [0, 1].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.1</td>
</tr>
<tr>
<td>lr_scheduler_step</td>
<td>The epochs at which to reduce the learning rate. As explained in the lr_scheduler_factor parameter, the learning rate is reduced by lr_scheduler_factor at these epochs. For example, if the value is set to &quot;10, 20&quot;, then the learning rate is reduced by lr_scheduler_factor after 10th epoch and again by lr_scheduler_factor after 20th epoch. The epochs are delimitedor &quot;,&quot;.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: string</td>
</tr>
<tr>
<td></td>
<td>Default value: no default value</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>The batch size for training. In a single-machine multi-GPU setting, each GPU handles mini_batch_size / num_gpu training samples. For the multi-machine training in dist_sync mode, the actual batch size is mini_batch_size * number of machines. See MXNet docs for more details.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 32</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| momentum       | The momentum for sgd and nag, ignored for other optimizers. 

**Optional**

Valid values: float. Range in [0, 1].

Default value: 0.9

| multi_label    | Flag to use for multi-label classification where each sample can be assigned multiple labels. Average accuracy across all classes is logged. 

**Optional**

Valid values: 0 or 1

Default value: 0

| num_layers     | Number of layers for the network. For data with large image size (for example, 224x224 - like ImageNet), we suggest selecting the number of layers from the set [18, 34, 50, 101, 152, 200]. For data with small image size (for example, 28x28 - like CIFAR), we suggest selecting the number of layers from the set [20, 32, 44, 56, 110]. The number of layers in each set is based on the ResNet paper. For transfer learning, the number of layers defines the architecture of base network and hence can only be selected from the set [18, 34, 50, 101, 152, 200]. 

**Optional**

Valid values: positive integer in [18, 34, 50, 101, 152, 200] or [20, 32, 44, 56, 110]

Default value: 152
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| **optimizer** | The optimizer type. For more details of the parameters for the optimizers, please refer to MXNet's API.  
  **Optional**  
  Valid values: One of sgd, adam, rmsprop, or nag.  
  - sgd: *Stochastic gradient descent*  
  - adam: *Adaptive momentum estimation*  
  - rmsprop: *Root mean square propagation*  
  - nag: *Nesterov accelerated gradient*  
  Default value: sgd |
| **precision_dtype** | The precision of the weights used for training. The algorithm can use either single precision (float32) or half precision (float16) for the weights. Using half-precision for weights results in reduced memory consumption.  
  **Optional**  
  Valid values: float32 or float16  
  Default value: float32 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>resize</td>
<td>The number of pixels in the shortest side of an image after resizing it for training. If the parameter is not set, then the training data is used without resizing. The parameter should be larger than both the width and height components of image_shape to prevent training failure.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong> when using image content types</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong> when using the RecordIO content type</td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: no default value</td>
</tr>
<tr>
<td>top_k</td>
<td>Reports the top-k accuracy during training. This parameter has to be greater than 1, since the top-1 training accuracy is the same as the regular training accuracy that has already been reported.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer larger than 1.</td>
</tr>
<tr>
<td></td>
<td>Default value: no default value</td>
</tr>
<tr>
<td>use_pretrained_model</td>
<td>Flag to use pre-trained model for training. If set to 1, then the pretrained model with the corresponding number of layers is loaded and used for training. Only the top FC layer are reinitialized with random weights. Otherwise, the network is trained from scratch.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 or 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
</tbody>
</table>
### Parameter Name | Description
--- | ---
use_weighted_loss | Flag to use weighted cross-entropy loss for multi-label classification (used only when `multi_label = 1`), where the weights are calculated based on the distribution of classes.

**Optional**

Valid values: 0 or 1
Default value: 0

weight_decay | The coefficient weight decay for sgd and nag, ignored for other optimizers.

**Optional**

Valid values: float. Range in [0, 1].
Default value: 0.0001

---

**Tune an Image Classification Model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics Computed by the Image Classification Algorithm**

The image classification algorithm is a supervised algorithm. It reports an accuracy metric that is computed during training. When tuning the model, choose this metric as the objective metric.
## Tunable Image Classification Hyperparameters

Tune an image classification model with the following hyperparameters. The hyperparameters that have the greatest impact on image classification objective metrics are: `mini_batch_size`, `learning_rate`, and `optimizer`. Tune the optimizer-related hyperparameters, such as `momentum`, `weight_decay`, `beta_1`, `beta_2`, `eps`, and `gamma`, based on the selected optimizer. For example, use `beta_1` and `beta_2` only when `adam` is the optimizer.

For more information about which hyperparameters are used in each optimizer, see [Image Classification Hyperparameters](#).

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>beta_1</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
<tr>
<td>beta_2</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
<tr>
<td>eps</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-8, MaxValue: 1.0</td>
</tr>
<tr>
<td>gamma</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-8, MaxValue: 0.999</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.5</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 8, MaxValue: 512</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------</td>
<td>--------------------------</td>
</tr>
<tr>
<td>momentum</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
<tr>
<td>optimizer</td>
<td>CategoricalParameterRanges</td>
<td>['sgd', 'adam', 'rmsprop', 'nag']</td>
</tr>
<tr>
<td>weight_decay</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
</tbody>
</table>

### Image Classification - TensorFlow

The Amazon SageMaker Image Classification - TensorFlow algorithm is a supervised learning algorithm that supports transfer learning with many pretrained models from the TensorFlow Hub. Use transfer learning to fine-tune one of the available pretrained models on your own dataset, even if a large amount of image data is not available. The image classification algorithm takes an image as input and outputs a probability for each provided class label. Training datasets must consist of images in .jpg, .jpeg, or .png format.

### Topics

- [How to use the SageMaker Image Classification - TensorFlow algorithm](#)
- [Input and output interface for the Image Classification - TensorFlow algorithm](#)
- [Amazon EC2 instance recommendation for the Image Classification - TensorFlow algorithm](#)
- [Image Classification - TensorFlow sample notebooks](#)
- [How Image Classification - TensorFlow Works](#)
- [TensorFlow Hub Models](#)
- [Image Classification - TensorFlow Hyperparameters](#)
- [Tune an Image Classification - TensorFlow model](#)

### How to use the SageMaker Image Classification - TensorFlow algorithm

You can use Image Classification - TensorFlow as an Amazon SageMaker built-in algorithm. The following section describes how to use Image Classification - TensorFlow with the SageMaker
Python SDK. For information on how to use Image Classification - TensorFlow from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

The Image Classification - TensorFlow algorithm supports transfer learning using any of the compatible pretrained TensorFlow Hub models. For a list of all available pretrained models, see TensorFlow Hub Models. Every pretrained model has a unique model_id. The following example uses MobileNet V2 1.00 224 (model_id: tensorflow-ic-imagenet-mobilenet-v2-100-224-classification-4) to fine-tune on a custom dataset. The pretrained models are all pre-downloaded from the TensorFlow Hub and stored in Amazon S3 buckets so that training jobs can run in network isolation. Use these pre-generated model training artifacts to construct a SageMaker Estimator.

First, retrieve the Docker image URI, training script URI, and pretrained model URI. Then, change the hyperparameters as you see fit. You can see a Python dictionary of all available hyperparameters and their default values with hyperparameters.retrieve_default. For more information, see Image Classification - TensorFlow Hyperparameters. Use these values to construct a SageMaker Estimator.

![Note]

Default hyperparameter values are different for different models. For larger models, the default batch size is smaller and the train_only_top_layer hyperparameter is set to "True".

This example uses the tf_flowers dataset, which contains five classes of flower images. We pre-downloaded the dataset from TensorFlow under the Apache 2.0 license and made it available with Amazon S3. To fine-tune your model, call .fit using the Amazon S3 location of your training dataset.

```python
from sagemaker import image_uris, model_uris, script_uris, hyperparameters
from sagemaker.estimator import Estimator

model_id, model_version = "tensorflow-ic-imagenet-mobilenet-v2-100-224-classification-4", "*
training_instance_type = "ml.p3.2xlarge"

# Retrieve the Docker image
train_image_uri =
image_uris.retrieve(model_id=model_id,model_version=model_version,image_scope="training",instance_type=training_instance_type)

# Call .fit using the Amazon S3 location of your training dataset
estimator.fit(train_image_uri)
```
# Retrieve the training script
train_source_uri = script_uris.retrieve(model_id=model_id, model_version=model_version,
    script_scope="training")

# Retrieve the pretrained model tarball for transfer learning
train_model_uri = model_uris.retrieve(model_id=model_id, model_version=model_version,
    model_scope="training")

# Retrieve the default hyper-parameters for fine-tuning the model
hyperparameters = hyperparameters.retrieve_default(model_id=model_id,
    model_version=model_version)

# [Optional] Override default hyperparameters with custom values
hyperparameters["epochs"] = "5"

# The sample training data is available in the following S3 bucket
training_data_bucket = f"jumpstart-cache-prod-{aws_region}"
training_data_prefix = "training-datasets/tf_flowers/"

training_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}"

output_bucket = sess.default_bucket()
output_prefix = "jumpstart-example-ic-training"
s3_output_location = f"s3://{output_bucket}/{output_prefix}/output"

# Create SageMaker Estimator instance
tf_ic_estimator = Estimator(
    role=aws_role,
    image_uri=train_image_uri,
    source_dir=train_source_uri,
    model_uri=train_model_uri,
    entry_point="transfer_learning.py",
    instance_count=1,
    instance_type=training_instance_type,
    max_run=360000,
    hyperparameters=hyperparameters,
    output_path=s3_output_location,
)

# Use S3 path of the training data to launch SageMaker TrainingJob
tf_ic_estimator.fit({"training": training_dataset_s3_path}, logs=True)
Input and output interface for the Image Classification - TensorFlow algorithm

Each of the pretrained models listed in TensorFlow Hub Models can be fine-tuned to any dataset with any number of image classes. Be mindful of how to format your training data for input to the Image Classification - TensorFlow model.

- **Training data input format:** Your training data should be a directory with as many subdirectories as the number of classes. Each subdirectory should contain images belonging to that class in .jpg, .jpeg, or .png format.

The following is an example of an input directory structure. This example dataset has two classes: roses and dandelion. The image files in each class folder can have any name. The input directory should be hosted in an Amazon S3 bucket with a path similar to the following: s3://bucket_name/input_directory/. Note that the trailing / is required.

```
input_directory
|-- roses
    |-- abc.jpg
    |-- def.jpg
|-- dandelion
    |-- ghi.jpg
    |-- jkl.jpg
```

Trained models output label mapping files that map class folder names to the indices in the list of output class probabilities. This mapping is in alphabetical order. For example, in the preceding example, the dandelion class is index 0 and the roses class is index 1.

After training, you have a fine-tuned model that you can further train using incremental training or deploy for inference. The Image Classification - TensorFlow algorithm automatically adds a pre-processing and post-processing signature to the fine-tuned model so that it can take in images as input and return class probabilities. The file mapping class indices to class labels is saved along with the models.

**Incremental training**

You can seed the training of a new model with artifacts from a model that you trained previously with SageMaker. Incremental training saves training time when you want to train a new model with the same or similar data.
You can only seed a SageMaker Image Classification - TensorFlow model with another Image Classification - TensorFlow model trained in SageMaker.

You can use any dataset for incremental training, as long as the set of classes remains the same. The incremental training step is similar to the fine-tuning step, but instead of starting with a pretrained model, you start with an existing fine-tuned model. For an example of incremental training with the SageMaker Image Classification - TensorFlow algorithm, see the [Introduction to SageMaker TensorFlow - Image Classification](https://docs.aws.amazon.com/AmazonSageMaker/latest/dg/tensorflow-image-classification-intro.html) sample notebook.

**Inference with the Image Classification - TensorFlow algorithm**

You can host the fine-tuned model that results from your TensorFlow Image Classification training for inference. Any input image for inference must be in .jpg, .jpeg, or .png format and be content type `application/x-image`. The Image Classification - TensorFlow algorithm resizes input images automatically.

Running inference results in probability values, class labels for all classes, and the predicted label corresponding to the class index with the highest probability encoded in JSON format. The Image Classification - TensorFlow model processes a single image per request and outputs only one line. The following is an example of a JSON format response:

```json
accept: application/json;verbose

{"probabilities": [prob_0, prob_1, prob_2, ...],
 "labels": [label_0, label_1, label_2, ...],
 "predicted_label": predicted_label}
```

If accept is set to `application/json`, then the model only outputs probabilities. For more information on training and inference with the Image Classification - TensorFlow algorithm, see the [Introduction to SageMaker TensorFlow - Image Classification](https://docs.aws.amazon.com/AmazonSageMaker/latest/dg/tensorflow-image-classification-intro.html) sample notebook.

**Amazon EC2 instance recommendation for the Image Classification - TensorFlow algorithm**

The Image Classification - TensorFlow algorithm supports all CPU and GPU instances for training, including:

- `ml.p2.xlarge`
We recommend GPU instances with more memory for training with large batch sizes. Both CPU (such as M5) and GPU (P2, P3, G4dn, or G5) instances can be used for inference.

**Image Classification - TensorFlow sample notebooks**

For more information about how to use the SageMaker Image Classification - TensorFlow algorithm for transfer learning on a custom dataset, see the [Introduction to SageMaker TensorFlow - Image Classification](#) notebook.

For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see [Amazon SageMaker Notebook Instances](#). After you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

**How Image Classification - TensorFlow Works**

The Image Classification - TensorFlow algorithm takes an image as input and classifies it into one of the output class labels. Various deep learning networks such as MobileNet, ResNet, Inception, and EfficientNet are highly accurate for image classification. There are also deep learning networks that are trained on large image datasets, such as ImageNet, which has over 11 million images and almost 11,000 classes. After a network is trained with ImageNet data, you can then fine-tune the network on a dataset with a particular focus to perform more specific classification tasks. The Amazon SageMaker Image Classification - TensorFlow algorithm supports transfer learning on many pretrained models that are available in the TensorFlow Hub.

According to the number of class labels in your training data, a classification layer is attached to the pretrained TensorFlow Hub model of your choice. The classification layer consists of a dropout layer, a dense layer, and a fully-connected layer with 2-norm regularizer that is initialized with random weights. The model has hyperparameters for the dropout rate of the dropout layer and the L2 regularization factor for the dense layer. You can then fine-tune either the entire network
(including the pretrained model) or only the top classification layer on new training data. With this method of transfer learning, training with smaller datasets is possible.

**TensorFlow Hub Models**

The following pretrained models are available to use for transfer learning with the Image Classification - TensorFlow algorithm.

The following models vary significantly in size, number of model parameters, training time, and inference latency for any given dataset. The best model for your use case depends on the complexity of your fine-tuning dataset and any requirements that you have on training time, inference latency, or model accuracy.

<table>
<thead>
<tr>
<th>Model Name</th>
<th>model_id</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>MobileNet V2 1.00 224</td>
<td>tensorflow-ic-imagenet-mobilenet-v2-100-224-classification-4</td>
<td>[TensorFlow Hub link]</td>
</tr>
<tr>
<td>MobileNet V2 0.75 224</td>
<td>tensorflow-ic-imagenet-mobilenet-v2-075-224-classification-4</td>
<td>[TensorFlow Hub link]</td>
</tr>
<tr>
<td>MobileNet V2 0.50 224</td>
<td>tensorflow-ic-imagenet-mobilenet-v2-050-224-classification-4</td>
<td>[TensorFlow Hub link]</td>
</tr>
<tr>
<td>MobileNet V2 0.35 224</td>
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<td>TensorFlow Hub link</td>
</tr>
</tbody>
</table>

**Image Classification - TensorFlow Hyperparameters**

Hyperparameters are parameters that are set before a machine learning model begins learning. The following hyperparameters are supported by the Amazon SageMaker built-in Image Classification - TensorFlow algorithm. See [Tune an Image Classification - TensorFlow model](#) for information on hyperparameter tuning.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>augmentation</td>
<td>Set to &quot;True&quot; to apply augmentation_random_flip , augmentation_random_rotation , and augmentation_random_zoom to the training data. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>augmentation_random_flip</td>
<td>Indicates which flip mode to use for data augmentation when augmentation is set to &quot;True&quot;. For more information, see RandomFlip in the TensorFlow documentation. Valid values: string, any of the following: (&quot;horizontal_and_vertical&quot; , &quot;vertical&quot; , or &quot;None&quot;). Default value: &quot;horizontal_and_vertical&quot;.</td>
</tr>
<tr>
<td>augmentation_random_rotation</td>
<td>Indicates how much rotation to use for data augmentation when augmentation is set to &quot;True&quot;. Values represent a fraction of $2\pi$. Positive values rotate counterclockwise while negative values rotate clockwise. 0 means no rotation. For more information, see RandomRotation in the TensorFlow documentation. Valid values: float, range: [-1.0, 1.0]. Default value: 0.2.</td>
</tr>
<tr>
<td>augmentation_random_zoom</td>
<td>Indicates how much vertical zoom to use for data augmentation when augmentation is set to &quot;True&quot;. Positive values zoom out while negative values zoom in. 0 means no zoom. For more information, see RandomZoom in the TensorFlow documentation. Valid values: float, range: [-1.0, 1.0]. Default value: 0.1.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
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<tr>
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</tr>
<tr>
<td>batch_size</td>
<td>The batch size for training. For training on instances with multiple GPUs, this batch size is used across the GPUs. Valid values: positive integer. Default value: 32.</td>
</tr>
<tr>
<td>beta_1</td>
<td>The beta1 for the &quot;adam&quot; optimizer. Represents the exponential decay rate for the first moment estimates. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.9.</td>
</tr>
<tr>
<td>beta_2</td>
<td>The beta2 for the &quot;adam&quot; optimizer. Represents the exponential decay rate for the second moment estimates. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.999.</td>
</tr>
<tr>
<td>binary_mode</td>
<td>When binary_mode is set to &quot;True&quot;, the model returns a single probability number for the positive class and can use additional eval_metric options. Use only for binary classification problems. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>dropout_rate</td>
<td>The dropout rate for the dropout layer in the top classification layer. Valid values: float, range: [0.0, 1.0]. Default value: 0.2</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping</td>
<td>Set to &quot;True&quot; to use early stopping logic during training. If &quot;False&quot;, early stopping is not used. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>early_stopping_min_delta</td>
<td>The minimum change needed to qualify as an improvement. An absolute change less than the value of early_stopping_min_delta does not qualify as improvement. Used only when early_stopping is set to &quot;True&quot;. Valid values: float, range: [0.0, 1.0]. Default value: 0.0.</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>The number of epochs to continue training with no improvement. Used only when early_stopping is set to &quot;True&quot;. Valid values: positive integer. Default value: 5.</td>
</tr>
<tr>
<td>epochs</td>
<td>The number of training epochs. Valid values: positive integer. Default value: 3.</td>
</tr>
<tr>
<td>epsilon</td>
<td>The epsilon for &quot;adam&quot;, &quot;rmsprop&quot;, &quot;adadelta&quot;, and &quot;adagrad&quot; optimizers. Usually set to a small value to avoid division by 0. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 1e-7.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| eval_metric                 | If `binary_mode` is set to "False", `eval_metric` can only be "accuracy". If `binary_mode` is "True", select any of the valid values. For more information, see [Metrics](https://www.tensorflow.org/api_docs/python/tf/metrics) in the TensorFlow documentation.  
Valid values: string, any of the following: ("accuracy", "precision", "recall", "auc", or "prc").  
Default value: "accuracy".                                                                                                                                 |
| image_resize_interpolation  | Indicates interpolation method used when resizing images. For more information, see [image.resize](https://www.tensorflow.org/api_docs/python/tf/imaging/image_resize) in the TensorFlow documentation.  
Valid values: string, any of the following: ("bilinear", "nearest", "bicubic", "area", "lanczos3", "lanczos5", "gaussian", or "mitchellcubic").  
Default value: "bilinear".                                                                                                                                 |
| initial_accumulator_value   | The starting value for the accumulators, or the per-parameter momentum values, for the "adagrad" optimizer. Ignored for other optimizers.  
Valid values: float, range: [0.0, 1.0].  
Default value: 0.0001.                                                                                                                                 |
| label_smoothing             | Indicates how much to relax the confidence on label values. For example, if `label_smoothing` is 0.1, then non-target labels are 0.1/num_classes and target labels are 0.9+0.1/num_classes.  
Valid values: float, range: [0.0, 1.0].  
Default value: 0.1.                                                                                                                                 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>The optimizer learning rate.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.001.</td>
</tr>
<tr>
<td>momentum</td>
<td>The momentum for &quot;sgd&quot;, &quot;nesterov&quot;, and &quot;rmsprop&quot; optimizers. Ignored for other optimizers.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.9.</td>
</tr>
<tr>
<td>optimizer</td>
<td>The optimizer type. For more information, see Optimizers in the TensorFlow documentation.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: (&quot;adam&quot;, &quot;sgd&quot;, &quot;nesterov&quot;, &quot;rmsprop&quot;, &quot;adagrad&quot;, &quot;adadelta&quot;).</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;adam&quot;.</td>
</tr>
<tr>
<td>regularizers_l2</td>
<td>The L2 regularization factor for the dense layer in the classification layer.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float, range: [0.0, 1.0].</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0001.</td>
</tr>
<tr>
<td>reinitialize_top_layer</td>
<td>If set to &quot;Auto&quot;, the top classification layer parameters are re-initialized during fine-tuning. For incremental training, top classification layer parameters are not re-initialized unless set to &quot;True&quot;.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string, any of the following: (&quot;Auto&quot;, &quot;True&quot; or &quot;False&quot;).</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;Auto&quot;.</td>
</tr>
</tbody>
</table>
### Parameter Name | Description
--- | ---
\(\rho\) | The discounting factor for the gradient of the "adadelta" and "rmsprop" optimizers. Ignored for other optimizers.
Valid values: float, range: \([0.0, 1.0]\).
Default value: 0.95.

\texttt{train\_only\_top\_layer} | If "True", only the top classification layer parameters are fine-tuned. If "False", all model parameters are fine-tuned.
Valid values: string, either: ("True" or "False").
Default value: "False".

---

**Tune an Image Classification - TensorFlow model**

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

**Metrics computed by the Image Classification - TensorFlow algorithm**

The image classification algorithm is a supervised algorithm. It reports an accuracy metric that is computed during training. When tuning the model, choose this metric as the objective metric.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>\texttt{validation:accuracy}</td>
<td>The ratio of the number of correct predictions to the total number of predictions made.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>
Tunable Image Classification - TensorFlow hyperparameters

Tune an image classification model with the following hyperparameters. The hyperparameters that have the greatest impact on image classification objective metrics are: `batch_size`, `learning_rate`, and `optimizer`. Tune the optimizer-related hyperparameters, such as `momentum`, `regularizers_l2`, `beta_1`, `beta_2`, and `eps` based on the selected optimizer. For example, use `beta_1` and `beta_2` only when `adam` is the optimizer.

For more information about which hyperparameters are used for each optimizer, see Image Classification - TensorFlow Hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 8, MaxValue: 512</td>
</tr>
<tr>
<td>beta_1</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
<tr>
<td>beta_2</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
<tr>
<td>eps</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-8, MaxValue: 1.0</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.5</td>
</tr>
<tr>
<td>momentum</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
<tr>
<td>optimizer</td>
<td>CategoricalParameterRanges</td>
<td>['sgd', 'adam', 'rmsprop', 'nesterov', 'adagrad', 'adadelta']</td>
</tr>
<tr>
<td>regularizers_l2</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
</tbody>
</table>
Object Detection - MXNet

The Amazon SageMaker Object Detection - MXNet algorithm detects and classifies objects in images using a single deep neural network. It is a supervised learning algorithm that takes images as input and identifies all instances of objects within the image scene. The object is categorized into one of the classes in a specified collection with a confidence score that it belongs to the class. Its location and scale in the image are indicated by a rectangular bounding box. It uses the Single Shot multibox Detector (SSD) framework and supports two base networks: VGG and ResNet. The network can be trained from scratch, or trained with models that have been pre-trained on the ImageNet dataset.

Topics

- Input/Output Interface for the Object Detection Algorithm
- EC2 Instance Recommendation for the Object Detection Algorithm
- Object Detection Sample Notebooks
- How Object Detection Works
- Object Detection Hyperparameters
- Tune an Object Detection Model
- Object Detection Request and Response Formats

Input/Output Interface for the Object Detection Algorithm

The SageMaker Object Detection algorithm supports both RecordIO (application/x-recordio) and image (image/png, image/jpeg, and application/x-image) content types for training in file mode and supports RecordIO (application/x-recordio) for training in pipe mode. However you can also train in pipe mode using the image files (image/png, image/jpeg, and application/x-image), without creating RecordIO files, by using the augmented manifest format. The recommended input format for the Amazon SageMaker object detection algorithms

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>train_only_top_layer</td>
<td>ContinuousParameterRanges</td>
<td>['True', 'False']</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>train_only_top_layer</td>
<td>ContinuousParameterRanges</td>
<td>['True', 'False']</td>
</tr>
</tbody>
</table>
is Apache MXNet RecordIO. However, you can also use raw images in .jpg or .png format. The algorithm supports only application/x-image for inference.

**Note**

To maintain better interoperability with existing deep learning frameworks, this differs from the protobuf data formats commonly used by other Amazon SageMaker algorithms.

See the [Object Detection Sample Notebooks](#) for more details on data formats.

### Train with the RecordIO Format

If you use the RecordIO format for training, specify both train and validation channels as values for the InputDataConfig parameter of the `CreateTrainingJob` request. Specify one RecordIO (.rec) file in the train channel and one RecordIO file in the validation channel. Set the content type for both channels to application/x-recordio. An example of how to generate RecordIO file can be found in the object detection sample notebook. You can also use tools from the [MXNet's GluonCV](#) to generate RecordIO files for popular datasets like the [PASCAL Visual Object Classes](#) and [Common Objects in Context (COCO)](#) .

### Train with the Image Format

If you use the image format for training, specify train, validation, train_annotation, and validation_annotation channels as values for the InputDataConfig parameter of `CreateTrainingJob` request. Specify the individual image data (.jpg or .png) files for the train and validation channels. For annotation data, you can use the JSON format. Specify the corresponding .json files in the train_annotation and validation_annotation channels. Set the content type for all four channels to image/png or image/jpeg based on the image type. You can also use the content type application/x-image when your dataset contains both .jpg and .png images. The following is an example of a .json file.

```json
{
    "file": "your_image_directory/sample_image1.jpg",
    "image_size": [
        {
            "width": 500,
            "height": 400,
            "depth": 3
        }
    ]
}
```
Each image needs a .json file for annotation, and the .json file should have the same name as the corresponding image. The name of above .json file should be "sample_image1.json". There are four properties in the annotation .json file. The property "file" specifies the relative path of the image file. For example, if your training images and corresponding .json files are stored in s3://your_bucket/train/sample_image and s3://your_bucket/train_annotation,
specify the path for your train and train_annotation channels as s3://your_bucket/train and s3://your_bucket/train_annotation, respectively.

In the .json file, the relative path for an image named sample_image1.jpg should be sample_image/sample_image1.jpg. The "image_size" property specifies the overall image dimensions. The SageMaker object detection algorithm currently only supports 3-channel images. The "annotations" property specifies the categories and bounding boxes for objects within the image. Each object is annotated by a "class_id" index and by four bounding box coordinates ("left", "top", "width", "height"). The "left" (x-coordinate) and "top" (y-coordinate) values represent the upper-left corner of the bounding box. The "width" (x-coordinate) and "height" (y-coordinate) values represent the dimensions of the bounding box. The origin (0, 0) is the upper-left corner of the entire image. If you have multiple objects within one image, all the annotations should be included in a single .json file. The "categories" property stores the mapping between the class index and class name. The class indices should be numbered successively and the numbering should start with 0. The "categories" property is optional for the annotation .json file.

**Train with Augmented Manifest Image Format**

The augmented manifest format enables you to do training in pipe mode using image files without needing to create RecordIO files. You need to specify both train and validation channels as values for the InputDataConfig parameter of the `CreateTrainingJob` request. While using the format, an S3 manifest file needs to be generated that contains the list of images and their corresponding annotations. The manifest file format should be in JSON Lines format in which each line represents one sample. The images are specified using the 'source-ref' tag that points to the S3 location of the image. The annotations are provided under the "AttributeNames" parameter value as specified in the `CreateTrainingJob` request. It can also contain additional metadata under the "AttributeNames" tag, but these are ignored by the algorithm. In the following example, the "AttributeNames" are contained in the list ["source-ref", "bounding-box"]:

```json
[
  {
    "source-ref": "s3://your_bucket/image1.jpg",
    "bounding-box": {
      "image_size": [{
        "width": 500,
        "height": 400,
        "depth": 3
      }],
      "annotations": [{
        "class_id": 0,
        "left": 111,
        "top": 134,
        "width": 61,
        "height": 128,
        "class_id": 5,
        "left": 161,
        "top": 250,
        "width": 80,
        "height": 50
      }],
      "bounding-box-metadata": {
        "class-map": {
          "0": "dog",
          "5": "horse"
        },
        "type": "groundtruth/object-detection"
      }
    }
  },
  {
    "source-ref": "s3://your_bucket/image2.jpg",
    "bounding-box": {
      "image_size": [{
        "width": 400,
        "height": 300,
        "depth": 3
      }],
      "annotations": [{
        "class_id": 1,
        "left": 100,
        "top": 120,
        "width": 43,
        "height": 78
      }],
      "bounding-box-metadata": {
        "class-map": {
          "1": "cat"
        },
        "type": "groundtruth/object-detection"
      }
    }
  }
]
```
The order of "AttributeNames" in the input files matters when training the Object Detection algorithm. It accepts piped data in a specific order, with image first, followed by annotations. So the "AttributeNames" in this example are provided with "source-ref" first, followed by "bounding-box". When using Object Detection with Augmented Manifest, the value of parameter RecordWrapperType must be set as "RecordIO".

For more information on augmented manifest files, see [Provide Dataset Metadata to Training Jobs with an Augmented Manifest File](#).

**Incremental Training**

You can also seed the training of a new model with the artifacts from a model that you trained previously with SageMaker. Incremental training saves training time when you want to train a new model with the same or similar data. SageMaker object detection models can be seeded only with another built-in object detection model trained in SageMaker.

To use a pretrained model, in the [CreateTrainingJob](#) request, specify the ChannelName as "model" in the InputDataConfig parameter. Set the ContentType for the model channel to application/x-sagemaker-model. The input hyperparameters of both the new model and the pretrained model that you upload to the model channel must have the same settings for the base_network and num_classes input parameters. These parameters define the network architecture. For the pretrained model file, use the compressed model artifacts (in .tar.gz format) output by SageMaker. You can use either RecordIO or image formats for input data.

For more information on incremental training and for instructions on how to use it, see [Use Incremental Training in Amazon SageMaker](#).

**EC2 Instance Recommendation for the Object Detection Algorithm**

The object detection algorithm supports P2, P3, G4dn, and G5 GPU instance families. We recommend using GPU instances with more memory for training with large batch sizes. You can run the object detection algorithm on multi-GPU and mult-machine settings for distributed training.

You can use both CPU (such as C5 and M5) and GPU (such as P3 and G4dn) instances for inference.

**Object Detection Sample Notebooks**

For a sample notebook that shows how to use the SageMaker Object Detection algorithm to train and host a model on the
Caltech Birds (CUB 200 2011) dataset using the Single Shot multibox Detector algorithm, see Amazon SageMaker Object Detection for Bird Species. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. The object detection example notebook using the Object Detection algorithm is located in the Introduction to Amazon Algorithms section. To open a notebook, click on its Use tab and select Create copy.

How Object Detection Works

The object detection algorithm identifies and locates all instances of objects in an image from a known collection of object categories. The algorithm takes an image as input and outputs the category that the object belongs to, along with a confidence score that it belongs to the category. The algorithm also predicts the object's location and scale with a rectangular bounding box. Amazon SageMaker Object Detection uses the Single Shot multibox Detector (SSD) algorithm that takes a convolutional neural network (CNN) pretrained for classification task as the base network. SSD uses the output of intermediate layers as features for detection.

Various CNNs such as VGG and ResNet have achieved great performance on the image classification task. Object detection in Amazon SageMaker supports both VGG-16 and ResNet-50 as a base network for SSD. The algorithm can be trained in full training mode or in transfer learning mode. In full training mode, the base network is initialized with random weights and then trained on user data. In transfer learning mode, the base network weights are loaded from pretrained models.

The object detection algorithm uses standard data augmentation operations, such as flip, rescale, and jitter, on the fly internally to help avoid overfitting.

Object Detection Hyperparameters

In the CreateTrainingJob request, you specify the training algorithm that you want to use. You can also specify algorithm-specific hyperparameters that are used to help estimate the parameters of the model from a training dataset. The following table lists the hyperparameters provided by Amazon SageMaker for training the object detection algorithm. For more information about how object training works, see How Object Detection Works.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_classes</td>
<td>The number of output classes. This parameter defines the dimensions of the network output and is typically set to the number of classes in the dataset.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>num_training_samples</td>
<td>The number of training examples in the input dataset.</td>
</tr>
<tr>
<td></td>
<td><strong>Note</strong></td>
</tr>
<tr>
<td></td>
<td>If there is a mismatch between this value and the number of samples in the training set, then the behavior of the lr_scheduler_step parameter will be undefined and distributed training accuracy may be affected.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>base_network</td>
<td>The base network architecture to use.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 'vgg-16' or 'resnet-50'</td>
</tr>
<tr>
<td></td>
<td>Default value: 'vgg-16'</td>
</tr>
<tr>
<td>early_stopping</td>
<td>True to use early stopping logic during training. False not to use it.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: True or False</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>early_stopping_min_epochs</td>
<td>The minimum number of epochs that must be run before the early stopping logic can be invoked. It is used only when early_stopping = True.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>The number of epochs to wait before ending training if no improvement, as defined by the early_stopping_tolerance hyperparameter, is made in the relevant metric. It is used only when early_stopping = True.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td>early_stopping_tolerance</td>
<td>The tolerance value that the relative improvement in validation:mAP, the mean average precision (mAP), is required to exceed to avoid early stopping. If the ratio of the change in the mAP divided by the previous best mAP is smaller than the early_stopping_tolerance value set, early stopping considers that there is no improvement. It is used only when early_stopping = True.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 \leq \text{float} \leq 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>image_shape</td>
<td>The image size for input images. We rescale the input image to a square image with this size. We recommend using 300 and 512 for better performance.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer ≥300</td>
</tr>
<tr>
<td></td>
<td>Default: 300</td>
</tr>
<tr>
<td>epochs</td>
<td>The number of training epochs.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default: 30</td>
</tr>
<tr>
<td>freeze_layer_pattern</td>
<td>The regular expression (regex) for freezing layers in the base network. For example, if we set freeze_layer_pattern = &quot;^(conv1_</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: string</td>
</tr>
<tr>
<td></td>
<td>Default: No layers frozen.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>kv_store</td>
<td>The weight update synchronization mode used for distributed training. The weights can be updated either synchronously or asynchronously across machines. Synchronous updates typically provide better accuracy than asynchronous updates but can be slower. See the <a href="#">Distributed Training MXNet tutorial</a> for details.</td>
</tr>
</tbody>
</table>

**Note**

This parameter is not applicable to single machine training.

**Optional**

Valid values: 'dist_sync' or 'dist_async'

- **'dist_sync'**: The gradients are synchronized after every batch with all the workers. With 'dist_sync', batch-size now means the batch size used on each machine. So if there are n machines and we use batch size b, then dist_sync behaves like a single machine with batch size n*b.

- **'dist_async'**: Performs asynchronous updates. The weights are updated whenever gradients are received from any machine and the weight updates are atomic. However, the order is not guaranteed.

**Default**: -
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>label_width</td>
<td>The force padding label width used to sync across training and validation data. For example, if one image in the data contains at most 10 objects, and each object's annotation is specified with 5 numbers, [class_id, left, top, width, height], then the label_width should be no smaller than (10*5 + header information length). The header information length is usually 2. We recommend using a slightly larger label_width for the training, such as 60 for this example.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Positive integer large enough to accommodate the largest annotation information length in the data.</td>
</tr>
<tr>
<td></td>
<td>Default: 350</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The initial learning rate.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float in (0, 1)</td>
</tr>
<tr>
<td></td>
<td>Default: 0.001</td>
</tr>
<tr>
<td>lr_scheduler_factor</td>
<td>The ratio to reduce learning rate. Used in conjunction with the lr_scheduler_step parameter defined as lr_new = lr_old * lr_scheduler_factor.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float in (0, 1)</td>
</tr>
<tr>
<td></td>
<td>Default: 0.1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>lr_scheduler_step</td>
<td>The epochs at which to reduce the learning rate. The learning rate is reduced by ( lr_scheduler_factor ) at epochs listed in a comma-delimited string: &quot;epoch1, epoch2, ...&quot;. For example, if the value is set to &quot;10, 20&quot; and the ( lr_scheduler_factor ) is set to 1/2, then the learning rate is halved after 10th epoch and then halved again after 20th epoch.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: string</td>
</tr>
<tr>
<td></td>
<td>Default: empty string</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>The batch size for training. In a single-machine multi-gpu setting, each GPU handles ( \text{mini_batch_size} / \text{num_gpu} ) training samples. For the multi-machine training in dist_sync mode, the actual batch size is ( \text{mini_batch_size} \times \text{number of machines} ). A large ( \text{mini_batch_size} ) usually leads to faster training, but it may cause out of memory problem. The memory usage is related to ( \text{mini_batch_size}, \text{image_shape}, \text{and base_network} ) architecture. For example, on a single p3.2xlarge instance, the largest ( \text{mini_batch_size} ) without an out of memory error is 32 with the base_network set to &quot;resnet-50&quot; and an ( \text{image_shape} ) of 300. With the same instance, you can use 64 as the ( \text{mini_batch_size} ) with the base network vgg-16 and an ( \text{image_shape} ) of 300.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default: 32</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>momentum</td>
<td>The momentum for sgd. Ignored for other optimizers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float in (0, 1]</td>
</tr>
<tr>
<td></td>
<td>Default: 0.9</td>
</tr>
<tr>
<td>nms_threshold</td>
<td>The non-maximum suppression threshold.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float in (0, 1]</td>
</tr>
<tr>
<td></td>
<td>Default: 0.45</td>
</tr>
<tr>
<td>optimizer</td>
<td>The optimizer types. For details on optimizer values, see <code>MXNet's API</code>.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: ['sgd', 'adam', 'rmsprop', 'adadelta']</td>
</tr>
<tr>
<td></td>
<td>Default: 'sgd'</td>
</tr>
<tr>
<td>overlap_threshold</td>
<td>The evaluation overlap threshold.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float in (0, 1]</td>
</tr>
<tr>
<td></td>
<td>Default: 0.5</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>use_pretrained_model</td>
<td>Indicates whether to use a pre-trained model for training. If set to 1, then the pre-trained model with corresponding architecture is loaded and used for training. Otherwise, the network is trained from scratch.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 or 1</td>
</tr>
<tr>
<td></td>
<td>Default: 1</td>
</tr>
<tr>
<td>weight_decay</td>
<td>The weight decay coefficient for sgd and rmsprop. Ignored for other optimizers.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: float in (0, 1)</td>
</tr>
<tr>
<td></td>
<td>Default: 0.0005</td>
</tr>
</tbody>
</table>

### Tune an Object Detection Model

*Automatic model tuning*, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see [Perform Automatic Model Tuning with SageMaker](#).

### Metrics Computed by the Object Detection Algorithm

The object detection algorithm reports on a single metric during training: `validation:mAP`. When tuning a model, choose this metric as the objective metric.
<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:mAP</td>
<td>Mean Average Precision (mAP) computed on the validation set.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

**Tunable Object Detection Hyperparameters**

Tune the Amazon SageMaker object detection model with the following hyperparameters. The hyperparameters that have the greatest impact on the object detection objective metric are: mini_batch_size, learning_rate, and optimizer.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-6, MaxValue: 0.5</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 8, MaxValue: 64</td>
</tr>
<tr>
<td>momentum</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
<tr>
<td>optimizer</td>
<td>CategoricalParameterRanges</td>
<td>['sgd', 'adam', 'rmsprop', 'adadelta']</td>
</tr>
<tr>
<td>weight_decay</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
</tbody>
</table>

**Object Detection Request and Response Formats**

**Request Format**

Query a trained model by using the model's endpoint. The endpoint takes .jpg and .png image formats with image/jpeg and image/png content-types.
Response Formats

The response is the class index with a confidence score and bounding box coordinates for all objects within the image encoded in JSON format. The following is an example of response .json file:

```
{
  "prediction": [
    [4.0, 0.86419455409049988, 0.3088374733924866, 0.07030484080314636, 0.7110607028007507, 0.9345266819000244],
    [0.0, 0.73376623392105103, 0.5714187026023865, 0.40427327156066895, 0.827075183391571, 0.9712159633636475],
    [4.0, 0.32643985450267792, 0.3677481412887573, 0.034883320331573486, 0.6318609714508057, 0.5967587828636169],
    [8.0, 0.22552496790885925, 0.6152569651603699, 0.5722782611846924, 0.882301390171051, 0.8985623121261597],
    [3.0, 0.42260299175977707, 0.019305512309074402, 0.08386176824569702, 0.3909356594085693, 0.5124796557426453]
  ]
}
```

Each row in this .json file contains an array that represents a detected object. Each of these object arrays consists of a list of six numbers. The first number is the predicted class label. The second number is the associated confidence score for the detection. The last four numbers represent the bounding box coordinates [xmin, ymin, xmax, ymax]. These output bounding box corner indices are normalized by the overall image size. Note that this encoding is different than that used by the input .json format. For example, in the first entry of the detection result, 0.3088374733924866 is the left coordinate (x-coordinate of upper-left corner) of the bounding box as a ratio of the overall image width, 0.07030484080314636 is the top coordinate (y-coordinate of upper-left corner) of the bounding box as a ratio of the overall image height, 0.7110607028007507 is the right coordinate (x-coordinate of lower-right corner) of the bounding box as a ratio of the overall image width, and 0.9345266819000244 is the bottom coordinate (y-coordinate of lower-right corner) of the bounding box as a ratio of the overall image height.

To avoid unreliable detection results, you might want to filter out the detection results with low confidence scores. In the object detection sample notebook, we provide examples of scripts that use a threshold to remove low confidence detections and to plot bounding boxes on the original images.

For batch transform, the response is in JSON format, where the format is identical to the JSON format described above. The detection results of each image is represented as a JSON file. For example:
For more details on training and inference, see the Object Detection Sample Notebooks.

**OUTPUT: JSON Response Format**

```json
{
  "image_size": [
    {
      "width": 500,
      "height": 400,
      "depth": 3
    }
  ],
  "annotations": [
    {
      "class_id": 0,
      "score": 0.943,
      "left": 111,
      "top": 134,
      "width": 61,
      "height": 128
    },
    {
      "class_id": 0,
      "score": 0.0013,
      "left": 161,
      "top": 250,
      "width": 79,
      "height": 143
    },
    {
      "class_id": 1,
      "score": 0.0133,
      "left": 101,
      "top": 185,
      "width": 42,
      "height": 130
    }
  ]
}
```
Object Detection - TensorFlow

The Amazon SageMaker Object Detection - TensorFlow algorithm is a supervised learning algorithm that supports transfer learning with many pretrained models from the TensorFlow Model Garden. Use transfer learning to fine-tune one of the available pretrained models on your own dataset, even if a large amount of image data is not available. The object detection algorithm takes an image as input and outputs a list of bounding boxes. Training datasets must consist of images in .jpg, .jpeg, or .png format.

Topics

- How to use the SageMaker Object Detection - TensorFlow algorithm
- Input and output interface for the Object Detection - TensorFlow algorithm
- Amazon EC2 instance recommendation for the Object Detection - TensorFlow algorithm
- Object Detection - TensorFlow sample notebooks
- How Object Detection - TensorFlow Works
- TensorFlow Models
- Object Detection - TensorFlow Hyperparameters
- Tune an Object Detection - TensorFlow model

How to use the SageMaker Object Detection - TensorFlow algorithm

You can use Object Detection - TensorFlow as an Amazon SageMaker built-in algorithm. The following section describes how to use Object Detection - TensorFlow with the SageMaker Python SDK. For information on how to use Object Detection - TensorFlow from the Amazon SageMaker Studio Classic UI, see SageMaker JumpStart.

The Object Detection - TensorFlow algorithm supports transfer learning using any of the compatible pretrained TensorFlow models. For a list of all available pretrained models, see TensorFlow Models. Every pretrained model has a unique model_id. The following example uses ResNet50 (model_id: tensorflow-od1-ssd-resnet50-v1-fpn-640x640-coco17-tpu-8) to fine-tune on a custom dataset. The pretrained models are all pre-downloaded from the TensorFlow Hub and stored in Amazon S3 buckets so that training jobs can run in network isolation. Use these pre-generated model training artifacts to construct a SageMaker Estimator.
First, retrieve the Docker image URI, training script URI, and pretrained model URI. Then, change the hyperparameters as you see fit. You can see a Python dictionary of all available hyperparameters and their default values with `hyperparameters.retrieve_default`. For more information, see [Object Detection - TensorFlow Hyperparameters](#). Use these values to construct a SageMaker Estimator.

**Note**

Default hyperparameter values are different for different models. For example, for larger models, the default number of epochs is smaller.

This example uses the **PennFudanPed** dataset, which contains images of pedestrians in the street. We pre-downloaded the dataset and made it available with Amazon S3. To fine-tune your model, call `.fit` using the Amazon S3 location of your training dataset.

```python
from sagemaker import image_uris, model_uris, script_uris, hyperparameters
from sagemaker.estimator import Estimator

model_id, model_version = "tensorflow-od1-ssd-resnet50-v1-fpn-640x640-coco17-tpu-8", "*

training_instance_type = "ml.p3.2xlarge"

# Retrieve the Docker image
train_image_uri =
    image_uris.retrieve(model_id=model_id, model_version=model_version, image_scope="training", instance_type=training_instance_type, region=None, framework=None)

# Retrieve the training script
train_source_uri = script_uris.retrieve(model_id=model_id, model_version=model_version, script_scope="training")

# Retrieve the pretrained model tarball for transfer learning
train_model_uri = model_uris.retrieve(model_id=model_id, model_version=model_version, model_scope="training")

# Retrieve the default hyperparameters for fine-tuning the model
hyperparameters = hyperparameters.retrieve_default(model_id=model_id, model_version=model_version)

# [Optional] Override default hyperparameters with custom values
hyperparameters["epochs"] = "5"
```
# Sample training data is available in this bucket

```python
training_data_bucket = f"jumpstart-cache-prod-{aws_region}"
training_data_prefix = "training-datasets/PennFudanPed_COCO_format/"

training_dataset_s3_path = f"s3://{training_data_bucket}/{training_data_prefix}"

output_bucket = sess.default_bucket()
output_prefix = "jumpstart-example-od-training"
s3_output_location = f"s3://{output_bucket}/{output_prefix}/output"
```

# Create an Estimator instance

```python
tf_od_estimator = Estimator(
    role=aws_role,
    image_uri=train_image_uri,
    source_dir=train_source_uri,
    model_uri=train_model_uri,
    entry_point="transfer_learning.py",
    instance_count=1,
    instance_type=training_instance_type,
    max_run=360000,
    hyperparameters=hyperparameters,
    output_path=s3_output_location,
)
```

# Launch a training job

```python
tf_od_estimator.fit(\"training": training_dataset_s3_path, logs=True)
```

For more information about how to use the SageMaker Object Detection - TensorFlow algorithm for transfer learning on a custom dataset, see the Introduction to SageMaker TensorFlow - Object Detection notebook.

**Input and output interface for the Object Detection - TensorFlow algorithm**

Each of the pretrained models listed in TensorFlow Models can be fine-tuned to any dataset with any number of image classes. Be mindful of how to format your training data for input to the Object Detection - TensorFlow model.

- **Training data input format**: Your training data should be a directory with an images subdirectory and an annotations.json file.
The following is an example of an input directory structure. The input directory should be hosted in an Amazon S3 bucket with a path similar to the following: s3://bucket_name/input_directory/. Note that the trailing / is required.

```
input_directory
  |--images
     |--abc.png
     |--def.png
     |--annotations.json
```

The annotations.json file should contain information for bounding boxes and their class labels in the form of a dictionary "images" and "annotations" keys. The value for the "images" key should be a list of dictionaries. There should be one dictionary for each image with the following information: `{"file_name": image_name, "height": height, "width": width, "id": image_id}`. The value for the "annotations" key should also be a list of dictionaries. There should be one dictionary for each bounding box with the following information: `{"image_id": image_id, "bbox": [xmin, ymin, xmax, ymax], "category_id": bbox_label}`.

After training, a label mapping file and trained model are saved to your Amazon S3 bucket.

**Incremental training**

You can seed the training of a new model with artifacts from a model that you trained previously with SageMaker. Incremental training saves training time when you want to train a new model with the same or similar data.

⚠️ **Note**


You can use any dataset for incremental training, as long as the set of classes remains the same. The incremental training step is similar to the fine-tuning step, but instead of starting with a pretrained model, you start with an existing fine-tuned model. For more information about how to use incremental training with the SageMaker Object Detection - TensorFlow, see the [Introduction to SageMaker TensorFlow - Object Detection](#) notebook.
Inference with the Object Detection - TensorFlow algorithm

You can host the fine-tuned model that results from your TensorFlow Object Detection training for inference. Any input image for inference must be in .jpg, .jpeg, or .png format and be content type application/x-image. The Object Detection - TensorFlow algorithm resizes input images automatically.

Running inference results in bounding boxes, predicted classes, and the scores of each prediction encoded in JSON format. The Object Detection - TensorFlow model processes a single image per request and outputs only one line. The following is an example of a JSON format response:

```json
accept: application/json;verbose

{"normalized_boxes":[[xmin1, xmax1, ymin1, ymax1],....],
  "classes":[classidx1, class_idx2,...],
  "scores":[score_1, score_2,...],
  "labels": [label1, label2, ...],
  "tensorflow_model_output":<original output of the model>}
```

If accept is set to application/json, then the model only outputs normalized boxes, classes, and scores.

Amazon EC2 instance recommendation for the Object Detection - TensorFlow algorithm

The Object Detection - TensorFlow algorithm supports all GPU instances for training, including:

- ml.p2.xlarge
- ml.p2.16xlarge
- ml.p3.2xlarge
- ml.p3.16xlarge

We recommend GPU instances with more memory for training with large batch sizes. Both CPU (such as M5) and GPU (P2 or P3) instances can be used for inference. For a comprehensive list of SageMaker training and inference instances across AWS Regions, see Amazon SageMaker Pricing.

Object Detection - TensorFlow sample notebooks

For more information about how to use the SageMaker Object Detection - TensorFlow algorithm for transfer learning on a custom dataset, see the Introduction to SageMaker TensorFlow - Object Detection notebook.
For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. To open a notebook, choose its Use tab and choose Create copy.

**How Object Detection - TensorFlow Works**

The Object Detection - TensorFlow algorithm takes an image as input and predicts bounding boxes and object labels. Various deep learning networks such as MobileNet, ResNet, Inception, and EfficientNet are highly accurate for object detection. There are also deep learning networks that are trained on large image datasets, such as Common Objects in Context (COCO), which has 328,000 images. After a network is trained with COCO data, you can then fine-tune the network on a dataset with a particular focus to perform more specific object detection tasks. The Amazon SageMaker Object Detection - TensorFlow algorithm supports transfer learning on many pretrained models that are available in the TensorFlow Model Garden.

According to the number of class labels in your training data, an object detection layer is attached to the pretrained TensorFlow model of your choice. You can then fine-tune either the entire network (including the pretrained model) or only the top classification layer on new training data. With this method of transfer learning, training with smaller datasets is possible.

**TensorFlow Models**

The following pretrained models are available to use for transfer learning with the Object Detection - TensorFlow algorithm.

The following models vary significantly in size, number of model parameters, training time, and inference latency for any given dataset. The best model for your use case depends on the complexity of your fine-tuning dataset and any requirements that you have on training time, inference latency, or model accuracy.

<table>
<thead>
<tr>
<th>Model Name</th>
<th>model_id</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>ResNet50 V1 FPN 640</td>
<td>tensorflow-od1-ssd-resnet50-v1-fpn-640x640-coco17-tpu-8</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td>Model Name</td>
<td>model_id</td>
<td>Source</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-----------------------------------------------</td>
<td>---------------------------------------------</td>
</tr>
<tr>
<td>EfficientDet D0 512</td>
<td>tensorflow-od1-ssd-efficientdet-d0-5</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td></td>
<td>12x512-coco17-tpu-8</td>
<td></td>
</tr>
<tr>
<td>EfficientDet D1 640</td>
<td>tensorflow-od1-ssd-efficientdet-d1-6</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td></td>
<td>40x640-coco17-tpu-8</td>
<td></td>
</tr>
<tr>
<td>EfficientDet D2 768</td>
<td>tensorflow-od1-ssd-efficientdet-d2-7</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td></td>
<td>68x768-coco17-tpu-8</td>
<td></td>
</tr>
<tr>
<td>EfficientDet D3 896</td>
<td>tensorflow-od1-ssd-efficientdet-d3-8</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td></td>
<td>96x896-coco17-tpu-32</td>
<td></td>
</tr>
<tr>
<td>MobileNet V1 FPN 640</td>
<td>tensorflow-od1-ssd-mobilenet-v1-fpn-640x640-coco17-tpu-8</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td>MobileNet V2 FPNLite 320</td>
<td>tensorflow-od1-ssd-mobilenet-v2-fpnlite-320x320-coco17-tpu-8</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td>MobileNet V2 FPNLite 640</td>
<td>tensorflow-od1-ssd-mobilenet-v2-fpnlite-640x640-coco17-tpu-8</td>
<td>TensorFlow Model Garden link</td>
</tr>
<tr>
<td>ResNet50 V1 FPN 1024</td>
<td>tensorflow-od1-ssd-resnet50-v1-fpn-1024x1024-coco17-tpu-8</td>
<td>TensorFlow Model Garden link</td>
</tr>
</tbody>
</table>
Object Detection - TensorFlow Hyperparameters

Hyperparameters are parameters that are set before a machine learning model begins learning. The following hyperparameters are supported by the Amazon SageMaker built-in Object Detection - TensorFlow algorithm. See [Tune an Object Detection - TensorFlow model](#) for information on hyperparameter tuning.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>batch_size</td>
<td>The batch size for training. Valid values: positive integer. Default value: 3.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>beta_1</td>
<td>The beta1 for the &quot;adam&quot; optimizer. Represents the exponential decay rate for the first moment estimates. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.9.</td>
</tr>
<tr>
<td>beta_2</td>
<td>The beta2 for the &quot;adam&quot; optimizer. Represents the exponential decay rate for the second moment estimates. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.999.</td>
</tr>
<tr>
<td>early_stopping</td>
<td>Set to &quot;True&quot; to use early stopping logic during training. If &quot;False&quot;, early stopping is not used. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
<tr>
<td>early_stopping_min_delta</td>
<td>The minimum change needed to qualify as an improvement. An absolute change less than the value of early_stopping_min_delta does not qualify as improvement. Used only when early_stopping is set to &quot;True&quot;. Valid values: float, range: [0.0, 1.0]. Default value: 0.0.</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>The number of epochs to continue training with no improvement. Used only when early_stopping is set to &quot;True&quot;. Valid values: positive integer. Default value: 5.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>epochs</td>
<td>The number of training epochs. Valid values: positive integer. Default value: 5 for smaller models, 1 for larger models.</td>
</tr>
<tr>
<td>epsilon</td>
<td>The epsilon for &quot;adam&quot;, &quot;rmsprop&quot;, &quot;adadelta&quot;, and &quot;adagrad&quot; optimizers. Usually set to a small value to avoid division by 0. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 1e-7.</td>
</tr>
<tr>
<td>initial_accumulator_value</td>
<td>The starting value for the accumulators, or the per-parameter momentum values, for the &quot;adagrad&quot; optimizer. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.1.</td>
</tr>
<tr>
<td>learning_rate</td>
<td>The optimizer learning rate. Valid values: float, range: [0.0, 1.0]. Default value: 0.001.</td>
</tr>
<tr>
<td>momentum</td>
<td>The momentum for the &quot;sgd&quot; and &quot;nesterov&quot; optimizers. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.9.</td>
</tr>
</tbody>
</table>

Use Built-in Algorithms
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>optimizer</td>
<td>The optimizer type. For more information, see Optimizers in the TensorFlow documentation. Valid values: string, any of the following: (&quot;adam&quot;, &quot;sgd&quot;, &quot;nesterov&quot;, &quot;rmsprop&quot;, &quot;adagrad&quot;, &quot;adadelta&quot;). Default value: &quot;adam&quot;.</td>
</tr>
<tr>
<td>reinitialize_top_layer</td>
<td>If set to &quot;Auto&quot;, the top classification layer parameters are re-initialized during fine-tuning. For incremental training, top classification layer parameters are not re-initialized unless set to &quot;True&quot;. Valid values: string, any of the following: (&quot;Auto&quot;, &quot;True&quot; or &quot;False&quot;). Default value: &quot;Auto&quot;.</td>
</tr>
<tr>
<td>rho</td>
<td>The discounting factor for the gradient of the &quot;adadelta&quot; and &quot;rmsprop&quot; optimizers. Ignored for other optimizers. Valid values: float, range: [0.0, 1.0]. Default value: 0.95.</td>
</tr>
<tr>
<td>train_only_on_top_layer</td>
<td>If &quot;True&quot;, only the top classification layer parameters are fine-tuned. If &quot;False&quot;, all model parameters are fine-tuned. Valid values: string, either: (&quot;True&quot; or &quot;False&quot;). Default value: &quot;False&quot;.</td>
</tr>
</tbody>
</table>

Tune an Object Detection - TensorFlow model

Automatic model tuning, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches
the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

For more information about model tuning, see Perform Automatic Model Tuning with SageMaker.

Metrics computed by the Object Detection - TensorFlow algorithm

Refer to the following chart to find which metrics are computed by the Object Detection - TensorFlow algorithm.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
<th>Regex Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:localization_loss</td>
<td>The localization loss for box prediction.</td>
<td>Minimize</td>
<td>Val_localization=([0-9\.]+)</td>
</tr>
</tbody>
</table>

Tunable Object Detection - TensorFlow hyperparameters

Tune an object detection model with the following hyperparameters. The hyperparameters that have the greatest impact on object detection objective metrics are: batch_size, learning_rate, and optimizer. Tune the optimizer-related hyperparameters, such as momentum, regularizers_l2, beta_1, beta_2, and eps based on the selected optimizer. For example, use beta_1 and beta_2 only when adam is the optimizer.

For more information about which hyperparameters are used for each optimizer, see Object Detection - TensorFlow Hyperparameters.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 8, MaxValue: 512</td>
</tr>
<tr>
<td>beta_1</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
<tr>
<td>beta_2</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.999</td>
</tr>
</tbody>
</table>
## Parameter Table

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>eps</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-8, MaxValue: 1.0</td>
</tr>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 1e-6, MaxValue: 0.5</td>
</tr>
<tr>
<td>momentum</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
<tr>
<td>optimizer</td>
<td>CategoricalParameterRanges</td>
<td>['sgd', 'adam', 'rmsprop', 'nesterov', 'adagrad', 'adadelta']</td>
</tr>
<tr>
<td>regularizers_l2</td>
<td>ContinuousParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
<tr>
<td>train_only_on_top_layer</td>
<td>CategoricalParameterRanges</td>
<td>['True', 'False']</td>
</tr>
<tr>
<td>initial_accumulator_value</td>
<td>CategoricalParameterRanges</td>
<td>MinValue: 0.0, MaxValue: 0.999</td>
</tr>
</tbody>
</table>

### Semantic Segmentation Algorithm

The SageMaker semantic segmentation algorithm provides a fine-grained, pixel-level approach to developing computer vision applications. It tags every pixel in an image with a class label from a predefined set of classes. Tagging is fundamental for understanding scenes, which is critical to an increasing number of computer vision applications, such as self-driving vehicles, medical imaging diagnostics, and robot sensing.

For comparison, the SageMaker **Image Classification - MXNet** is a supervised learning algorithm that analyzes only whole images, classifying them into one of multiple output categories. The **Object Detection - MXNet** is a supervised learning algorithm that detects and classifies all instances...
of an object in an image. It indicates the location and scale of each object in the image with a rectangular bounding box.

Because the semantic segmentation algorithm classifies every pixel in an image, it also provides information about the shapes of the objects contained in the image. The segmentation output is represented as a grayscale image, called a segmentation mask. A segmentation mask is a grayscale image with the same shape as the input image.

The SageMaker semantic segmentation algorithm is built using the MXNet Gluon framework and the Gluon CV toolkit. It provides you with a choice of three built-in algorithms to train a deep neural network. You can use the Fully-Convolutional Network (FCN) algorithm, Pyramid Scene Parsing (PSP) algorithm, or DeepLabV3.

Each of the three algorithms has two distinct components:

- The **backbone** (or encoder)—A network that produces reliable activation maps of features.
- The **decoder**—A network that constructs the segmentation mask from the encoded activation maps.

You also have a choice of backbones for the FCN, PSP, and DeepLabV3 algorithms: ResNet50 or ResNet101. These backbones include pretrained artifacts that were originally trained on the ImageNet classification task. You can fine-tune these backbones for segmentation using your own data. Or, you can initialize and train these networks from scratch using only your own data. The decoders are never pretrained.

To deploy the trained model for inference, use the SageMaker hosting service. During inference, you can request the segmentation mask either as a PNG image or as a set of probabilities for each class for each pixel. You can use these masks as part of a larger pipeline that includes additional downstream image processing or other applications.

**Topics**

- Semantic Segmentation Sample Notebooks
- Input/Output Interface for the Semantic Segmentation Algorithm
- EC2 Instance Recommendation for the Semantic Segmentation Algorithm
- Semantic Segmentation Hyperparameters
- Tuning a Semantic Segmentation Model
**Semantic Segmentation Sample Notebooks**

For a sample Jupyter notebook that uses the SageMaker semantic segmentation algorithm to train a model and deploy it to perform inferences, see the Semantic Segmentation Example. For instructions on how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances.

To see a list of all of the SageMaker samples, create and open a notebook instance, and choose the SageMaker Examples tab. The example semantic segmentation notebooks are located under Introduction to Amazon algorithms. To open a notebook, choose its Use tab, and choose Create copy.

**Input/Output Interface for the Semantic Segmentation Algorithm**

SageMaker semantic segmentation expects the customer's training dataset to be on Amazon Simple Storage Service (Amazon S3). Once trained, it produces the resulting model artifacts on Amazon S3. The input interface format for the SageMaker semantic segmentation is similar to that of most standardized semantic segmentation benchmarking datasets. The dataset in Amazon S3 is expected to be presented in two channels, one for train and one for validation using four directories, two for images and two for annotations. Annotations are expected to be uncompressed PNG images. The dataset might also have a label map that describes how the annotation mappings are established. If not, the algorithm uses a default. It also supports the augmented manifest image format (application/x-image) for training in Pipe input mode straight from Amazon S3. For inference, an endpoint accepts images with an image/jpeg content type.

**How Training Works**

The training data is split into four directories: train, train_annotation, validation, and validation_annotation. There is a channel for each of these directories. The dataset also expected to have one label_map.json file per channel for train_annotation and validation_annotation respectively. If you don't provide these JSON files, SageMaker provides the default set label map.

The dataset specifying these files should look similar to the following example:

```text
s3://bucket_name
  |- train
      | - 0000.jpg
      | - coffee.jpg
```
Every JPG image in the train and validation directories has a corresponding PNG label image with the same name in the train_annotation and validation_annotation directories. This naming convention helps the algorithm to associate a label with its corresponding image during training. The train, train_annotation, validation, and validation_annotation channels are mandatory. The annotations are single-channel PNG images. The format works as long as the metadata (modes) in the image helps the algorithm read the annotation images into a single-channel 8-bit unsigned integer. For more information on our support for modes, see the Python Image Library documentation. We recommend using the 8-bit pixel, true color P mode.

The image that is encoded is a simple 8-bit integer when using modes. To get from this mapping to a map of a label, the algorithm uses one mapping file per channel, called the label map. The label map is used to map the values in the image with actual label indices. In the default label map, which is provided by default if you don’t provide one, the pixel value in an annotation matrix (image) directly index the label. These images can be grayscale PNG files or 8-bit indexed PNG files. The label map file for the unscaled default case is the following:

```
{
    "scale": "1"
}
```

To provide some contrast for viewing, some annotation software scales the label images by a constant amount. To support this, the SageMaker semantic segmentation algorithm provides a rescaling option to scale down the values to actual label values. When scaling down doesn’t convert the value to an appropriate integer, the algorithm defaults to the greatest integer less than
or equal to the scale value. The following code shows how to set the scale value to rescale the label values:

```json
{
   "scale": "3"
}
```

The following example shows how this "scale" value is used to rescale the encoded_label values of the input annotation image when they are mapped to the mapped_label values to be used in training. The label values in the input annotation image are 0, 3, 6, with scale 3, so they are mapped to 0, 1, 2 for training:

```python
encoded_label = [0, 3, 6]
mapped_label = [0, 1, 2]
```

In some cases, you might need to specify a particular color mapping for each class. Use the map option in the label mapping as shown in the following example of a label_map file:

```json
{
   "map": {
      "0": 5,
      "1": 0,
      "2": 2
   }
}
```

This label mapping for this example is:

```python
encoded_label = [0, 5, 2]
mapped_label = [1, 0, 2]
```

With label mappings, you can use different annotation systems and annotation software to obtain data without a lot of preprocessing. You can provide one label map per channel. The files for a label map in the label_map channel must follow the naming conventions for the four directory structure. If you don't provide a label map, the algorithm assumes a scale of 1 (the default).

**Training with the Augmented Manifest Format**

The augmented manifest format enables you to do training in Pipe mode using image files without needing to create RecordIO files. The augmented manifest file contains data objects and should be
in JSON Lines format, as described in the `CreateTrainingJob` request. Each line in the manifest is an entry containing the Amazon S3 URI for the image and the URI for the annotation image.

Each JSON object in the manifest file must contain a `source-ref` key. The `source-ref` key should contain the value of the Amazon S3 URI to the image. The labels are provided under the `AttributeNames` parameter value as specified in the `CreateTrainingJob` request. It can also contain additional metadata under the `metadata` tag, but these are ignored by the algorithm. In the example below, the `AttributeNames` are contained in the list of image and annotation references ["source-ref", "city-streets-ref"]. These names must have -ref appended to them. When using the Semantic Segmentation algorithm with Augmented Manifest, the value of the `RecordWrapperType` parameter must be "RecordIO" and value of the `ContentType` parameter must be application/x-recordio.

```json
{"source-ref": "S3 bucket location", "city-streets-ref": "S3 bucket location", "city-streets-metadata": {"job-name": "label-city-streets"} }
```

For more information on augmented manifest files, see [Provide Dataset Metadata to Training Jobs with an Augmented Manifest File](#).

### Incremental Training

You can also seed the training of a new model with a model that you trained previously using SageMaker. This incremental training saves training time when you want to train a new model with the same or similar data. Currently, incremental training is supported only for models trained with the built-in SageMaker Semantic Segmentation.

To use your own pre-trained model, specify the `ChannelName` as "model" in the `InputDataConfig` for the `CreateTrainingJob` request. Set the `ContentType` for the model channel to application/x-sagemaker-model. The backbone, algorithm, `crop_size`, and `num_classes` input parameters that define the network architecture must be consistently specified in the input hyperparameters of the new model and the pre-trained model that you upload to the model channel. For the pretrained model file, you can use the compressed (.tar.gz) artifacts from SageMaker outputs. You can only use Image formats for input data. For more information on incremental training and for instructions on how to use it, see [Use Incremental Training in Amazon SageMaker](#).

### Produce Inferences

To query a trained model that is deployed to an endpoint, you need to provide an image and an `AcceptType` that denotes the type of output required. The endpoint takes JPEG images with an
image/jpeg content type. If you request an AcceptType of image/png, the algorithm outputs a PNG file with a segmentation mask in the same format as the labels themselves. If you request an accept type of application/x-recordio-protobuf, the algorithm returns class probabilities encoded in recordio-protobuf format. The latter format outputs a 3D tensor where the third dimension is the same size as the number of classes. This component denotes the probability of each class label for each pixel.

**EC2 Instance Recommendation for the Semantic Segmentation Algorithm**

The SageMaker semantic segmentation algorithm only supports GPU instances for training, and we recommend using GPU instances with more memory for training with large batch sizes. The algorithm can be trained using P2, P3, G4dn, or G5 instances in single machine configurations.

For inference, you can use either CPU instances (such as C5 and M5) and GPU instances (such as P3 and G4dn) or both. For information about the instance types that provide varying combinations of CPU, GPU, memory, and networking capacity for inference, see [Amazon SageMaker ML Instance Types](#).

**Semantic Segmentation Hyperparameters**

The following tables list the hyperparameters supported by the Amazon SageMaker semantic segmentation algorithm for network architecture, data inputs, and training. You specify Semantic Segmentation for training in the AlgorithmName of the CreateTrainingJob request.

**Network Architecture Hyperparameters**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>backbone</td>
<td>The backbone to use for the algorithm's encoder component.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: resnet-50, resnet-101</td>
</tr>
<tr>
<td></td>
<td>Default value: resnet-50</td>
</tr>
<tr>
<td>use_pretrained_model</td>
<td>Whether a pretrained model is to be used for the backbone.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: True, False</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Default value: True</td>
<td></td>
</tr>
</tbody>
</table>

**algorithm**

The algorithm to use for semantic segmentation.

*Optional*

Valid values:

- fcn: [Fully-Convolutional Network (FCN) algorithm](#)
- psp: [Pyramid Scene Parsing (PSP) algorithm](#)
- deeplab: [DeepLab V3 algorithm](#)

Default value: fcn

---

**Data Hyperparameters**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_classes</td>
<td>The number of classes to segment.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 2 ≤ positive integer ≤ 254</td>
</tr>
<tr>
<td>num_training_samples</td>
<td>The number of samples in the training data. The algorithm uses this value to set up the learning rate scheduler.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td>base_size</td>
<td>Defines how images are rescaled before cropping. Images are rescaled such that the long size length is set to base_size multiplied by a random number from 0.5 to 2.0, and the short size is computed to preserve the aspect ratio.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
</tbody>
</table>
### Training Hyperparameters

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>early_stopping</td>
<td>Whether to use early stopping logic during training.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: True, False</td>
</tr>
<tr>
<td></td>
<td>Default value: False</td>
</tr>
<tr>
<td>early_stopping_min_epochs</td>
<td>The minimum number of epochs that must be run.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td>early_stopping_patience</td>
<td>The number of epochs that meet the tolerance for lower performance before the algorithm enforces an early stop.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
</tbody>
</table>

**Parameter Name**

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valid values: positive integer &gt; 16</td>
</tr>
<tr>
<td>Default value: 520</td>
</tr>
<tr>
<td>Optional</td>
</tr>
<tr>
<td>Valid values: positive integer &gt; 16</td>
</tr>
<tr>
<td>Default value: 240</td>
</tr>
<tr>
<td>Parameter Name</td>
</tr>
<tr>
<td>-----------------------------</td>
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<tr>
<td></td>
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<tr>
<td>early_stopping_tolerance</td>
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<tr>
<td>epochs</td>
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<td></td>
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<tr>
<td></td>
</tr>
<tr>
<td>gamma1</td>
</tr>
<tr>
<td></td>
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<tr>
<td></td>
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<tr>
<td></td>
</tr>
<tr>
<td>gamma2</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Parameter Name</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>learning_rate</td>
</tr>
</tbody>
</table>
| lr_scheduler    | The shape of the learning rate schedule that controls its decrease over time. **Optional** Valid values:  
  - **step**: A stepwise decay, where the learning rate is reduced (multiplied) by the $\text{lr_scheduler_factor}$ after epochs specified by $\text{lr_scheduler_step}$.  
  - **poly**: A smooth decay using a polynomial function.  
  - **cosine**: A smooth decay using a cosine function.  
 Default value: poly |
<p>| lr_scheduler_factor | If $\text{lr_scheduler}$ is set to step, the ratio by which to reduce (multiply) the $\text{learning_rate}$ after each of the epochs specified by the $\text{lr_scheduler_step}$. Otherwise, ignored. <strong>Optional</strong> Valid values: $0 \leq \text{float} \leq 1$ Default value: 0.1 |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>lr_scheduler_step</td>
<td>A comma delimited list of the epochs after which the learning_rate is reduced (multiplied) by an lr_scheduler_factor. For example, if the value is set to &quot;10, 20&quot;, then the learning_rate is reduced by lr_scheduler_factor after the 10th epoch and again by this factor after 20th epoch.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditionally Required</strong> if lr_scheduler is set to step. Otherwise, ignored.</td>
</tr>
<tr>
<td></td>
<td>Valid values: string</td>
</tr>
<tr>
<td></td>
<td>Default value: (No default, as the value is required when used.)</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>The batch size for training. Using a large mini_batch_size usually results in faster training, but it might cause you to run out of memory. Memory usage is affected by the values of the mini_batch_size and image_shape parameters, and the backbone architecture.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: positive integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 16</td>
</tr>
<tr>
<td>momentum</td>
<td>The momentum for the sgd optimizer. When you use other optimizers, the semantic segmentation algorithm ignores this parameter.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: 0 &lt; float ≤ 1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.9</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| **optimizer**  | The type of optimizer. For more information about an optimizer, choose the appropriate link:  
  - adam: *Adaptive momentum estimation*  
  - adagrad: *Adaptive gradient descent*  
  - nag: *Nesterov accelerated gradient*  
  - rmsprop: *Root mean square propagation*  
  - sgd: *Stochastic gradient descent*  
  
  **Optional**  
  Valid values: adam, adagrad, nag, rmsprop, sgd  
  Default value: sgd |
| **syncbn**     | If set to True, the batch normalization mean and variance are computed over all the samples processed across the GPUs.  
  
  **Optional**  
  Valid values: True, False  
  Default value: False |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| validation_mini_batch_size | The batch size for validation. A large mini_batch_size usually results in faster training, but it might cause you to run out of memory. Memory usage is affected by the values of the mini_batch_size and image_shape parameters, and the backbone architecture.  
- To score the validation on the entire image without cropping the images, set this parameter to 1. Use this option if you want to measure performance on the entire image as a whole.  

**Note**  
Setting the validation_mini_batch_size parameter to 1 causes the algorithm to create a new network model for every image. This might slow validation and training.  

- To crop images to the size specified in the crop_size parameter, even during evaluation, set this parameter to a value greater than 1.  

**Optional**  
Valid values: positive integer  
Default value: 16 |
| weight_decay       | The weight decay coefficient for the sgd optimizer. When you use other optimizers, the algorithm ignores this parameter.  

**Optional**  
Valid values: 0 < float < 1  
Default value: 0.0001 |
Tuning a Semantic Segmentation Model

Automatic model tuning, also known as hyperparameter tuning, finds the best version of a model by running many jobs that test a range of hyperparameters on your dataset. You choose the tunable hyperparameters, a range of values for each, and an objective metric. You choose the objective metric from the metrics that the algorithm computes. Automatic model tuning searches the hyperparameters chosen to find the combination of values that result in the model that optimizes the objective metric.

Metrics Computed by the Semantic Segmentation Algorithm

The semantic segmentation algorithm reports two validation metrics. When tuning hyperparameter values, choose one of these metrics as the objective.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Optimization Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>validation:mIOU</td>
<td>The area of the intersection of the predicted segmentation and the ground truth divided by the area of union between them for images in the validation set. Also known as the Jaccard Index.</td>
<td>Maximize</td>
</tr>
<tr>
<td>validation:pixel_accuracy</td>
<td>The percentage of pixels that are correctly classified in images from the validation set.</td>
<td>Maximize</td>
</tr>
</tbody>
</table>

Tunable Semantic Segmentation Hyperparameters

You can tune the following hyperparameters for the semantic segmentation algorithm.

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Parameter Type</th>
<th>Recommended Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>learning_rate</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-4, MaxValue: 1e-1</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Parameter Type</td>
<td>Recommended Ranges</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------</td>
<td>----------------------------</td>
</tr>
<tr>
<td>mini_batch_size</td>
<td>IntegerParameterRanges</td>
<td>MinValue: 1, MaxValue: 128</td>
</tr>
<tr>
<td>momentum</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 0.9, MaxValue: 0.999</td>
</tr>
<tr>
<td>optimizer</td>
<td>CategoricalParameterRanges</td>
<td>['sgd', 'adam', 'adadelta']</td>
</tr>
<tr>
<td>weight_decay</td>
<td>ContinuousParameterRange</td>
<td>MinValue: 1e-5, MaxValue: 1e-3</td>
</tr>
</tbody>
</table>

**Use Reinforcement Learning with Amazon SageMaker**

Reinforcement learning (RL) combines fields such as computer science, neuroscience, and psychology to determine how to map situations to actions to maximize a numerical reward signal. This notion of a reward signal in RL stems from neuroscience research into how the human brain makes decisions about which actions maximize reward and minimize punishment. In most situations, humans are not given explicit instructions on which actions to take, but instead must learn both which actions yield the most immediate rewards, and how those actions influence future situations and consequences.

The problem of RL is formalized using Markov decision processes (MDPs) that originate from dynamical systems theory. MDPs aim to capture high-level details of a real problem that a learning agent encounters over some period of time in attempting to achieve some ultimate goal. The learning agent should be able to determine the current state of its environment and identify possible actions that affect the learning agent’s current state. Furthermore, the learning agent’s goals should correlate strongly to the state of the environment. A solution to a problem formulated in this way is known as a reinforcement learning method.

**What are the differences between reinforcement, supervised, and unsupervised learning paradigms?**

Machine learning can be divided into three distinct learning paradigms: supervised, unsupervised, and reinforcement.
In supervised learning, an external supervisor provides a training set of labeled examples. Each example contains information about a situation, belongs to a category, and has a label identifying the category to which it belongs. The goal of supervised learning is to generalize in order to predict correctly in situations that are not present in the training data.

In contrast, RL deals with interactive problems, making it infeasible to gather all possible examples of situations with correct labels that an agent might encounter. This type of learning is most promising when an agent is able to accurately learn from its own experience and adjust accordingly.

In unsupervised learning, an agent learns by uncovering structure within unlabeled data. While a RL agent might benefit from uncovering structure based on its experiences, the sole purpose of RL is to maximize a reward signal.

Topics

• Why is Reinforcement Learning Important?
• Markov Decision Process (MDP)
• Key Features of Amazon SageMaker RL
• Reinforcement Learning Sample Notebooks
• Sample RL Workflow Using Amazon SageMaker RL
• RL Environments in Amazon SageMaker
• Distributed Training with Amazon SageMaker RL
• Hyperparameter Tuning with Amazon SageMaker RL

Why is Reinforcement Learning Important?

RL is well-suited for solving large, complex problems, such as supply chain management, HVAC systems, industrial robotics, game artificial intelligence, dialog systems, and autonomous vehicles. Because RL models learn by a continuous process of receiving rewards and punishments for every action taken by the agent, it is possible to train systems to make decisions under uncertainty and in dynamic environments.

Markov Decision Process (MDP)

RL is based on models called Markov Decision Processes (MDPs). An MDP consists of a series of time steps. Each time step consists of the following:
Environment

Defines the space in which the RL model operates. This can be either a real-world environment or a simulator. For example, if you train a physical autonomous vehicle on a physical road, that would be a real-world environment. If you train a computer program that models an autonomous vehicle driving on a road, that would be a simulator.

State

Specifies all information about the environment and past steps that is relevant to the future. For example, in an RL model in which a robot can move in any direction at any time step, the position of the robot at the current time step is the state, because if we know where the robot is, it isn't necessary to know the steps it took to get there.

Action

What the agent does. For example, the robot takes a step forward.

Reward

A number that represents the value of the state that resulted from the last action that the agent took. For example, if the goal is for a robot to find treasure, the reward for finding treasure might be 5, and the reward for not finding treasure might be 0. The RL model attempts to find a strategy that optimizes the cumulative reward over the long term. This strategy is called a policy.

Observation

Information about the state of the environment that is available to the agent at each step. This might be the entire state, or it might be just a part of the state. For example, the agent in a chess-playing model would be able to observe the entire state of the board at any step, but a robot in a maze might only be able to observe a small portion of the maze that it currently occupies.

Typically, training in RL consists of many episodes. An episode consists of all of the time steps in an MDP from the initial state until the environment reaches the terminal state.

Key Features of Amazon SageMaker RL

To train RL models in SageMaker RL, use the following components:


• An RL environment. You can use custom environments, open-source environments, or commercial environments. For information, see RL Environments in Amazon SageMaker.

The following diagram shows the RL components that are supported in SageMaker RL.
Reinforcement Learning Sample Notebooks

For complete code examples, see the reinforcement learning sample notebooks in the SageMaker Examples repository.

Sample RL Workflow Using Amazon SageMaker RL

The following example describes the steps for developing RL models using Amazon SageMaker RL.

1. **Formulate the RL problem**—First, formulate the business problem into an RL problem. For example, auto scaling enables services to dynamically increase or decrease capacity depending on conditions that you define. Currently, this requires setting up alarms, scaling policies, thresholds, and other manual steps. To solve this with RL, we define the components of the Markov Decision Process:
   a. **Objective**—Scale instance capacity so that it matches the desired load profile.
   b. **Environment**—A custom environment that includes the load profile. It generates a simulated load with daily and weekly variations and occasional spikes. The simulated system has a delay between when new resources are requested and when they become available for serving requests.
   c. **State**—The current load, number of failed jobs, and number of active machines.
   d. **Action**—Remove, add, or keep the same number of instances.
   e. **Reward**—A positive reward for successful transactions and a high penalty for failing transactions beyond a specified threshold.

2. **Define the RL environment**—The RL environment can be the real world where the RL agent interacts or a simulation of the real world. You can connect open source and custom environments developed using Gym interfaces and commercial simulation environments such as MATLAB and Simulink.

3. **Define the presets**—The presets configure the RL training jobs and define the hyperparameters for the RL algorithms.

4. **Write the training code**—Write training code as a Python script and pass the script to a SageMaker training job. In your training code, import the environment files and the preset files, and then define the main() function.

5. **Train the RL Model**—Use the SageMaker RLEstimator in the Amazon SageMaker Python SDK to start an RL training job. If you are using local mode, the training job runs on the notebook instance. When you use SageMaker for training, you can select GPU or CPU.
instances. Store the output from the training job in a local directory if you train in local mode, or on Amazon S3 if you use SageMaker training.

The RLEstimator requires the following information as parameters.

a. The source directory where the environment, presets, and training code are uploaded.
b. The path to the training script.
c. The RL toolkit and deep learning framework you want to use. This automatically resolves to the Amazon ECR path for the RL container.
d. The training parameters, such as the instance count, job name, and S3 path for output.
e. Metric definitions that you want to capture in your logs. These can also be visualized in CloudWatch and in SageMaker notebooks.

6. **Visualize training metrics and output**—After a training job that uses an RL model completes, you can view the metrics you defined in the training jobs in CloudWatch. You can also plot the metrics in a notebook by using the Amazon SageMaker Python SDK analytics library. Visualizing metrics helps you understand how the performance of the model as measured by the reward improves over time.

**Note**

If you train in local mode, you can’t visualize metrics in CloudWatch.

7. **Evaluate the model**—Checkpointed data from the previously trained models can be passed on for evaluation and inference in the checkpoint channel. In local mode, use the local directory. In SageMaker training mode, you need to upload the data to S3 first.

8. **Deploy RL models**—Finally, deploy the trained model on an endpoint hosted on SageMaker containers or on an edge device by using AWS IoT Greengrass.

For more information on RL with SageMaker, see [Using RL with the SageMaker Python SDK](#).

**RL Environments in Amazon SageMaker**

Amazon SageMaker RL uses environments to mimic real-world scenarios. Given the current state of the environment and an action taken by the agent or agents, the simulator processes the impact of the action, and returns the next state and a reward. Simulators are useful in cases where it is not safe to train an agent in the real world (for example, flying a drone) or if the RL algorithm takes a long time to converge (for example, when playing chess).
The following diagram shows an example of the interactions with a simulator for a car racing game.

![Diagram showing interactions with a simulator for a car racing game]

The simulation environment consists of an agent and a simulator. Here, a convolutional neural network (CNN) consumes images from the simulator and generates actions to control the game controller. With multiple simulations, this environment generates training data of the form \(\text{state}_t, \text{action}, \text{state}_{t+1}, \text{reward}_{t+1}\). Defining the reward is not trivial and impacts the RL model quality. We want to provide a few examples of reward functions, but would like to make it user-configurable.

**Topics**

- [Use OpenAI Gym Interface for Environments in SageMaker RL](#)
- [Use Open-Source Environments](#)
- [Use Commercial Environments](#)

**Use OpenAI Gym Interface for Environments in SageMaker RL**

To use OpenAI Gym environments in SageMaker RL, use the following API elements. For more information about OpenAI Gym, see [Gym Documentation](#).

- `env.action_space`—Defines the actions the agent can take, specifies whether each action is continuous or discrete, and specifies the minimum and maximum if the action is continuous.
- `env.observation_space`—Defines the observations the agent receives from the environment, as well as minimum and maximum for continuous observations.
- `env.reset()`—Initializes a training episode. The `reset()` function returns the initial state of the environment, and the agent uses the initial state to take its first action. The action is
then sent to `step()` repeatedly until the episode reaches a terminal state. When `step()` returns `done = True`, the episode ends. The RL toolkit re-initializes the environment by calling `reset()`.

- **step()**—Takes the agent action as input and outputs the next state of the environment, the reward, whether the episode has terminated, and an `info` dictionary to communicate debugging information. It is the responsibility of the environment to validate the inputs.

- **`env.render()`**—Used for environments that have visualization. The RL toolkit calls this function to capture visualizations of the environment after each call to the `step()` function.

### Use Open-Source Environments

You can use open-source environments, such as EnergyPlus and RoboSchool, in SageMaker RL by building your own container. For more information about EnergyPlus, see [https://energyplus.net/](https://energyplus.net/). For more information about RoboSchool, see [https://github.com/openai/roboschool](https://github.com/openai/roboschool). The HVAC and RoboSchool examples in the [SageMaker examples repository](https://github.com/aws/sagemaker-examples) show how to build a custom container to use with SageMaker RL:

### Use Commercial Environments

You can use commercial environments, such as MATLAB and Simulink, in SageMaker RL by building your own container. You need to manage your own licenses.

### Distributed Training with Amazon SageMaker RL

Amazon SageMaker RL supports multi-core and multi-instance distributed training. Depending on your use case, training and/or environment rollout can be distributed. For example, SageMaker RL works for the following distributed scenarios:

- **Single training instance and multiple rollout instances of the same instance type.** For an example, see the Neural Network Compression example in the [SageMaker examples repository](https://github.com/aws/sagemaker-examples).

- **Single trainer instance and multiple rollout instances, where different instance types for training and rollouts.** For an example, see the AWS DeepRacer / AWS RoboMaker example in the [SageMaker examples repository](https://github.com/aws/sagemaker-examples).

- **Single trainer instance that uses multiple cores for rollout.** For an example, see the Roboschool example in the [SageMaker examples repository](https://github.com/aws/sagemaker-examples). This is useful if the simulation environment is light-weight and can run on a single thread.

- **Multiple instances for training and rollouts.** For an example, see the Roboschool example in the [SageMaker examples repository](https://github.com/aws/sagemaker-examples).
Hyperparameter Tuning with Amazon SageMaker RL

You can run a hyperparameter tuning job to optimize hyperparameters for Amazon SageMaker RL. The Roboschool example in the sample notebooks in the SageMaker examples repository shows how you can do this with RL Coach. The launcher script shows how you can abstract parameters from the Coach preset file and optimize them.

Run your local code as a SageMaker training job

You can run your local machine learning (ML) Python code as a large single-node Amazon SageMaker training job or as multiple parallel jobs. You can do this by annotating your code with an @remote decorator, as shown in the following code example. Distributed training (across multiple instances) are not supported with remote functions.

```python
@remote(**settings)
def divide(x, y):
    return x / y
```

The SageMaker Python SDK will automatically translate your existing workspace environment and any associated data processing code and datasets into a SageMaker training job that runs on the SageMaker training platform. You can also activate a persistent cache feature, which will further reduce job start latency by caching previously downloaded dependency packages. This reduction in job latency is greater than the reduction in latency from using SageMaker managed warm pools alone. For more information, see Using persistent cache.

Note

Distributed training jobs are not supported by remote functions.

The following sections show how to annotate your local ML code with an @remote decorator and tailor your experience for your use case. This includes customizing your environment and integrating with SageMaker Experiments.

Topics

- Set up your environment
- Invoking a function
Set up your environment

Choose one of the following three options to set up your environment.

Run your code from Amazon SageMaker Studio Classic

You can annotate and run your local ML code from SageMaker Studio Classic by creating a SageMaker Notebook and attaching any image available on SageMaker Studio Classic image. The following instructions help you create a SageMaker Notebook, install the SageMaker Python SDK, and annotate your code with the decorator.

1. Create a SageMaker Notebook and attach an image in SageMaker Studio Classic as follows:
   a. Follow the instructions in Launch Amazon SageMaker Studio Classic in the Amazon SageMaker Developer Guide.
   b. Select Studio from the left navigation pane. This opens a new window.
   c. In the Get Started dialog box, select a user profile from the down arrow. This opens a new window.
   d. Select Open Studio Classic.
   e. Select Open Launcher from the main working area. This opens a new page.
   f. Select Create notebook from the main working area.
   g. Select Base Python 3.0 from the down arrow next to Image in the Change environment dialog box.

The @remote decorator automatically detects the image attached to the SageMaker Studio Classic notebook and uses it to run the SageMaker training job. If image_uri is specified either as an argument in the decorator or in the configuration file, then the value specified in image_uri will be used instead of the detected image.
For more information about how to create a notebook in SageMaker Studio Classic, see the Create a Notebook from the File Menu section in Create or Open an Amazon SageMaker Studio Classic Notebook.

For a list of available images, see Supported Docker images.

2. Install the SageMaker Python SDK.

To annotate your code with the @remote function inside a SageMaker Studio Classic Notebook, you must have the SageMaker Python SDK installed. Install the SageMaker Python SDK, as shown in the following code example.

```
!pip install sagemaker
```

3. Use @remote decorator to run functions in a SageMaker training job.

To run your local ML code, first create a dependencies file to instruct SageMaker where to locate your local code. To do so, follow these steps:

a. From the SageMaker Studio Classic Launcher main working area, in Utilities and files, choose Text file. This opens a new tab with a text file called untitled.txt.

For more information about the SageMaker Studio Classic user interface (UI), see Amazon SageMaker Studio Classic UI Overview.

b. Rename untitled.txt to requirements.txt.

c. Add all the dependencies required for the code along with the SageMaker library to requirements.txt.

A minimal code example for requirements.txt for the example divide function is provided in the following section, as follows.

```
sagemaker
```

d. Run your code with the remote decorator by passing the dependencies file, as follows.

```
from sagemaker.remote_function import remote

@remote(instance_type="ml.m5.xlarge", dependencies='./requirements.txt')
def divide(x, y):
  return x / y
```
divide(2, 3.0)

For additional code examples, see the sample notebook quick_start.ipynb.

If you’re already running a SageMaker Studio Classic notebook, and you install the Python SDK as instructed in 2. Install the SageMaker Python SDK, you must restart your kernel. For more information, see Use the SageMaker Studio Classic Notebook Toolbar in the Amazon SageMaker Developer Guide.

Run your code from an Amazon SageMaker notebook

You can annotate your local ML code from a SageMaker notebook instance. The following instructions show how to create a notebook instance with a custom kernel, install the SageMaker Python SDK, and annotate your code with the decorator.

1. Create a notebook instance with a custom conda kernel.

You can annotate your local ML code with an @remote decorator to use inside of a SageMaker training job. First you must create and customize a SageMaker notebook instance to use a kernel with Python version 3.7 or higher, up to 3.10.x. To do so, follow these steps:

b. In the left navigation panel, choose Notebook to expand its options.
c. Choose Notebook Instances from the expanded options.
d. Choose the Create Notebook Instance button. This opens a new page.
e. For Notebook instance name, enter a name with a maximum of 63 characters and no spaces. Valid characters: A-Z, a-z, 0-9, and :+=@_%- (hyphen).
f. In the Notebook instance settings dialog box, expand the right arrow next to Additional Configuration.
g. Under Lifecycle configuration - optional, expand the down arrow and select Create a new lifecycle configuration. This opens a new dialog box.
h. Under Name, enter a name for your configuration setting.
i. In the Scripts dialog box, in the Start notebook tab, replace the existing contents of the text box with the following script.

```bash
#!/bin/bash
```

Set up your environment
set -e

sudo -u ec2-user -i <<'EOF'
unset SUDO_UID
WORKING_DIR=/home/ec2-user/SageMaker/custom-miniconda/
source "$WORKING_DIR/miniconda/bin/activate"
for env in $WORKING_DIR/miniconda/envs/\*; do
    BASENAME=$(basename "$env")
    source activate "$BASENAME"
    python -m ipykernel install --user --name "$BASENAME" --display-name "Custom ($BASENAME)"
done
EOF

echo "Restarting the Jupyter server.."
# restart command is dependent on current running Amazon Linux and JupyterLab
CURR_VERSION_AL=$(cat /etc/system-release)
CURR_VERSION_JS=$(jupyter --version)
if [[ $CURR_VERSION_JS == *"jupyter_core : 4.9.1"* ]] && [[ $CURR_VERSION_AL == *" release 2018"* ]]; then
    sudo initctl restart jupyter-server --no-wait
else
    sudo systemctl --no-block restart jupyter-server.service
fi

j. In the **Scripts** dialog box, in the **Create notebook** tab, replace the existing contents of the text box with the following script.

```bash
#!/bin/bash

set -e

sudo -u ec2-user -i <<'EOF'
unset SUDO_UID
# Install a separate conda installation via Miniconda
WORKING_DIR=/home/ec2-user/SageMaker/custom-miniconda
mkdir -p "$WORKING_DIR"
wget https://repo.anaconda.com/miniconda/Miniconda3-4.6.14-Linux-x86_64.sh -O "$WORKING_DIR/miniconda.sh"
bash "$WORKING_DIR/miniconda.sh" -b -u -p "$WORKING_DIR/miniconda"
rm -rf "$WORKING_DIR/miniconda.sh"
# Create a custom conda environment
EOF
```
source "${WORKING_DIR}/miniconda/bin/activate"
KERNEL_NAME="custom_python310"
PYTHON="3.10"
conda create --yes --name "${KERNEL_NAME}" python="${PYTHON}" pip
conda activate "${KERNEL_NAME}"
pip install --quiet ipykernel
# Customize these lines as necessary to install the required packages
EOF

k. Choose the **Create configuration** button on the bottom right of the window.
l. Choose the **Create notebook instance** button on the bottom right of the window.
m. Wait for the notebook instance **Status** to change from **Pending** to **InService**.

2. Create a Jupyter notebook in the notebook instance.

The following instructions show how to create a Jupyter notebook using Python 3.10 in your newly created SageMaker instance.

a. After the notebook instance **Status** from the previous step is **InService**, do the following:
   i. Select **Open Jupyter** under **Actions** in the row containing your newly created notebook instance **Name**. This opens a new Jupyter server.

b. In the Jupyter server, select **New** from the top right menu.

c. From the down arrow, select **conda_custom_python310**. This creates a new Jupyter notebook that uses a Python 3.10 kernel. This new Jupyter notebook can now be used similarly to a local Jupyter notebook.

3. Install the SageMaker Python SDK.

After your virtual environment is running, install the SageMaker Python SDK by using the following code example.

```bash
!pip install sagemaker
```

4. Use an @remote decorator to run functions in a SageMaker training job.

When you annotate your local ML code with an @remote decorator inside the SageMaker notebook, SageMaker training will automatically interpret the function of your code and run it as a SageMaker training job. Set up your notebook by doing the following:

a. Select the kernel name in the notebook menu from the SageMaker notebook instance that you created in step 1, **Create a SageMaker Notebook instance with a custom kernel**.
For more information, see Change an Image or a Kernel.

b. From the down arrow, choose the a custom conda kernel that uses a version of Python that is 3.7 or higher.

As an example, selecting conda_custom_python310 chooses the kernel for Python 3.10.

c. Choose Select.

d. Wait for the kernel's status to show as idle, which indicates that the kernel has started.

e. In the Jupyter Server Home, select New from the top right menu.

f. Next to the down arrow, select Text file. This creates a new text file called untitled.txt.

g. Rename untitled.txt to requirements.txt and add any dependencies required for the code along with sagemaker.

h. Run your code with the remote decorator by passing the dependencies file as shown below.

```python
from sagemaker.remote_function import remote

@remote(instance_type="ml.m5.xlarge", dependencies='./requirements.txt')
def divide(x, y):
    return x / y

divide(2, 3.0)
```

See the sample notebook quick_start.ipnyb for additional code examples.

Run your code from within your local IDE

You can annotate your local ML code with an @remote decorator inside your preferred local IDE. The following steps show the necessary prerequisites, how to install the Python SDK, and how to annotate your code with the @remote decorator.

1. Install prerequisites by setting up the AWS Command Line Interface (AWS CLI) and creating a role, as follows:
   - Onboard to a SageMaker domain following the instructions in the AWS CLI Prerequisites section of Set Up Amazon SageMaker Prerequisites.
   - Create an IAM role following the Create execution role section of SageMaker Roles.

2. Create a virtual environment by using either PyCharm or conda and using Python version 3.7 or higher, up to 3.10.x.
• Set up a virtual environment using PyCharm as follows:
  a. Select **File** from the main menu.
  b. Choose **New Project**.
  c. Choose **Conda** from the down arrow under **New environment using**.
  d. In the field for **Python version** use the down arrow to select a version of Python that is 3.7 or above. You can go up to 3.10.x from the list.

![PyCharm New Project dialog](image)

• If you have Anaconda installed, you can set up a virtual environment using conda, as follows:
  • Open an Anaconda prompt terminal interface.
  • Create and activate a new conda environment using a Python version of 3.7 or higher, up to 3.10x. The following code example shows how to create a conda environment using Python version 3.10.

```
conda create -n sagemaker_jobs_quick_start python=3.10 pip
conda activate sagemaker_jobs_quick_start
```

3. Install the SageMaker Python SDK.
To package your code from your preferred IDE, you must have a virtual environment set up using Python 3.7 or higher, up to 3.10x. You also need a compatible container image. Install the SageMaker Python SDK using the following code example.

```
pip install sagemaker
```

4. Wrap your code inside the `@remote` decorator. The SageMaker Python SDK will automatically interpret the function of your code and run it as a SageMaker training job. The following code examples show how to import the necessary libraries, set up a SageMaker session, and annotate a function with the `@remote` decorator.

You can run your code by either providing the dependencies needed directly, or by using dependencies from the active conda environment.

- **To provide the dependencies directly, do the following:**
  - Create a `requirements.txt` file in the working directory that the code resides in.
  - Add all of the dependencies required for the code along with the SageMaker library. The following section provides a minimal code example for `requirements.txt` for the example `divide` function.

```
sagemaker
```

- **Run your code with the `@remote` decorator by passing the dependencies file. In the following code example, replace the IAM role name with an AWS Identity and Access Management (IAM) role ARN that you would like SageMaker to use to run your job.**

```
import boto3
import sagemaker
from sagemaker.remote_function import remote

sm_session =
    sagemaker.Session(boto_session=boto3.session.Session(region_name="us-west-2"))
settings = dict(
    sagemaker_session=sm_session,
    role=<The IAM role name>,
    instance_type="ml.m5.xlarge",
    dependencies='./requirements.txt'
)
@remote(**settings)
```
def divide(x, y):
    return x / y

if __name__ == "__main__":
    print(divide(2, 3.0))

• To use dependencies from the active conda environment, use the value `auto_capture` for the dependencies parameter, as shown in the following.

```python
import boto3
import sagemaker
from sagemaker.remote_function import remote

sm_session = sagemaker.Session(boto_session=boto3.session.Session(region_name="us-west-2"))
settings = dict(
    sagemaker_session=sm_session,
    role=<The IAM role name>,
    instance_type="ml.m5.xlarge",
    dependencies="auto_capture"
)

@remote(**settings)
def divide(x, y):
    return x / y

if __name__ == "__main__":
    print(divide(2, 3.0))
```

⚠️ Note

You can also implement the previous code inside a Jupyter notebook. PyCharm Professional Edition supports Jupyter natively. For more guidance, see Jupyter notebook support in PyCharm's documentation.

**Invoking a function**

To invoke a function inside the @remote decorator, use either of the following methods:
**Use an @remote decorator to invoke a function.**

**Use the RemoteExecutor API to invoke a function.**

If you use the @remote decorator method to invoke a function, the training job will wait for the function to complete before starting a new task. However, if you use the RemoteExecutor API, you can run more than one job in parallel. The following sections show both ways of invoking a function.

**Use an @remote decorator to invoke a function**

You can use the @remote decorator to annotate a function. SageMaker will transform the code inside the decorator into a SageMaker training job. The training job will then invoke the function inside the decorator and wait for the job to complete. The following code example shows how to import the required libraries, start a SageMaker instance, and annotate a matrix multiplication with the @remote decorator.

```python
from sagemaker.remote_function import remote
import numpy as np

@remote(instance_type="ml.m5.large")
def matrix_multiply(a, b):
    return np.matmul(a, b)

a = np.array([[1, 0],
              [0, 1]])
b = np.array([1, 2])

assert (matrix_multiply(a, b) == np.array([1,2])).all()
```

The decorator is defined as follows.

```python
def remote(*,
        **kwarg):
    ...
```

When you invoke a decorated function, SageMaker Python SDK loads any exceptions raised by an error into local memory. In the following code example, the first call to the divide function completes successfully and the result is loaded into local memory. In the second call to the divide function, the code returns an error and this error is loaded into local memory.
from sagemaker.remote_function import remote
import pytest

@remote()
def divide(a, b):
    return a/b

# the underlying job is completed successfully
# and the function return is loaded
assert divide(10, 5) == 2

# the underlying job fails with "AlgorithmError"
# and the function exception is loaded into local memory
with pytest.raises(ZeroDivisionError):
    divide(10, 0)

⚠️ Note
The decorated function is run as a remote job. If the thread is interrupted, the underlying job will not be stopped.

How to change the value of a local variable

The decorator function is run on a remote machine. Changing a non-local variable or input arguments inside a decorated function will not change the local value.

In the following code example, a list and a dict are appended inside the decorator function. This does not change when the decorator function is invoked.

```python
a = []

@remote
def func():
    a.append(1)

# when func is invoked, a in the local memory is not modified
func()
func()
func()

# a stays as []
```
To change the value of a local variable declared inside of a decorator function, return the variable from the function. The following code example shows that the value of a local variable is changed when it is returned from the function.

```python
a = {"key-1": "value-1"}

@remote
def func(a):
    a["key-2"] = "value-2"
    return a

a = func(a)

-> {"key-1": "value-1", "key-2": "value-2"}
```

**Data serialization and deserialization**

When you invoke a remote function, SageMaker automatically serializes your function arguments during the input and output stages. Function arguments and returns are serialized using `cloudpickle`. SageMaker supports serializing the following Python objects and functions.

- Built-in Python objects including dicts, lists, floats, ints, strings, boolean values and tuples
- Numpy arrays
- Pandas Dataframes
- Scikit-learn datasets and estimators
- PyTorch models
- TensorFlow models
- The Booster class for XGBoost
The following can be used with some limitations.

- Dask DataFrames
- The XGBoost Dmatrix class
- TensorFlow datasets and subclasses
- PyTorch models

The following section contains best practices for using the previous Python classes with some limitations in your remote function, information about where SageMaker stores your serialized data and how to manage access to it.

**Best practices for Python classes with limited support for remote data serialization**

You can use the Python classes listed in this section with limitations. The next sections discuss best practices for how to use the following Python classes.

- Dask DataFrames
- The XGBoost DMatrix class
- TensorFlow datasets and subclasses
- PyTorch models

**Best practices for Dask**

Dask is an open-source library used for parallel computing in Python. This section shows the following.

- How to pass a Dask DataFrame into your remote function
- How to convert summary statistics from a Dask DataFrame into a Pandas DataFrame

**How to pass a Dask DataFrame into your remote function**

Dask DataFrames are often used to process large datasets because they can hold datasets that require more memory than is available. This is because a Dask DataFrame does not load your local data into memory. If you pass a Dask DataFrame as a function argument to your remote function, Dask may pass a reference to the data in your local disk or cloud storage, instead of the data itself. The following code shows an example of passing a Dask DataFrame inside your remote function that will operate on an empty DataFrame.
# Do not pass a Dask DataFrame to your remote function as follows

def clean(df: dask.DataFrame):
    cleaned = df[...]

Dask will load the data from the Dask DataFrame into memory only when you use the DataFrame. If you want to use a Dask DataFrame inside a remote function, provide the path to the data. Then Dask will read the dataset directly from the data path that you specify when the code runs.

The following code example shows how to use a Dask DataFrame inside the remote function clean. In the code example, raw_data_path is passed to clean instead of the Dask DataFrame. When the code runs, the dataset is read directly from the location of an Amazon S3 bucket specified in raw_data_path. Then the persist function keeps the dataset in memory to facilitate the subsequent random_split function and written back to the output data path in an S3 bucket using Dask DataFrame API functions.

```python
import dask.dataframe as dd

@remote(
    instance_type='ml.m5.24xlarge',
    volume_size=300,
    keep_alive_period_in_seconds=600)
#pass the data path to your remote function rather than the Dask DataFrame itself
def clean(raw_data_path: str, output_data_path: str, split_ratio: list[float]):
    df = dd.read_parquet(raw_data_path) #pass the path to your DataFrame
    cleaned = df[\(\text{df.column\_a} \geq 1\) \& \(\text{df.column\_a} < 5\)]\n    .drop(['\text{column\_b}', '\text{column\_c}'], axis=1)\n    .persist() #keep the data in memory to facilitate the following random_split operation

    train_df, test_df = cleaned.random_split(split_ratio, random_state=10)

    train_df.to_parquet(os.path.join(output_data_path, 'train'))
    test_df.to_parquet(os.path.join(output_data_path, 'test'))

    clean("s3://my-bucket/raw/", "s3://my-bucket/cleaned/", split_ratio=[0.7, 0.3])
```

**How to convert summary statistics from a Dask DataFrame into a Pandas DataFrame**

Summary statistics from a Dask DataFrame can be converted into a Pandas DataFrame by invoking the compute method as shown in the following example code. In the example, the S3 bucket
contains a large Dask DataFrame that cannot fit into memory or into a Pandas dataframe. In the following example, a remote function scans the data set and returns a Dask DataFrame containing the output statistics from `describe` to a Pandas DataFrame.

```python
executor = RemoteExecutor(
    instance_type='ml.m5.24xlarge',
    volume_size=300,
    keep_alive_period_in_seconds=600)

future = executor.submit(lambda: dd.read_parquet("s3://my-bucket/raw/").describe().compute())

future.result()
```

**Best practices for the XGBoost DMatrix class**

DMatrix is an internal data structure used by XGBoost to load data. A DMatrix object can't be pickled in order to move easily between compute sessions. Directly passing DMatrix instances will fail with a `SerializationError`.

**How to pass a data object to your remote function and train with XGBoost**

To convert a Pandas DataFrame into a DMatrix instance and use it for training in your remote function, pass it directly to the remote function as shown in the following code example.

```python
import xgboost as xgb

@remote
def train(df, params):
    # Convert a pandas dataframe into a DMatrix DataFrame and use it for training
    dtrain = DMatrix(df)
    return xgb.train(dtrain, params)
```

**Best practices for TensorFlow datasets and sub-classes**

TensorFlow datasets and subclasses are internal objects used by TensorFlow to load data during training. TensorFlow datasets and subclasses can't be pickled in order to move easily between compute sessions. Directly passing Tensorflow datasets or subclasses will fail with a `SerializationError`. Use the Tensorflow I/O APIs to load data from the storage, as shown in the following code example.
import tensorflow as tf
import tensorflow_io as tfio

@remote
def train(data_path: str, params):
    dataset = tf.data.TextLineDataset(tf.data.Dataset.list_files(f"{data_path}/*.txt"))
    ...
    train("s3://my-bucket/data", {"})

**Best practices for PyTorch models**

PyTorch models are serializable and can be passed between your local environment and remote function. If your local environment and remote environment have different device types, such as (GPUs and CPUs), you cannot return a trained model to your local environment. For example, if the following code is developed in a local environment without GPUs but run in an instance with GPUs, returning the trained model directly will lead to a `DeserializationError`.

```python
# Do not return a model trained on GPUs to a CPU-only environment as follows

@remote(instance_type='ml.g4dn.xlarge')
def train(...):
    if torch.cuda.is_available():
        device = torch.device("cuda")
    else:
        device = torch.device("cpu") # a device without GPU capabilities

    model = Net().to(device)

    # train the model
    ...

    return model

model = train(...) #returns a DeserializationError if run on a device with GPU
```

To return a model trained in a GPU environment to one that contains only CPU capabilities, use the PyTorch model I/O APIs directly as shown in the code example below.

```python
import s3fs
```
Where SageMaker stores your serialized data

When you invoke a remote function, SageMaker automatically serializes your function arguments and return values during the input and output stages. This serialized data is stored under a root directory in your S3 bucket. You specify the root directory, `<s3_root_uri>`, in a configuration file. The parameter `job_name` is automatically generated for you.

Under the root directory, SageMaker creates a `<job_name>` folder, which holds your current work directory, serialized function, the arguments for your serialized function, results and any exceptions that arose from invoking the serialized function.

Under `<job_name>`, the directory `workdir` contains a zipped archive of your current working directory. The zipped archive includes any Python files in your working directory and the `requirements.txt` file, which specifies any dependencies needed to run your remote function.
The following is an example of the folder structure under an S3 bucket that you specify in your configuration file.

```
<s3_root_uri>/  # specified by s3_root_uri or S3RootUri
  <job_name>/  # automatically generated for you
    workdir/workspace.zip  # archive of the current working directory (workdir)
    function/  # serialized function
    arguments/  # serialized function arguments
    results/  # returned output from the serialized function including the model
    exception/  # any exceptions from invoking the serialized function
```

The root directory that you specify in your S3 bucket is not meant for long term storage. The serialized data are tightly tied to the Python version and machine learning (ML) framework version that were used during serialization. If you upgrade the Python version or ML framework, you may not be able to use your serialized data. Instead, do the following.

- Store your model and model artifacts in a format that is agnostic to your Python version and ML framework.
- If you upgrade your Python or ML framework, access your model results from your long-term storage.

⚠️ **Important**

To delete your serialized data after a specified amount of time, set a [lifetime configuration](#) on your S3 bucket.

💡 **Note**

Files that are serialized with the Python [pickle](#) module can be less portable than other data formats including CSV, Parquet and JSON. Be wary of loading pickled files from unknown sources.

For more information about what to include in a configuration file for a remote function, see [Configuration File](#).
Access to your serialized data

Administrators can provide settings for your serialized data, including its location and any encryption settings in a configuration file. By default, the serialized data are encrypted with an AWS Key Management Service (AWS KMS) Key. Administrators can also restrict access to the root directory that you specify in your configuration file with a bucket policy. The configuration file can be shared and used across projects and jobs. For more information, see Configuration File.

Use the RemoteExecutor API to invoke a function

You can use the RemoteExecutor API to invoke a function. SageMaker Python SDK will transform the code inside the RemoteExecutor call into a SageMaker training job. The training job will then invoke the function as an asynchronous operation and return a future. If you use the RemoteExecutor API, you can run more than one training job in parallel. For more information about futures in Python, see Futures.

The following code example shows how to import the required libraries, define a function, start a SageMaker instance, and use the API to submit a request to run 2 jobs in parallel.

```python
from sagemaker.remote_function import RemoteExecutor
def matrix_multiply(a, b):
    return np.matmul(a, b)

a = np.array([[1, 0],
              [0, 1]])
b = np.array([1, 2])

with RemoteExecutor(max_parallel_job=2, instance_type="ml.m5.large") as e:
    future = e.submit(matrix_multiply, a, b)

assert (future.result() == np.array([1, 2])).all()
```

The RemoteExecutor class is an implementation of the concurrent.futures.Executor library.

The following code example shows how to define a function and call it using the RemoteExecutor API. In this example, the RemoteExecutor will submit 4 jobs in total, but only 2 in parallel. The last two jobs will reuse the clusters with minimal overhead.

```python
from sagemaker.remote_function.client import RemoteExecutor
```
def divide(a, b):
    return a/b

with RemoteExecutor(max_parallel_job=2, keep_alive_period_in_seconds=60) as e:
    futures = [e.submit(divide, a, 2) for a in [3, 5, 7, 9]]

for future in futures:
    print(future.result())

The `max_parallel_job` parameter only serves as a rate limiting mechanism without optimizing compute resource allocation. In the previous code example, `RemoteExecutor` doesn’t reserve compute resources for the two parallel jobs before any jobs are submitted. For more information about `max_parallel_job` or other parameters for the @remote decorator, see Remote function classes and methods specification.

**Future class for the RemoteExecutor API**

A future class is a public class that represents the return function from the training job when it is invoked asynchronously. The future class implements the `concurrent.futures.Future` class. This class can be used to do operations on the underlying job and load data into memory.

**Configuration file**

The Amazon SageMaker Python SDK supports setting of default values for AWS infrastructure primitive types. After administrators configure these defaults, they are automatically passed when SageMaker Python SDK calls supported APIs. The arguments for the decorator function can be put inside of configuration files. This is so that you can separate settings that are related to the infrastructure from the code base. For more information about parameters and arguments for the remote function and methods, see Remote function classes and methods specification.

You can set infrastructure settings for the network configuration, IAM roles, Amazon S3 folder for input, output data, and tags inside the configuration file. The configuration file can be used when invoking a function using either the @remote decorator or the RemoteExecutor API.

An example configuration file that defines the dependencies, resources, and other arguments follows. This example configuration file is used to invoke a function that is initiated either using the @remote decorator or the RemoteExecutor API.
PythonSDK:

Modules:

RemoteFunction:

  Dependencies: 'path/to/requirements.txt'
  EnableInterContainerTrafficEncryption: true
  EnvironmentVariables: {'EnvVarKey': 'EnvVarValue'}
  ImageUri: '366666666666.dkr.ecr.us-west-2.amazonaws.com/my-image:latest'
  IncludeLocalWorkDir: true
  CustomFileFilter:
    IgnoreNamePatterns:
    - '*.ipynb'
    - 'data'
  InstanceType: 'ml.m5.large'
  JobCondaEnvironment: 'your_conda_env'

PreExecutionCommands:

  - 'command_1'
  - 'command_2'

PreExecutionScript: 'path/to/script.sh'

RoleArn: 'arn:aws:iam::366666666666:role/MyRole'

S3KmsKeyId: 'yourkmskeyid'

S3RootUri: 's3://my-bucket/my-project'

VpcConfig:

  SecurityGroupIds:
  - 'sg123'
  Subnets:
  - 'subnet-1234'

Tags: [{'Key': 'yourTagKey', 'Value': 'yourTagValue'}]

VolumeKmsKeyId: 'yourkmskeyid'

The @remote decorator and RemoteExecutor will look for Dependencies in the following configuration files:

- An admin-defined configuration file.
- A user-defined configuration file.

The default locations for these configuration files depend on, and are relative to, your environment. The following code example returns the default location of your admin and user configuration files. These commands must be run in the same environment where you're using the SageMaker Python SDK.

```python
import os
```

```python
# Configuration file
```
from platformdirs import site_config_dir, user_config_dir

# Prints the location of the admin config file
print(os.path.join(site_config_dir("sagemaker"), "config.yaml"))

# Prints the location of the user config file
print(os.path.join(user_config_dir("sagemaker"), "config.yaml"))

You can override the default locations of these files by setting the SAGEMAKER_ADMIN_CONFIG_OVERRIDE and SAGEMAKER_USER_CONFIG_OVERRIDE environment variables for the admin-defined and user-defined configuration file paths, respectively.

If a key exists in both the admin-defined and user-defined configuration files, the value in the user-defined file will be used.

**Customize your runtime environment**

You can customize your runtime environment to use your preferred local integrated development environments (IDEs), SageMaker notebooks, or SageMaker Studio Classic notebooks to write your ML code. SageMaker will help package and submit your functions and its dependencies as a SageMaker training job. This allows you to access the capacity of the SageMaker training server to run your training jobs.

Both the remote decorator and the RemoteExecutor methods to invoke a function allow users to define and customize their runtime environment. You can use either a requirements.txt file or a conda environment YAML file.

To customize a runtime environment using both a conda environment YAML file and a requirements.txt file, refer to the following code example.

```python
# specify a conda environment inside a yaml file
@remote(instance_type="ml.m5.large",
        image_uri="my_base_python:latest",
        dependencies="./environment.yml")
def matrix_multiply(a, b):
    return np.matmul(a, b)

# use a requirements.txt file to import dependencies
@remote(instance_type="ml.m5.large",
        image_uri="my_base_python:latest",
        dependencies='./requirements.txt')
def matrix_multiply(a, b):
```
```
return np.matmul(a, b)
```

Alternatively, you can set dependencies to `auto_capture` to let the SageMaker Python SDK capture the installed dependencies in the active conda environment. The following are required for `auto_capture` to work reliably:

- You must have an active conda environment. We recommend not using the base conda environment for remote jobs so that you can reduce potential dependency conflicts. Not using the base conda environment also allows for faster environment setup in the remote job.
- You must not have any dependencies installed using pip with a value for the parameter `--extra-index-url`.
- You must not have any dependency conflicts between packages installed with conda and packages installed with pip in the local development environment.
- Your local development environment must not contain operating system-specific dependencies that are not compatible with Linux.

In case `auto_capture` does not work, we recommend that you pass in your dependencies as a `requirement.txt` or `conda environment.yaml` file, as described in the first coding example in this section.

**Container image compatibility**

The following table shows a list of SageMaker training images that are compatible with the `@remote` decorator.

<table>
<thead>
<tr>
<th>Name</th>
<th>Python Version</th>
<th>Image URI - CPU</th>
<th>Image URI - GPU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data Science</td>
<td>3.7(py37)</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
</tr>
<tr>
<td>Name</td>
<td>Python Version</td>
<td>Image URI - CPU</td>
<td>Image URI - GPU</td>
</tr>
<tr>
<td>-----------------------</td>
<td>----------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Data Science 2.0</td>
<td>3.8(py38)</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
</tr>
<tr>
<td>Data Science 3.0</td>
<td>3.10(py310)</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
</tr>
<tr>
<td>Base Python 2.0</td>
<td>3.8(py38)</td>
<td>Python SDK selects this image when it detects that development environment is using Python 3.8 runtime. Otherwise Python SDK automatically selects this image when used as SageMaker Studio Classic Notebook kernel image.</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as SageMaker Studio Classic Notebook kernel image.</td>
</tr>
<tr>
<td>Name</td>
<td>Python Version</td>
<td>Image URI - CPU</td>
<td>Image URI - GPU</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------</td>
<td>-------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Base Python 3.0</td>
<td>3.10(py310)</td>
<td>Python SDK selects this image when it detects that development environment is using Python 3.8 runtime. Otherwise Python SDK automatically selects this image when used as SageMaker Studio Classic Notebook kernel image.</td>
<td>For SageMaker Studio Classic Notebooks only. Python SDK automatically selects the image URI when used as Studio Classic Notebook kernel image.</td>
</tr>
<tr>
<td>DLC-Tensorflow 2.11.0 for SageMaker training</td>
<td>3.9(py39)</td>
<td>763104351884.dkr.ecr.&lt;region&gt;.amazonaws.com/tensorflow-training:2.11.0-cpu-py39-ubuntu20.04-sagemaker</td>
<td>763104351884.dkr.ecr.&lt;region&gt;.amazonaws.com/tensorflow-training:2.11.0-gpu-py39-cu112-ubuntu20.04-sagemaker</td>
</tr>
<tr>
<td>Name</td>
<td>Python Version</td>
<td>Image URI - CPU</td>
<td>Image URI - GPU</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>----------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>DLC-PyTorch 2.0.0 for SageMaker training</td>
<td>3.10(py310)</td>
<td>763104351884.dkr&lt;region&gt;.amazon aws.com/pytorch-training:2.0.0-cpu-py310-ubuntu20.04-sagemaker</td>
<td>763104351884.dkr&lt;region&gt;.amazon aws.com/pytorch-training:2.0.0-gpu-py310-cu118-ubuntu20.04-sagemaker</td>
</tr>
<tr>
<td>Name</td>
<td>Python Version</td>
<td>Image URI - CPU</td>
<td>Image URI - GPU</td>
</tr>
<tr>
<td>------</td>
<td>----------------</td>
<td>-----------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>DLC-PyTorch 1.11.0 for SageMaker training</td>
<td>3.8(py38)</td>
<td>763104351884.dkr.ecr.&lt;region&gt;.amazonaws.com/pytorch-training:1.11.0-cpu-py38-ubuntu20.04-sagemaker</td>
<td>763104351884.dkr.ecr.&lt;region&gt;.amazonaws.com/pytorch-training:1.11.0-gpu-py38-cu113-ubuntu20.04-sagemaker</td>
</tr>
</tbody>
</table>

**Note**

To run jobs locally using AWS Deep Learning Containers (DLC) images, use the image URIs found in the [DLC documentation](#). The DLC images do not support the `auto_capture` value for dependencies.

Jobs with [SageMaker Distribution in SageMaker Studio](#) run in a container as a non-root user named `sagemaker-user`. This user needs full permission to access `/opt/ml` and `/tmp`. Grant this permission by adding `sudo chmod -R 777 /opt/ml /tmp` to the `pre_execution_commands` list, as shown in the following snippet:

```python
@remote(pre_execution_commands=["sudo chmod -R 777 /opt/ml /tmp"])
def func():
    pass
```

You can also run remote functions with your custom images. For compatibility with remote functions, custom images should be built with Python version 3.7.x-3.10.x. The following is a minimal Dockerfile example showing you how to use a Docker image with Python 3.10.

```text
FROM python:3.10
```
To create conda environments in your image and use it to run jobs, set the environment variable SAGEMAKER_JOB_CONDA_ENV to the conda environment name. If your image has the SAGEMAKER_JOB_CONDA_ENV value set, the remote function cannot create a new conda environment during the training job runtime. Refer to the following Dockerfile example that uses a conda environment with Python version 3.10.

```bash
FROM continuumio/miniconda3:4.12.0

ENV SHELL=/bin/bash \
    CONDA_DIR=/opt/conda \
    SAGEMAKER_JOB_CONDA_ENV=sagemaker-job-env

RUN conda create -n $SAGEMAKER_JOB_CONDA_ENV \
    && conda install -n $SAGEMAKER_JOB_CONDA_ENV python=3.10 -y \
    && conda clean --all -f -y 
```

For SageMaker to use mamba to manage your Python virtual environment in the container image, install the mamba toolkit from miniforge. To use mamba, add the following code example to your Dockerfile. Then, SageMaker will detect the mamba availability at runtime and use it instead of conda.

```bash
#Mamba Installation
RUN curl -L -O "https://github.com/conda-forge/miniforge/releases/latest/download/Mambaforge-Linux-x86_64.sh" \
    && bash Mambaforge-Linux-x86_64.sh -b -p "/opt/conda" \
    && /opt/conda/bin/conda init bash
```

Using a custom conda channel on an Amazon S3 bucket is not compatible with mamba when using a remote function. If you choose to use mamba, make sure you are not using a custom conda channel on Amazon S3. For more information, see the Prerequisites section under Custom conda repository using Amazon S3.

The following is a complete Dockerfile example showing how to create a compatible Docker image.

```bash
FROM python:3.10

RUN apt-get update -y \
    # Needed for awscli to work
```
The resulting image from running the previous Dockerfile example can also be used as a SageMaker Studio Classic kernel image.

Logging parameters and metrics with Amazon SageMaker Experiments

This guide show how to log parameters and metrics with Amazon SageMaker Experiments. A SageMaker experiment consists of runs, and each run consists of all the inputs, parameters, configurations and results for a single model training interaction.

You can log parameters and metrics from a remote function using either the @remote decorator or the RemoteExecutor API.

To log parameters and metrics from a remote function, choose one of the following methods:

- Instantiate a SageMaker experiment run inside a remote function using Run from the SageMaker Experiments library. For more information, see Create an Amazon SageMaker Experiment.
Use the `load_run` function inside a remote function from the SageMaker Experiments library. This will load a Run instance that is declared outside of the remote function.

The following sections show how to create and track lineage with SageMaker experiment runs by using the previous listed methods. The sections also describe cases that are not supported by SageMaker training.

**Use the @remote decorator to integrate with SageMaker Experiments**

You can either instantiate an experiment in SageMaker, or load a current SageMaker experiment from inside a remote function. The following sections show you how to use either method.

**Create an experiment with SageMaker Experiments**

You can create an experiment run in SageMaker experiment. To do this you pass your experiment name, run name, and other parameters into your remote function.

The following code example imports the name of your experiment, the name of the run, and the parameters to log during each run. The parameters `param_1` and `param_2` are logged over time inside a training loop. Common parameters may include batch size or epochs. In this example, the metrics `metric_a` and `metric_b` are logged for a run over time inside a training loop. Other common metrics may include accuracy or loss.

```python
from sagemaker.remote_function import remote
from sagemaker.experiments.run import Run

# Define your remote function
@remote
def train(value_1, value_2, exp_name, run_name):
    ...
    ...
    # Creates the experiment
    with Run(
        experiment_name=exp_name,
        run_name=run_name,
    ) as run:
        ...
        # Define values for the parameters to log
        run.log_parameter("param_1", value_1)
        run.log_parameter("param_2", value_2)
        ...
```

Logging parameters and metrics with Amazon SageMaker Experiments
# Define metrics to log
run.log_metric("metric_a", 0.5)
run.log_metric("metric_b", 0.1)

# Invoke your remote function
train(1.0, 2.0, "my-exp-name", "my-run-name")

## Load current SageMaker Experiments with a job initiated by the @remote decorator

Use the `load_run()` function from the SageMaker Experiments library to load the current run object from the run context. You can also use the `load_run()` function within your remote function. Load the run object initialized locally by the `with` statement on the run object as shown in the following code example.

```python
from sagemaker.experiments.run import Run, load_run

# Define your remote function
@remote
def train(value_1, value_2):
    ...
    ...
    with load_run() as run:
        run.log_metric("metric_a", value_1)
        run.log_metric("metric_b", value_2)

# Invoke your remote function
with Run(
    experiment_name="my-exp-name",
    run_name="my-run-name",
) as run:
    train(0.5, 1.0)
```

## Load a current experiment run within a job initiated with the RemoteExecutor API

You can also load a current SageMaker experiment run if your jobs were initiated with the RemoteExecutor API. The following code example shows how to use RemoteExecutor API with the SageMaker Experiments `load_run` function. You do this to load a current SageMaker experiment run and capture metrics in the job submitted by RemoteExecutor.
from sagemaker.experiments.run import Run, load_run

def square(x):
    with load_run() as run:
        result = x * x
        run.log_metric("result", result)
    return result

with RemoteExecutor(
    max_parallel_job=2,
    instance_type="ml.m5.large"
) as e:
    with Run(
        experiment_name="my-exp-name",
        run_name="my-run-name",
    ):
        future_1 = e.submit(square, 2)

Unsupported uses for SageMaker Experiments while annotating your code with an @remote decorator

SageMaker does not support passing a Run type object to an @remote function or using global Run objects. The following examples show code that will throw a SerializationError.

The following code example attempts to pass a Run type object to an @remote decorator, and it generates an error.

@remote
def func(run: Run):
    run.log_metrics("metric_a", 1.0)

with Run(...) as run:
    func(run) ---> SerializationError caused by NotImplementedError

The following code example attempts to use a global run object instantiated outside of the remote function. In the code example, the train() function is defined inside the with Run context, referencing a global run object from within. When train() is called, it generates an error.

with Run(...) as run:
    @remote
def train(metric_1, value_1, metric_2, value_2):
    run.log_parameter(metric_1, value_1)
    run.log_parameter(metric_2, value_2)

train("p1", 1.0, "p2", 0.5) ---> SerializationError caused by NotImplementedError

Using modular code with the @remote decorator

You can organize your code into modules for ease of workspace management during development and still use the @remote function to invoke a function. You can also replicate the local modules from your development environment to the remote job environment. To do so, set the parameter include_local_workdir to True, as shown in the following code example.

@remote(
    include_local_workdir=True,
)

Note

The @remote decorator and parameter must appear in the main file, rather than in any of the dependent files.

When include_local_workdir is set to True, SageMaker packages all of the Python scripts while maintaining the directory structure in the process' current directory. It also makes the dependencies available in the job's working directory.

For example, suppose your Python script which processes the MNIST dataset is divided into a main.py script and a dependent pytorch_mnist.py script. main.py calls the dependent script. Also, the main.py script contains code to import the dependency as shown.

from mnist_impl.pytorch_mnist import ...

The main.py file must also contain the @remote decorator, and it must set the include_local_workdir parameter to True.

The include_local_workdir parameter by default includes all the Python scripts in the directory. You can customize which files you want to upload to the job by using this parameter in
Amazon SageMaker

Developer Guide

conjunction with the custom_file_filter parameter. You can either pass a function that ﬁlters
job dependencies to be uploaded to S3, or a CustomFileFilter object that speciﬁes the local
directories and ﬁles to ignore in the remote function. You can use custom_file_filter only if
include_local_workdir is set to True—otherwise the parameter is ignored.
The following example uses CustomFileFilter to ignore all notebook ﬁles and folders or ﬁles
named data when uploading ﬁles to S3.
@remote(
include_local_workdir=True,
custom_file_filter=CustomFileFilter(
ignore_pattern_names=[ # files or directories to ignore
"*.ipynb", # all notebook files
"data", # folter or file named data
]
)
)

The following example demonstrates how you can package an entire workspace.
@remote(
include_local_workdir=True,
custom_file_filter=CustomFileFilter(
ignore_pattern_names=[] # package whole workspace
)
)

The following example shows how you can use a function to ﬁlter ﬁles.
import os
def my_filter(path: str, files: List[str]) -> List[str]:
to_ignore = []
for file in files:
if file.endswith(".txt") or file.endswith(".ipynb"):
to_ignore.append(file)
return to_ignore
@remote(
include_local_workdir=True,
custom_file_filter=my_filter
Using modular code with the @remote decorator

3118


Best practices in structuring your working directory

The following best practices suggest how you can organize your directory structure while using the @remote decorator in your modular code.

- Put the @remote decorator in a file that resides at the root level directory of the workspace.
- Structure the local modules at the root level.

The following example image shows the recommended directory structure. In this example structure, the main.py script is located at the root level directory.

```
### config.yaml
### data/
### main.py <----------------- @remote used here
### mnist_impl
# ### __pycache__/
# # ### pytorch_mnist.cpython-310.pyc
# ### pytorch_mnist.py <-------- dependency of main.py
### requirements.txt
```

The following example image shows a directory structure that will result in inconsistent behavior when it is used to annotate your code with an @remote decorator.

In this example structure, the main.py script that contains the @remote decorator is not located at the root level directory. The following structure is NOT recommended.

```
### config.yaml
### entrypoint
# ### data
# ### main.py <----------------- @remote used here
### mnist_impl
# ### __pycache__/
# # ### pytorch_mnist.cpython-310.pyc
# ### pytorch_mnist.py <-------- dependency of main.py
### requirements.txt
```
Private repository for runtime dependencies

You can use pre-execution commands or script to configure a dependency manager like pip or conda in your job environment. To achieve network isolation, use either of these options to redirect your dependency managers to access your private repositories and run remote functions within a VPC. The pre-execution commands or script will run before your remote function runs. You can define them with the `@remote` decorator, the `RemoteExecutor` API, or within a configuration file.

The following sections show you how to access a private Python Package Index (PyPI) repository managed with AWS CodeArtifact. The sections also show how to access a custom conda channel hosted on Amazon Simple Storage Service (Amazon S3).

**How to use a custom PyPI repository managed with AWS CodeArtifact**

To use CodeArtifact to manage a custom PyPI repository, the following prerequisites are required:

- Your private PyPI repository should already have been created. You can utilize AWS CodeArtifact to create and manage your private package repositories. To learn more about CodeArtifact, see the [CodeArtifact User Guide](https://docs.aws.amazon.com/codeartifact/latest/userguide/).

- Your VPC should have access to your CodeArtifact repository. To allow a connection from your VPC to your CodeArtifact repository, you must do the following:
  - Create VPC endpoints for CodeArtifact.
  - Create an Amazon S3 gateway endpoint for your VPC, which allows CodeArtifact to store package assets.

The following pre-execution command example shows how to configure pip in the SageMaker training job to point to your CodeArtifact repository. For more information, see [Configure and use pip with CodeArtifact](https://docs.aws.amazon.com/codeartifact/latest/userguide/).

```bash
# use a requirements.txt file to import dependencies
@remote(
    instance_type="ml.m5.large",
    image_uri = "my_base_python:latest",
    dependencies = './requirements.txt',
    pre_execution_commands=[
        "aws codeartifact login --tool pip --domain my-org --domain-owner <000000000000> --repository my-codeartifact-python-repo --endpoint-url https://vpce-xxxxx.api.codeartifact.us-east-1.vpce.amazonaws.com"
    ]
```
How to use a custom conda channel hosted on Amazon S3

To use Amazon S3 to manage a custom conda repository, the following prerequisites are required:

- Your private conda channel must already be set up in your Amazon S3 bucket, and all dependent packages must be indexed and uploaded to your Amazon S3 bucket. For instructions on how to index your conda packages, see Creating custom channels.
- Your VPC should have access to the Amazon S3 bucket. For more information, see Endpoints for Amazon S3.
- The base conda environment in your job image should have boto3 installed. To check your environment, enter the following in your Anaconda prompt to check that boto3 appears in the resulting generated list.

```
conda list -n base
```

- You job image should be installed with conda, not mamba. To check your environment, ensure that the previous code prompt does not return mamba.

The following pre-execution commands example shows how to configure conda in the SageMaker training job to point to your private channel on Amazon S3. The pre-execution commands removes the defaults channel and adds custom channels to a .condarc conda configuration file.

```python
# specify your dependencies inside a conda yaml file
@remote(
    instance_type="ml.m5.large"
    image_uri = "my_base_python:latest",
    dependencies = "./environment.yml",
    pre_execution_commands=[
        "conda config --remove channels 'defaults'"
        "conda config --add channels 's3://my_bucket/my-conda-repository/conda-forge/'",
        "conda config --add channels 's3://my_bucket/my-conda-repository/main/'"
    ]
)

def matrix_multiply(a, b):
    return np.matmul(a, b)
```
Example notebooks

You can transform a training code in an existing workspace environment and any associated data processing code and datasets into a SageMaker training job. The following notebooks show you how to customize your environment, job settings, and more for an image classification problem, using the XGBoost algorithm and Hugging Face.

The quick_start notebook contains the following code examples:

- How to customize your job settings with a configuration file.
- How to invoke Python functions as jobs, asynchronously.
- How to customize the job runtime environment by bringing in additional dependencies.
- How to use local dependencies with the @remote function method.

The following notebooks provide additional code examples for different ML problems types and implementations.

- To see code examples to use the @remote decorator for an image classification problem, open the pytorch_mnist.ipynb notebook. This classification problem recognizes handwritten digits using the Modified National Institute of Standards and Technology (MNIST) sample dataset.
- To see code examples for using the @remote decorator for the previous image classification problem with a script, see the Pytorch MNIST sample script, train.py.
- To see how the XGBoost algorithm implemented with an @remote decorator: Open the xgboost_abalone.ipynb notebook.
- To see how Hugging Face is integrated with an @remote decorator: Open the huggingface.ipynb notebook.

Manage Machine Learning with Amazon SageMaker Experiments

Amazon SageMaker Experiments is a capability of Amazon SageMaker that lets you create, manage, analyze, and compare your machine learning experiments.

Experimentation in machine learning
Machine learning is an iterative process. You need to experiment with multiple combinations of data, algorithms, and parameters, all while observing the impact of incremental changes on model accuracy. Over time, this iterative experimentation can result in thousands of model training runs and model versions. This makes it hard to track the best performing models and their input configurations. It's also difficult to compare active experiments with past experiments to identify opportunities for further incremental improvements. Use SageMaker Experiments to organize, view, analyze, and compare iterative ML experimentation to gain comparative insights and track your best performing models.

**Manage ML experimentation with SageMaker Experiments**

SageMaker Experiments automatically tracks the inputs, parameters, configurations, and results of your iterations as *runs*. You can assign, group, and organize these runs into *experiments*. SageMaker Experiments is integrated with Amazon SageMaker Studio Classic, providing a visual interface to browse your active and past experiments, compare runs on key performance metrics, and identify the best performing models. SageMaker Experiments tracks all of the steps and artifacts that went into creating a model, and you can quickly revisit the origins of a model when you are troubleshooting issues in production, or auditing your models for compliance verifications.

Use SageMaker Experiments to view, manage, analyze, and compare both custom experiments that you programmatically create and experiments automatically created from SageMaker jobs.

**Supported AWS Regions**

SageMaker Experiments is generally available in all AWS commercial Regions where Amazon SageMaker Studio Classic is available, except the China Regions.

**Topics**

- [Create an Amazon SageMaker Experiment](#)
- [View, search, and compare experiment runs](#)
- [SageMaker integrations](#)
- [Example notebooks for Amazon SageMaker Experiments](#)
- [Monitor experiment training metrics with AWS CloudTrail](#)
- [Clean Up Amazon SageMaker Experiment Resources](#)
- [Additional supported SDK](#)
- [Experiments FAQs](#)
- [Search Using the Amazon SageMaker Console and API](#)
Create an Amazon SageMaker Experiment

Create an Amazon SageMaker experiment to track your machine learning (ML) workflows with a few lines of code from your preferred development environment. You can then browse your experiments, create visualizations for analysis, and find the best performing model. You can also integrate SageMaker Experiments into your SageMaker training script using the SageMaker Python SDK.

Overview

The following components make up the building blocks of an experiment in Amazon SageMaker.

- **experiment**: An experiment is a collection of runs. When you initialize a run in your training loop, you include the name of the experiment that the run belongs to. Experiment names must be unique within your AWS account.

- **Run**: A run consists of all the inputs, parameters, configurations, and results for one interaction of model training. Initialize an experiment run for tracking a training job with `Run.init()`.

  **Note**
  
  We recommend that you initialize a Run object in a Jupyter Notebook, and create the SageMaker job for your experiment within the context of this Run object initialization. To refer to this Run object in script mode, use the `load_run()` operation. For examples, see [Example notebooks for Amazon SageMaker Experiments](#).

- **load_run**: To run your experiments in script mode, refer to an initialized Run object with `load_run()`. If an experiment for a run exists, `load_run` returns the experiment context. Generally, you use `load_run` with no arguments to track metrics, parameters, and artifacts within a SageMaker training or processing job script.

  ```python
  # Load run from a local script passing experiment and run names
  with load_run(experiment_name=experiment_name, run_name=run_name) as run:
  ```

```
• **log_parameter**: Log parameters for a run, such as batch size or epochs, over time in a training loop with `run.log_parameter()`. `log_parameter` records a single name-value pair in a run. You can use `run.log_parameters()` to log multiple parameters. If called multiple times within a run for a parameter of the same name, `log_parameter` overwrites any previous value. The name must be a string and the value must be either a string, integer, or float.

```python
# Log a single parameter
run.log_parameter("param1", "value1")

# Log multiple parameters
run.log_parameters(
    {
        "param2": "value2",
        "param3": "value3"
    }
)
```

• **log_metric**: Log metrics for a run, such as accuracy or loss, over time in a training loop with `run.log_metric()`. `log_metric` records a name-value pair where the name is a string and the value is an integer or float. To declare the frequency of logging over the course of the run, define a step value. You can then visualize these metrics in the Studio Classic Experiments UI. For more information, see [View, search, and compare experiment runs](#).

```python
# Log a metric over the course of a run
run.log_metric(name="Final_loss", value=finalloss)

# Log a metric over the course of a run at each epoch
run.log_metric(name="test:loss", value=loss, step=epoch)
```

• **log_artifact**: Log any input or output artifacts related to a run with `run.log_artifact()`. Log artifacts such as S3 URIs, datasets, models, and more for your experiment to help you keep track of artifacts across multiple runs. `is_output` is `True` by default. To record the artifact as an input artifact instead of an output artifact, set `is_output` to `False`.

```python
# Track a string value as an input or output artifact
```
run.log_artifact(name="training_data", value="data.csv" is_output=False)

- log_file: Log any input or output files related to a run, such as training or test data, and store them in Amazon S3 with run.log_file(). is_output is True by default. To record the file as an input artifact instead of an output artifact, set is_output to False.

# Upload a local file to S3 and track it as an input or output artifact
run.log_file("training_data.csv", name="training_data", is_output=False)

For more information on initializing a Run object, see Experiments in the SageMaker Python SDK documentation. For information on visualizing logged experiment data and automatic logging, see View, search, and compare experiment runs.

Create an experiment with the SageMaker Python SDK

The following section demonstrates how to create an Amazon SageMaker Experiment using the SageMaker Python SDK. This example uses the Run class to track a Keras model in a notebook environment. The Keras Callback class provides an operation on_epoch_end which emits metrics at the end of each epoch. First, define a Callback class.

```
class ExperimentCallback(keras.callbacks.Callback):
    """ ""
    def __init__(self, run, model, x_test, y_test):
        """Save params in constructor""
        self.run = run
        self.model = model
        self.x_test = x_test
        self.y_test = y_test

    def on_epoch_end(self, epoch, logs=None):
        """ ""
        keys = list(logs.keys())
        for key in keys:
            run.log_metric(name=key, value=logs[key], step=epoch)
            print("Epoch: {}\n{} -> {}").format(epoch, key, logs[key])
```

Next, train the Keras model in a notebook environment and track it as an experiment.
This example carries out jobs sequentially. To run SageMaker jobs asynchronously, you may need to increase your resource limit.

```python
from sagemaker.experiments import Run

# The run name is an optional argument to `run.init`.
with Run(experiment_name = 'my-experiment') as run:

    # Define values for the parameters to log
    run.log_parameter("batch_size", batch_size)
    run.log_parameter("epochs", epochs)
    run.log_parameter("dropout", 0.5)

    # Define input artifacts
    run.log_file('datasets/input_train.npy', is_output = False)
    run.log_file('datasets/input_test.npy', is_output = False)
    run.log_file('datasets/input_train_labels.npy', is_output = False)
    run.log_file('datasets/input_test_labels.npy', is_output = False)

    # Train locally
    model.fit(
        x_train,
        y_train,
        batch_size=batch_size,
        epochs=epochs,
        validation_split=0.1,
        callbacks = [ExperimentCallback(run, model, x_test, y_test)]
    )

    score = model.evaluate(x_test, y_test, verbose=0)
    print("Test loss:", score[0])
    print("Test accuracy:", score[1])

    # Define metrics to log
    run.log_metric(name = "Final Test Loss", value = score[0])
    run.log_metric(name = "Final Test Accuracy", value = score[1])
```

For more code examples and example notebooks, see Example notebooks for Amazon SageMaker Experiments.
Create an experiment using SageMaker script mode

You can use SageMaker script mode to write your own code to train a model and track it as an experiment. When creating an experiment with script mode, use `load_run()`.

```python
# Make sure that you have the latest version of the SageMaker Python SDK
import os
os.system("pip install -U sagemaker")

# Import additional requirements
import boto3
from sagemaker.session import Session
from sagemaker.experiments.run import load_run

# Define training script
if __name__ == "__main__":
    session = Session(boto3.session.Session(region_name=args.region))
    with load_run(sagemaker_session=session) as run:
        # Define values for the parameters to log
        run.log_parameters({
            "batch_size": batch_size,
            "epochs": epochs,
            "dropout": 0.5
        })
        # Define input artifacts
        run.log_file('datasets/input_train.npy', is_output = False)
        run.log_file('datasets/input_test.npy', is_output = False)
        run.log_file('datasets/input_train_labels.npy', is_output = False)
        run.log_file('datasets/input_test_labels.npy', is_output = False)

        # Train the model
        model.fit(
            x_train,
            y_train,
            batch_size=batch_size,
            epochs=epochs,
            validation_split=0.1,
            callbacks = [ExperimentCallback(run, model, x_test, y_test)]
        )

        score = model.evaluate(x_test, y_test, verbose=0)
        print("Test loss:", score[0])
        print("Test accuracy:", score[1])
```
# Define metrics to log
run.log_metric(name = "Final Test Loss", value = score[0])
run.log_metric(name = "Final Test Accuracy", value = score[1])

For more code examples and example notebooks on using Amazon SageMaker Experiments in SageMaker script mode, see [Track experiments for SageMaker training jobs using script mode](#).

For more information on script mode, see [Use script mode in a supported framework](#). You can also define custom metrics in script mode by specifying a name and regular expression for each metric that a tuning job monitors. See [Use a custom algorithm for training](#) for more information.

**View your experiment in Studio**

⚠️ **Important**
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).

To view your experiments, you must do so through Studio Classic, which you can access through the updated Studio experience.

Within Studio, choose **Experiments** on the lefthand navigation pane. Then, choose **View Studio Classic**. For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#). For more information on getting started with the updated Studio experience, see [Amazon SageMaker Studio](#).
View your experiment in Studio Classic

⚠️ Important
As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

To view the experiment in Studio Classic, in the left sidebar, choose Experiments.

Select the name of the experiment to view all associated runs. It might take a moment for the list to refresh and display a new experiment or experiment run. You can click Refresh to update the page. Your experiment list should look similar to the following:
To view the runs that make up your experiment, select the experiment name. For more information, see [View, search, and compare experiment runs](#).

**View unassigned runs**

All SageMaker jobs, including training jobs, processing jobs, and transform jobs, correspond to runs and create Run objects by default. If you launch these jobs without explicitly associating them with an experiment, the resulting runs are unassigned and can be viewed in the [Unassigned runs](#) section of the Studio Classic Experiments UI.
To clean up the resources you created, see Clean Up Amazon SageMaker Experiment Resources.

View, search, and compare experiment runs

An Amazon SageMaker experiment consists of multiple run groups with a related objective. A run group consists of one or more runs, such as a data preprocessing job and a training job.

To view your experiments, you must do so through Studio Classic. For an overview of the Studio Classic user interface, see Amazon SageMaker Studio Classic UI Overview. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
Amazon SageMaker Studio Classic provides an experiments browser that you can use to view lists of experiments and runs. You can choose one of these entities to view detailed information about the entity or choose multiple entities for comparison. You can filter the list of experiments by entity name, type, and tags.

To view experiments and runs

1. To view the experiment in Studio Classic, in the left sidebar, choose **Experiments**.

Select the name of the experiment to view all associated runs. You can search experiments by typing directly into the **Search** bar or filtering for experiment type. You can also choose which columns to display in your experiment or run list.

It might take a moment for the list to refresh and display a new experiment or experiment run. You can click **Refresh** to update the page. Your experiment list should look similar to the following:

2. In the experiments list, double-click an experiment to display a list of the runs in the experiment.
## Note

Experiment runs that are automatically created by SageMaker jobs and containers are visible in the Experiments Studio Classic UI by default. To hide runs created by SageMaker jobs for a given experiment, choose the settings icon and toggle **Show jobs**.

3. Double-click a run to display information about a specific run.

In the **Overview** pane, choose any of the following headings to see available information about each run:

- **Metrics** – Metrics that are logged during a run.
- **Charts** – Build your own charts to compare runs.
- **Output artifacts** – Any resulting artifacts of the experiment run and the artifact locations in Amazon S3.
- **Bias reports** – Pre-training or post-training bias reports generated using Clarify.
- **Explainability** – Explainability reports generated using Clarify.
- **Debugs** – A list of debugger rules and any issues found.

**Compare and analyze runs**

To analyze experiment runs, select the experiment of your choice in the Amazon SageMaker Studio Classic Experiments UI and then select the runs that you want to compare. You must select between 1 and 20 runs. After you have your runs selected, choose **Analyze** in the upper right-hand corner.

**To compare experiment runs:**

1. After navigating to the experiment of your choice, select all the runs that you want to compare. You must choose more than 1 and less than 20 runs to analyze.
2. Choose **Analyze** in the upper right-hand corner.
3. Visualize the comparative metrics of multiple experiment runs in a histogram, line chart, scatter plot, or bar chart. To add a chart, choose **Add Chart**, select values for your chart axes, and choose **Create**.

![Image of experiment runs comparison](image-url)
You can update, download, or delete existing charts.

Log charts

Logging charts and visualizations is available for classification models. You can log a confusion matrix, receiver operating characteristics, or precision and recall graphs.

Log and visualize metrics with the following Python SDK methods:

- `log_confusion_matrix`: Records a confusion matrix artifact that you can view in the Charts section of the Run Overview in Studio Classic.
- `log_roc_curve`: Records a receiver operating characteristic artifact that you can view in the Charts section of the Run Overview in Studio Classic.
- `log_precision_recall`: Records a precision recall graph that you can view in the Charts section of the Run Overview in Studio Classic.

An automatically logged precision recall record creates a chart similar to the following:
Amazon SageMaker integrations

Amazon SageMaker Experiments is integrated with a number of SageMaker features. Certain SageMaker jobs automatically create experiments. You can view and manage SageMaker Clarify bias reports or SageMaker Debugger output tensors for specific experiment runs directly in the Studio Classic Experiments UI.

- **Automatic experiment creation**
- **Bias and explainability reports**
- **Debugging**

**Automatic experiment creation**

Amazon SageMaker automatically creates experiments when running Autopilot jobs, hyperparameter optimization (HPO) jobs, or Pipeline executions. You can view these experiments in the Studio Classic Experiments UI.
Autopilot

Amazon SageMaker Experiments is integrated with Amazon SageMaker Autopilot. When you perform an Autopilot job, SageMaker Experiments creates an experiment for that job as well as runs for each of the different combinations of the available run components, parameters, and artifacts. You can find these runs in the SageMaker Experiments UI by filtering for the run type Autopilot. For more information, see Automate model development with Amazon SageMaker Autopilot.

HPO

Amazon SageMaker Experiments is integrated with HPO jobs. An HPO job automatically creates Amazon SageMaker experiments, runs, and components for each training job that it completes. You can find these runs in the SageMaker Experiments UI by filtering for the run type HPO. For more information, see Tune Multiple Algorithms with Hyperparameter Optimization to Find the Best Model.

Pipelines

Amazon SageMaker Model Building Pipelines is closely integrated with Amazon SageMaker Experiments. By default, when SageMaker Pipelines creates and executes a pipeline, experiments, runs, and components are created if they do not already exist. You can find these runs in the SageMaker Experiments UI by filtering for the run type Pipelines. For more information, see Amazon SageMaker Experiments Integration.

Bias and explainability reports

Manage SageMaker Clarify bias and explainability reports for experiment runs directly through Studio Classic. To view reports, find and select the name of the experiment run of your choice in Studio Classic. Choose Bias reports to see any Clarify bias reports associated with the experiment run.
Choose **Explanations** to see any Clarify explainability reports associated with the experiment run.
You can generate pre-training or post-training bias reports that analyze bias in datasets or model predictions using labels and bias metrics with SageMaker Clarify. You can also use SageMaker Clarify to generate explainability reports that document model behavior for global or local data samples. For more information, see Amazon SageMaker Clarify Bias Detection and Model Explainability.

**Debugging**

You can debug model training progress with Amazon SageMaker Debugger and view debug output tensors in the Studio Classic Experiments UI. Choose the name of the run associated with the Debugger report and choose Debugger.

Then, choose the training job name to view the associated Amazon SageMaker Debugger dashboard.
For more information, see [Debug Training Jobs Using Amazon SageMaker Debugger](#).

**Example notebooks for Amazon SageMaker Experiments**

The following tutorials demonstrate how to track runs for various model training experiments. You can view the resulting experiments in Studio Classic after running the notebooks. To clean up the resources created by a notebook, see [Clean Up Amazon SageMaker Experiment Resources](#). For a tutorial that showcases additional features of Studio, see [Amazon SageMaker Studio Classic Tour](#).

**Track experiments in a notebook environment**

To learn more about tracking experiments in a notebook environment, see the following example notebooks:

- [Track an experiment while training a Keras model locally](#)
- [Track an experiment while training a Pytorch model locally or in your notebook](#)
Track bias and explainability for your experiments with SageMaker Clarify

For a step-by-step guide on tracking bias and explainability for your experiments, see the following example notebook:

- [Fairness and Explainability with SageMaker Clarify](#)

Track experiments for SageMaker training jobs using script mode

For more information about tracking experiments for SageMaker training jobs, see the following example notebooks:

- [Run a SageMaker Experiment with Pytorch Distributed Data Parallel - MNIST Handwritten Digits Classification](#)
- [Track an experiment while training a Pytorch model with a SageMaker Training Job](#)
- [Train a TensorFlow model with a SageMaker training job and track it using SageMaker Experiments](#)

Monitor experiment training metrics with AWS CloudTrail

The training metrics for Amazon SageMaker Experiments are integrated with AWS CloudTrail, a service that provides a record of actions taken by a user, role, or an AWS service. CloudTrail captures all API calls for `BatchPutMetrics` as events. SageMaker automatically calls `BatchPutMetrics` when you create an experiment run using the SageMaker SDK for Python. AWS CloudTrail captures data related to calls for resource type `AWS::SageMaker::ExperimentTrialComponent`.

### Note

In the Studio Classic Experiments UI, trials are referred to as *run groups* and trial components are referred to as *runs*.

When you create an experiment run, you can also configure the continuous delivery of CloudTrail events to an Amazon S3 bucket. Use CloudTrail to monitor all ingested training metrics for an experiment run, including information such as the metric name, the training step of the recorded metric, the timestamp, and the metric value. CloudTrail events also include the experiment
run ARN, the ID of the account that created the run, and the resource type, which should be
AWS::SageMaker::ExperimentTrialComponent.

To monitor BatchPutMetrics API calls as CloudTrail events, you must first set up the logging
of data plane API activity in CloudTrail. See Logging data events for trails for more information.
For granular control over which API calls you want to selectively log and pay for, you can filter
CloudTrail events by resource type. Specify AWS::SageMaker::ExperimentTrialComponent
as a resource type to monitor calls to the BatchPutMetrics API. For more information, see
DataResource in the AWS CloudTrail API reference. To learn more about CloudTrail, see the AWS
CloudTrail User Guide.

For an in-depth explanation of how Amazon SageMaker works with AWS CloudTrail, see Log
Amazon SageMaker API Calls with AWS CloudTrail.

The following is an example CloudTrail event for a training metric in an experiment run:

```json
{
  ...
  "eventTime": "2022-12-14T21:53:41Z",
  "eventSource": "metrics-sagemaker.amazonaws.com",
  "eventName": "BatchPutMetrics",
  "awsRegion": "us-east-1",
  "sourceIPAddress": "192.0.2.0",
  "userAgent": "aws-cli/2.7.25 Python/3.9.11 Linux/5.4.214-134.408.amzn2int.x86_64 x86_64.amzn.2 prompt/off command/sm-metrics.batch-put-metrics",
  "requestParameters": {
    "trialComponentName": "trial-component-name",
    "metricData": [
      {
        "metricName": "foo",
        "timestamp": 1670366870000,
        "step": 101,
        "value": 0.9
      }
    ]
  },
  ...
  "resources": [
    {
      "accountId": "abcdef01234567890",
      "type": "AWS::SageMaker::ExperimentTrialComponent",
    }
  ]
}
```
Clean Up Amazon SageMaker Experiment Resources

To avoid incurring unnecessary charges, delete the Amazon SageMaker Experiment resources you no longer need. You can't delete Experiment resources through the SageMaker Management Console or the Amazon SageMaker Studio Classic UI. This topic shows you how to clean up these resources using the SageMaker Python SDK, Boto3, and the Experiments SDK.

Topics
- Clean Up Using the SageMaker Python SDK (Recommended)
- Clean Up Using the Python SDK (Boto3)
- Clean Up Using the Experiments SDK

Clean Up Using the SageMaker Python SDK (Recommended)

To clean up using the SageMaker Python SDK

```python
from sagemaker.experiments.experiment import Experiment

exp = Experiment.load(experiment_name=experiment_name, sagemaker_session=sm_session)
exp._delete_all(action="--force")
```

Clean Up Using the Python SDK (Boto3)

To clean up using Boto 3

```python
import boto3
sm = boto3.Session().client('sagemaker')

def cleanup_boto3(experiment_name):
```

Define cleanup_boto3
trials = sm.list_trials(ExperimentName=experiment_name)['TrialSummaries']
print('TrialNames:')
for trial in trials:
    trial_name = trial['TrialName']
    print(f"\n{trial_name}\")

components_in_trial = sm.list_trial_components(TrialName=trial_name)
print('TrialComponentNames:')
for component in components_in_trial['TrialComponentSummaries']:
    component_name = component['TrialComponentName']
    print(f'\t{component_name}\")
sm.disassociate_trial_component(TrialComponentName=component_name,
    TrialName=trial_name)
try:
    # comment out to keep trial components
    sm.delete_trial_component(TrialComponentName=component_name)
except:
    # component is associated with another trial
    continue
    # to prevent throttling
    time.sleep(.5)
sm.delete_trial(TrialName=trial_name)
sm.delete_experiment(ExperimentName=experiment_name)
print(f'\nExperiment {experiment_name} deleted')

Call cleanup_boto3

# Use experiment name not display name
experiment_name = "experiment-name"
cleanup_boto3(experiment_name)

Clean Up Using the Experiments SDK

To clean up using the Experiments SDK

import sys
!{sys.executable} -m pip install sagemaker-experiments

import time
from smexperiments.experiment import Experiment
from smexperiments.trial import Trial
Define cleanup_sme_sdk

```python
def cleanup_sme_sdk(experiment):
    for trial_summary in experiment.list_trials():
        trial = Trial.load(trial_name=trial_summary.trial_name)
        for trial_component_summary in trial.list_trial_components():
            tc = TrialComponent.load(
                trial_component_name=trial_component_summary.trial_component_name)
            trial.remove_trial_component(tc)
            try:
                # comment out to keep trial components
                tc.delete()
            except:
                # tc is associated with another trial
                continue
            # to prevent throttling
            time.sleep(.5)
            trial.delete()
        trial.delete()
        experiment_name = experiment.experiment_name
        experiment.delete()
        print(f"\nExperiment {experiment_name} deleted")
```

Call cleanup_sme_sdk

```python
experiment_to_cleanup = Experiment.load(
    # Use experiment name not display name
    experiment_name="experiment-name")
cleanup_sme_sdk(experiment_to_cleanup)
```

Additional supported SDK

⚠️ Important

As of v2.123.0, SageMaker Experiments is now fully integrated with the SageMaker Python SDK and you no longer need to use the separate SageMaker Experiments SDK. We recommend creating an experiment with sagemaker.experiments.run rather than the following smexperiments module.
The following section describes how to create a SageMaker Experiment with the SageMaker Experiments SDK.

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

---

**Create an Amazon SageMaker Experiment with the SageMaker Experiments SDK**

Create an Amazon SageMaker experiment to track your SageMaker training, processing, and transform jobs.

The following procedure shows you how to create a SageMaker experiment for a SageMaker training, processing, or transform job. Steps labeled as (Studio Classic) describe how to view the experiment in Amazon SageMaker Studio Classic. You don't have to run the experiment in Studio Classic to view the experiment in Studio Classic.

1. Import the `sys` module to install the SDKs.

   ```py
   import sys
   ```

2. (Optional) The Amazon SageMaker Python SDK, comes preinstalled in Amazon SageMaker Studio Classic. If you plan to run your code outside Studio Classic, install the SageMaker Python SDK.

   ```sh
   !{sys.executable} -m pip install sagemaker
   ```

3. Install the SageMaker Experiments Python SDK.

   ```sh
   !{sys.executable} -m pip install sagemaker-experiments
   ```

4. Import modules.

   ```py
   import time
   from time import strftime
   ```
import sagemaker

from smexperiments.experiment import Experiment
from smexperiments.trial import Trial
from smexperiments.trial_component import TrialComponent
from smexperiments.tracker import Tracker

5. Get the execution role and create the SageMaker session.

role = sagemaker.get_execution_role()
sm_sess = sagemaker.session.Session()

6. Create a SageMaker experiment. The experiment name must be unique in your account.

   **Note**

   The tags parameter is optional. You can search for the tag using Studio Classic, the SageMaker console, and the SDK. Tags can also be applied to trials and trial components.

   create_date = strftime("%Y-%m-%d-%H-%M-%S")
   demo_experiment = Experiment.create(experiment_name = "DEMO-{}".format(create_date),
                                          description = "Demo experiment",
                                          tags = [{"Key": 'demo-experiments', 'Value':
                                                                  'demo1'}])

7. (Studio Classic) To view the experiment in Amazon SageMaker Studio Classic, in the left sidebar, choose the **Experiments**.

   After the code runs, the experiment list contains the new experiment. It might take a moment for the list to refresh and display the experiment. The filter on the experiment tag is also displayed. Only experiments that have a matching tag are displayed. Your list should look similar to the following:
8. Create a trial for the experiment. The trial name must be unique in your account.

```python
demo_trial = Trial.create(trial_name = "DEMO-{}".format(create_date),
                        experiment_name = demo_experiment.experiment_name,
                        tags = [{'Key': 'demo-trials', 'Value': 'demo1'}])
```

9. Create a trial component as part of the trial. The trial component is the SageMaker job.

Add the `ExperimentConfig` parameter to the appropriate method. The SageMaker jobs listed in the following table are supported.

<table>
<thead>
<tr>
<th>Job</th>
<th>SageMaker Python SDK method</th>
<th>Boto3 method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training</td>
<td><code>Estimator.fit</code></td>
<td><code>CreateTrainingJob</code></td>
</tr>
<tr>
<td>Processing</td>
<td><code>Processor.run</code></td>
<td><code>CreateProcessingJob</code></td>
</tr>
<tr>
<td>Transform</td>
<td><code>Transformer.transform</code></td>
<td><code>CreateTransformJob</code></td>
</tr>
</tbody>
</table>

Additional supported SDK

3149
The following examples are for a training job. The Tags parameter adds a tag to the trial component. ExperimentName isn’t specified because the trial was associated with the experiment when the trial was created in an earlier step.

**Using the SageMaker Python SDK**

```python
sagemaker.estimator.Estimator(
    ...,  
    sagemaker_session = sm_sess,
    tags = [{'Key': 'demo-jobs', 'Value': 'demo2'}])

estimator.fit(
    ...,  
    experiment_config = {
        # "ExperimentName"
        "TrialName" : demo_trial.trial_name,
        "TrialComponentDisplayName" : "TrainingJob",
    }
)
```

**Using Boto3**

```python
create_training_job(
    ...,  
    "ExperimentConfig": {
        # "ExperimentName"
        "TrialName" : demo_trial.trial_name,
        "TrialComponentDisplayName" : "TrainingJob",
    },
    "Tags": [{'Key': 'demo-jobs', 'Value': 'demo2'}])
```

10. *(Studio Classic)* In the experiment list, double-click the experiment to display a list of the trials in the experiment. In the Studio Classic UI, trials are referred to as *run groups* and trial components are referred to as *runs*. Your list should look similar to the following:
11. (Studio Classic) To view information about the experiment, trial, and job (trial component), see View, search, and compare experiment runs.

To clean up the resources you created, see Clean Up Amazon SageMaker Experiment Resources.

**Experiments FAQs**

Refer to the following FAQ items for answers to commonly asked questions about SageMaker Experiments.

**Q. What is the recommended method to create an experiment?**

A: Experiments are a collection of runs aimed at finding the best model to solve a problem. To initialize a run within an experiment, use the SageMaker Python SDK Run class. For more examples, see Create an Amazon SageMaker Experiment.

**Q. Can I create an experiment using SageMaker script mode?**

Yes. You can create experiments using SageMaker script mode. In the Jupyter notebook or Python file you are using to define your estimator, initialize a run using the Run class. Within the run, launch an estimator with your custom entry point script. Within that entry point script, use the
load_run method to initialize the run you defined within the entry point script and log your metrics. For in-depth examples, see Track experiments for SageMaker training jobs using script mode.

Q. What SageMaker jobs automatically create experiments?

SageMaker Hyperparameter Optimization (HPO) jobs (also known as tuning jobs) automatically create experiments to track all the training jobs launched during a hyperparameter search. All other SageMaker jobs create unassigned runs unless launched from within an experiment.

Q. What kind of SageMaker jobs can I create an experiment for?

You can use SageMaker Experiments to track metrics from training jobs, processing jobs, and transform jobs.

Q. Why do I see experiments and runs in the Experiments Studio Classic UI that I did not create using the SageMaker Python SDK?

Experiment runs that are automatically created by SageMaker jobs and containers are visible in the Experiments Studio Classic UI by default. To hide runs created by SageMaker jobs for a given experiment, choose the settings icon and toggle Show jobs.

Q. Is the SageMaker Experiments SDK still supported?

Yes, the SageMaker Experiments SDK is still supported. However, as of v2.123.0, SageMaker Experiments is fully integrated with the SageMaker Python SDK. We recommend using the SageMaker Python SDK to create experiments and runs. For more information, see Create an Amazon SageMaker Experiment.

Q. Can I use distributed training with my experiments?

A: Yes. However, metrics for distributed training can be logged only at the epoch level. Be sure that you only log metrics generated by the leader node, as shown in the following example:

```python
... if rank == 0:
    test_loss, correct, target, pred = test(model, test_loader, device, tracker)
    logger.info("Test Average loss: {:.4f}, Test Accuracy: {:.0f}%\n".format(
```
```
    test_loss, test_accuracy)
    )
    run.log_metric(name = "train_loss", value = loss.item(), step = epoch)
    run.log_metric(name = "test_loss", value = test_loss, step = epoch)
    run.log_metric(name = "test_accuracy", value = test_accuracy, step = epoch)
    ...
```

For more information, see the [Run a SageMaker Experiment with Pytorch Distributed Data Parallel - MNIST Handwritten Digits Classification](#) example notebook.

**Q. What are unassigned runs?**

A: All jobs in SageMaker (training jobs, processing jobs, transform jobs) correspond to runs. When launching these jobs, TrialComponents are created by default. TrialComponents map directly to runs. If these jobs are launched without being explicitly associated with an experiment or run, they are created as unassigned runs.

**Q. Do I need to pass the experiment run context to the training script when running a SageMaker training job?**

A: Yes. You need to load the run context into the training script, along with the SageMaker session information.

```python
from sagemaker.session import Session
from sagemaker.experiments.run import load_run

session = Session(boto3.session.Session(region_name=args.region))

with load_run(sagemaker_session=session) as run:
    run.log_parameters(
        {"num_train_samples": len(train_set.data), "num_test_samples": len(test_set.data)}
    )
```

**Q. How do I add a new run to an experiment analysis?**

A: If you already created a comparison for your experiment and want to add a new run to analyze, select all the runs from your previous analysis as well as the new run and choose **Analyze**. If you don't see your new run in the resulting analysis page, then refresh the Studio Classic browser. Note that refreshing your Studio Classic browser may impact your other open tabs.
Search Using the Amazon SageMaker Console and API

Developing a machine learning model typically requires extensive experimenting with different datasets, algorithms, and hyperparameter values. To manage up to thousands of machine learning model experiments, use the search capabilities in SageMaker.

You can use SageMaker search to:

- Organize, find, and evaluate training jobs using properties, hyperparameters, performance metrics, or any metadata.
- Find the best performing model by reviewing training job and model metrics, such as training loss or validation accuracy.
- Trace a model's lineage to the training job and its related resources, such as the training datasets.

This topic covers searching from the SageMaker console and the SageMaker API.

Topics

- Organize, Find, and Evaluate Training Jobs (Console)
- Find and Evaluate Training Jobs (API)
- Verify the Datasets Used by Your Training Jobs
- Trace Model Lineage

Organize, Find, and Evaluate Training Jobs (Console)

To organize training jobs, assign one or more tags to them.

To find a specific training job, model, or resource, use model tracking to search on keywords assigned to any searchable items. *Searchable items* include training jobs, models, hyperparameters, metadata, tags, and URLs. To refine your tracking results, you can search using multiple criteria.

To choose the best model for deployment, evaluate how all models performed against one or more metrics. You can use model tracking results to list, sort, and evaluate the performance of the models in your experiments.

Topics

- Use Tags to Track Training Jobs (Console)
- Find Training Jobs (Console)
Use Tags to Track Training Jobs (Console)

To group training jobs, create tags with descriptive keys and a value. For example, create tag keys for: project, owner, customer, and industry.

Add tags to training jobs (console)

1. Open the Amazon SageMaker console.
2. In the navigation pane, choose Training jobs and Create training job.
3. Scroll to the bottom of the page and enter a key and value for the tag.

4. To add another tag, choose Add tag, and add another key-value pair.

Find Training Jobs (Console)

You can search for training jobs using a variety of job attributes. Note that some search parameters appear only if you have created a training job with that attribute. For example, Tags appears only if you have added a tag for a training job.

To find training jobs (console)

1. Open the Amazon SageMaker console.
2. In the navigation pane, choose Search.
3. Add Parameters.
   a. In the search box, enter a parameter and choose a parameter type, for example TrainingJobName.
b. Choose a conditional operation. For numeric values, use operators such as **is equals to**, **lesser than**, or **or greater than**. For text-based values, use operators such as **equals to** or **contains**.

c. Enter a value for the parameter.

4. (Optional) To refine your search, add additional search criteria. Choose **Add row** and enter the parameter values.

5. Choose **Search**.

---

**Evaluate Models (Console)**

To evaluate a model's performance, review its metadata, hyperparameters, and metrics. To highlight metrics, adjust the view to show only metrics and important hyperparameters.

**To evaluate a model (console)**

1. Open the [Amazon SageMaker console](https://console.aws.amazon.com/sagemaker/home).  
2. In the navigation pane, choose **Search** and search for training jobs by specifying relevant parameters. The results are displayed in a table.

```
<table>
<thead>
<tr>
<th>HyperParameter</th>
<th>HyperParameter</th>
<th>Metric</th>
<th>Metric</th>
<th>Metric</th>
<th>Metric</th>
</tr>
</thead>
<tbody>
<tr>
<td>mini_batch_size</td>
<td>predictor_type</td>
<td>trainbinary_f_beta</td>
<td>progress</td>
<td>trainobjective_loss</td>
<td>binary_classification_accuracy</td>
</tr>
<tr>
<td>300</td>
<td>binary_classifier</td>
<td>0.966639518737795</td>
<td>100</td>
<td>0.02381423674523804</td>
<td>0.9934399724006653</td>
</tr>
<tr>
<td>100</td>
<td>binary_classifier</td>
<td>0.9652714135262634</td>
<td>100</td>
<td>0.0235049128553240967</td>
<td>0.993179976940155</td>
</tr>
<tr>
<td>200</td>
<td>binary_classifier</td>
<td>0.9647442698478699</td>
<td>100</td>
<td>0.023529807378053665</td>
<td>0.99308000199508667</td>
</tr>
</tbody>
</table>
```

3. Open the preferences window by choosing the settings icon in the search results table.

4. To show or hide a hyperparameter or metric, turn it on or off by choosing **Hyperparameter** or **Metric**.

5. Make necessary changes, then choose **Update view**.

6. After viewing metrics and important hyperparameters, you can compare and contrast the result. Then, you can choose the best model to host or investigate the models that are performing poorly.
Find and Evaluate Training Jobs (API)

To the find and evaluate training jobs or to get suggestions for items used in experiments that are searchable, you can use the Search API.

Topics

- Find Training Jobs (API)
- Evaluate Models (API)
- Get Suggestions for a Search (API)

Find Training Jobs (API)

To find training jobs, create a search parameter using the search_params parameter. Then use the search function in the smclient subprocess in the AWS SDK for Python (Boto3).

The following example shows how to use the Search API to find training jobs.

```python
import boto3

search_params={
    "MaxResults": 10,
    "Resource": "TrainingJob",
    "SearchExpression": {
        "Filters":[
            {
                "Name": "Tags.Project",
                "Operator": "Equals",
                "Value": "Project_Binary_Classifier"
            }
        ],
        "SortBy": "Metrics.train:binary_classification_accuracy",
        "SortOrder": "Descending"
    }
}

smclient = boto3.client(service_name='sagemaker')
results = smclient.search(**search_params)
```

Evaluate Models (API)

To evaluate models, run a search as described in Find Training Jobs (API), review model metrics, then, use the AWS SDK for Python (Boto3) to create a table and plot it.

The following example shows how to evaluate models and to display the results in a table.
import pandas

headers=["Training Job Name", "Training Job Status", "Batch Size", "Binary Classification Accuracy"]
rows=[]
for result in results['Results']:
    trainingJob = result['TrainingJob']
    metrics = trainingJob['FinalMetricDataList']
    rows.append([trainingJob['TrainingJobName'],
                  trainingJob['TrainingJobStatus'],
                  trainingJob['HyperParameters']['mini_batch_size'],
                  metrics[[x['MetricName'] for x in
                       metrics].index('train:binary_classification_accuracy')]["Value"]
                  ])

df = pandas.DataFrame(data=rows,columns=headers)

from IPython.display import display, HTML
display(HTML(df.to_html()))

Get Suggestions for a Search (API)

To get suggestions for a search, use the GetSearchSuggestions API.

The following example for AWS SDK for Python (Boto3) is a get_search_suggestions request for items containing linear.

```
search_suggestion_params=
    "Resource": "TrainingJob",
    "SuggestionQuery": {
        "PropertyNameQuery": {
            "PropertyNameHint": "linear"
        }
    }
```

The following is an example response for a get_search_suggestions request.

```
{
    'PropertyNameSuggestions': [["PropertyName":
        'hyperparameters.linear_init_method'],
       [{'PropertyName': 'hyperparameters.linear_init_value'},
        {'PropertyName': 'hyperparameters.linear_init_sigma'}],
```
After getting search suggestions, you can use one of the property names in a search.

**Verify the Datasets Used by Your Training Jobs**

You can use model tracking capability to verify which datasets were used in training, where holdout datasets were used, and other details about training jobs. For example, use model tracking capability to verify that a specific dataset was used in a training job for an audit or to verify compliance.

To check whether a specific dataset was used in a training job, you search for the URL to its location in Amazon Simple Storage Service (Amazon S3). Model tracking capability returns the training jobs that used the dataset that you specify. If your search doesn't return the dataset (the result is empty), the dataset wasn't used in a training job. An empty result confirms, for example, that a holdout dataset wasn't used.

**Trace Model Lineage**

You can use model tracking capability to get information about the lineage of training jobs and the model resources that were used for them, including the dataset, algorithm, hyperparameters, and metrics. For example, if you find that the performance of a hosted model has declined, you can review its training job and the resources it used to determine what's causing the problem.

**Topics**

- [Trace Model Lineage (Console)]
- [Trace Model Lineage (API)]

**Trace Model Lineage (Console)**

**To trace a model's lineage (console)**

1. Open the [Amazon SageMaker console].
2. In the navigation pane, choose **Endpoints**, and choose the relevant endpoint.
3. Scroll to the **Endpoint configuration settings** section. This section lists all of the model versions deployed at the endpoint, with a hyperlink to the training job that created each.
Trace Model Lineage (API)

To trace a model's lineage, get the model's name, then use it to search for training jobs.

The following example shows how to trace a model's lineage using the API.

```python
# Get the name of model deployed at endpoint
endpoint_config = smclient.describe_endpoint_config(EndpointConfigName=endpointName)
model_name = endpoint_config['ProductionVariants'][0]['ModelName']

# Get the model's name
model = smclient.describe_model(ModelName=model_name)

# Search the training job by the location of model artifacts in Amazon S3
search_params = {
    "MaxResults": 1,
    "Resource": "TrainingJob",
    "SearchExpression": {
        "Filters": [
            {
                "Name": "ModelArtifacts.S3ModelArtifacts",
                "Operator": "Equals",
                "Value": model['PrimaryContainer']['ModelDataUrl']
            }
        ]
    }
}
results = smclient.search(**search_params)
```

After finding the training job, you can review the resources used to train the model.

Perform Automatic Model Tuning with SageMaker

Amazon SageMaker automatic model tuning (AMT), also known as hyperparameter tuning, finds the best version of a model by running many training jobs on your dataset. To do this, AMT uses the algorithm and ranges of hyperparameters that you specify. It then chooses the hyperparameter values that creates a model that performs the best, as measured by a metric that you choose.

For example, suppose that you want to solve a binary classification problem on a marketing dataset. Your goal is to maximize the area under the curve (AUC) metric of the algorithm by training an XGBoost Algorithm model. You want to find which values for the eta, alpha, min_child_weight, and max_depth hyperparameters that will train the best model. Specify a range of values for these hyperparameters. Then, SageMaker hyperparameter tuning searches
within these ranges to find a combination of values that creates a training job that creates a model with the highest AUC. To conserve resources or meet a specific model quality expectation, you can also set up completion criteria to stop tuning after the criteria have been met.

You can use SageMaker AMT with built-in algorithms, custom algorithms, or SageMaker pre-built containers for machine learning frameworks.

SageMaker AMT can use an Amazon EC2 Spot instance to optimize costs when running training jobs. For more information, see Use Managed Spot Training in Amazon SageMaker.

Before you start using hyperparameter tuning, you should have a well-defined machine learning problem, including the following:

- A dataset
- An understanding of the type of algorithm that you need to train
- A clear understanding of how you measure success

Prepare your dataset and algorithm so that they work in SageMaker and successfully run a training job at least once. For information about setting up and running a training job, see Get started.

Topics

- How Hyperparameter Tuning Works
- Define metrics and environment variables
- Define Hyperparameter Ranges
- Track and set completion criteria for your tuning job
- Tune Multiple Algorithms with Hyperparameter Optimization to Find the Best Model
- Example: Hyperparameter Tuning Job
- Stop Training Jobs Early
- Run a Warm Start Hyperparameter Tuning Job
- Resource Limits for Automatic Model Tuning
- Best Practices for Hyperparameter Tuning

How Hyperparameter Tuning Works

When you build complex machine learning systems like deep learning neural networks, exploring all of the possible combinations is impractical. Hyperparameter tuning can accelerate your
productivity by trying many variations of a model. It looks for the best model automatically by focusing on the most promising combinations of hyperparameter values within the ranges that you specify. To get good results, you must choose the right ranges to explore.

Use the API reference guide to understand how to interact with hyperparameter tuning. The examples on this page can be found in the HyperParameterTuningJobConfig and HyperbandStrategyConfig APIs.

Grid Search

When using grid search, hyperparameter tuning chooses combinations of values from the range of categorical values that you specify when you create the job. Only categorical parameters are supported when using the grid search strategy. You do not need to specify the MaxNumberOfTrainingJobs. The number of training jobs created by the tuning job will be automatically calculated to be the total number of distinct categorical combinations possible. If specified, the value of MaxNumberOfTrainingJobs should equal the total number of distinct categorical combinations possible.

Random Search

When using random search, hyperparameter tuning chooses a random combination of values from within the ranges that you specify for hyperparameters for each training job it launches. Because the choice of hyperparameter values doesn't depend on the results of previous training jobs, you can run the maximum number of concurrent training jobs without affecting the performance of the tuning.

For an example notebook that uses random search, see the Random search and hyperparameter scaling with SageMaker XGBoost and Automatic Model Tuning notebook.

Bayesian Optimization

Bayesian optimization treats hyperparameter tuning like a regression problem. Given a set of input features (the hyperparameters), hyperparameter tuning optimizes a model for the metric that
you choose. To solve a regression problem, hyperparameter tuning makes guesses about which hyperparameter combinations are likely to get the best results, and runs training jobs to test these values. After testing a set of hyperparameter values, hyperparameter tuning uses regression to choose the next set of hyperparameter values to test.

Hyperparameter tuning uses an Amazon SageMaker implementation of Bayesian optimization.

When choosing the best hyperparameters for the next training job, hyperparameter tuning considers everything that it knows about this problem so far. Sometimes it chooses a combination of hyperparameter values close to the combination that resulted in the best previous training job to incrementally improve performance. This allows hyperparameter tuning to exploit the best known results. Other times, it chooses a set of hyperparameter values far removed from those it has tried. This allows it to explore the range of hyperparameter values to try to find new areas that are not yet well understood. The explore/exploit trade-off is common in many machine learning problems.

For more information about Bayesian optimization, see the following:

**Basic Topics on Bayesian Optimization**

- [A Tutorial on Bayesian Optimization of Expensive Cost Functions, with Application to Active User Modeling and Hierarchical Reinforcement Learning](#)
- [Practical Bayesian Optimization of Machine Learning Algorithms](#)
- [Taking the Human Out of the Loop: A Review of Bayesian Optimization](#)

**Speeding up Bayesian Optimization**

- [Google Vizier: A Service for Black-Box Optimization](#)
- [Learning Curve Prediction with Bayesian Neural Networks](#)
- [Speeding up automatic hyperparameter optimization of deep neural networks by extrapolation of learning curves](#)

**Advanced Modeling and Transfer Learning**

- [Scalable Hyperparameter Transfer Learning](#)
- [Bayesian Optimization with Tree-structured Dependencies](#)
- [Bayesian Optimization with Robust Bayesian Neural Networks](#)
Hyperband

Hyperband is a multi-fidelity based tuning strategy that dynamically reallocates resources. Hyperband uses both intermediate and final results of training jobs to re-allocate epochs to well-utilized hyperparameter configurations and automatically stops those that underperform. It also seamlessly scales to using many parallel training jobs. These features can significantly speed up hyperparameter tuning over random search and Bayesian optimization strategies.

Hyperband should only be used to tune iterative algorithms that publish results at different resource levels. For example, Hyperband can be used to tune a neural network for image classification which publishes accuracy metrics after every epoch.

For more information about Hyperband, see the following links:

- Hyperband: A Novel Bandit-Based Approach to Hyperparameter Optimization
- Massively Parallel Hyperparameter Tuning
- BOHB: Robust and Efficient Hyperparameter Optimization at Scale
- Model-based Asynchronous Hyperparameter and Neural Architecture Search

Hyperband with early stopping

Training jobs can be stopped early when they are unlikely to improve the objective metric of the hyperparameter tuning job. This can help reduce compute time and avoid overfitting your model. Hyperband uses an advanced internal mechanism to apply early stopping. Thus, the parameter TrainingJobEarlyStoppingType in the HyperParameterTuningJobConfig API must be set to OFF when using the Hyperband internal early stopping feature.

⚠️ Note

Hyperparameter tuning might not improve your model. It is an advanced tool for building machine solutions. As such, it should be considered part of the scientific development process.
Define metrics and environment variables

A tuning job optimizes hyperparameters for training jobs that it launches by using a metric to evaluate performance. This guide shows how to define metrics so that you can use a custom algorithm for training, or use a built-in algorithm from Amazon SageMaker. This guide also shows how to specify environment variables during an Automatic model tuning (AMT) job.

Define metrics

Amazon SageMaker hyperparameter tuning parses your machine learning algorithm's stdout and stderr streams to find metrics, such as loss or validation-accuracy. The metrics show how well the model is performing on the dataset.

The following sections describe how to use two types of algorithms for training: built-in and custom.

Use a built-in algorithm for training

If you use one of the SageMaker built-in algorithms, metrics are already defined for you. In addition, built-in algorithms automatically send metrics to hyperparameter tuning for optimization. These metrics are also written to Amazon CloudWatch logs. For more information, see Log Amazon SageMaker Events with Amazon CloudWatch.

For the objective metric for the tuning job, choose one of the metrics that the built-in algorithm emits. For a list of available metrics, see the model tuning section for the appropriate algorithm in Use Amazon SageMaker Built-in Algorithms or Pre-trained Models.

You can choose up to 40 metrics to monitor in your tuning job. Select one of those metrics to be the objective metric. The hyperparameter tuning job returns the training job that performed the best against the objective metric.

Note

Hyperparameter tuning automatically sends an additional hyperparameter _tuning_objective_metric to pass your objective metric to the tuning job for use during training.
Use a custom algorithm for training

This section shows how to define your own metrics to use your own custom algorithm for training. When doing so, make sure that your algorithm writes at least one metric to stderr or stdout. Hyperparameter tuning parses these streams to find algorithm metrics that show how well the model is performing on the dataset.

You can define custom metrics by specifying a name and regular expression for each metric that your tuning job monitors. Then, pass these metric definitions to the CreateHyperParameterTuningJob API in the TrainingJobDefinition parameter in the MetricDefinitions field of AlgorithmSpecification.

The following shows sample output from a log written to stderr or stdout by a training algorithm.

GAN_loss=0.138318;  Scaled_reg=2.654134; disc:[-0.017371,0.102429] real 93.3% gen 0.0% 
disc-combined=0.000000; disc_train_loss=1.374587;  Loss = 16.020744; Iteration 0 took 0.704s;  Elapsed=0s

The following code example shows how to use regular expressions in Python (regex). This is used to search the sample log output and capture the numeric values of four different metrics.

```
[{
  "Name": "ganloss",
  "Regex": "GAN_loss=(.*?);",
},
{
  "Name": "disc-combined",
  "Regex": "disc-combined=(.*?);",
},
{
  "Name": "discloss",
  "Regex": "disc_train_loss=(.*?);",
},
{
  "Name": "loss",
  "Regex": "Loss = (.*?);",
}]
```

In regular expressions, parenthesis () are used to group parts of the regular expression together.
• For the loss metric that is defined in the code example, the expression (\(.*?\)); captures any character between the exact text "Loss=" and the first semicolon (\(;) character.

• The character \(\cdot\) instructs the regular expression to match any character.

• The character \(\ast\) means to match zero or more characters.

• The character \(\?\) means capture only until the first instance of the \(\;\) character.

The loss metric defined in the code sample will capture Loss = 16.020744 from the sample output.

Choose one of the metrics that you define as the objective metric for the tuning job. If you are using the SageMaker API, specify the value of the name key in the HyperParameterTuningJobObjective field of the HyperParameterTuningJobConfig parameter that you send to the CreateHyperParameterTuningJob operation.

Specify environment variables

SageMaker AMT optimizes hyperparameters within a tuning job to find the best parameters for model performance. You can use environment variables to configure your tuning job to change its behavior. You can also use environment variables that you used during training inside your tuning job.

If you want to use an environment variable from your tuning job or specify a new environment variable, input a string value for Environment within the SageMaker HyperParameterTrainingJobDefinition API. Pass this training job definition to the CreateHyperParameterTuningJob API.

For example, the environment variable SM_LOG_LEVEL can be set to the following values to tailor the output from a Python container.

```
NOTSET=0
DEBUG=10
INFO=20
WARN=30
ERROR=40
CRITICAL=50
```

As an example, to set the log level to 10 to debug your container logs, set the environment variable inside the HyperParameterTrainingJobDefinition, as follows.
Define Hyperparameter Ranges

This guide shows how to use SageMaker APIs to define hyperparameter ranges. It also provides a list of hyperparameter scaling types that you can use.

Choosing hyperparameters and ranges significantly affects the performance of your tuning job. Hyperparameter tuning finds the best hyperparameter values for your model by searching over a range of values that you specify for each tunable hyperparameter. You can also specify up to 100 static hyperparameters that do not change over the course of the tuning job. You can use up to 100 hyperparameters in total (static + tunable). For guidance on choosing hyperparameters and ranges, see Best Practices for Hyperparameter Tuning. You can also use autotune to find optimal tuning job settings. For more information, see the following Autotune section.

### Note
SageMaker Automatic Model Tuning (AMT) may add additional hyperparameters(s) that contribute to the limit of 100 total hyperparameters. Currently, to pass your objective metric to the tuning job for use during training, SageMaker adds _tuning_objective_metric automatically.
**Static hyperparameters**

Use static hyperparameters for the following cases: For example, you can use AMT to tune your model using `param1` (a tunable parameter) and `param2` (a static parameter). If you do, then use a search space for `param1` that lies between two values, and pass `param2` as a static hyperparameter, as follows.

```
param1: ["range_min","range_max"]
param2: "static_value"
```

Static hyperparameters have the following structure:

```
"StaticHyperParameters": {
    "objective" : "reg:squarederror",
    "dropout_rate": "0.3"
}
```

You can use the Amazon SageMaker API to specify key value pairs in the `StaticHyperParameters` field of the `HyperParameterTrainingJobDefinition` parameter that you pass to the `CreateHyperParameterTuningJob` operation.

**Dynamic hyperparameters**

You can use the SageMaker API to define [hyperparameter ranges](hyperparameter_ranges). Specify the names of hyperparameters and ranges of values in the `ParameterRanges` field of the `HyperParameterTuningJobConfig` parameter that you pass to the `CreateHyperParameterTuningJob` operation.

The `ParameterRanges` field has three subfields: categorical, integer, and continuous. You can define up to 30 total (categorical + integer + continuous) tunable hyperparameters to search over.

**Note**

Each categorical hyperparameter can have at most 30 different values.

Dynamic hyperparameters have the following structure:

```
"ParameterRanges": {
```
If you create a tuning job with a Grid strategy, you can only specify categorical values. You don't need to provide the `MaxNumberOfTrainingJobs`. This value is inferred from the total number of configurations that can be produced from your categorical parameters. If specified, the value of `MaxNumberOfTrainingJobs` should be equal to the total number of distinct categorical combinations possible.

**Autotune**

To save time and resources searching for hyperparameter ranges, resources or objective metrics, autotune can automatically guess optimal values for some hyperparameter fields. Use autotune to find optimal values for the following fields:

- **ParameterRanges** – The names and ranges of hyperparameters that a tuning job can optimize.
- **ResourceLimits** – The maximum resources to be used in a tuning job. These resources can include the maximum number of training jobs, maximum runtime of a tuning job, and the maximum number of training jobs that can be run at the same time.
• **TrainingJobEarlyStoppingType** – A flag that stops a training job if a job is not significantly improving against an objective metric. Defaults to enabled. For more information, see [Stop Training Jobs Early](#).

• **RetryStrategy** – The number of times to retry a training job. Non-zero values for RetryStrategy can increase the likelihood that your job will complete successfully.

• **Strategy** – Specifies how hyperparameter tuning chooses the combinations of hyperparameter values to use for the training job that it launches.

• **ConvergenceDetected** – A flag to indicate that Automatic Model Tuning (AMT) has detected model convergence.

To use autotune, do the following:

1. Specify the hyperparameter and an example value in the `AutoParameters` field of the `ParameterRanges` API.
2. Enable autotune.

AMT will determine if your hyperparameters and example values are eligible for autotune. Hyperparameters that can be used in autotune are automatically assigned to the appropriate parameter range type. Then, AMT uses `ValueHint` to select an optimal range for you. You can use the `DescribeHyperParameterTrainingJob` API to view these ranges.

The following example shows you how to configure a tuning job that uses autotune. In the configuration example, the hyperparameter `max_depth` has `ValueHint` containing an example value of 4.

```
config = {
    'Autotune': {'Mode': 'Enabled'},
    'HyperParameterTuningJobName': 'my-autotune-job',
    'HyperParameterTuningJobConfig': {
        'HyperParameterTuningJobObjective': {'Type': 'Minimize', 'MetricName': 'validation:rmse'},
        'ResourceLimits': {'MaxNumberOfTrainingJobs': 5, 'MaxParallelTrainingJobs': 1},
        'ParameterRanges': {
            'AutoParameters': [
                {'Name': 'max_depth', 'ValueHint': '4'}
            ]
        }
    }
}
```
Continuing the previous example, a tuning job is created after the previous configuration is included in a call to the CreateHyperParameterTuningJob API. Then, autotune converts the max_depth hyperparameter in AutoParameters to the hyperparameter IntegerParameterRanges. The following response from a DescribeHyperParameterTrainingJob API shows that the optimal IntegerParameterRanges for max_depth are between 2 and 8.

```json
{
    'HyperParameterTuningJobName': 'my_job',
    'HyperParameterTuningJobConfig': {
        'ParameterRanges': {
            'IntegerParameterRanges': [
                {'Name': 'max_depth', 'MinValue': '2', 'MaxValue': '8'},
            ],
        },
    'TrainingJobDefinition': {
        ...
    },
    'Autotune': {'Mode': 'Enabled'}
}
```

**Hyperparameter scaling types**

For integer and continuous hyperparameter ranges, you can choose the scale that you want hyperparameter tuning to use. For example, to search the range of values, you can specify a value for the ScalingType field of the hyperparameter range. You can choose from the following hyperparameter scaling types:

**Auto**

SageMaker hyperparameter tuning chooses the best scale for the hyperparameter.

**Linear**

Hyperparameter tuning searches the values in the hyperparameter range by using a linear scale. Typically, you choose this if the range of all values from the lowest to the highest is relatively
small (within one order of magnitude). Uniformly searching values from the range provides a reasonable exploration of the entire range.

Logarithmic

Hyperparameter tuning searches the values in the hyperparameter range by using a logarithmic scale.

Logarithmic scaling works only for ranges that have values greater than 0.

Choose logarithmic scaling when you're searching a range that spans several orders of magnitude.

For example, if you're tuning a Tune a linear learner model model, and you specify a range of values between .0001 and 1.0 for the learning_rate hyperparameter, consider the following:

Searching uniformly on a logarithmic scale gives you a better sample of the entire range than searching on a linear scale would. This is because searching on a linear scale would, on average, devote 90 percent of your training budget to only the values between .1 and 1.0. As a result, that leaves only 10 percent of your training budget for the values between .0001 and .1.

ReverseLogarithmic

Hyperparameter tuning searches the values in the hyperparameter range by using a reverse logarithmic scale. Reverse logarithmic scaling is supported only for continuous hyperparameter ranges. It is not supported for integer hyperparameter ranges.

Choose reverse logarithmic scaling when you are searching a range that is highly sensitive to small changes that are very close to 1.

Reverse logarithmic scaling works only for ranges that are entirely within the range 0<=x<1.0.

For an example notebook that uses hyperparameter scaling, see these Amazon SageMaker hyperparameter examples on GitHub.

**Track and set completion criteria for your tuning job**

You can use completion criteria to instruct Automatic model tuning (AMT) to stop your tuning job if certain conditions are met. With these conditions, you can set a minimum model performance or maximum number of training jobs that don't improve when evaluated against the objective metric. You can also track the progress of your tuning job and decide to let it continue or to stop it
manually. This guide shows you how to set completion criteria, check the progress of and stop your tuning job manually.

**Set completion criteria for your tuning job**

During hyperparameter optimization, a tuning job will launch several training jobs inside a loop. The tuning job will do the following.

- Check your training jobs for completion and update statistics accordingly
- Decide what combination of hyperparameters to evaluate next.

AMT will continuously check the training jobs that were launched from your tuning job to update statistics. These statistics include tuning job runtime and best training job. Then, AMT determines whether it should stop the job according to your completion criteria. You can also check these statistics and stop your job manually. For more information about stopping a job manually, see the [Stopping your tuning job manually](#) section.

As an example, if your tuning job meets your objective, you can stop tuning early to conserve resources or ensure model quality. AMT checks your job performance against your completion criteria and stops the tuning job if any have been met.

You can specify the following kinds of completion criteria:

- **MaxNumberOfTrainingJobs** – The maximum number of training jobs to be run before tuning is stopped.
- **MaxNumberOfTrainingJobsNotImproving** – The maximum number of training jobs that do not improve performance against the objective metric from the current best training job. As an example, if the best training job returned an objective metric that had an accuracy of 90%, and **MaxNumberOfTrainingJobsNotImproving** is set to 10. In this example, tuning will stop after 10 training jobs fail to return an accuracy higher than 90%.
- **MaxRuntimeInSeconds** – The upper limit of wall clock time in seconds of how long a tuning job can run.
- **TargetObjectiveMetricValue** – The value of the objective metric against which the tuning job is evaluated. Once this value is met, AMT stops the tuning job.
- **CompleteOnConvergence** – A flag to stop tuning after an internal algorithm determines that the tuning job is unlikely to improve more than 1% over the objective metric from the best training job.
Selecting completion criteria

You can choose one or multiple completion criteria to stop your hyperparameter tuning job after a condition has been meet. The following instructions show you how to select completion criteria and how to decide which is the most appropriate for your use case.

- Use `MaxNumberOfTrainingJobs` in the `ResourceLimits` API to set an upper limit for the number of training jobs that can be run before your tuning job is stopped. Start with a large number and adjust it based on model performance against your tuning job objective. Most users input values of around 50 or more training jobs to find an optimal hyperparameter configuration. Users looking for higher levels of model performance will use 200 or more training jobs.

- Use `MaxNumberOfTrainingJobsNotImproving` in the `BestObjectiveNotImproving` API field to stop training if model performance fails to improve after a specified number of jobs. Model performance is evaluated against an objective function. After the `MaxNumberOfTrainingJobsNotImproving` is met, AMT will stop the tuning job. Tuning jobs tend to make the most progress in the beginning of the job. Improving model performance against an objective function will require a larger number of training jobs towards the end of tuning. Select a value for `MaxNumberOfTrainingJobsNotImproving` by checking the performance of similar training jobs against your objective metric.

- Use `MaxRuntimeInSeconds` in the `ResourceLimits` API to set an upper limit for the amount of wall clock time that the tuning job may take. Use this field to meet a deadline by which the tuning job must complete or to limit compute resources.

To get an estimated total compute time in seconds for a tuning job, use the following formula:

\[
\text{Estimated max compute time in seconds} = \text{MaxRuntimeInSeconds} \times \text{MaxParallelTrainingJobs} \times \text{MaxInstancesPerTrainingJob}
\]

**Note**

The actual duration of a tuning job may deviate slightly from the value specified in this field.

- Use `TargetObjectiveMetricValue` in the `TuningJobCompletionCriteria` API to stop your tuning job. You stop the tuning job after any training job that is launched by the tuning job reaches this objective metric value. Use this field if your use case depends on reaching a specific performance level, rather than spending compute resources to find the best possible model.
• Use CompleteOnConvergence in the TuningJobCompletionCriteria API to stop a tuning job after AMT has detected that the tuning job has converged, and is unlikely to make further significant progress. Use this field when it is not clear what values for any of the other completion criteria should be used. AMT determines convergence based on an algorithm developed and tested on a wide range of diverse benchmarks. A tuning job is defined to have converged when none of the training jobs return significant improvement (1% or less). Improvement is measured against the objective metric returned by the highest performing job, so far.

**Combining different completion criteria**

You can also combine any of the different completion criteria in the same tuning job. AMT will stop the tuning job when any one of the completion criteria is met. For example, if you want to tune your model until it meets an objective metric, but don't want to keep tuning if your job has converged, use the following guidance.

• Specify TargetObjectiveMetricValue in the TuningJobCompletionCriteria API to set a target objective metrics value to reach.
• Set CompleteOnConvergence to Enabled to stop a tuning job if AMT has determined that model performance is unlikely to improve.

**Track tuning job progress**

You can use the DescribeHyperParameterTuningJob API to track the progress of your tuning job at any time while it is running. You don't have to specify completion criteria to obtain tracking information for your tuning job. Use the following fields to obtain statistics about your tuning job.

• BestTrainingJob – An object that describes the best training job obtained so far, evaluated against your objective metric. Use this field to check your current model performance and the value of the objective metric of this best training job.
• ObjectiveStatusCounters – An object that specifies the total number of training jobs completed in a tuning job. To estimate average duration of a tuning job, use ObjectiveStatusCounters and the total runtime of a tuning job. You can use the average duration to estimate how much longer your tuning job will run.
• ConsumedResources – The total resources, such as RunTimeInSeconds, consumed by your tuning job. Compare ConsumedResources, found in the DescribeHyperParameterTuningJob API, against BestTrainingJob in the same API. You can also compare ConsumedResources
against the response from the `ListTrainingJobsForHyperParameterTuningJob` API to assess if your tuning job is making satisfactory progress given the resources being consumed.

- **TuningJobCompletionDetails** – Tuning job completion information that includes the following:
  - The timestamp of when convergence is detected if the job has converged.
  - The number of training jobs that have not improved model performance. Model performance is evaluated against the objective metric from the best training job.

Use the tuning job completion criteria to assess how likely your tuning job is to improve your model performance. Model performance is evaluated against the best objective metric if it ran to completion.

**Stopping your tuning job manually**

You can determine if you should let the tuning job run until it completes or if you should stop the tuning job manually. To determine this, use the information returned by the parameters in the `DescribeHyperParameterTuningJob` API, as shown in the previous Tracking tuning job progress section. As an example, if your model performance does not improve after several training jobs complete, you may choose to stop the tuning job. Model performance is evaluated against the best objective metric.

To stop the tuning job manually, use the `StopHyperParameterTuningJob` API and provide the name of the tuning job to be stopped.

**Tune Multiple Algorithms with Hyperparameter Optimization to Find the Best Model**

To create a new hyperparameter optimization (HPO) job with Amazon SageMaker that tunes multiple algorithms, you must provide job settings that apply to all of the algorithms to be tested and a training definition for each of these algorithms. You must also specify the resources you want to use for the tuning job.

- The **job settings** to configure include warm starting, early stopping, and the tuning strategy. Warm starting and early stopping are available only when tuning a single algorithm.
- The **training job definition** to specify the name, algorithm source, objective metric, and the range of values, when required, to configure the set of hyperparameter values for each training job. It configures the channels for data inputs, data output locations, and any checkpoint storage locations for each training job. The definition also configures the resources to deploy...
for each training job, including instance types and counts, managed spot training, and stopping conditions.

- The **tuning job resources**: to deploy, including the maximum number of concurrent training jobs that a hyperparameter tuning job can run concurrently and the maximum number of training jobs that the hyperparameter tuning job can run.

**Get Started**

You can create a new hyperparameter tuning job, clone a job, add, or edit tags to a job from the console. You can also use the search feature to find jobs by their name, creation time, or status. Alternatively, you can also hyperparameter tuning jobs with the SageMaker API.

- **In the console**: To create a new job, open the Amazon SageMaker console at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/), choose **Hyperparameter tuning jobs** from the **Training** menu, and then choose **Create hyperparameter tuning job**. Then following the configuration steps to create a training job for each algorithm that you want to use. These steps are documented in the [Create a Hyperparameter Optimization Tuning Job for One or More Algorithms (Console)](https://console.aws.amazon.com/sagemaker/) topic.

⚠️ **Note**

When you start the configuration steps, note that the warm start and early stopping features are not available to use with multi-algorithm HPO. If you want to use these features, you can only tune a single algorithm at a time.

- **With the API**: For instructions on using the SageMaker API to create a hyperparameter tuning job, see [Example: Hyperparameter Tuning Job](https://console.aws.amazon.com/sagemaker/). When you call `CreateHyperParameterTuningJob` to tune multiple algorithms, you must provide a list of training definitions using `TrainingJobDefinitions` instead of specifying a single `TrainingJobDefinition`. You must provide job settings that apply to all of the algorithms to be tested and a training definition for each of these algorithms. You must also specify the resources that you want to use for the tuning job. Choose only one of these definition types depending on the number of algorithms that are being tuned.

**Topics**

- [Create a Hyperparameter Optimization Tuning Job for One or More Algorithms (Console)](https://console.aws.amazon.com/sagemaker/)
Create a Hyperparameter Optimization Tuning Job for One or More Algorithms (Console)

This guide shows you how to create a new hyperparameter optimization (HPO) tuning job for one or more algorithms. To create an HPO job, define the settings for the tuning job, and create training job definitions for each algorithm being tuned. Next, configure the resources for and create the tuning job. The following sections provide details about how to complete each step. We provide an example of how to tune multiple algorithms using the SageMaker SDK for Python client at the end of this guide.

Components of a tuning job

An HPO tuning job contains the following three components:

- Tuning job settings
- Training job definitions
- Tuning job configuration

The way that these components are included in your HPO tuning job depends on whether your tuning job contains one or multiple training algorithms. The following guide describes each of the components and gives an example of both types of tuning jobs.

Tuning job settings

Your tuning job settings are applied across all of the algorithms in the HPO tuning job. Warm start and early stopping are available only when you're tuning a single algorithm. After you define the job settings, you can create individual training definitions for each algorithm or variation that you want to tune.

Warm start

If you cloned this job, you can use the results from a previous tuning job to improve the performance of this new tuning job. This is the warm start feature, and it's only available when tuning a single algorithm. With the warm start option, you can choose up to five previous hyperparameter tuning jobs to use. Alternatively, you can use transfer learning to add additional data to the parent tuning job. When you select this option, you choose one previous tuning job as the parent.
Early stopping

To reduce compute time and avoid overfitting your model, you can stop training jobs early. Early stopping is helpful when the training job is unlikely to improve the current best objective metric of the hyperparameter tuning job. Like warm start, this feature is only available when tuning a single algorithm. This is an automatic feature without configuration options, and it's disabled by default. For more information about how early stopping works, the algorithms that support it, and how to use it with your own algorithms, see Stop Training Jobs Early.

Tuning strategy

Tuning strategy can be either random, Bayesian, or Hyperband. These selections specify how automatic tuning algorithms search specified hyperparameter ranges that are selected in a later step. Random search chooses random combinations of values from the specified ranges and can be run sequentially or in parallel. Bayesian optimization chooses values based on what is likely to get the best result according to the known history of previous selections. Hyperband uses a multi-fidelity strategy that dynamically allocates resources toward well-utilized jobs and automatically stops those that underperform. The new configuration that starts after stopping other configurations is chosen randomly.

Hyperband can only be used with iterative algorithms, or algorithms that run steps in iterations, such as XGBoost or Random Cut Forest. Hyperband can't be used with non-iterative algorithms, such as decision trees or k-Nearest Neighbors. For more information about search strategies, see How Hyperparameter Tuning Works.

Note

Hyperband uses an advanced internal mechanism to apply early stopping. Therefore, when you use the Hyperband internal early stopping feature, the parameter TrainingJobEarlyStoppingType in the HyperParameterTuningJobConfig API must be set to OFF.
Tags

To help you manage tuning jobs, you can enter tags as key-value pairs to assign metadata to tuning jobs. Values in the key-value pair are not required. You can use the key without values. To see the keys associated with a job, choose the Tags tab on the details page for tuning job. For more information about using tags for tuning jobs, see Manage Hyperparameter Tuning and Training Jobs.

Training job definitions

To create a training job definition, you must configure the algorithm and parameters, define the data input and output, and configure resources. Provide at least one TrainingJobDefinition for each HPO tuning job. Each training definition specifies the configuration for an algorithm.

To create several definitions for your training job, you can clone a job definition. Cloning a job can save time because it copies all of the job settings, including data channels and Amazon S3 storage locations for output artifacts. You can edit a cloned job to change what you need for your use case.

Topics

- Configure algorithm and parameters
- Define data input and output
- Configure training job resources
- Add or clone a training job

Configure algorithm and parameters

The following list describes what you need to configure the set of hyperparameter values for each training job.

- A name for your tuning job
- Permission to access services
- Parameters for any algorithm options
- An objective metric
- The range of hyperparameter values, when required

Name
Provide a unique name for the training definition.

**Permissions**

Amazon SageMaker requires permissions to call other services on your behalf. Choose an AWS Identity and Access Management (IAM) role, or let AWS create a role with the AmazonSageMakerFullAccess IAM policy attached.

**Optional security settings**

The network isolation setting prevents the container from making any outbound network calls. This is required for AWS Marketplace machine learning offerings.

You can also choose to use a virtual private cloud (VPC).

ℹ️ **Note**

Inter-container encryption is only available when you create a job definition from the API.

**Algorithm options**

You can choose built-in algorithms, your own algorithm, your own container with an algorithm, or you can subscribe to an algorithm from AWS Marketplace.

- If you choose a built-in algorithm, it has the Amazon Elastic Container Registry (Amazon ECR) image information pre-populated.
- If you choose your own container, you must specify the (Amazon ECR) image information. You can select the input mode for the algorithm as file or pipe.
- If you plan to supply your data using a CSV file from Amazon S3, you should select the file.

**Metrics**

When you choose a built-in algorithm, metrics are provided for you. If you choose your own algorithm, you must define your metrics. You can define up to 20 metrics for your tuning job to monitor. You must choose one metric as the objective metric. For more information about how to define a metric for a tuning job, see [Define metrics](#).

**Objective metric**
To find the best training job, set an objective metric and whether to maximize or minimize it. After the training job is complete, you can view the tuning job detail page. The detail page provides a summary of the best training job that is found using this objective metric.

**Hyperparameter configuration**

When you choose a built-in algorithm, the default values for its hyperparameters are set for you, using ranges that are optimized for the algorithm that's being tuned. You can change these values as you see fit. For example, instead of a range, you can set a fixed value for a hyperparameter by setting the parameter's type to `static`. Each algorithm has different required and optional parameters. For more information, see [Best Practices for Hyperparameter Tuning](#) and [Define Hyperparameter Ranges](#).

**Define data input and output**

Each training job definition for a tuning job must configure the channels for data inputs, data output locations, and optionally, any checkpoint storage locations for each training job.

**Input data configuration**

Input data is defined by channels. Each channel its own source location (Amazon S3 or Amazon Elastic File System), compression, and format options. You can define up to 20 channels of input sources. If the algorithm that you choose supports multiple input channels, you can specify those, too. For example, when you use the [XGBoost churn prediction notebook](#), you can add two channels: train and validation.

**Checkpoint configuration**

Checkpoints are periodically generated during training. For the checkpoints to be saved, you must choose an Amazon S3 location. Checkpoints are used in metrics reporting, and are also used to resume managed spot training jobs. For more information, see [Use checkpoints in Amazon SageMaker](#).

**Output data configuration**

Define an Amazon S3 location for the artifacts of the training job to be stored. You have the option of adding encryption to the output using an AWS Key Management Service (AWS KMS) key.

**Configure training job resources**

Each training job definition for a tuning job must configure the resources to deploy, including instance types and counts, managed spot training, and stopping conditions.
Resource configuration

Each training definition can have a different resource configuration. You choose the instance type and number of nodes.

Managed spot training

You can save computer costs for jobs if you have flexibility in start and end times by allowing SageMaker to use spare capacity to run jobs. For more information, see Use Managed Spot Training in Amazon SageMaker.

Stopping condition

The stopping condition specifies the maximum duration that's allowed for each training job.

Add or clone a training job

After you create a training job definition for a tuning job, you will return to the Training Job Definition(s) panel. This panel is where you can create additional training job definitions to train additional algorithms. You can select the Add training job definition and work through the steps to define a training job again.

Alternatively, to replicate an existing training job definition and edit it for the new algorithm, choose Clone from the Action menu. The clone option can save time because it copies all of the job’s settings, including the data channels and Amazon S3 storage locations. For more information about cloning, see Manage Hyperparameter Tuning and Training Jobs.

Tuning job configuration

Resource Limits

You can specify the maximum number of concurrent training jobs that a hyperparameter tuning job can run concurrently (10 at most). You can also specify the maximum number of training jobs that the hyperparameter tuning job can run (500 at most). The number of parallel jobs should not exceed the number of nodes that you have requested across all of your training definitions. The total number of jobs can’t exceed the number of jobs that your definitions are expected to run.

Review the job settings, the training job definitions, and the resource limits. Then select Create hyperparameter tuning job.
HPO tuning job example

To run a hyperparameter optimization (HPO) training job, first create a training job definition for each algorithm that's being tuned. Next, define the tuning job settings and configure the resources for the tuning job. Finally, run the tuning job.

If your HPO tuning job contains a single training algorithm, the SageMaker tuning function will call the HyperparameterTuner API directly and pass in your parameters. If your HPO tuning job contains multiple training algorithms, your tuning function will call the create function of the HyperparameterTuner API. The create function tells the API to expect a dictionary containing one or more estimators.

In the following section, code examples show how to tune a job containing either a single training algorithm or multiple algorithms using the SageMaker Python SDK.

Create training job definitions

When you create a tuning job that includes multiple training algorithms, your tuning job configuration will include the estimators and metrics and other parameters for your training jobs. Therefore, you need to create the training job definition first, and then configure your tuning job.

The following code example shows how to retrieve two SageMaker containers containing the built-in algorithms XGBoost and Linear Learner. If your tuning job contains only one training algorithm, omit one of the containers and one of the estimators.

```python
import sagemaker
from sagemaker import image_uris
from sagemaker.estimator import Estimator

ts = sagemaker.Session()
region = sess.boto_region_name
role = sagemaker.get_execution_role()

bucket = sess.default_bucket()
prefix = "sagemaker/multi-algo-hpo"

# Define the training containers and initialize the estimators
xgb_container = image_uris.retrieve("xgboost", region, "latest")
ll_container = image_uris.retrieve("linear-learner", region, "latest")

xgb_estimator = Estimator(
```
xgb_container,
role=role,
inference_count=1,
inference_type="ml.m4.xlarge",
output_path='s3://{}/{}xgb_output'.format(bucket, prefix),
sagemaker_session=sess,
)

ll_estimator = Estimator(
  ll_container,
  role,
inference_count=1,
inference_type="ml.c4.xlarge",
output_path='s3://{}/{}ll_output'.format(bucket, prefix),
sagemaker_session=sess,
)

# Set static hyperparameters
ll_estimator.set_hyperparameters(predictor_type="binary_classifier")
xgb_estimator.set_hyperparameters(
  eval_metric="auc",
  objective="binary:logistic",
  num_round=100,
  rate_drop=0.3,
  tweedie_variance_power=1.4,
)

Next, define your input data by specifying the training, validation, and testing datasets, as shown in the following code example. This example shows how to tune multiple training algorithms.


training_data = sagemaker.inputs.TrainingInput(
  s3_data="s3://{}//{}train".format(bucket, prefix), content_type="csv"
)
validation_data = sagemaker.inputs.TrainingInput(
  s3_data="s3://{}//{}validate".format(bucket, prefix), content_type="csv"
)
test_data = sagemaker.inputs.TrainingInput(
  s3_data="s3://{}//{}test".format(bucket, prefix), content_type="csv"
)

train_inputs = {
  "estimator-1": {
    "train": training_data,
    "validate": validation_data,
    "test": test_data,
  },
If your tuning algorithm contains only one training algorithm, your train_inputs should contain only one estimator.

You must upload the inputs for the training, validation, and training datasets to your Amazon S3 bucket before you use those in an HPO tuning job.

**Define resources and settings for your tuning job**

This section shows how to initialize a tuner, define resources, and specify job settings for your tuning job. If your tuning job contains multiple training algorithms, these settings are applied to all of the algorithms that are contained inside your tuning job. This section provides two code examples to define a tuner. The code examples show you how to optimize a single training algorithm followed by an example of how to tune multiple training algorithms.

**Tune a single training algorithm**

The following code example shows how to initialize a tuner and set hyperparameter ranges for one SageMaker built-in algorithm, XGBoost.

```python
from sagemaker.tuner import HyperparameterTuner
from sagemaker.parameter import ContinuousParameter, IntegerParameter

hyperparameter_ranges = {
    "max_depth": IntegerParameter(1, 10),
    "eta": ContinuousParameter(0.1, 0.3),
}

objective_metric_name = "validation:accuracy"

tuner = HyperparameterTuner(
    xgb_estimator,
    objective_metric_name,
    hyperparameter_ranges,
)```
Tune multiple training algorithms

Each training job requires different configurations, and these are specified using a dictionary. The following code example shows how to initialize a tuner with configurations for two SageMaker built-in algorithms, XGBoost and Linear Learner. The code example also shows how to set a tuning strategy and other job settings, such as the compute resources for the tuning job. The following code example uses `metric_definitions_dict`, which is optional.

```python
from sagemaker.tuner import HyperparameterTuner
from sagemaker.parameter import ContinuousParameter, IntegerParameter

# Initialize your tuner
tuner = HyperparameterTuner.create(
    estimator_dict={
        "estimator-1": xgb_estimator,
        "estimator-2": ll_estimator,
    },
    objective_metric_name_dict={
        "estimator-1": "validation:auc",
        "estimator-2": "test:binary_classification_accuracy",
    },
    hyperparameter_ranges_dict={
        "estimator-1": {"eta": ContinuousParameter(0.1, 0.3)},
        "estimator-2": {"learning_rate": ContinuousParameter(0.1, 0.3)},
    },
    metric_definitions_dict={
        "estimator-1": [
            {"Name": "validation:auc", "Regex": "Overall test accuracy: (.*)\.;"}
        ],
        "estimator-2": [
            {
                "Name": "test:binary_classification_accuracy",
                "Regex": "Overall test accuracy: (.*)\.;",
            }
        ],
    },
    strategy="Bayesian",
    max_jobs=10,
)
```
Run your HPO tuning job

Now you can run your tuning job by passing your training inputs to the `fit` function of the `HyperparameterTuner` class. The following code example shows how to pass the `train_inputs` parameter, that is defined in a previous code example, to your tuner.

```python
max_parallel_jobs=3,
)

Run your HPO tuning job

```
tuner.fit(inputs=train_inputs, include_cls_metadata ={}, estimator_kwargs ={})
```

Manage Hyperparameter Tuning and Training Jobs

A tuning job can contain many training jobs and creating and managing these jobs and their definitions can become a complex and onerous task. SageMaker provides tools to help facilitate the management of these jobs. Tuning jobs you have run can be accessed from the Amazon SageMaker console at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/). Select Hyperparameter tuning job from the Training menu to see the list. This page is also where you start the procedure to create a new tuning job by selecting Create hyperparameter tuning job.

To see the training jobs run a part of a tuning job, select one of the hyperparameter tuning jobs from the list. The tabs on the tuning job page allow you to inspect the training jobs, their definitions, the tags and configuration used for the tuning job, and the best training job found during tuning. You can select the best training job or any of the other training jobs that belong to the tuning job to see all of their settings. From here you can create a model that uses the hyperparameter values found by a training job by selecting Create Model or you can clone the training job by selecting Clone.

Cloning

You can save time by cloning a training job that belongs to a hyperparameter tuning job. Cloning copies all of the job's settings, including data channels, S3 storage locations for output artifacts. You can do this for training jobs you have already run from the tuning job page, as just described, or when you are creating additional training job definitions while creating a hyperparameter tuning job, as described in Add or clone a training job step of that procedure.

Tagging

Automatic Model Tuning launches multiple training jobs within a single parent tuning job to discover the ideal weighting of model hyperparameters. Tags can be added to the parent tuning
job as described in the Components of a tuning job section and these tags are then propagated to the individual training jobs underneath. Customers can use these tags for purposes, such as cost allocation or access control. To add tags using the SageMaker SDK, use AddTags API. For more information about using tagging for AWS resources, see Tagging AWS resources.

Example: Hyperparameter Tuning Job

This example shows how to create a new notebook for configuring and launching a hyperparameter tuning job. The tuning job uses the XGBoost Algorithm to train a model to predict whether a customer will enroll for a term deposit at a bank after being contacted by phone.

You use the low-level SDK for Python (Boto3) to configure and launch the hyperparameter tuning job, and the AWS Management Console to monitor the status of hyperparameter tuning jobs. You can also use the Amazon SageMaker high-level Amazon SageMaker Python SDK to configure, run, monitor, and analyze hyperparameter tuning jobs. For more information, see https://github.com/aws/sagemaker-python-sdk.

Prerequisites

To run the code in this example, you need

- An AWS account and an administrator user
- An Amazon S3 bucket for storing your training dataset and the model artifacts created during training
- A running SageMaker notebook instance

Topics

- Create a Notebook Instance
- Get the Amazon SageMaker Boto 3 Client
- Get the SageMaker Execution Role
- Use an Amazon S3 bucket for input and output
- Download, Prepare, and Upload Training Data
- Configure and Launch a Hyperparameter Tuning Job
- Clean up
Create a Notebook Instance

Create a Jupyter notebook that contains a pre-installed environment with the default Anaconda installation and Python3.

To create a Jupyter notebook

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Open a running notebook instance, by choosing Open next to its name. The Jupyter notebook server page appears:

3. To create a notebook, choose Files, New, and conda_python3.
4. Name the notebook.

Next Step

Get the Amazon SageMaker Boto 3 Client

Get the Amazon SageMaker Boto 3 Client

Import Amazon SageMaker Python SDK, AWS SDK for Python (Boto3), and other Python libraries. In a new Jupyter notebook, paste the following code to the first cell:

```python
import sagemaker
import boto3

import numpy as np
    # For performing matrix operations
import pandas as pd
    # For manipulating tabular data
from time import gmtime, strftime
import os

region = boto3.Session().region_name
```
The preceding code cell defines `region` and `smclient` objects that you will use to call the built-in XGBoost algorithm and set the SageMaker hyperparameter tuning job.

**Next Step**

**Get the SageMaker Execution Role**

Get the execution role for the notebook instance. This is the IAM role that you created for your notebook instance.

To find the ARN of the IAM execution role attached to a notebook instance:

2. On the left navigation pane, choose **Notebook** then **Notebook instances**.
3. From the list of notebooks, select the notebook that you want to view.
4. The ARN is in the **Permissions and encryption** section.

Alternatively, [Amazon SageMaker Python SDK](https://boto3.amazonaws.com/v1/documentation/api/latest/reference/services/sagemaker.html) users can retrieve the ARN of the execution role attached to their user profile or a notebook instance by running the following code:

```python
from sagemaker import get_execution_role

role = get_execution_role()
print(role)
```


**Next Step**

**Use an Amazon S3 bucket for input and output**

Set up a S3 bucket to upload training datasets and save training output data for your hyperparameter tuning job.
To use a default S3 bucket

Use the following code to specify the default S3 bucket allocated for your SageMaker session. `prefix` is the path within the bucket where SageMaker stores the data for the current training job.

```python
sess = sagemaker.Session()
bucket = sess.default_bucket()  # Set a default S3 bucket
prefix = 'DEMO-automatic-model-tuning-xgboost-dm'
```

To use a specific S3 bucket (Optional)

If you want to use a specific S3 bucket, use the following code and replace the strings to the exact name of the S3 bucket. The name of the bucket must contain `sagemaker`, and be globally unique. The bucket must be in the same AWS Region as the notebook instance that you use for this example.

```python
bucket = "sagemaker-your-preferred-s3-bucket"

sess = sagemaker.Session(
    default_bucket = bucket
)
```

Note

The name of the bucket doesn't need to contain `sagemaker` if the IAM role that you use to run the hyperparameter tuning job has a policy that gives the `S3FullAccess` permission.

Next Step

Download, Prepare, and Upload Training Data

Download, Prepare, and Upload Training Data

For this example, you use a training dataset of information about bank customers that includes the customer's job, marital status, and how they were contacted during the bank's direct marketing campaign. To use a dataset for a hyperparameter tuning job, you download it, transform the data, and then upload it to an Amazon S3 bucket.
For more information about the dataset and the data transformation that the example performs, see the `hpo_xgboost_direct_marketing_sagemaker_APIs` notebook in the **Hyperparameter Tuning** section of the **SageMaker Examples** tab in your notebook instance.

### Download and Explore the Training Dataset

To download and explore the dataset, run the following code in your notebook:

```bash
!unzip -o bank-additional.zip
data = pd.read_csv('./bank-additional/bank-additional-full.csv', sep=';')
pd.set_option('display.max_columns', 500)  # Make sure we can see all of the columns
pd.set_option('display.max_rows', 5)      # Keep the output on one page
data
```

### Prepare and Upload Data

Before creating the hyperparameter tuning job, prepare the data and upload it to an S3 bucket where the hyperparameter tuning job can access it.

Run the following code in your notebook:

```python
data['no_previous_contact'] = np.where(data['pdays'] == 999, 1, 0)  # Indicator variable to capture when pdays takes a value of 999
data['not_working'] = np.where(np.in1d(data['job'], ['student', 'retired', 'unemployed']), 1, 0)  # Indicator for individuals not actively employed
model_data = pd.get_dummies(data)  # Convert categorical variables to sets of indicators
model_data = model_data.drop(['duration', 'emp.var.rate', 'cons.price.idx', 'cons.conf.idx', 'euribor3m', 'nr.employed'], axis=1)
train_data, validation_data, test_data = np.split(model_data.sample(frac=1, random_state=1729), [int(0.7 * len(model_data)), int(0.9*len(model_data))])
pd.concat([train_data['y_yes'], train_data.drop(['y_no', 'y_yes'], axis=1)]).to_csv('train.csv', index=False, header=False)
pd.concat([validation_data['y_yes'], validation_data.drop(['y_no', 'y_yes'], axis=1)]).to_csv('validation.csv', index=False, header=False)
pd.concat([test_data['y_yes'], test_data.drop(['y_no', 'y_yes'], axis=1)]).to_csv('test.csv', index=False, header=False)
```
Next Step

Configure and Launch a Hyperparameter Tuning Job

Configure and Launch a Hyperparameter Tuning Job

A hyperparameter is a high-level parameter that influences the learning process during model training. To get the best model predictions, you can optimize a hyperparameter configuration or set hyperparameter values. The process of finding an optimal configuration is called hyperparameter tuning. To configure and launch a hyperparameter tuning job, complete the steps in these guides.

Topics

- Settings for the hyperparameter tuning job
- Configure the training jobs
- Name and launch the hyperparameter tuning job
- Monitor the Progress of a Hyperparameter Tuning Job
- View the Status of the Training Jobs
- View the Best Training Job

Settings for the hyperparameter tuning job

To specify settings for the hyperparameter tuning job, define a JSON object when you create the tuning job. Pass this JSON object as the value of the HyperParameterTuningJobConfig parameter to the CreateHyperParameterTuningJob API.

In this JSON object, specify the following:

In this JSON object, you specify:

- HyperParameterTuningJobObjective – The objective metric used to evaluate the performance of the training job launched by the hyperparameter tuning job.
- ParameterRanges – The range of values that a tunable hyperparameter can use during optimization. For more information, see Define Hyperparameter Ranges
- **RandomSeed** – A value used to initialize a pseudo-random number generator. Setting a random seed will allow the hyperparameter tuning search strategies to produce more consistent configurations for the same tuning job (optional).

- **ResourceLimits** – The maximum number of training and parallel training jobs that the hyperparameter tuning job can use.

**Note**

If you use your own algorithm for hyperparameter tuning, rather than a SageMaker built-in algorithm, you must define metrics for your algorithm. For more information, see [Define metrics](#).

The following code example shows how to configure a hyperparameter tuning job using the built-in XGBoost algorithm. The code example shows how to define ranges for the eta, alpha, min_child_weight, and max_depth hyperparameters. For more information about these and other hyperparameters see [XGBoost Parameters](#).

In this code example, the objective metric for the hyperparameter tuning job finds the hyperparameter configuration that maximizes `validation:auc`. SageMaker built-in algorithms automatically write the objective metric to CloudWatch Logs. The following code example also shows how to set a RandomSeed.

```python
tuning_job_config = {
    "ParameterRanges": {
        "CategoricalParameterRanges": [],
        "ContinuousParameterRanges": [
            {
                "MaxValue": "1",
                "MinValue": "0",
                "Name": "eta"
            },
            {
                "MaxValue": "2",
                "MinValue": "0",
                "Name": "alpha"
            },
            {
                "MaxValue": "10",
                "MinValue": "0",
                "Name": "max_depth"
            }
        ]
    }
}
```

Example: Hyperparameter Tuning Job
Configure the training jobs

The hyperparameter tuning job will launch training jobs to find an optimal configuration of hyperparameters. These training jobs should be configured using the SageMaker CreateHyperParameterTuningJob API.

To configure the training jobs, define a JSON object and pass it as the value of the TrainingJobDefinition parameter inside CreateHyperParameterTuningJob.

In this JSON object, you can specify the following:

- **AlgorithmSpecification** – The registry path of the Docker image containing the training algorithm and related metadata. To specify an algorithm, you can use your own custom built algorithm inside a Docker container or a SageMaker built-in algorithm (required).
- **InputDataConfig** – The input configuration, including the ChannelName, ContentType, and data source for your training and test data (required).
- **InputDataConfig** – The input configuration, including the ChannelName, ContentType, and data source for your training and test data (required).
• The storage location for the algorithm's output. Specify the S3 bucket where you want to store the output of the training jobs.

• RoleArn – The Amazon Resource Name (ARN) of an AWS Identity and Access Management (IAM) role that SageMaker uses to perform tasks. Tasks include reading input data, downloading a Docker image, writing model artifacts to an S3 bucket, writing logs to Amazon CloudWatch Logs, and writing metrics to Amazon CloudWatch (required).

• StoppingCondition – The maximum runtime in seconds that a training job can run before being stopped. This value should be greater than the time needed to train your model (required).

• MetricDefinitions – The name and regular expression that defines any metrics that the training jobs emit. Define metrics only when you use a custom training algorithm. The example in the following code uses a built-in algorithm, which already has metrics defined. For information about defining metrics (optional), see Define metrics.

• TrainingImage – The Docker container image that specifies the training algorithm (optional).

• StaticHyperParameters – The name and values of hyperparameters that are not tuned in the tuning job (optional).

The following code example sets static values for the eval_metric, num_round, objective, rate_drop, and tweedie_variance_power parameters of the XGBoost Algorithm built-in algorithm.

SageMaker Python SDK v1

```python
from sagemaker.amazon.amazon_estimator import get_image_uri
training_image = get_image_uri(region, 'xgboost', repo_version='1.0-1')

s3_input_train = 's3://{}/{}//train'.format(bucket, prefix)
s3_input_validation = 's3://{}/{}//validation/'.format(bucket, prefix)

training_job_definition = {
    "AlgorithmSpecification": {
        "TrainingImage": training_image,
        "TrainingInputMode": "File"
    },
    "InputDataConfig": [
        {
            "ChannelName": "train",
            "CompressionType": "None",
            "ContentType": "csv"
        }
    ]
}
```
"DataSource": {
    "S3DataSource": {
        "S3DataDistributionType": "FullyReplicated",
        "S3DataType": "S3Prefix",
        "S3Uri": s3_input_train
    }
},
{
    "ChannelName": "validation",
    "CompressionType": "None",
    "ContentType": "csv",
    "DataSource": {
        "S3DataSource": {
            "S3DataDistributionType": "FullyReplicated",
            "S3DataType": "S3Prefix",
            "S3Uri": s3_input_validation
        }
    }
},
"OutputDataConfig": {
    "S3OutputPath": "s3://{}/{}/output".format(bucket,prefix)
},
"ResourceConfig": {
    "InstanceCount": 2,
    "InstanceType": "ml.c4.2xlarge",
    "VolumeSizeInGB": 10
},
"RoleArn": role,
"StaticHyperParameters": {
    "eval_metric": "auc",
    "num_round": "100",
    "objective": "binary:logistic",
    "rate_drop": "0.3",
    "tweedie_variance_power": "1.4"
},
"StoppingCondition": {
    "MaxRuntimeInSeconds": 43200
}
training_image = sagemaker.image_uris.retrieve('xgboost', region, '1.0-1')

s3_input_train = 's3://{}//{}/train'.format(bucket, prefix)
s3_input_validation = 's3://{}//{}/validation/'.format(bucket, prefix)

training_job_definition = {
    "AlgorithmSpecification": {
        "TrainingImage": training_image,
        "TrainingInputMode": "File"
    },
    "InputDataConfig": [
        {
            "ChannelName": "train",
            "CompressionType": "None",
            "ContentType": "csv",
            "DataSource": {
                "S3DataSource": {
                    "S3DataDistributionType": "FullyReplicated",
                    "S3DataType": "S3Prefix",
                    "S3Uri": s3_input_train
                }
            }
        },
        {
            "ChannelName": "validation",
            "CompressionType": "None",
            "ContentType": "csv",
            "DataSource": {
                "S3DataSource": {
                    "S3DataDistributionType": "FullyReplicated",
                    "S3DataType": "S3Prefix",
                    "S3Uri": s3_input_validation
                }
            }
        }
    ],
    "OutputDataConfig": {
        "S3OutputPath": "s3://{}//{}/output".format(bucket, prefix)
    },
    "ResourceConfig": {
        "InstanceCount": 2,
        "InstanceType": "ml.c4.2xlarge",
    }
}

Example: Hyperparameter Tuning Job
Name and launch the hyperparameter tuning job

After you configure the hyperparameter tuning job, you can launch it by calling the `CreateHyperParameterTuningJob` API. The following code example uses `tuning_job_config` and `training_job_definition`. These were defined in the previous two code examples to create a hyperparameter tuning job.

```python

tuning_job_name = "MyTuningJob"
smclient.create_hyper_parameter_tuning_job(HyperParameterTuningJobName =
tuning_job_name,
    HyperParameterTuningJobConfig =
    tuning_job_config,
    TrainingJobDefinition =
    training_job_definition)
```

Monitor the Progress of a Hyperparameter Tuning Job

To monitor the progress of a hyperparameter tuning job and the training jobs that it launches, use the Amazon SageMaker console.

Topics

- View the Status of the Hyperparameter Tuning Job
View the Status of the Hyperparameter Tuning Job

To view the status of the hyperparameter tuning job

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Hyperparameter tuning jobs.

3. In the list of hyperparameter tuning jobs, check the status of the hyperparameter tuning job you launched. A tuning job can be:
   - Completed—The hyperparameter tuning job successfully completed.
   - InProgress—The hyperparameter tuning job is in progress. One or more training jobs are still running.
   - Failed—The hyperparameter tuning job failed.
   - Stopped—The hyperparameter tuning job was manually stopped before it completed. All training jobs that the hyperparameter tuning job launched are stopped.
   - Stopping—The hyperparameter tuning job is in the process of stopping.

View the Status of the Training Jobs

To view the status of the training jobs that the hyperparameter tuning job launched

1. In the list of hyperparameter tuning jobs, choose the job that you launched.
2. Choose Training jobs.

3. View the status of each training job. To see more details about a job, choose it in the list of training jobs. To view a summary of the status of all of the training jobs that the hyperparameter tuning job launched, see Training job status counter.

A training job can be:

- Completed—The training job successfully completed.
- InProgress—The training job is in progress.
- Stopped—The training job was manually stopped before it completed.
- Failed (Retryable)—The training job failed, but can be retried. A failed training job can be retried only if it failed because an internal service error occurred.
- Failed (Non-retryable)—The training job failed and can't be retried. A failed training job can't be retried when a client error occurs.

**Note**

Hyperparameter tuning jobs can be stopped and the underlying resources **deleted**, but the jobs themselves cannot be deleted.
View the Best Training Job

A hyperparameter tuning job uses the objective metric that each training job returns to evaluate training jobs. While the hyperparameter tuning job is in progress, the best training job is the one that has returned the best objective metric so far. After the hyperparameter tuning job is complete, the best training job is the one that returned the best objective metric.

To view the best training job, choose **Best training job**.

To deploy the best training job as a model that you can host at a SageMaker endpoint, choose **Create model**.

Next Step

Clean up

Clean up

To avoid incurring unnecessary charges, when you are done with the example, use the AWS Management Console to delete the resources that you created for it.

Note

If you plan to explore other examples, you might want to keep some of these resources, such as your notebook instance, S3 bucket, and IAM role.
1. Open the SageMaker console at https://console.aws.amazon.com/sagemaker/ and delete the notebook instance. Stop the instance before deleting it.

2. Open the Amazon S3 console at https://console.aws.amazon.com/s3/ and delete the bucket that you created to store model artifacts and the training dataset.

3. Open the IAM console at https://console.aws.amazon.com/iam/ and delete the IAM role. If you created permission policies, you can delete them, too.

4. Open the Amazon CloudWatch console at https://console.aws.amazon.com/cloudwatch/ and delete all of the log groups that have names starting with /aws/sagemaker/.

Stop Training Jobs Early

Stop the training jobs that a hyperparameter tuning job launches early when they are not improving significantly as measured by the objective metric. Stopping training jobs early can help reduce compute time and helps you avoid overfitting your model. To configure a hyperparameter tuning job to stop training jobs early, do one of the following:

- If you are using the AWS SDK for Python (Boto3), set the TrainingJobEarlyStoppingType field of the HyperParameterTuningJobConfig object that you use to configure the tuning job to AUTO.
- If you are using the Amazon SageMaker Python SDK, set the early_stopping_type parameter of the HyperParameterTuner object to Auto.
- In the Amazon SageMaker console, in the Create hyperparameter tuning job workflow, under Early stopping, choose Auto.

For a sample notebook that demonstrates how to use early stopping, see https://github.com/awslabs/amazon-sagemaker-examples/blob/master/hyperparameter_tuning/image_classification_early_stopping/hpo_image_classification_early_stopping.ipynb or open the hpo_image_classification_early_stopping.ipynb notebook in the Hyperparameter Tuning section of the SageMaker Examples in a notebook instance. For information about using sample notebooks in a notebook instance, see Example Notebooks.

How Early Stopping Works

When you enable early stopping for a hyperparameter tuning job, SageMaker evaluates each training job the hyperparameter tuning job launches as follows:
After each epoch of training, get the value of the objective metric.

Compute the running average of the objective metric for all previous training jobs up to the same epoch, and then compute the median of all of the running averages.

If the value of the objective metric for the current training job is worse (higher when minimizing or lower when maximizing the objective metric) than the median value of running averages of the objective metric for previous training jobs up to the same epoch, SageMaker stops the current training job.

**Algorithms That Support Early Stopping**

To support early stopping, an algorithm must emit objective metrics for each epoch. The following built-in SageMaker algorithms support early stopping:

- **LightGBM**
- **CatBoost**
- **AutoGluon-Tabular**
- **TabTransformer**
- **Linear Learner Algorithm**—Supported only if you use `objective_loss` as the objective metric.
- **XGBoost Algorithm**
- **Image Classification - MXNet**
- **Object Detection - MXNet**
- **Sequence-to-Sequence Algorithm**
- **IP Insights**

**Note**

This list of built-in algorithms that support early stopping is current as of December 13, 2018. Other built-in algorithms might support early stopping in the future. If an algorithm emits a metric that can be used as an objective metric for a hyperparameter tuning job (preferably a validation metric), then it supports early stopping.
To use early stopping with your own algorithm, you must write your algorithms such that it emits the value of the objective metric after each epoch. The following list shows how you can do that in different frameworks:

**TensorFlow**

Use the `tf.keras.callbacks.ProgbarLogger` class. For information, see the `tf.keras.callbacks.ProgbarLogger API`.

**MXNet**

Use the `mxnet.callback.LogValidationMetricsCallback`. For information, see the `mxnet.callback APIs`.

**Chainer**

Extend chainer by using the `extensions.Evaluator` class. For information, see the `chainer.training.extensions.Evaluator API`.

**PyTorch and Spark**

There is no high-level support. You must explicitly write your training code so that it computes objective metrics and writes them to logs after each epoch.

### Run a Warm Start Hyperparameter Tuning Job

Use warm start to start a hyperparameter tuning job using one or more previous tuning jobs as a starting point. The results of previous tuning jobs are used to inform which combinations of hyperparameters to search over in the new tuning job. Hyperparameter tuning uses either Bayesian or random search to choose combinations of hyperparameter values from ranges that you specify. For more information, see [How Hyperparameter Tuning Works](#). Using information from previous hyperparameter tuning jobs can help increase the performance of the new hyperparameter tuning job by making the search for the best combination of hyperparameters more efficient.

**Note**

Warm start tuning jobs typically take longer to start than standard hyperparameter tuning jobs, because the results from the parent jobs have to be loaded before the job can start. The increased time depends on the total number of training jobs launched by the parent jobs.
Reasons to consider warm start include the following:

- To gradually increase the number of training jobs over several tuning jobs based on results after each iteration.
- To tune a model using new data that you received.
- To change hyperparameter ranges that you used in a previous tuning job, change static hyperparameters to tunable, or change tunable hyperparameters to static values.
- You stopped a previous hyperparameter job early or it stopped unexpectedly.

**Topics**

- Types of Warm Start Tuning Jobs
- Warm Start Tuning Restrictions
- Warm Start Tuning Sample Notebook
- Create a Warm Start Tuning Job

**Types of Warm Start Tuning Jobs**

There are two different types of warm start tuning jobs:

**IDENTICAL_DATA_AND_ALGORITHM**

The new hyperparameter tuning job uses the same input data and training image as the parent tuning jobs. You can change the hyperparameter ranges to search and the maximum number of training jobs that the hyperparameter tuning job launches. You can also change hyperparameters from tunable to static, and from static to tunable, but the total number of static plus tunable hyperparameters must remain the same as it is in all parent jobs. You cannot use a new version of the training algorithm, unless the changes in the new version do not affect the algorithm itself. For example, changes that improve logging or adding support for a different data format are allowed.

Use identical data and algorithm when you use the same training data as you used in a previous hyperparameter tuning job, but you want to increase the total number of training jobs or change ranges or values of hyperparameters.

When you run an warm start tuning job of type IDENTICAL_DATA_AND_ALGORITHM, there is an additional field in the response to DescribeHyperParameterTuningJob named OverallBestTrainingJob. The value of this field is the TrainingJobSummary for the training
job with the best objective metric value of all training jobs launched by this tuning job and all parent jobs specified for the warm start tuning job.

TRANSFER_LEARNING

The new hyperparameter tuning job can include input data, hyperparameter ranges, maximum number of concurrent training jobs, and maximum number of training jobs that are different than those of its parent hyperparameter tuning jobs. You can also change hyperparameters from tunable to static, and from static to tunable, but the total number of static plus tunable hyperparameters must remain the same as it is in all parent jobs. The training algorithm image can also be a different version from the version used in the parent hyperparameter tuning job. When you use transfer learning, changes in the dataset or the algorithm that significantly affect the value of the objective metric might reduce the usefulness of using warm start tuning.

Warm Start Tuning Restrictions

The following restrictions apply to all warm start tuning jobs:

- A tuning job can have a maximum of 5 parent jobs, and all parent jobs must be in a terminal state (Completed, Stopped, or Failed) before you start the new tuning job.
- The objective metric used in the new tuning job must be the same as the objective metric used in the parent jobs.
- The total number of static plus tunable hyperparameters must remain the same between parent jobs and the new tuning job. Because of this, if you think you might want to use a hyperparameter as tunable in a future warm start tuning job, you should add it as a static hyperparameter when you create a tuning job.
- The type of each hyperparameter (continuous, integer, categorical) must not change between parent jobs and the new tuning job.
- The number of total changes from tunable hyperparameters in the parent jobs to static hyperparameters in the new tuning job, plus the number of changes in the values of static hyperparameters cannot be more than 10. For example, if the parent job has a tunable categorical hyperparameter with the possible values red and blue, you change that hyperparameter to static in the new tuning job, that counts as 2 changes against the allowed total of 10. If the same hyperparameter had a static value of red in the parent job, and you change the static value to blue in the new tuning job, it also counts as 2 changes.
- Warm start tuning is not recursive. For example, if you create MyTuningJob3 as a warm start tuning job with MyTuningJob2 as a parent job, and MyTuningJob2 is itself an warm start
tuning job with a parent job MyTuningJob1, the information that was learned when running MyTuningJob1 is not used for MyTuningJob3. If you want to use the information from MyTuningJob1, you must explicitly add it as a parent for MyTuningJob3.

- The training jobs launched by every parent job in a warm start tuning job count against the 500 maximum training jobs for a tuning job.

- Hyperparameter tuning jobs created before October 1, 2018 cannot be used as parent jobs for warm start tuning jobs.

**Warm Start Tuning Sample Notebook**

For a sample notebook that shows how to use warm start tuning, see https://github.com/awslabs/amazon-sagemaker-examples/blob/master/hyperparameter_tuning/image_classification_warmstart/hpo_image_classification_warmstart.ipynb. For instructions how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Example Notebooks. Once you have created a notebook instance and opened it, select the SageMaker Examples tab to see a list of all the SageMaker samples. The warm start tuning example notebook is located in the Hyperparameter tuning section, and is named hpo_image_classification_warmstart.ipynb. To open a notebook, click on its Use tab and select Create copy.

**Create a Warm Start Tuning Job**

You can use either the low-level AWS SDK for Python (Boto 3) or the high-level SageMaker Python SDK to create a warm start tuning job.

**Topics**

- Create a Warm Start Tuning Job (Low-level SageMaker API for Python (Boto 3))
- Create a Warm Start Tuning Job (SageMaker Python SDK)

**Create a Warm Start Tuning Job (Low-level SageMaker API for Python (Boto 3))**

To use warm start tuning, you specify the values of a HyperParameterTuningJobWarmStartConfig object, and pass that as the WarmStartConfig field in a call to CreateHyperParameterTuningJob.
The following code shows how to create a `HyperParameterTuningJobWarmStartConfig` object and pass it to `CreateHyperParameterTuningJob` job by using the low-level SageMaker API for Python (Boto 3).

Create the `HyperParameterTuningJobWarmStartConfig` object:

```python
warm_start_config = {
    "ParentHyperParameterTuningJobs" : [
    {"HyperParameterTuningJobName" : 'MyParentTuningJob'}
    ],
    "WarmStartType" : "IdenticalDataAndAlgorithm"
}
```

Create the warm start tuning job:

```python
smclient = boto3.Session().client('sagemaker')
smclient.create_hyper_parameter_tuning_job(HyperParameterTuningJobName =
    'MyWarmStartTuningJob',
    HyperParameterTuningJobConfig = tuning_job_config, # See notebook for tuning configuration
    TrainingJobDefinition = training_job_definition, # See notebook for job definition
    WarmStartConfig = warm_start_config)
```

**Create a Warm Start Tuning Job (SageMaker Python SDK)**

To use the [Amazon SageMaker Python SDK](https://github.com/aws/sagemaker-python-sdk#sagemaker-automatic-model-tuning) to run a warm start tuning job, you:

- Specify the parent jobs and the warm start type by using a `WarmStartConfig` object.
- Pass the `WarmStartConfig` object as the value of the `warm_start_config` argument of a `HyperparameterTuner` object.
- Call the `fit` method of the `HyperparameterTuner` object.


This example uses an estimator that uses the `Image Classification - MXNet` algorithm for training. The following code sets the hyperparameter ranges that the warm start tuning job searches within to find the best combination of values. For information about setting hyperparameter ranges, see [Define Hyperparameter Ranges](https://github.com/aws/sagemaker-python-sdk#sagemaker-automatic-model-tuning).
hyperparameter_ranges = {'learning_rate': ContinuousParameter(0.0, 0.1),
   'momentum': ContinuousParameter(0.0, 0.99)}

The following code configures the warm start tuning job by creating a WarmStartConfig object.

```python
from sagemaker.tuner import WarmStartConfig, WarmStartTypes

parent_tuning_job_name = "MyParentTuningJob"
warm_start_config = WarmStartConfig(warm_start_type=WarmStartTypes.IDENTICAL_DATA_AND_ALGORITHM,
                                        parents={parent_tuning_job_name})
```

Now set the values for static hyperparameters, which are hyperparameters that keep the same value for every training job that the warm start tuning job launches. In the following code, `imageclassification` is an estimator that was created previously.

```python
imageclassification.set_hyperparameters(num_layers=18,
                                        image_shape='3,224,224',
                                        num_classes=257,
                                        num_training_samples=15420,
                                        mini_batch_size=128,
                                        epochs=30,
                                        optimizer='sgd',
                                        top_k='2',
                                        precision_dtype='float32',
                                        augmentation_type='crop')
```

Now create the `HyperparameterTuner` object and pass the `WarmStartConfig` object that you previously created as the `warm_start_config` argument.

```python
tuner_warm_start = HyperparameterTuner(imageclassification,
                                        'validation:accuracy',
                                        hyperparameter_ranges,
                                        objective_type='Maximize',
                                        max_jobs=10,
                                        max_parallel_jobs=2,
                                        base_tuning_job_name='warmstart',
                                        warm_start_config=warm_start_config)
```

Finally, call the `fit` method of the `HyperparameterTuner` object to launch the warm start tuning job.
tuner_warm_start.fit(
    {'train': s3_input_train, 'validation': s3_input_validation},
    include_cls_metadata=False)

## Resource Limits for Automatic Model Tuning

SageMaker sets the following default limits for resources used by automatic model tuning:

<table>
<thead>
<tr>
<th>Resource</th>
<th>Regions</th>
<th>Default limits</th>
<th>Can be increased to</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of parallel (concurrent) hyperparameter tuning jobs</td>
<td>All</td>
<td>100</td>
<td>N/A</td>
</tr>
<tr>
<td>Number of hyperparameters that can be searched *</td>
<td>All</td>
<td>30</td>
<td>N/A</td>
</tr>
<tr>
<td>Number of metrics defined per hyperparameter tuning job</td>
<td>All</td>
<td>20</td>
<td>N/A</td>
</tr>
<tr>
<td>Number of parallel training jobs per hyperparameter tuning job</td>
<td>All</td>
<td>10</td>
<td>100</td>
</tr>
<tr>
<td>[Bayesian optimization] Number of training jobs per hyperparameter tuning job</td>
<td>All</td>
<td>750</td>
<td>N/A</td>
</tr>
<tr>
<td>[Random search] Number of training</td>
<td>All</td>
<td>750</td>
<td>10000</td>
</tr>
<tr>
<td>Resource</td>
<td>Regions</td>
<td>Default limits</td>
<td>Can be increased to</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>---------</td>
<td>----------------</td>
<td>---------------------</td>
</tr>
<tr>
<td>jobs per hyperparameter tuning job</td>
<td>All</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>[Hyperband] Number of training jobs per hyperparameter tuning job</td>
<td>All</td>
<td>750</td>
<td>N/A</td>
</tr>
<tr>
<td>[Grid] Number of training jobs per hyperparameter tuning job, either specified explicitly or inferred from the search space</td>
<td>All</td>
<td>750</td>
<td>N/A</td>
</tr>
<tr>
<td>Maximum run time for a hyperparameter tuning job</td>
<td>All</td>
<td>30 days</td>
<td>N/A</td>
</tr>
</tbody>
</table>

* Each categorical hyperparameter can have at most 30 different values.

**Resource limit example**

When you plan hyperparameter tuning jobs, you also have to take into account the limits on training resources. For information about the default resource limits for SageMaker training jobs, see [SageMaker Limits](#). Every concurrent training instance on which all of your hyperparameter tuning jobs run counts against the total number of training instances allowed. For example, if you run 10 concurrent hyperparameter tuning jobs, each of those hyperparameter tuning jobs runs 100 total training jobs and 20 concurrent training jobs. Each of those training jobs runs on one `ml.m4.xlarge` instance. The following limits apply:

- Number of concurrent hyperparameter tuning jobs: You don't need to increase the limit, because 10 tuning jobs is below the limit of 100.
- Number of training jobs per hyperparameter tuning job: You don't need to increase the limit, because 100 training jobs is below the limit of 750.
• Number of concurrent training jobs per hyperparameter tuning job: You need to request a limit increase to 20, because the default limit is 10.

• SageMaker training ml.m4.xlarge instances: You need to request a limit increase to 200, because you have 10 hyperparameter tuning jobs, each of which is running 20 concurrent training jobs. The default limit is 20 instances.

• SageMaker training total instance count: You need to request a limit increase to 200, because you have 10 hyperparameter tuning jobs, each of which is running 20 concurrent training jobs. The default limit is 20 instances.

To request a quota increase:

1. Open the [AWS Support Center](https://aws.amazon.com/support/) page, sign in if necessary, and then choose [Create case](https://aws.amazon.com/support/ tinha/index.jsp).  
2. On the [Create case](https://aws.amazon.com/support/ tinha) page, choose [Service limit increase](https://aws.amazon.com/support/ tinha/service-limit-increase/).  
4. On the [Requests](https://aws.amazon.com/support/ tinha/case/requests/) panel for Request 1, select the [Region](https://aws.amazon.com/support/ tinha/case/region/), the resource [Limit](https://aws.amazon.com/support/ tinha/case/limit/) to increase and the [New Limit value](https://aws.amazon.com/support/ tinha/case/new-limit-value/) you are requesting. Select [Add another request](https://aws.amazon.com/support/ tinha/case/add-another-request/) if you have additional requests for quota increases.
5. In the **Case description** panel, provide a description of your use case.

6. In the **Contact options** panel, select your preferred **Contact methods (Web, Chat or Phone)** and then choose **Submit**.

**Best Practices for Hyperparameter Tuning**

Hyperparameter optimization (HPO) is not a fully-automated process. To improve optimization, follow these best practices for hyperparameter tuning.

**Topics**

- [Choosing a tuning strategy](#)
Choosing a tuning strategy

For large jobs, using the Hyperband tuning strategy can reduce computation time. Hyperband has an early stopping mechanism to stop under-performing jobs. Hyperband can also reallocate resources towards well-utilized hyperparameter configurations and run parallel jobs. For smaller training jobs using less runtime, use either random search or Bayesian optimization.

Use Bayesian optimization to make increasingly informed decisions about improving hyperparameter configurations in the next run. Bayesian optimization uses information gathered from prior runs to improve subsequent runs. Because of its sequential nature, Bayesian optimization cannot massively scale.

Use random search to run a large number of parallel jobs. In random search, subsequent jobs do not depend on the results from prior jobs and can be run independently. Compared to other strategies, random search is able to run the largest number of parallel jobs.

Use grid search to reproduce results of a tuning job, or if simplicity and transparency of the optimization algorithm are important. You can also use grid search to explore the entire hyperparameter search space evenly. Grid search methodically searches through every hyperparameter combination to find optimal hyperparameter values. Unlike grid search, Bayesian optimization, random search and Hyperband all draw hyperparameters randomly from the search space. Because grid search analyzes every combination of hyperparameters, optimal hyperparameter values will be identical between tuning jobs that use the same hyperparameters.

Choosing the number of hyperparameters

During optimization, the computational complexity of a hyperparameter tuning job depends on the following:

- The number of hyperparameters
- The range of values that Amazon SageMaker has to search
Although you can simultaneously specify up to 30 hyperparameters, limiting your search to a smaller number can reduce computation time. Reducing computation time allows SageMaker to converge more quickly to an optimal hyperparameter configuration.

**Choosing hyperparameter ranges**

The range of values that you choose to search can adversely affect hyperparameter optimization. For example, a range that covers every possible hyperparameter value can lead to large compute times and a model that doesn't generalize well to unseen data. If you know that using a subset of the largest possible range is appropriate for your use case, consider limiting the range to that subset.

**Using the correct scales for hyperparameters**

During hyperparameter tuning, SageMaker attempts to infer if your hyperparameters are log-scaled or linear-scaled. Initially, SageMaker assumes linear scaling for hyperparameters. If hyperparameters are log-scaled, choosing the correct scale will make your search more efficient. You can also select Auto for ScalingType in the `CreateHyperParameterTuningJob` API if you want SageMaker to detect the scale for you.

**Choosing the best number of parallel training jobs**

You can use the results of previous trials to improve the performance of subsequent trials. Choose the largest number of parallel jobs that would provide a meaningful incremental result that is also within your region and account compute constraints. Use the `MaxParallelTrainingJobs` field to limit the number of training jobs that a hyperparameter tuning job can launch in parallel. For more information, see Running multiple HPO jobs in parallel on Amazon SageMaker.

**Running training jobs on multiple instances**

When a training job runs on multiple machines in distributed mode, each machine emits an objective metric. HPO can only use one of these emitted objective metrics to evaluate model performance. In distributed mode, HPO uses the objective metric that was reported by the last running job across all instances.

**Using a random seed to reproduce hyperparameter configurations**

You can specify an integer as a random seed for hyperparameter tuning and use that seed during hyperparameter generation. Later, you can use the same seed to reproduce hyperparameter
configurations that are consistent with your previous results. For random search and Hyperband strategies, using the same random seed can provide up to 100% reproducibility of the previous hyperparameter configuration for the same tuning job. For Bayesian strategy, using the same random seed will improve reproducibility for the same tuning job.

**Refine data during training with Amazon SageMaker smart sifting**

Amazon SageMaker smart sifting is in preview release and is subject to change.

SageMaker smart sifting is a capability of SageMaker Training that helps improve the efficiency of your training datasets and reduce total training time and cost.

Modern deep learning models such as large language models (LLMs) or vision transformer models often require massive datasets to achieve acceptable accuracy. For example, LLMs often require trillions of tokens or petabytes of data to converge. The growing size of training datasets, along with the size of state-of-the-art models, can increase the compute time and cost of model training.

Invariably, samples in a dataset do not contribute equally to the learning process during model training. A significant proportion of computational resources provisioned during training might be spent on processing easy samples that do not contribute substantially to the overall accuracy of a model. Ideally, training datasets would only include samples that are actually improving the model convergence. Filtering out less helpful data can reduce training time and compute cost. However, identifying less helpful data can be challenging and risky. It is practically difficult to identify which samples are less informative before training, and model accuracy can be impacted if the wrong samples or too many samples are excluded.

Smart sifting of data with Amazon SageMaker can help reduce training time and cost by improving data efficiency. The SageMaker smart sifting algorithm evaluates the loss value of each data during the data loading stage of a training job and excludes samples which are less informative to the model. By using refined data for training, the total time and cost of training your model is reduced by eliminating unnecessary forward and backward passes on non-improving data. Therefore, there is minimal or no impact on the accuracy of the model.

SageMaker smart sifting is available through SageMaker Training Deep Learning Containers (DLCs) and supports PyTorch workloads via the `PyTorch DataLoader`. Just a few lines of code change
are needed to implement SageMaker smart sifting and you do not need to change your existing training or data processing workflows.

**How SageMaker smart sifting works**

The goal of SageMaker smart sifting is to sift through your training data during the training process and only feed more informative samples to the model. During typical training with PyTorch, data is iteratively sent in batches to the training loop and to accelerator devices (such as GPUs or Trainium chips) by the **PyTorch DataLoader**. SageMaker smart sifting is implemented at this data loading stage and is thus independent of any upstream data pre-processing in your training pipeline. SageMaker smart sifting uses your model and its user-specified loss function to do an evaluative forward pass of each data sample as it is loaded. Samples that return *low-loss* values have less of an impact on the model's learning and are thus excluded from training, because it is already *easy* for the model to make the right prediction about them with high confidence. Meanwhile, those relatively high-loss samples are what the model still needs to learn, so these are kept for training. A key input you can set for SageMaker smart sifting is the proportion of data to exclude. For example, by setting the proportion to 25%, samples distributed in the lowest quartile of the distribution of loss (taken from a user-specified number of previous samples) are excluded from training. High-loss samples are accumulated in a refined data batch. The refined data batch is sent to the training loop (forward and backward pass), and the model learns and trains on the refined data batch.

The following diagram shows an overview of how the SageMaker smart sifting algorithm is designed.
In short, SageMaker smart sifting operates during training as data is loaded. The SageMaker smart sifting algorithm runs loss calculation over the batches, and sifts non-improving data out before the forward and backward pass of each iteration. The refined data batch is then used for the forward and backward pass.

SageMaker smart sifting works for PyTorch-based training jobs with classic distributed data parallelism, which makes model replicas on each GPU worker and performs AllReduce. It works with PyTorch DDP and the SageMaker distributed data parallel library.
Supported frameworks and AWS Regions

Before using SageMaker smart sifting data loader, check if your framework of choice is supported, that the instance types are available in your AWS account, and that your AWS account is in one of the supported AWS Regions.

Supported Frameworks

SageMaker smart sifting supports the following deep learning frameworks and is available through AWS Deep Learning Containers.

Topics

- PyTorch

PyTorch

<table>
<thead>
<tr>
<th>Framework</th>
<th>Framework version</th>
<th>Deep Learning Container URI</th>
</tr>
</thead>
<tbody>
<tr>
<td>PyTorch</td>
<td>2.1.0</td>
<td>763104351884.dkr.ecr.region.amazonaws.com/pytorch-training:2.1.0-gpu-py310-cu121-ubuntu20.04-sagemaker</td>
</tr>
</tbody>
</table>

For more information about the pre-built containers, see SageMaker Framework Containers in the AWS Deep Learning Containers GitHub repository.

AWS Regions

The containers packaged with the SageMaker smart sifting library are available in the AWS Regions where AWS Deep Learning Containers are in service.
Instance types

You can use SageMaker smart sifting for any PyTorch training jobs on any instance types. We recommend that you use P4d, P4de, or P5 instances.

Apply SageMaker smart sifting to your training script

The SageMaker smart sifting library is packaged in the SageMaker framework DLCs as a complementary library. It provides a filtering logic against training samples that have relatively lower impact on model training, and your model can reach the desired model accuracy with fewer training samples when compared to the model training with full data samples.

PyTorch

These instructions demonstrate how to enable SageMaker smart sifting with your training script.

1. Configure the SageMaker smart sifting interface.

   The SageMaker smart sifting library implements a relative-threshold loss-based sampling technique that helps filter out samples with lower impact on reducing the loss value. The SageMaker smart sifting algorithm calculates the loss value of every input data sample using a forward pass, and calculates its relative percentile against the loss values of preceding data.

   The following two parameters are what you need to specify to the RelativeProbabilisticSiftConfig class for creating a sifting configuration object.

   • Specify the proportion of data that should be used for training to the beta_value parameter.
   • Specify the number of samples used in the comparison with the loss_history_length parameter.

   The following code example demonstrates setting up an object of the RelativeProbabilisticSiftConfig class.

```
from smart_sifting.sift_config.sift_configs import (
    RelativeProbabilisticSiftConfig
    LossConfig
    SiftingBaseConfig
)
```
sift_config=RelativeProbabilisticSiftConfig(
    beta_value=0.5,
    loss_history_length=500,
    loss_based_sift_config=LossConfig(
        sift_config=SiftingBaseConfig(sift_delay=0)
    )
)

For more information about the loss_based_sift_config parameter and related classes, see the section called “SageMaker smart sifting configuration modules” in the SageMaker smart sifting Python SDK reference section.

The sift_config object in the preceding code example is used in step 4 for setting up the SiftingDataloader class.

2. (Optional) Configure a SageMaker smart sifting batch transform class.

Different training use cases require different training data formats. Given the variety of data formats, the SageMaker smart sifting algorithm needs to identify how to perform sifting on a particular batch. To address this, SageMaker smart sifting provides a batch transform module that helps convert batches into standardized formats that it can efficiently sift.

a. SageMaker smart sifting handles batch transform of training data in the following formats: Python lists, dictionaries, tuples, and tensors. For these data formats, SageMaker smart sifting automatically handles the batch data format conversion, and you can skip the rest of this step. If you skip this step, in step 4 for configuring SiftingDataloader, leave the batch_transforms parameter of SiftingDataloader to its default value, which is None.

b. If your dataset is not in these format, you should proceed to the rest of this step to create a custom batch transform using SiftingBatchTransform.

In cases in which your dataset isn’t in one of the supported formats by SageMaker smart sifting, you might run into errors. Such data format errors can be resolved by adding the batch_format_index or batch_transforms parameter to the SiftingDataloader class, which you set up in step 4. The following shows example errors due to an incompatible data format and resolutions for them.

<table>
<thead>
<tr>
<th>Error Message</th>
<th>Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Issue Description</td>
<td>Error Description</td>
</tr>
<tr>
<td>----------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Batches of type {type(batch)} are not supported by default.</td>
<td>This error indicates the batch format is not supported by default. You should implement a custom batch transform class, and use this by specifying it to the batch_transforms parameter of the SiftingDataloader class.</td>
</tr>
<tr>
<td>Unable to index the batch of type {type(batch)}</td>
<td>This error indicates the batch object cannot be indexed normally. User must implement a custom batch transform and pass this using the batch_transforms parameter.</td>
</tr>
<tr>
<td>Batch size {batch_size} does not match dimension 0 or dimension 1 sizes</td>
<td>This error occurs when the provided batch size does not match the 0th or 1st dimensions of the batch. User must implement a custom batch transform and pass this using the batch_transforms parameter.</td>
</tr>
<tr>
<td>Both dimension 0 and dimension 1 match batch size</td>
<td>This error indicates that since multiple dimensions match the provided batch size, more information is required to sift the batch. The user can provide the batch_format_index parameter to indicate if the batch is indexable by sample or feature. Users may also implement a custom batch transform, but this is more work than required.</td>
</tr>
</tbody>
</table>

To resolve the aforementioned issues, you need to create a custom batch transform class using the SiftingBatchTransform module. A batch transform class should consist of a pair of transform and reverse-transform functions. The function pair converts your data format to a format that SageMaker smart sifting algorithm can process. After you create a batch transform class, the class returns a SiftingBatch object that you'll pass to the SiftingDataloader class in step 4.
The following are examples of custom batch transform classes of the SiftingBatchTransform module.

- An example of a custom list batch transform implementation with SageMaker smart sifting for cases where the dataloader chunk has inputs, masks, and labels.

```python
from typing import Any
import torch
from smart_sifting.data_model.data_model_interface import SiftingBatchTransform
from smart_sifting.data_model.list_batch import ListBatch

class ListBatchTransform(SiftingBatchTransform):
    def transform(self, batch: Any):
        inputs = batch[0].tolist()
        labels = batch[-1].tolist()  # assume the last one is the list of labels
        return ListBatch(inputs, labels)

    def reverse_transform(self, list_batch: ListBatch):
        a_batch = [torch.tensor(list_batch.inputs),
                   torch.tensor(list_batch.labels)]
        return a_batch
```

- An example of a custom list batch transform implementation with SageMaker smart sifting for cases where no labels are needed for reverse transformation.

```python
class ListBatchTransformNoLabels(SiftingBatchTransform):
    def transform(self, batch: Any):
        return ListBatch(batch[0].tolist())

    def reverse_transform(self, list_batch: ListBatch):
        a_batch = [torch.tensor(list_batch.inputs)]
        return a_batch
```

- An example of a custom tensor batch implementation with SageMaker smart sifting for cases where the data loader chunk has inputs, masks, and labels.

```python
from typing import Any
```
from smart_sifting.data_model.data_model_interface import SiftingBatchTransform
from smart_sifting.data_model.tensor_batch import TensorBatch

class TensorBatchTransform(SiftingBatchTransform):
    def transform(self, batch: Any):
        a_tensor_batch = TensorBatch(
            batch[0], batch[-1]
        )  # assume the last one is the list of labels
        return a_tensor_batch

    def reverse_transform(self, tensor_batch: TensorBatch):
        a_batch = [tensor_batch.inputs, tensor_batch.labels]
        return a_batch

After you create a SiftingBatchTransform-implemented batch transform class, you use this class in step 4 for setting up the SiftingDataloader class. The rest of this guide assumes that a ListBatchTransform class is created. In step 4, this class is passed to the batch_transforms.

3. Create a class for implementing the SageMaker smart sifting Loss interface. This tutorial assumes that the class is named SiftingImplementedLoss. While setting up this class, we recommend that you use the same loss function in the model training loop. Go through the following substeps for creating a SageMaker smart sifting Loss implemented class.

   a. SageMaker smart sifting calculates a loss value for each training data sample, as opposed to calculating a single loss value for a batch. To ensure that SageMaker smart sifting uses the same loss calculation logic, create a smart-sifting-implemented loss function using the SageMaker smart sifting Loss module that uses your loss function and calculates loss per training sample.

   Tip

   SageMaker smart sifting algorithm runs on every data sample, not on the entire batch, so you should add an initialization function to set the PyTorch loss function without any reduction strategy.
self.loss = torch.nn.CrossEntropyLoss(reduction='none')

This is also shown in the following code example.

b. Define a loss function that accepts the original_batch (or transformed_batch if you have set up a batch transform in step 2) and the PyTorch model. Using the specified loss function with no reduction, SageMaker smart sifting runs a forward pass for each data sample to evaluate its loss value.

The following code is an example of a smart-sifting-implemented Loss interface named SiftingImplementedLoss.

```python
from typing import Any
import torch
import torch.nn as nn
from torch import Tensor
from smart_sifting.data_model.data_model_interface import SiftingBatch
from smart_sifting.loss.abstract_sift_loss_module import Loss

model = ... # a PyTorch model based on torch.nn.Module

class SiftingImplementedLoss(Loss):
    # You should add the following initialization function
    # to calculate loss per sample, not per batch.
    def __init__(self):
        self.loss_no_reduction = torch.nn.CrossEntropyLoss(reduction='none')

    def loss(self, model: torch.nn.Module, transformed_batch: SiftingBatch, original_batch: Any = None, ) -> torch.Tensor:
        device = next(model.parameters()).device
        batch = [t.to(device) for t in original_batch] # use this if you use original batch and skipped step 2
        # batch = [t.to(device) for t in transformed_batch] # use this if you transformed batches in step 2
```
Before the training loop hits the actual forward pass, this sifting loss calculation is done during the data loading phase of fetching a batch in each iteration. The individual loss value is then compared to previous loss values, and its relative percentile is estimated per the object of RelativeProbabilisticSiftConfig you have set up in step 1.

4. Wrap the PyTorch data loader by the SageMaker SiftingDataloader class.

Finally, use all the SageMaker smart sifting implemented classes you configured in the previous steps to the SageMaker SiftingDataloader configuration class. This class is a wrapper for PyTorch DataLoader. By wrapping PyTorch DataLoader, SageMaker smart sifting is registered to run as part of data loading in each iteration of a PyTorch training job. The following code example demonstrates implementing SageMaker data sifting to a PyTorch DataLoader.

```python
from smart_sifting.dataloader.sift_dataloader import SiftingDataloader
from torch.utils.data import DataLoader

train_dataloader = DataLoader(...) # PyTorch data loader

# Wrap the PyTorch data loader by SiftingDataloader
train_dataloader = SiftingDataloader(
    sift_config=sift_config, # config object of RelativeProbabilisticSiftConfig
    orig_dataloader=train_dataloader,
    batch_transforms=ListBatchTransform(), # Optional, this is the custom class from step 2
    loss_impl=SiftingImplementedLoss(), # PyTorch loss function wrapped by the Sifting Loss interface
    model=model,
    log_batch_data=False
)
```

**Hugging Face Transformers**

There are two ways to implement the SageMaker smart sifting into the Transformers Trainer class.

---

Apply SageMaker smart sifting to your training script
**Note**

If you use one of the DLCs for PyTorch with the SageMaker smart sifting package installed, note that you need to install the transformers library. You can install additional packages by extending the DLCs or passing requirements.txt to the training job launcher class for PyTorch ([sagemaker.pytorch.PyTorch](https://sagemaker.pytorch.org)) in the SageMaker Python SDK.

**Simple setup**

The simplest way to implement SageMaker smart sifting into the Transformers `Trainer` class is to use the `enable_sifting` function. This function accepts an existing `Trainer` object, and wraps the existing `DataLoader` object with `SiftingDataloader`. You can continue using the same training object. See the following example usage.

```python
from smart_sifting.integrations.trainer import enable_sifting
from smart_sifting.loss.abstract_sift_loss_module import Loss
from smart_sifting.sift_config.sift_configs import (RelativeProbabilisticSiftConfig,
LossConfig,
SiftingBaseConfig)

class SiftingImplementedLoss(Loss):
    def loss(self, model, transformed_batch, original_batch):
        loss_fct = MSELoss(reduction="none")  # make sure to set reduction to "none"
        logits = model.bert(**original_batch)
        return loss_fct(logits, original_batch.get("labels"))

sift_config = RelativeProbabilisticSiftConfig(
    beta_value=0.5,
    loss_history_length=500,
    loss_based_sift_config=LossConfig(
        sift_config=SiftingBaseConfig(sift_delay=0)
    )
)

trainer = Trainer(...)  
enable_sifting(trainer, sift_config, loss=SiftingImplementedLoss())  # updates the trainer with Sifting Loss and config
```
The `SiftingDataloader` class is an iterable data loader. The exact size of the resulting dataset is not known beforehand due to the random sampling during sifting. As a result, the Hugging Face Trainer expects the `max_steps` training argument. Note that this argument overrides the epoch configuration parameter `num_train_epochs`. If your original data loader was also iterable, or your training uses `max_steps` and a single epoch, then the `SiftingDataloader` performs the same as the existing dataloader. If the original dataloader was not iterable or `max_steps` was not provided, the Hugging Face Trainer might throw an error message similar to the following.

```
args.max_steps must be set to a positive value if dataloader does not have a length, was -1
```

To address this, the `enable_sifting` function provides an optional `set_epochs` parameter. This enables training with epochs, using the number of epochs provided by `num_train_epochs` argument of the Trainer class, and sets `max_steps` to the maximum system integer, allowing training to progress until the specified epochs have completed.

**Custom setup**

For a custom integration of the SageMaker smart sifting data loader, you can utilize a custom Hugging Face Trainer class. Within any subclass of Trainer, the `get_train_dataloader()` function can be overridden to return an object of the `SiftingDataloader` class instead. For cases with existing custom trainers, this approach might be less intrusive but requires code changes than the simple setup option. The following is an example implementation of SageMaker smart sifting into a custom Hugging Face Trainer class.

```python
from smart_sifting.sift_config.sift_configs import (  
    RelativeProbabilisticSiftConfig  
    LossConfig  
    SiftingBaseConfig  
)
from smart_sifting.dataloader.sift_dataloader import SiftingDataloader  
from smart_sifting.loss.abstract_sift_loss_module import Loss  
from smart_sifting.data_model.data_model_interface import SiftingBatch,  
    SiftingBatchTransform  
from smart_sifting.data_model.list_batch import ListBatch  

class SiftingListBatchTransform(SiftingBatchTransform):
    def transform(self, batch: Any):
```

Apply SageMaker smart sifting to your training script
inputs = batch[0].tolist()
labels = batch[-1].tolist()  # assume the last one is the list of labels
return ListBatch(inputs, labels)

def reverse_transform(self, list_batch: ListBatch):
a_batch = [torch.tensor(list_batch.inputs), torch.tensor(list_batch.labels)]
return a_batch

class SiftingImplementedLoss():
    # You should add the following initialization function
    # to calculate loss per sample, not per batch.
    def __init__(self):
        self.celoss = torch.nn.CrossEntropyLoss(reduction='none')

    def loss(
        self,
        model: torch.nn.Module,
        transformed_batch: SiftingBatch,
        original_batch: Any = None,
    ) -> torch.Tensor:
        device = next(model.parameters()).device
        batch = [t.to(device) for t in original_batch]

        # compute loss
        outputs = model(batch)
        return self.celoss(outputs.logits, batch[2])

class SiftingImplementedTrainer(Trainer):
    def get_train_dataloader(self):
        dl = super().get_train_dataloader()

        sift_config = RelativeProbabilisticSiftConfig(
            beta_value=0.5,
            loss_history_length=500,
            loss_based_sift_config=LossConfig(
                sift_config=SiftingBaseConfig(sift_delay=0)
            )
        )

        return SiftingDataloader(
            sift_config=sift_config,
            orig_dataloader=dl,
            batch_transforms=SiftingListBatchTransform(),
            loss_impl=SiftingImplementedLoss(),
        )
Using the wrapped `Trainer` class, create an object of it as follows.

```python
trainer = SiftingImplementedTrainer(
    model=model,
    args=training_args,
    train_dataset=small_train_dataset,
    eval_dataset=small_eval_dataset
)

trainer.train()
```

### Best practices, considerations, and troubleshooting

#### Best practices

- Smart sifting of data on SageMaker uses additional forward passes to analyze and filter your training data. In turn, there are fewer backward passes as less impactful data is excluded from your training job. Because of this, models which have long or expensive backward passes see the greatest efficiency gains when using smart sifting. Meanwhile, if your model's forward pass takes longer than its backward pass, overhead could increase total training time. To measure the time spent by each pass, you can run a pilot training job and collect logs that record the time on the processes. Also consider using SageMaker Profiler that provides profiling tools and UI application. To learn more, see [Use Amazon SageMaker Profiler to profile activities on AWS compute resources](#).

- SageMaker smart sifting supports PyTorch model training with traditional data parallelism and distributed data parallelism, which makes model replicas in all GPU workers and uses the `AllReduce` operation. It doesn't work with model parallelism techniques, including sharded data parallelism.

- Because SageMaker smart sifting works for data parallelism jobs, make sure that the model you train fits in each GPU memory.

- SageMaker smart sifting runs on individual data in batches during data loading, so make sure that you set the reduction strategy of the PyTorch loss function to "none" for non-reduction. With reduction set to "mean" or "sum", the loss function returns a single loss value, which leads SageMaker smart sifting not working properly.
Troubleshooting

If you run into an error, you can use the following list to try to troubleshoot the issue. If you need further support, reach out to the SageMaker team at <sm-smart-sifting-feedback@amazon.com>.

Exceptions from the SageMaker smart sifting library

Use the following reference of exceptions raised by the SageMaker smart sifting library to troubleshoot errors and identify causes.

<table>
<thead>
<tr>
<th>Exception Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiftConfigValidationException</td>
<td>Thrown from the SageMaker smart sifting library in case of any missing Config key or unsupported value type for Sift Key</td>
</tr>
<tr>
<td>UnsupportedDataFormatException</td>
<td>Thrown from the SageMaker smart sifting library in case of any unsupported DataFormat for Sifting logic</td>
</tr>
<tr>
<td>LossImplementationNotProvidedException</td>
<td>Thrown in case of missing or not implementing Loss interface</td>
</tr>
</tbody>
</table>

Security in SageMaker smart sifting

Because the SageMaker smart sifting library runs processes of removing less valuable training samples, it requires full access to training datasets as they are produced by the data loader. This access is not different than the access already provided to PyTorch in normal training scenario.

SageMaker smart sifting has built-in logging with security implications. By default, SageMaker smart sifting logs are only application-level logs containing metrics, latencies, and user errors or warnings. Users can, however, choose to enable verbose logs, which log full batch data to show which samples were removed from a given batch. These logs are emitted using Python loggers and are not uploaded or stored anywhere by the library. In the case of automatic log uploading to CloudWatch or similar services, please note that using verbose logs may result in sensitive training data being uploaded off of the training instance.
Beyond the aforementioned logging, SageMaker smart sifting does not have any network functionality nor does it interact with the local file system. User data is stored as in-memory objects for the entirety of the time it is used by the library.

**SageMaker smart sifting Python SDK reference**

This page provides a reference of Python modules you need for applying SageMaker smart sifting to your training script.

**SageMaker smart sifting configuration modules**

```python
class smart_sifting.sift_config.sift_configs.RelativeProbabilisticSiftConfig()
```

The SageMaker smart sifting configuration class.

**Parameters**

- `beta_value` (float) – A beta (constant) value for calculating the probability of selecting a sample for training based on the percentile of the loss in the history of loss values. Lowering the beta value results in a lower percentage of data sifted, and raising it results in a higher percentage of data sifted. There’s no minimum or maximum value for the beta value, beside that it must be a positive value. Refer to the following reference table for sifting rates with respect to `beta_value`.

<table>
<thead>
<tr>
<th><code>beta_value</code></th>
<th>Proportion of data kept (%)</th>
<th>Proportion of data sifted out (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1</td>
<td>90.91</td>
<td>9.01</td>
</tr>
<tr>
<td>0.25</td>
<td>80</td>
<td>20</td>
</tr>
<tr>
<td>0.5</td>
<td>66.67</td>
<td>33.33</td>
</tr>
<tr>
<td>1</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>2</td>
<td>33.33</td>
<td>66.67</td>
</tr>
<tr>
<td>3</td>
<td>25</td>
<td>75</td>
</tr>
</tbody>
</table>
• **loss_history_length (int)** – The number of previous training losses to store for the relative threshold loss based sampling.

• **loss_based_sift_config (dict or a LossConfig object)** – Specify a LossConfig object that returns the SageMaker smart sifting Loss interface configuration.

```python
class smart_sifting.sift_config.sift_configs.LossConfig()
```

The configuration class for the `loss_based_sift_config` parameter of the `RelativeProbabilisticSiftConfig` class.

**Parameters**

- **sift_config (dict or a SiftingBaseConfig object)** – Specify a SiftingBaseConfig object that returns a sifting base configuration dictionary.

```python
class smart_sifting.sift_config.sift_configs.SiftingBaseConfig()
```

The configuration class for the `sift_config` parameter of `LossConfig`.

**Parameters**

- **sift_delay (int)** – The number of training steps to wait for before start sifting. We recommend that you start sifting after all the layers in the model has sufficient view of the training data. The default value is 1000.

- **repeat_delay_per_epoch (bool)** – Specify whether to delay sifting every epoch. The default value is False.

**SageMaker smart sifting data batch transform modules**

```python
class smart_sifting.data_model.data_model_interface.SiftingBatchTransform
```

A SageMaker smart sifting Python module for defining how to perform batch transform. Using this, you can set up a batch transform class that converts the data format of your training data to `SiftingBatch` format, which SageMaker smart sifting can sift and accumulate into a sifted batch.
class smart_sifting.data_model.data_model_interface.SiftingBatch

An interface to define a batch data type that can be sifted and accumulated.

class smart_sifting.data_model.list_batch.ListBatch

A module for keeping track of a list batch for sifting.

class smart_sifting.data_model.tensor_batch.TensorBatch

A module for keeping track of a tensor batch for sifting.

**SageMaker smart sifting loss implementation module**

class smart_sifting.loss.abstract_sift_loss_module.Loss

A wrapper module for registering the SageMaker smart sifting interface to the loss function of a PyTorch-based model.

**SageMaker smart sifting data loader wrapper module**

class smart_sifting.dataloader.sift_dataloader.SiftingDataloader

A wrapper module for registering the SageMaker smart sifting interface to the data loader of a PyTorch-based model.

The Main Sifting Dataloader iterator sifts out training samples from a dataloader based on a sift configuration

**Parameters**

- `sift_config` (dict or a RelativeProbabilisticSiftConfig object) – A RelativeProbabilisticSiftConfig object.
- `orig_dataloader` (a PyTorch DataLoader object) – Specify the PyTorch DataLoader object to be wrapped.
- `batch_transforms` (a SiftingBatchTransform object) – (Optional) If your data format is not supported by the SageMaker smart sifting library’s default transform, you need to create a batch transform class using the SiftingBatchTransform module. This parameter is to pass the batch transform class that SiftingDataloader can run to convert the data for SageMaker smart sifting algorithm can accept.
• model (a PyTorch model object) – The original PyTorch model
• loss_impl (a sifting loss function of
  smart_sifting.loss.abstract_sift_loss_module.Loss) – A sifting loss function that is configured with the Loss module and wraps the PyTorch loss function.
• log_batch_data (bool) – Specify whether to log batch data. If set to True, SageMaker smart sifting logs the details of the batches that are kept or sifted. We recommend that you turn it on only for a pilot training job. When logging is on, the samples are loaded to GPU and transferred to CPU, which introduces overhead. The default value is False.

**SageMaker smart sifting release notes**

See the following release notes to track the latest updates for the SageMaker smart sifting capability.

**SageMaker smart sifting release notes: November 29, 2023**

**New Features**

• Launched the Amazon SageMaker smart sifting library at AWS re:Invent 2023.

**Migration to AWS Deep Learning Containers**

• The SageMaker smart sifting library passed integration testing and is available in AWS Deep Learning Containers. To find a complete list of the pre-built containers with the SageMaker smart sifting library, see the section called “Supported frameworks and AWS Regions”.

**Debug and improve model performance**

The essence of training machine learning models, deep learning neural networks, transformer models is in achieving stable model convergence, and as such, state-of-the-art models have millions, billions, or trillions of model parameters. The number of operations to update the gigantic number of model parameters during each iteration can easily become astronomical. To identify model convergence issues, it is important to be able to access the model parameters, activations, and gradients computed during optimization processes.

Amazon SageMaker provides two debugging tools to help identify such convergence issues and gain visibility into your models.
Amazon SageMaker with TensorBoard

To offer a greater compatibility with the open-source community tools within the SageMaker Training platform, SageMaker hosts TensorBoard as an application in SageMaker domain. You can bring your training jobs to SageMaker and keep using the TensorBoard summary writer to collect the model output tensors. Because TensorBoard is implemented into SageMaker domain, it also gives you more options to manage user profiles under the SageMaker domain in your AWS account, and provides fine control over the user profiles by granting access to specific actions and resources. To learn more, see the section called “Use TensorBoard”.

Amazon SageMaker Debugger

Amazon SageMaker Debugger is a capability of SageMaker that provides tools to register hooks to callbacks to extract model output tensors and save them in Amazon Simple Storage Service. It provides built-in rules for detecting model convergence issues, such as overfitting, saturated activation functions, vanishing gradients, and more. You can also set up the built-in rules with Amazon CloudWatch Events and AWS Lambda for taking automated actions against detected issues, and set up Amazon Simple Notification Service to receive email or text notifications. To learn more, see the section called “Use SageMaker Debugger”.

Topics

- Use TensorBoard to debug and analyze training jobs in Amazon SageMaker
- Use Amazon SageMaker Debugger to debug and improve model performance
- Access a training container through AWS Systems Manager for remote debugging
- Release notes for debugging capabilities of Amazon SageMaker

Use TensorBoard to debug and analyze training jobs in Amazon SageMaker

Amazon SageMaker with TensorBoard is a capability of Amazon SageMaker that brings the visualization tools of TensorBoard to SageMaker, integrated with SageMaker Training and domain. It provides options to administer your AWS account and users belonging to the account through SageMaker domain, to give the domain users access to the TensorBoard data with appropriate permissions to Amazon S3, and help the domain users perform model debugging tasks using the TensorBoard visualization plugins. SageMaker with TensorBoard is extended with the SageMaker Data Manager plugin, with which domain users can access a number of training jobs in one place within the TensorBoard application.
**Note**

This feature is for training and debugging deep learning models using the PyTorch or TensorFlow framework.

---

**For data scientists**

Training large models can have scientific problems that require data scientists to debug and resolve them in order to improve model convergence and stabilize gradient descent processes.

When you encounter model training issues, such as loss not converging, or vanishing or exploding weights and gradients, you need to access tensor data to dive deep and analyze the model parameters, scalars, and any custom metrics. Using SageMaker with TensorBoard, you can visualize model output tensors extracted from training jobs. As you experiment with different models, multiple training runs, and model hyperparameters, you can select multiple training jobs in TensorBoard and compare them in one place.

**For administrators**

Through the TensorBoard landing page in the SageMaker console or [SageMaker domain](https://aws.amazon.com/sagemaker/), you can manage TensorBoard application users if you are an administrator of an AWS account or SageMaker domain. Each domain user can access their own TensorBoard application given the granted permissions. As a SageMaker domain administrator and domain user, you can create and delete the TensorBoard application given the permission level you have.

**Supported frameworks and AWS Regions**

This feature supports the following machine learning frameworks and AWS Regions.

**Frameworks**

- PyTorch
- TensorFlow
- Hugging Face Transformers

**AWS Regions**

- US East (N. Virginia) (us-east-1)
• US East (Ohio) (us-east-2)
• US West (Oregon) (us-west-2)
• Europe (Frankfurt) (eu-central-1)
• Europe (Ireland) (eu-west-1)

**Note**

Amazon SageMaker with TensorBoard runs the TensorBoard application on an ml.r5.large instance and incurs charges after the SageMaker free tier or the free trial period of the feature. For more information, see [Amazon SageMaker Pricing](#).

**Prerequisites**

The following list shows the prerequisites to start using SageMaker with TensorBoard.

• A SageMaker domain that's set up with Amazon VPC in your AWS account.

  For instructions on setting up a domain, see [Onboard to Amazon SageMaker domain using quick setup](#). You also need to add domain user profiles for individual users to access the TensorBoard on SageMaker. For more information, see [Add and remove SageMaker domain user profiles](#).

• The following list is the minimum set of permissions for using TensorBoard on SageMaker.
  
  • sagemaker:CreateApp
  • sagemaker:DeleteApp
  • sagemaker:DescribeTrainingJob
  • sagemaker:Search
  • s3:GetObject
  • s3:ListBucket

**Prepare a training job with a TensorBoard output data configuration**

A typical training job for deep learning in SageMaker consists of two main steps: preparing a training script and configuring a SageMaker Training job launcher. In this section, you can check the required changes to collect TensorBoard-compatible data from SageMaker Training.
Step 1: Modify your training script

Make sure you determine which output tensors and scalars to collect, and modify code lines in your training script using any of the following tools: TensorBoardX, TensorFlow Summary Writer, PyTorch Summary Writer, or SageMaker Debugger.

Also make sure that you specify the TensorBoard data output path as the log directory (`log_dir`) for callback in the training container.

For more information about callbacks per framework, see the following resources.

- For PyTorch, use `torch.utils.tensorboard.SummaryWriter`. See also the **Using TensorBoard in PyTorch** and **Log scalars** sections in the **PyTorch tutorials**. Alternatively, you can use **TensorBoardX Summary Writer**.

  ```python
  LOG_DIR="/opt/ml/output/tensorboard"
  tensorboard_callback=torch.utils.tensorboard.writer.SummaryWriter(log_dir=LOG_DIR)
  ```

- For TensorFlow, use the native callback for TensorBoard, `tf.keras.callbacks.TensorBoard`.

  ```python
  LOG_DIR="/opt/ml/output/tensorboard"
  tensorboard_callback=tf.keras.callbacks.TensorBoard(
                  log_dir=LOG_DIR, histogram_freq=1)
  ```

- For Transformers with PyTorch, you can use `transformers.integrations.TensorBoardCallback`.

  For Transformers with TensorFlow, use the `tf.keras.tensorboard.callback`, and pass that to the keras callback in transformers.

Tip

You can also use a different container local output path. However, in **Step 2: Construct a SageMaker training launcher with TensorBoard data configuration**, you must map the paths correctly for SageMaker to successfully search the local path and save the TensorBoard data to the S3 output bucket.

- For guidance on modifying training scripts using the SageMaker Debugger Python library, see the section called “Step 1: Adapt Your Training Script to Register a Hook”.
Step 2: Construct a SageMaker training launcher with TensorBoard data configuration

Use the `sagemaker.debugger.TensorBoardOutputConfig` while configuring a SageMaker framework estimator. This configuration API maps the S3 bucket you specify for saving TensorBoard data with the local path in the training container (`/opt/ml/output/tensorboard`). Pass the object of the module to the `tensorboard_output_config` parameter of the estimator class. The following code snippet shows an example of preparing a TensorFlow estimator with the TensorBoard output configuration parameter.

```python
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import TensorBoardOutputConfig

# Set variables for training job information, # such as s3_out_bucket and other unique tags.
...

LOG_DIR = '/opt/ml/output/tensorboard'

output_path = os.path.join(
    's3_output_bucket', 'sagemaker-output', 'date_str', 'your-training_job_name'
)

tensorboard_output_config = TensorBoardOutputConfig(
    s3_output_path=os.path.join(output_path, 'tensorboard'),
    container_local_output_path=LOG_DIR
)

estimator = TensorFlow(
    entry_point='train.py',

Use TensorBoard
```
source_dir="src",
    role=role,
    image_uri=image_uri,
    instance_count=1,
    instance_type="ml.c5.xlarge",
    base_job_name="your-training_job_name",
    tensorboard_output_config=tensorboard_output_config,
    hyperparameters=hyperparameters
)

How to access TensorBoard on SageMaker

You can access TensorBoard by two methods: programmatically using the sagemaker.interactive_apps.tensorboard module that generates an unsigned or a presigned URL, or using the TensorBoard landing page in the SageMaker console. After you open TensorBoard, SageMaker runs the TensorBoard plugin and automatically finds all training job output data in TensorBoard-compatible file format.

Topics

• Open TensorBoard using the sagemaker.interactive_apps.tensorboard module
• Open TensorBoard using the get_app_url function as an estimator class method
• Open TensorBoard through the SageMaker console

Open TensorBoard using the sagemaker.interactive_apps.tensorboard module

The sagemaker.interactive_apps.tensorboard module provides a function called get_app_url that generates unsigned or presigned URLs to open the TensorBoard application in any environment in SageMaker or Amazon EC2. This is to provide a unified experience for both Studio Classic and non-Studio Classic users. For the Studio environment, you can open TensorBoard by running the get_app_url() function as it is, or you can also specify a job name to start tracking as the TensorBoard application opens. For non-Studio Classic environments, you can open TensorBoard by providing your domain and user profile information to the utility function. With this functionality, regardless of where or how you run training code and launch training jobs, you can directly access TensorBoard by running the get_app_url function in your Jupyter notebook or terminal.
Note

This functionality is available in the SageMaker Python SDK v2.184.0 and later. To use this functionality, make sure that you upgrade the SDK by running `pip install sagemaker --upgrade`.

Topics

- Option 1: For SageMaker Studio Classic
- Option 2: For non-Studio Classic environments

Option 1: For SageMaker Studio Classic

If you are using SageMaker Studio Classic, you can directly open the TensorBoard application or retrieve an unsigned URL by running the `get_app_url` function as follows. As you are already within the Studio Classic environment and signed in as a domain user, `get_app_url()` generates unsigned URL because it is not necessary to authenticate again.

To open the TensorBoard application

The following code automatically opens the TensorBoard application from the unsigned URL that the `get_app_url()` function returns in the your environment's default web browser.

```python
from sagemaker.interactive_apps import tensorboard
region = "us-west-2"
app = tensorboard.TensorBoardApp(region)
app.get_app_url(
    training_job_name="your-training_job_name" # Optional. Specify the job name to track a specific training job
)
```

To retrieve an unsigned URL and open the TensorBoard application manually

The following code prints an unsigned URL that you can copy to a web browser and open the TensorBoard application.

```python
from sagemaker.interactive_apps import tensorboard
```
region = "us-west-2"
app = tensorboard.TensorBoardApp(region)
print("Navigate to the following URL:")
print(
    app.get_app_url(
        training_job_name="your-training_job_name", # Optional. Specify the name of the
        job to track.
        open_in_default_web_browser=False # Set to False to print the URL to
        terminal.
    )
)

Note that if you run the preceding two code samples outside the SageMaker Studio Classic
environment, the function will return a URL to the TensorBoard landing page in the SageMaker
console, because these do not have sign-in information to your domain and user profile. For
creating a presigned URL, see Option 2 in the following section.

**Option 2: For non-Studio Classic environments**

If you use non-Studio Classic environments, such as SageMaker Notebook instance or Amazon EC2,
and want to open TensorBoard directly from the environment you are in, you need to generate
a URL presigned with your domain and user profile information. A *presigned* URL is a URL that's
signed in to Amazon SageMaker Studio Classic while the URL is being created with your domain
and user profile, and therefore granted access to all of the domain applications and files associated
with your domain. To open TensorBoard through a presigned URL, use the `get_app_url` function
with your domain and user profile name as follows.

Note that this option requires the domain user to have the
`sagemaker:CreatePresignedDomainUrl` permission. Without the permission, the domain user
will receive an exception error.

⚠️ **Important**

Do not share any presigned URLs. The `get_app_url` function creates presigned URLs,
which automatically authenticates with your domain and user profile and gives access to
any applications and files associated with your domain.
app.get_app_url(
    training_job_name="your-training_job_name", # Optional. Specify the name of the
    job to track.
    create_presigned_domain_url=True, # Required to be set to True for
    domain_id="your-domain-id", # Required if creating a presigned
    URL (create_presigned_domain_url=True).
    user_profile_name="your-user-profile-name", # Required if creating a presigned
    URL (create_presigned_domain_url=True).
    open_in_default_web_browser=False, # Optional. Set to False to print
    the URL to terminal.
    optional_create_presigned_url_kwargs={} # Optional. Add any additional args
    for Boto3 create_presigned_domain_url
)

Tip

The get_app_url function runs the
SageMaker.Client.create_presigned_domain_url API in the AWS SDK for Python
(Boto3) in the backend. As the Boto3 create_presigned_domain_url API creates
presigned domain URLs that expire in 300 seconds by default, presigned TensorBoard
application URLs also expire in 300 seconds. If you want to extend the expiration time, pass
the ExpiresInSeconds argument to the optional_create_presigned_url_kwargs argument of the get_app_url function as follows.

optional_create_presigned_url_kwargs={"ExpiresInSeconds": 1500}

Note

If any of your input passed to the arguments of get_app_url is invalid, the function
outputs a URL to the TensorBoard landing page instead of opening the TensorBoard
application. The output message would be similar to the following.

Navigate to the following URL:
Open TensorBoard using the `get_app_url` function as an estimator class method

If you are in the process of running a training job using the estimator class of the SageMaker Python SDK and have an active object of the estimator class, you can also access the `get_app_url` function as a class method of the estimator class. Open the TensorBoard application or retrieve an unsigned URL by running the `get_app_url` method as follows. The `get_app_url` class method pulls the training job name from the estimator and opens the TensorBoard application with the specified job.

⚠️ Note

This functionality is available in the SageMaker Python SDK v2.184.0 and later. To use this functionality, make sure that you upgrade the SDK by running `pip install sagemaker --upgrade`.

Topics

- Option 1: For SageMaker Studio Classic
- Option 2: For non-Studio Classic environments

Option 1: For SageMaker Studio Classic

To open the TensorBoard application

The following code automatically opens the TensorBoard application from the unsigned URL that the `get_app_url()` method returns in the your environment's default web browser.

```python
estimator.get_app_url(
    app_type=SupportedInteractiveAppTypes.TENSORBOARD # Required.
)
```

To retrieve an unsigned URL and open the TensorBoard application manually

The following code prints an unsigned URL that you can copy to a web browser and open the TensorBoard application.

```python
print(estimator.get_app_url())
```
Note that if you run the preceding two code samples outside the SageMaker Studio Classic environment, the function will return a URL to the TensorBoard landing page in the SageMaker console, because these do not have sign-in information to your domain and user profile. For creating a presigned URL, see Option 2 in the following section.

**Option 2: For non-Studio Classic environments**

If you use non-Studio Classic environments, such as SageMaker Notebook instance and Amazon EC2, and want to generate a presigned URL to open the TensorBoard application, use the `get_app_url` method with your domain and user profile information as follows.

Note that this option requires the domain user to have the `sagemaker:CreatePresignedDomainUrl` permission. Without the permission, the domain user will receive an exception error.

**Important**

Do not share any presigned URLs. The `get_app_url` function creates presigned URLs, which automatically authenticates with your domain and user profile and gives access to any applications and files associated with your domain.
Open TensorBoard through the SageMaker console

You can also use the SageMaker console UI to open the TensorBoard application. There are two options to open the TensorBoard application through the SageMaker console.

Topics

- Option 1: Launch TensorBoard from the domain details page
- Option 2: Launch TensorBoard from the TensorBoard landing page

Option 1: Launch TensorBoard from the domain details page

Navigate to the domain details page

The following procedure shows how to navigate to the domain details page.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. From the list of domains, select the domain in which you want to launch the TensorBoard application.

Launch a user profile application

The following procedure shows how to launch a Studio Classic application that is scoped to a user profile.

1. On the domain details page, choose the User profiles tab.
2. Identify the user profile for which you want to launch the Studio Classic application.
3. Choose Launch for your selected user profile, then choose TensorBoard.

Option 2: Launch TensorBoard from the TensorBoard landing page

The following procedure describes how to launch a TensorBoard application from the TensorBoard landing page.
1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.

2. On the left navigation pane, choose TensorBoard.

3. Under Get started, select the domain in which you want to launch the Studio Classic application. If your user profile only belongs to one domain, you do not see the option for selecting a domain.

4. Select the user profile for which you want to launch the Studio Classic application. If there is no user profile in the domain, choose Create user profile. For more information, see Add and Remove User Profiles.

5. Choose Open TensorBoard.

The following screenshot shows the location of TensorBoard in the left navigation pane of the SageMaker console and the SageMaker with TensorBoard landing page in the main pane.

Access and visualize training output data in TensorBoard

You can conduct an online or offline analysis by loading collected output tensors from S3 buckets paired with training jobs during or after training.

When you open the TensorBoard application, TensorBoard opens with the SageMaker Data Manager tab. The following screenshot shows the full view of the SageMaker Data Manager tab in the TensorBoard application.
In the **SageMaker Data Manager** tab, you can select any training job and load TensorBoard-compatible training output data from Amazon S3.

1. In the **Search training jobs** section, use the filters to narrow down the list of training jobs you want to find, load, and visualize.

2. In the **List of training jobs** section, use the check boxes to choose training jobs from which you want to pull data and visualize for debugging.

3. Choose **Add selected jobs**. The selected jobs should appear in the **Tracked training jobs** section, as shown in the following screenshot.
The SageMaker Data Manager tab only shows training jobs configured with the TensorBoardOutputConfig parameter. Make sure you have configured the SageMaker estimator with this parameter. For more information, see Step 2: Construct a SageMaker training launcher with TensorBoard data configuration.

The visualization tabs might not appear if you are using SageMaker with TensorBoard for the first time or no data is loaded from a previous use. After adding training jobs and waiting for a few seconds, refresh the viewer by choosing the clockwise circular arrow on the upper-right corner. The visualization tabs should appear after the job data are successfully loaded. You can also set to auto-refresh using the Settings button next to the refresh button in the upper right corner.

Explore training output data visualized in TensorBoard

In the graphics tabs, you can see the list of the loaded training jobs in the left pane. You can also use the check boxes of the training jobs to show or hide visualizations. The TensorBoard dynamic
plugins are activated dynamically depending on how you have set your training script to include summary writers and pass callbacks for tensor and scalar collection, and therefore the graphics tabs also appear dynamically. The following screenshots show example views of each tab with visualization of two training jobs that collected metrics for time series, scalar, graph, distribution, and histogram plugins.

The **TIME SERIES** tab view

![Time Series Tab View](image)

The **SCALARS** tab view

![Scalars Tab View](image)
The GRAPHS tab view
The DISTRIBUTIONS tab view

The HISTOGRAMS tab view
Delete unused TensorBoard applications

After you are done with monitoring and experimenting with jobs in TensorBoard, shut the TensorBoard application down.

1. Open the SageMaker console.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Choose your domain.
5. Choose your user profile.
7. Choose Yes, delete app.
8. Type delete in the text box, then choose Delete.
9. A blue message should appear at the top of the screen: default is being deleted.

Considerations

Consider the following when using SageMaker with TensorBoard.
• You cannot share the TensorBoard applications for collaboration purposes because SageMaker domain does not allow application sharing among users. Users can share the output tensors saved in an S3 bucket, if they have access to the bucket.

• The visualization plugins might not appear when you first launch the TensorBoard application. After you select training jobs in the SageMaker Data Manager plugin, the TensorBoard application loads the TensorBoard data and populates the visualization plugins.

• The TensorBoard applications automatically shuts down after 1 hour of inactivity. If you want to shut the application down when you are done using it, make sure to manually shut down TensorBoard to avoid paying for the instance hosting it. For instructions on deleting the application, see *Delete unused TensorBoard applications*.

### Use Amazon SageMaker Debugger to debug and improve model performance

Debug model output tensors from machine learning training jobs in real time and detect non-converging issues using Amazon SageMaker Debugger.

### Amazon SageMaker Debugger Features

A machine learning (ML) training job can have problems such as overfitting, saturated activation functions, and vanishing gradients, which can compromise model performance.

SageMaker Debugger provides tools to debug training jobs and resolve such problems to improve the performance of your model. Debugger also offers tools to send alerts when training anomalies are found, take actions against the problems, and identify the root cause of them by visualizing collected metrics and tensors.

SageMaker Debugger supports the Apache MXNet, PyTorch, TensorFlow, and XGBoost frameworks. For more information about available frameworks and versions supported by SageMaker Debugger, see *Supported Frameworks and Algorithms*.
The high-level Debugger workflow is as follows:

1. Modify your training script with the `sagemaker-debugger` Python SDK if needed.
2. Configure a SageMaker training job with SageMaker Debugger.
   - Configure using the SageMaker Estimator API (for Python SDK).
   - Configure using the SageMaker `CreateTrainingJob` request (for Boto3 or CLI).
   - Configure custom training containers with SageMaker Debugger.
3. Start a training job and monitor training issues in real time.
   - List of Debugger Built-in Rules.
4. Get alerts and take prompt actions against the training issues.
   - Receive texts and emails and stop training jobs when training issues are found using Debugger Built-in Actions for Rules.
   - Set up your own actions using Amazon CloudWatch Events and AWS Lambda.
5. Explore deep analysis of the training issues.
   - For debugging model output tensors, see Visualize Debugger Output Tensors in TensorBoard.
6. Fix the issues, consider the suggestions provided by Debugger, and repeat steps 1–5 until you optimize your model and achieve target accuracy.

The SageMaker Debugger developer guide walks you through the following topics.

**Topics**

- Supported Frameworks and Algorithms
Supported Frameworks and Algorithms

The following table shows SageMaker machine learning frameworks and algorithms supported by Debugger.

<table>
<thead>
<tr>
<th>SageMaker-supported frameworks and algorithms</th>
<th>Debugging output tensors</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TensorFlow</strong></td>
<td><strong>AWS TensorFlow deep learning containers</strong> 1.15.4 or later</td>
</tr>
<tr>
<td><strong>PyTorch</strong></td>
<td><strong>AWS PyTorch deep learning containers</strong> 1.5.0 or later</td>
</tr>
<tr>
<td><strong>MXNet</strong></td>
<td><strong>AWS MXNet deep learning containers</strong> 1.6.0 or later</td>
</tr>
<tr>
<td><strong>XGBoost</strong></td>
<td>1.0-1, 1.2-1, 1.3-1</td>
</tr>
<tr>
<td><strong>SageMaker generic estimator</strong></td>
<td><strong>Custom training containers</strong> (available for TensorFlow, PyTorch, MXNet, and XGBoost with manual hook registration)</td>
</tr>
</tbody>
</table>

- **Debugging output tensors** – Track and debug model parameters, such as weights, gradients, biases, and scalar values of your training job. Available deep learning frameworks are Apache MXNet, TensorFlow, PyTorch, and XGBoost.
Important
For the TensorFlow framework with Keras, SageMaker Debugger deprecates the zero code change support for debugging models built using the `tf.keras` modules of TensorFlow 2.6 and later. This is due to breaking changes announced in the TensorFlow 2.6.0 release note. For instructions on how to update your training script, see the section called “TensorFlow”.

Important
From PyTorch v1.12.0 and later, SageMaker Debugger deprecates the zero code change support for debugging models. This is due to breaking changes that cause SageMaker Debugger to interfere with the `torch.jit` functionality. For instructions on how to update your training script, see the section called “PyTorch”.

If the framework or algorithm that you want to train and debug is not listed in the table, go to the AWS Discussion Forum and leave feedback on SageMaker Debugger.

AWS Regions

Amazon SageMaker Debugger is available in all regions where Amazon SageMaker is in service except the following region.

- Asia Pacific (Jakarta): ap-southeast-3

To find if Amazon SageMaker is in service in your AWS Region, see AWS Regional Services.

Use Debugger with Custom Training Containers

Bring your training containers to SageMaker and gain insights into your training jobs using Debugger. Maximize your work efficiency by optimizing your model on Amazon EC2 instances using the monitoring and debugging features.
For more information about how to build your training container with the sagemaker-debugger client library, push it to the Amazon Elastic Container Registry (Amazon ECR), and monitor and debug, see Use Debugger with Custom Training Containers.

Debugger Open-Source GitHub Repositories

Debugger APIs are provided through the SageMaker Python SDK and designed to construct Debugger hook and rule configurations for the SageMaker CreateTrainingJob and DescribeTrainingJob API operations. The sagemaker-debugger client library provides tools to register hooks and access the training data through its trial feature, all through its flexible and powerful API operations. It supports the machine learning frameworks TensorFlow, PyTorch, MXNet, and XGBoost on Python 3.6 and later.

For direct resources about the Debugger and sagemaker-debugger API operations, see the following links:

- [The Amazon SageMaker Python SDK documentation](#)
- [The Amazon SageMaker Python SDK - Debugger APIs](#)
- [The sagemaker-debugger Python SDK documentation](#) for the Amazon SageMaker Debugger open source client library
- [The sagemaker-debugger PyPI](#)

If you use the SDK for Java to conduct SageMaker training jobs and want to configure Debugger APIs, see the following references:

- [Amazon SageMaker Debugger API Operations](#)
- [Configure Debugger Using Amazon SageMaker API](#)

Amazon SageMaker Debugger Architecture

This topic walks you through a high-level overview of the Amazon SageMaker Debugger workflow.

Debugger supports profiling functionality for performance optimization to identify computation issues, such as system bottlenecks and underutilization, and to help optimize hardware resource utilization at scale.
Debugger's debugging functionality for *model optimization* is about analyzing non-converging training issues that can arise while minimizing the loss functions using optimization algorithms, such as gradient descent and its variations.

The following diagram shows the architecture of SageMaker Debugger. The blocks with bold boundary lines are what Debugger manages to analyze your training job.
Debugger stores the following data from your training jobs in your secured Amazon S3 bucket:

- **Output tensors** – Collections of scalars and model parameters that are continuously updated during the forward and backward passes while training ML models. The output tensors include...
scalar values (accuracy and loss) and matrices (weights, gradients, input layers, and output layers).

Note

By default, Debugger monitors and debugs SageMaker training jobs without any Debugger-specific parameters configured in SageMaker estimators. Debugger collects system metrics every 500 milliseconds and basic output tensors (scalar outputs such as loss and accuracy) every 500 steps. It also runs the ProfilerReport rule to analyze the system metrics and aggregate the Studio Debugger insights dashboard and a profiling report. Debugger saves the output data in your secured Amazon S3 bucket.

The Debugger built-in rules run on processing containers, which are designed to evaluate machine learning models by processing the training data collected in your S3 bucket (see Process Data and Evaluate Models). The built-in rules are fully managed by Debugger. You can also create your own rules customized to your model to watch for any issues you want to monitor.

Get Started with Debugger Tutorials

The following topics walk you through tutorials from the basics to advanced use cases of monitoring, profiling, and debugging SageMaker training jobs using Debugger. Explore the Debugger features and learn how you can debug and improve your machine learning models efficiently by using Debugger.

Topics

- Debugger Tutorial Videos
- Debugger Example Notebooks
- Debugger Advanced Demos and Visualization

Debugger Tutorial Videos

The following videos provide a tour of Amazon SageMaker Debugger capabilities using SageMaker Studio and SageMaker notebook instances.

Topics

- Debug Models with Amazon SageMaker Debugger in Studio
• Deep Dive on Amazon SageMaker Debugger and SageMaker Model Monitor

Debug Models with Amazon SageMaker Debugger in Studio

*Julien Simon, AWS Technical Evangelist | Length: 14 minutes 17 seconds*

This tutorial video demonstrates how to use Amazon SageMaker Debugger to capture and inspect debugging information from a training model. The example training model used in this video is a simple convolutional neural network (CNN) based on Keras with the TensorFlow backend. SageMaker in a TensorFlow framework and Debugger enable you to build an estimator directly using the training script and debug the training job.

**Debug Models with Amazon SageMaker Debugger (part 1)**

You can find the example notebook in the video in [this Studio Demo repository](#) provided by the author. You need to clone the debugger.ipynb notebook file and the mnist_keras_tf.py training script to your SageMaker Studio or a SageMaker notebook instance. After you clone the two files, specify the path `keras_script_path` to the `mnist_keras_tf.py` file inside the debugger.ipynb notebook. For example, if you cloned the two files in the same directory, set it as `keras_script_path = "mnist_keras_tf.py"`.

**Deep Dive on Amazon SageMaker Debugger and SageMaker Model Monitor**

*Julien Simon, AWS Technical Evangelist | Length: 44 minutes 34 seconds*

This video session explores advanced features of Debugger and SageMaker Model Monitor that help boost productivity and the quality of your models. First, this video shows how to detect and fix training issues, visualize tensors, and improve models with Debugger. Next, at 22:41, the video shows how to monitor models in production and identify prediction issues such as missing features or data drift using SageMaker Model Monitor. Finally, it offers cost optimization tips to help you make the most of your machine learning budget.

**Debug Models with Debugger (part 2)**

You can find the example notebook in the video in [this AWS Dev Days 2020 repository](#) offered by the author.
Debugger Example Notebooks

*SageMaker Debugger example notebooks* are provided in the `aws/amazon-sagemaker-examples` repository. The Debugger example notebooks walk you through basic to advanced use cases of debugging and profiling training jobs.

We recommend that you run the example notebooks on SageMaker Studio or a SageMaker Notebook instance because most of the examples are designed for training jobs in the SageMaker ecosystem, including Amazon EC2, Amazon S3, and Amazon SageMaker Python SDK.

To clone the example repository to SageMaker Studio, follow the instructions at *Amazon SageMaker Studio Tour*.

To find the examples in a SageMaker Notebook instance, follow the instructions at *SageMaker Notebook Instance Example Notebooks*.

⚠️ Important

To use the new Debugger features, you need to upgrade the SageMaker Python SDK and the SMDebug client library. In your iPython kernel, Jupyter Notebook, or JupyterLab environment, run the following code to install the latest versions of the libraries and restart the kernel.

```python
import sys
import IPython
!{sys.executable} -m pip install -U sagemaker smdebug
IPython.Application.instance().kernel.do_shutdown(True)
```

Debugger Example Notebooks for Profiling Training Jobs

The following list shows Debugger example notebooks introducing Debugger's adaptability to monitor and profile training jobs for various machine learning models, datasets, and frameworks.

<table>
<thead>
<tr>
<th>Notebook Title</th>
<th>Framework</th>
<th>Model</th>
<th>Dataset</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amazon SageMaker</td>
<td>TensorFlow</td>
<td>Keras</td>
<td>Cifar-10</td>
<td>This notebook provides an introduction to interact</td>
</tr>
<tr>
<td>Notebook Title</td>
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<td>Dataset</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-----------</td>
<td>-------</td>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td>Debugger Profiling Data Analysis</td>
<td>TensorFlow</td>
<td>1-D Convolutional Neural Network</td>
<td>IMDB dataset</td>
<td>Interactive analysis of profiled data captured by SageMaker Debugger. Explore the full functionality of the SMDebug interactive analysis tools.</td>
</tr>
<tr>
<td>Profile machine learning training with Amazon SageMaker Debugger</td>
<td>TensorFlow</td>
<td>ResNet50</td>
<td>Cifar-10</td>
<td>Profile a TensorFlow 1-D CNN for sentiment analysis of IMDB data that consists of movie reviews labeled as having positive or negative sentiment. Explore the Studio Debugger insights and Debugger profiling report.</td>
</tr>
<tr>
<td>Profiling PyTorch ResNet model training with various distributed training settings</td>
<td>PyTorch</td>
<td>ResNet50</td>
<td>Cifar-10</td>
<td>Run PyTorch training jobs with various distributed training settings, monitor system resource utilization, and profile model performance using Debugger.</td>
</tr>
</tbody>
</table>
Debugger Example Notebooks for Analyzing Model Parameters

The following list shows Debugger example notebooks introducing Debugger's adaptability to debug training jobs for various machine learning models, datasets, and frameworks.

<table>
<thead>
<tr>
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<th>Framework</th>
<th>Model</th>
<th>Dataset</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amazon SageMaker Debugger - Use built-in rule</td>
<td>TensorFlow</td>
<td>Convolutional Neural Network</td>
<td>MNIST</td>
<td>Use the Amazon SageMaker Debugger built-in rules for debugging a TensorFlow model.</td>
</tr>
<tr>
<td>Amazon SageMaker Debugger - Tensorflow 2.1</td>
<td>TensorFlow</td>
<td>ResNet50</td>
<td>Cifar-10</td>
<td>Use the Amazon SageMaker Debugger hook configuration and built-in rules for debugging a model with the Tensorflow 2.1 framework.</td>
</tr>
<tr>
<td>Visualizing Debugging Tensors of MXNet training</td>
<td>MXNet</td>
<td>Gluon Convolutional Neural Network</td>
<td>Fashion MNIST</td>
<td>Run a training job and configure SageMaker Debugger to store all tensors from this job, then visualize those tensors in a notebook.</td>
</tr>
<tr>
<td>Enable Spot Training with Amazon SageMaker Debugger</td>
<td>MXNet</td>
<td>Gluon Convolutional Neural Network</td>
<td>Fashion MNIST</td>
<td>Learn how Debugger collects tensor data from a training job on a spot instance, and how to use the Debugger built-in rules with managed spot training.</td>
</tr>
<tr>
<td>Explain an XGBoost model that predicts an individual</td>
<td>XGBoost</td>
<td>XGBoost Regression</td>
<td>Adult Census dataset</td>
<td>Learn how to use the Debugger hook and built-in rules for collecting and visualizing tensor data from an XGBoost regression model,</td>
</tr>
</tbody>
</table>
To find advanced visualizations of model parameters and use cases, see the next topic at Debugger Advanced Demos and Visualization.

**Debugger Advanced Demos and Visualization**

The following demos walk you through advanced use cases and visualization scripts using Debugger.

**Topics**

- [Train and Tune Your Models with Amazon SageMaker Experiments and Debugger](#)
- [Using SageMaker Debugger to Monitor a Convolutional Autoencoder Model Training](#)
- [Using SageMaker Debugger to Monitor Attentions in BERT Model Training](#)
- [Using SageMaker Debugger to Visualize Class Activation Maps in Convolutional Neural Networks (CNNs)](#)

**Train and Tune Your Models with Amazon SageMaker Experiments and Debugger**

*Dr. Nathalie Rauschmayr, AWS Applied Scientist | Length: 49 minutes 26 seconds*

**Train and Prune Models with SageMaker Experiments and Debugger**

Find out how Amazon SageMaker Experiments and Debugger can simplify the management of your training jobs. Amazon SageMaker Debugger provides transparent visibility into training jobs and saves training metrics into your Amazon S3 bucket. SageMaker Experiments enables you to call the training information as *trials* through SageMaker Studio and supports visualization of the training job. This helps you keep model quality high while reducing less important parameters based on importance rank.

This video demonstrates a *model pruning* technique that makes pre-trained ResNet50 and AlexNet models lighter and affordable while keeping high standards for model accuracy.
SageMaker Estimator trains those algorithms supplied from the PyTorch model zoo in an AWS Deep Learning Containers with PyTorch framework, and Debugger extracts training metrics from the training process.

The video also demonstrates how to set up a Debugger custom rule to watch the accuracy of a pruned model, to trigger an Amazon CloudWatch event and an AWS Lambda function when the accuracy hits a threshold, and to automatically stop the pruning process to avoid redundant iterations.

Learning objectives are as follows:

- Learn how to use SageMaker to accelerate ML model training and improve model quality.
- Understand how to manage training iterations with SageMaker Experiments by automatically capturing input parameters, configurations, and results.
- Discover how Debugger makes the training process transparent by automatically capturing real-time tensor data from metrics such as weights, gradients, and activation outputs of convolutional neural networks.
- Use CloudWatch to trigger Lambda when Debugger catches issues.
- Master the SageMaker training process using SageMaker Experiments and Debugger.

You can find the notebooks and training scripts used in this video from SageMaker Debugger PyTorch Iterative Model Pruning.

The following image shows how the iterative model pruning process reduces the size of AlexNet by cutting out the 100 least significant filters based on importance rank evaluated by activation outputs and gradients.

The pruning process reduced the initial 50 million parameters to 18 million. It also reduced the estimated model size from 201 MB to 73 MB.
You also need to track model accuracy, and the following image shows how you can plot the model pruning process to visualize changes in model accuracy based on the number of parameters in SageMaker Studio.
In SageMaker Studio, choose the **Experiments** tab, select a list of tensors saved by Debugger from the pruning process, and then compose a **Trial Component List** panel. Select all ten iterations and then choose **Add chart** to create a **Trial Component Chart**. After you decide on a model to deploy, choose the trial component and choose a menu to perform an action or choose **Deploy model**.
Note

To deploy a model through SageMaker Studio using the following notebook example, add a line at the end of the `train` function in the `train.py` script.

```python
# In the train.py script, look for the train function in line 58.
def train(epochs, batch_size, learning_rate):
    ...
    print('acc:{:.4f}'.format(correct/total))
    hook.save_scalar("accuracy", correct/total, sm_metric=True)

    # Add the following code to line 128 of the train.py script to save the pruned models
    # under the current SageMaker Studio model directory
    torch.save(model.state_dict(), os.environ['SM_MODEL_DIR'] + '/model.pt')
```

Using SageMaker Debugger to Monitor a Convolutional Autoencoder Model Training

This notebook demonstrates how SageMaker Debugger visualizes tensors from an unsupervised (or self-supervised) learning process on a MNIST image dataset of handwritten numbers.

The training model in this notebook is a convolutional autoencoder with the MXNet framework. The convolutional autoencoder has a bottleneck-shaped convolutional neural network that consists of an encoder part and a decoder part.

The encoder in this example has two convolution layers to produce compressed representation (latent variables) of the input images. In this case, the encoder produces a latent variable of size (1, 20) from an original input image of size (28, 28) and significantly reduces the size of data for training by 40 times.

The decoder has two deconvolutional layers and ensures that the latent variables preserve key information by reconstructing output images.

The convolutional encoder powers clustering algorithms with smaller input data size and the performance of clustering algorithms such as k-means, k-NN, and t-Distributed Stochastic Neighbor Embedding (t-SNE).
This notebook example demonstrates how to visualize the latent variables using Debugger, as shown in the following animation. It also demonstrates how the t-SNE algorithm classifies the latent variables into ten clusters and projects them into a two-dimensional space. The scatter plot color scheme on the right side of the image reflects the true values to show how well the BERT model and t-SNE algorithm organize the latent variables into the clusters.

Using SageMaker Debugger to Monitor Attentions in BERT Model Training

Bidirectional Encode Representations from Transformers (BERT) is a language representation model. As the name of model reflects, the BERT model builds on transfer learning and the Transformer model for natural language processing (NLP).

The BERT model is pre-trained on unsupervised tasks such as predicting missing words in a sentence or predicting the next sentence that naturally follows a previous sentence. The training data contains 3.3 billion words (tokens) of English text, from sources such as Wikipedia and electronic books. For a simple example, the BERT model can give a high attention to appropriate verb tokens or pronoun tokens from a subject token.

The pre-trained BERT model can be fine-tuned with an additional output layer to achieve state-of-the-art model training in NLP tasks, such as automated responses to questions, text classification, and many others.
Debugger collects tensors from the fine-tuning process. In the context of NLP, the weight of neurons is called *attention*.

This notebook demonstrates how to use the [pre-trained BERT model from the GluonNLP model zoo](https://gluon-nlp.mxnet.io/hub.html) on the Stanford Question and Answering dataset and how to set up SageMaker Debugger to monitor the training job.

Plotting *attention scores* and individual neurons in the query and key vectors can help to identify causes of incorrect model predictions. With SageMaker Debugger, you can retrieve the tensors and plot the *attention-head view* in real time as training progresses and understand what the model is learning.

The following animation shows the attention scores of the first 20 input tokens for ten iterations in the training job provided in the notebook example.
Using SageMaker Debugger to Visualize Class Activation Maps in Convolutional Neural Networks (CNNs)

This notebook demonstrates how to use SageMaker Debugger to plot class activation maps for image detection and classification in convolutional neural networks (CNNs). In deep learning, a convolutional neural network (CNN or ConvNet) is a class of deep neural networks, most commonly applied to analyzing visual imagery. One of the applications that adopts the class activation maps is self-driving cars, which require instantaneous detection and classification of images such as traffic signs, roads, and obstacles.

In this notebook, the PyTorch ResNet model is trained on the German Traffic Sign Dataset, which contains more than 40 classes of traffic-related objects and more than 50,000 images in total.

During the training process, SageMaker Debugger collects tensors to plot the class activation maps in real time. As shown in the animated image, the class activation map (also called as a saliency map) highlights regions with high activation in red color.

Using tensors captured by Debugger, you can visualize how the activation map evolves during the model training. The model starts by detecting the edge on the lower-left corner at the beginning of the training job. As the training progresses, the focus shifts to the center and detects the speed limit sign, and the model successfully predicts the input image as Class 3, which is a class of speed limit 60km/h signs, with a 97% confidence level.
Debug Training Jobs Using Amazon SageMaker Debugger

To prepare your training script and run training jobs with SageMaker Debugger to debug model training progress, you follow the typical two-step process: modify your training script using the sagemaker-debugger Python SDK, and construct a SageMaker estimator using the SageMaker Python SDK. Go through the following topics to learn how to use SageMaker Debugger's debugging functionality.

Topics

- **Step 1: Adapt Your Training Script to Register a Hook**
- **Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK**
- **SageMaker Debugger Interactive Report for XGBoost**
- **Action on Amazon SageMaker Debugger Rules**
- **Visualize Amazon SageMaker Debugger Output Tensors in TensorBoard**

### Step 1: Adapt Your Training Script to Register a Hook

Amazon SageMaker Debugger comes with a client library called the sagemaker-debugger Python SDK. The sagemaker-debugger Python SDK provides tools for adapting your training script before training and analysis tools after training. In this page, you'll learn how to adapt your training script using the client library.

The sagemaker-debugger Python SDK provides wrapper functions that help register a hook to extract model tensors, without altering your training script. To get started with collecting model output tensors and debug them to find training issues, make the following modifications in your training script.

**Tip**

While you're following this page, use the sagemaker-debugger open source SDK documentation for API references.

Topics

- **Adapt Your PyTorch Training Script**
- **Adapt Your TensorFlow Training Script**
Adapt Your PyTorch Training Script

To start collecting model output tensors and debug training issues, make the following modifications to your PyTorch training script.

For PyTorch 1.12.0

If you bring a PyTorch training script, you can run the training job and extract model output tensors with a few additional code lines in your training script. You need to use the hook APIs in the sagemaker-debugger client library. Walk through the following instructions that break down the steps with code examples.

1. Create a hook.

(Recommended) For training jobs within SageMaker

```python
import smdebug.pytorch as smd
hook=smd.get_hook(create_if_not_exists=True)
```

When you launch a training job in the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK” with any of the DebuggerHookConfig, TensorBoardConfig, or Rules in your estimator, SageMaker adds a JSON configuration file to your training instance that is picked up by the get_hook function. Note that if you do not include any of the configuration APIs in your estimator, there will be no configuration file for the hook to find, and the function returns None.

(Optional) For training jobs outside SageMaker

If you run training jobs in local mode, directly on SageMaker Notebook instances, Amazon EC2 instances, or your own local devices, use smd.Hook class to create a hook. However, this approach can only store the tensor collections and usable for TensorBoard visualization. SageMaker Debugger’s built-in Rules don’t work with the local mode because the Rules require SageMaker ML training instances and S3 to store outputs from the remote instances in real time. The smd.get_hook API returns None in this case.

If you want to create a manual hook to save tensors in local mode, use the following code snippet with the logic to check if the smd.get_hook API returns None and create a manual hook using the smd.Hook class. Note that you can specify any output directory in your local machine.
import smdebug.pytorch as smd
hook = smd.get_hook(create_if_not_exists=True)

if hook is None:
    hook = smd.Hook(
        out_dir='/path/to/your/local/output/',
        export_tensorboard=True
    )

2. Wrap your model with the hook's class methods.

The hook.register_module() method takes your model and iterates through each layer, looking for any tensors that match with regular expressions that you’ll provide through the configuration in the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK”. The collectable tensors through this hook method are weights, biases, activations, gradients, inputs, and outputs.

    hook.register_module(model)

**Tip**

If you collect the entire output tensors from a large deep learning model, the total size of those collections can exponentially grow and might cause bottlenecks. If you want to save specific tensors, you can also use the hook.save_tensor() method. This method helps you pick the variable for the specific tensor and save to a custom collection named as you want. For more information, see step 7 of this instruction.

3. Warp the loss function with the hook’s class methods.

The hook.register_loss method is to wrap the loss function. It extracts loss values every save_interval that you’ll set during configuration in the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK”, and saves them to the "losses" collection.

    hook.register_loss(loss_function)

4. Add hook.set_mode(ModeKeys.TRAIN) in the train block. This indicates the tensor collection is extracted during the training phase.
def train():
    ...
    hook.set_mode(ModeKeys.TRAIN)

5. Add `hook.set_mode(ModeKeys.EVAL)` in the validation block. This indicates the tensor collection is extracted during the validation phase.

def validation():
    ...
    hook.set_mode(ModeKeys.EVAL)

6. Use `hook.save_scalar()` to save custom scalars. You can save scalar values that aren't in your model. For example, if you want to record the accuracy values computed during evaluation, add the following line of code below the line where you calculate accuracy.

hook.save_scalar("accuracy", accuracy)

Note that you need to provide a string as the first argument to name the custom scalar collection. This is the name that'll be used for visualizing the scalar values in TensorBoard, and can be any string you want.

7. Use `hook.save_tensor()` to save custom tensors. Similarly to `hook.save_scalar()`, you can save additional tensors, defining your own tensor collection. For example, you can extract input image data that are passed into the model and save as a custom tensor by adding the following code line, where "images" is an example name of the custom tensor, `image_inputs` is an example variable for the input image data.

hook.save_tensor("images", image_inputs)

Note that you must provide a string to the first argument to name the custom tensor. `hook.save_tensor()` has the third argument `collections_to_write` to specify the tensor collection to save the custom tensor. The default is `collections_to_write="default"`. If you don't explicitly specify the third argument, the custom tensor is saved to the "default" tensor collection.

After you have completed adapting your training script, proceed to the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK”.

Use SageMaker Debugger
Adapt Your TensorFlow Training Script

To start collecting model output tensors and debug training issues, make the following modifications to your TensorFlow training script.

Create a hook for training jobs within SageMaker

```python
import smdebug.tensorflow as smd

hook=smd.get_hook(hook_type="keras", create_if_not_exists=True)
```

This creates a hook when you start a SageMaker training job. When you launch a training job in the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK” with any of the DebuggerHookConfig, TensorBoardConfig, or Rules in your estimator, SageMaker adds a JSON configuration file to your training instance that is picked up by the `smd.get_hook` method. Note that if you do not include any of the configuration APIs in your estimator, there will be no configuration file for the hook to find, and the function returns None.

(Optional) Create a hook for training jobs outside SageMaker

If you run training jobs in local mode, directly on SageMaker Notebook instances, Amazon EC2 instances, or your own local devices, use `smd.Hook` class to create a hook. However, this approach can only store the tensor collections and usable for TensorBoard visualization. SageMaker Debugger’s built-in Rules don’t work with the local mode. The `smd.get_hook` method also returns None in this case.

If you want to create a manual hook, use the following code snippet with the logic to check if the hook returns None and create a manual hook using the `smd.Hook` class.

```python
import smdebug.tensorflow as smd

hook=smd.get_hook(hook_type="keras", create_if_not_exists=True)

if hook is None:
    hook=smd.KerasHook(
        out_dir='path/to/your/local/output/',
        export_tensorboard=True
    )
```

After adding the hook creation code, proceed to the following topic for TensorFlow Keras.
Note

SageMaker Debugger currently supports TensorFlow Keras only.

Register the hook in your TensorFlow Keras training script

The following procedure walks you through how to use the hook and its methods to collect output scalars and tensors from your model and optimizer.

1. Wrap your Keras model and optimizer with the hook's class methods.

   The hook.register_model() method takes your model and iterates through each layer, looking for any tensors that match with regular expressions that you'll provide through the configuration in the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK”. The collectable tensors through this hook method are weights, biases, and activations.

   ```python
   model = tf.keras.Model(...)  
   hook.register_model(model)
   ```

2. Wrap the optimizer by the hook.wrap_optimizer() method.

   ```python
   optimizer = tf.keras.optimizers.Adam(...)  
   optimizer = hook.wrap_optimizer(optimizer)
   ```


   To collect tensors from the model, such as the input and output tensors of each layer, you must run the training in eager mode. Otherwise, SageMaker Debugger will not be able to collect the tensors. However, other tensors, such as model weights, biases, and the loss, can be collected without explicitly running in eager mode.

   ```python
   model.compile(  
       loss="categorical_crossentropy",  
       optimizer=optimizer,  
       metrics=["accuracy"],  
       # Required for collecting tensors of each layer  
       run_eagerly=True
   )
   ```
4. Register the hook to the `tf.keras.Model.fit()` method.

To collect the tensors from the hooks that you registered, add `callbacks=[hook]` to the Keras `model.fit()` class method. This will pass the `sagemaker-debugger` hook as a Keras callback.

```python
model.fit(
    X_train, Y_train,
    batch_size=batch_size,
    epochs=epoch,
    validation_data=(X_valid, Y_valid),
    shuffle=True,
    callbacks=[hook]
)
```

5. TensorFlow 2.x provides only symbolic gradient variables that do not provide access to their values. To collect gradients, wrap `tf.GradientTape` by the `hook.wrap_tape()` method, which requires you to write your own training step as follows.

```python
def training_step(model, dataset):
    with hook.wrap_tape(tf.GradientTape()) as tape:
        pred=model(data)
        loss_value=loss_fn(labels, pred)
        grads=tape.gradient(loss_value, model.trainable_variables)
        optimizer.apply_gradients(zip(grads, model.trainable_variables))
```

By wrapping the tape, the `sagemaker-debugger` hook can identify output tensors such as gradients, parameters, and losses. Wrapping the tape ensures that the `hook.wrap_tape()` method around functions of the tape object, such as `push_tape()`, `pop_tape()`, `gradient()`, will set up the writers of SageMaker Debugger and save tensors that are provided as input to `gradient()` (trainable variables and loss) and output of `gradient()` (gradients).

**Note**

To collect with a custom training loop, make sure that you use eager mode. Otherwise, SageMaker Debugger is not able to collect any tensors.

For a full list of actions that the `sagemaker-debugger` hook APIs offer to construct hooks and save tensors, see [Hook Methods](#) in the `sagemaker-debugger Python SDK documentation`. 
After you have completed adapting your training script, proceed to the section called “Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK”.

**Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK**

To configure a SageMaker estimator with SageMaker Debugger, use Amazon SageMaker Python SDK and specify Debugger-specific parameters. To fully utilize the debugging functionality, there are three parameters you need to configure: debugger_hook_config, tensorboard_output_config, and rules.

⚠️ **Important**

Before constructing and running the estimator fit method to launch a training job, make sure that you adapt your training script following the instructions at the section called “Step 1: Adapt Your Training Script to Register a Hook”.

**Construct a SageMaker Estimator with Debugger-specific parameters**

The code examples in this section show how to construct a SageMaker estimator with the Debugger-specific parameters.

⚠️ **Note**

The following code examples are templates for constructing the SageMaker framework estimators and not directly executable. You need to proceed to the next sections and configure the Debugger-specific parameters.

**PyTorch**

```python
# An example of constructing a SageMaker PyTorch estimator
import boto3
import sagemaker
from sagemaker.pytorch import PyTorch
from sagemaker.debugger import CollectionConfig, DebuggerHookConfig, Rule, rule_configs

session=boto3.session.Session()
region=session.region_name
```
debugger_hook_config=DebuggerHookConfig(...)  
rules=[  
    Rule.sagemaker(rule_configs.built_in_rule())  
]  
estimator=PyTorch(  
    entry_point="directory/to/your_training_script.py",  
    role=sagemaker.get_execution_role(),  
    base_job_name="debugger-demo",  
    instance_count=1,  
    instance_type="ml.p3.2xlarge",  
    framework_version="1.12.0",  
    py_version="py37",  

    # Debugger-specific parameters  
    debugger_hook_config=debugger_hook_config,  
    rules=rules  
)  
estimator.fit(wait=False)

TensorFlow

# An example of constructing a SageMaker TensorFlow estimator
import boto3
import sagemaker
from sagemaker.tensorflow import TensorFlow  
from sagemaker.debugger import CollectionConfig, DebuggerHookConfig, Rule, rule_configs

session=boto3.session.Session()  
region=session.region_name

designer_hook_config=DebuggerHookConfig(...)  
rules=[  
    Rule.sagemaker(rule_configs.built_in_rule()),  
    ProfilerRule.sagemaker(rule_configs.BuiltinRule())  
]  
estimator=TensorFlow(  
    entry_point="directory/to/your_training_script.py",  
    role=sagemaker.get_execution_role(),  
    base_job_name="debugger-demo",  
    instance_count=1,  
    instance_type="ml.p3.2xlarge",  
    framework_version="1.12.0",  
    py_version="py37",  
    debugger_hook_config=debugger_hook_config,  
    rules=rules  
)  
estimator.fit(wait=False)
base_job_name="debugger-demo",
instance_count=1,
instance_type="ml.p3.2xlarge",
framework_version="2.9.0",
py_version="py39",

# Debugger-specific parameters
debugger_hook_config=debugger_hook_config,
rules=rules
)

estimator.fit(wait=False)

MXNet

# An example of constructing a SageMaker MXNet estimator
import sagemaker
from sagemaker.mxnet import MXNet
from sagemaker.debugger import CollectionConfig, DebuggerHookConfig, Rule,
rule_configs

debugger_hook_config=DebuggerHookConfig(...)  
rules=[
     Rule.sagemaker(rule_configs.built_in_rule())
]

estimator=MXNet(
    entry_point="directory/to/your_training_script.py",
    role=sagemaker.get_execution_role(),
    base_job_name="debugger-demo",
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="1.7.0",
    py_version="py37",

    # Debugger-specific parameters
debugger_hook_config=debugger_hook_config,
    rules=rules
)

estimator.fit(wait=False)
XGBoost

# An example of constructing a SageMaker XGBoost estimator
import sagemaker
from sagemaker.xgboost.estimator import XGBoost
from sagemaker.debugger import CollectionConfig, DebuggerHookConfig, Rule,
   rule_configs

   debugger_hook_config=DebuggerHookConfig(...)  
   rules=[
       Rule.sagemaker(rule_configs.built_in_rule())
   ]
   
estimator=XGBoost(
       entry_point="directory/to/your_training_script.py",
       role=sagemaker.get_execution_role(),
       base_job_name="debugger-demo",
       instance_count=1,
       instance_type="ml.p3.2xlarge",
       framework_version="1.5-1",

       # Debugger-specific parameters
       debugger_hook_config=debugger_hook_config,
       rules=rules)

   estimator.fit(wait=False)

Generic estimator

# An example of constructing a SageMaker generic estimator using the XGBoost
algorithm base image
import boto3
import sagemaker
from sagemaker.estimator import Estimator
from sagemaker import image_uris
from sagemaker.debugger import CollectionConfig, DebuggerHookConfig, Rule,
   rule_configs

   debugger_hook_config=DebuggerHookConfig(...)  
   rules=[
       Rule.sagemaker(rule_configs.built_in_rule())
   ]

```python
region=boto3.Session().region_name
xgboost_container=sagemaker.image_uris.retrieve("xgboost", region, "1.5-1")

estimator=Estimator(
    role=sagemaker.get_execution_role(),
    image_uri=xgboost_container,
    base_job_name="debugger-demo",
    instance_count=1,
    instance_type="ml.m5.2xlarge",

    # Debugger-specific parameters
    debugger_hook_config=debugger_hook_config,
    rules=rules
)

estimator.fit(wait=False)
```

Configure the following parameters to activate SageMaker Debugger:

- **debugger_hook_config** (an object of `DebuggerHookConfig`) – Required to activate the hook in the adapted training script during the section called “Step 1: Adapt Your Training Script to Register a Hook”, configure the SageMaker training launcher (estimator) to collect output tensors from your training job, and save the tensors into your secured S3 bucket or local machine. To learn how to configure the `debugger_hook_config` parameter, see [Configure SageMaker Debugger to Save Tensors](#).

- **rules** (a list of `Rule` objects) – Configure this parameter to activate SageMaker Debugger built-in rules that you want to run in real time. The built-in rules are logics that automatically debug the training progress of your model and find training issues by analyzing the output tensors saved in your secured S3 bucket. To learn how to configure the `rules` parameter, see [Configure Debugger Built-in Rules](#). To find a complete list of built-in rules for debugging output tensors, see the section called “Debugger Rule”. If you want to create your own logic to detect any training issues, see the section called “Create Custom Rules”.

**Note**

The built-in rules are available only through SageMaker training instances. You cannot use them in local mode.
• `tensorboard_output_config` (an object of `TensorBoardOutputConfig`) – Configure SageMaker Debugger to collect output tensors in the TensorBoard-compatible format and save to your S3 output path specified in the `TensorBoardOutputConfig` object. To learn more, see the section called “Visualize Debugger Output Tensors in TensorBoard”.

**Note**

The `tensorboard_output_config` must be configured with the `debugger_hook_config` parameter, which also requires you to adapt your training script by adding the `sagemaker-debugger` hook.

**Note**

SageMaker Debugger securely saves output tensors in subfolders of your S3 bucket. For example, the format of the default S3 bucket URI in your account is `s3://sagemaker-<region>-<12digit_account_id>/<base-job-name>/<debugger-subfolders>/`. There are two subfolders created by SageMaker Debugger: `debug-output`, and `rule-output`. If you add the `tensorboard_output_config` parameter, you'll also find `tensorboard-output` folder.

See the following topics to find more examples of how to configure the Debugger-specific parameters in detail.

**Topics**

- [Configure SageMaker Debugger to Save Tensors](#)
- [Configure Debugger Built-in Rules](#)
- [Turn Off Debugger](#)
- [Useful SageMaker Estimator Classmethods for Debugger](#)

### Configure SageMaker Debugger to Save Tensors

*Tensors* are data collections of updated parameters from the backward and forward pass of each training iteration. SageMaker Debugger collects the output tensors to analyze the state of a training job. SageMaker Debugger's `CollectionConfig` and `DebuggerHookConfig` API
operations provide methods for grouping tensors into *collections* and saving them to a target S3 bucket.

**Note**

After properly configured and activated, SageMaker Debugger saves the output tensors in a default S3 bucket, unless otherwise specified. The format of the default S3 bucket URI is `s3://sagemaker-<region>-<12digit_account_id>/<training-job-name>/debug-output/`.

While constructing a SageMaker estimator, activate SageMaker Debugger by specifying the `debugger_hook_config` parameter. The following steps include examples of how to set up the `debugger_hook_config` using the `CollectionConfig` and `DebuggerHookConfig` API operations to pull tensors out of your training jobs and save them.

**Configure Tensor Collections Using the CollectionConfig API**

Use the `CollectionConfig` API operation to configure tensor collections. Debugger provides pre-built tensor collections that cover a variety of regular expressions (regex) of parameters if using Debugger-supported deep learning frameworks and machine learning algorithms. As shown in the following example code, add the built-in tensor collections you want to debug.

```python
from sagemaker debugger import CollectionConfig

collection_configs = [
    CollectionConfig(name="weights"),
    CollectionConfig(name="gradients")
]
```

The preceding collections set up the Debugger hook to save the tensors every 500 steps based on the default "save_interval" value.

For a full list of available Debugger built-in collections, see [Debugger Built-in Collections](#).

If you want to customize the built-in collections, such as changing the save intervals and tensor regex, use the following `CollectionConfig` template to adjust parameters.

```python
from sagemaker debugger import CollectionConfig
```
collection_configs=[
    CollectionConfig(
        name="tensor_collection",
        parameters={
            "key_1": "value_1",
            "key_2": "value_2",
            ...
            "key_n": "value_n"
        }
    )
]

For more information about available parameter keys, see `CollectionConfig` in the Amazon SageMaker Python SDK. For example, the following code example shows how you can adjust the save intervals of the "losses" tensor collection at different phases of training: save loss every 100 steps in training phase and validation loss every 10 steps in validation phase.

```python
from sagemaker.debugger import CollectionConfig
collection_configs=[
    CollectionConfig(
        name="losses",
        parameters={
            "train.save_interval": "100",
            "eval.save_interval": "10"
        }
    )
]
```

Tip

This tensor collection configuration object can be used for both `DebuggerHookConfig` and `Rule` API operations.

Configure the DebuggerHookConfig API to Save Tensors

Use the `DebuggerHookConfig` API to create a debugger_hook_config object using the collection_configs object you created in the previous step.

```python
from sagemaker.debugger import DebuggerHookConfig
```
Debugger saves the model training output tensors into the default S3 bucket. The format of the default S3 bucket URI is `s3://sagemaker-<region>-<12digit_account_id>/<training-job-name>/debug-output/`.

If you want to specify an exact S3 bucket URI, use the following code example:

```python
from sagemaker.debugger import DebuggerHookConfig
debuger_hook_config=DebuggerHookConfig(
    s3_output_path="specify-your-s3-bucket-uri",
    collection_configs=collection_configs
)
```

For more information, see DebuggerHookConfig in the Amazon SageMaker Python SDK.

**Example Notebooks and Code Samples to Configure Debugger Hook**

The following sections provide notebooks and code examples of how to use Debugger hook to save, access, and visualize output tensors.

**Topics**

- Tensor Visualization Example Notebooks
- Save Tensors Using Debugger Built-in Collections
- Save Tensors Using Debugger Modified Built-in Collections
- Save Tensors Using Debugger Custom Collections

**Tensor Visualization Example Notebooks**

The following two notebook examples show advanced use of Amazon SageMaker Debugger for visualizing tensors. Debugger provides a transparent view into training deep learning models.

- Interactive Tensor Analysis in SageMaker Studio Notebook with MXNet

  This notebook example shows how to visualize saved tensors using Amazon SageMaker Debugger. By visualizing the tensors, you can see how the tensor values change while training
deep learning algorithms. This notebook includes a training job with a poorly configured neural network and uses Amazon SageMaker Debugger to aggregate and analyze tensors, including gradients, activation outputs, and weights. For example, the following plot shows the distribution of gradients of a convolutional layer that is suffering from a vanishing gradient problem.

![Distribution of Gradients](image)

This notebook also illustrates how a good initial hyperparameter setting improves the training process by generating the same tensor distribution plots.

- **Visualizing and Debugging Tensors from MXNet Model Training**

This notebook example shows how to save and visualize tensors from an MXNet Gluon model training job using Amazon SageMaker Debugger. It illustrates that Debugger is set to save all tensors to an Amazon S3 bucket and retrieves ReLu activation outputs for the visualization. The following figure shows a three-dimensional visualization of the ReLu activation outputs. The color scheme is set to blue to indicate values close to 0 and yellow to indicate values close to 1.
In this notebook, the `TensorPlot` class imported from `tensor_plot.py` is designed to plot convolutional neural networks (CNNs) that take two-dimensional images for inputs. The `tensor_plot.py` script provided with the notebook retrieves tensors using Debugger and visualizes the CNN. You can run this notebook on SageMaker Studio to reproduce the tensor visualization and implement your own convolutional neural network model.

- **Real-time Tensor Analysis in a SageMaker Notebook with MXNet**

  This example guides you through installing required components for emitting tensors in an Amazon SageMaker training job and using the Debugger API operations to access those tensors while training is running. A gluon CNN model is trained on the Fashion MNIST dataset. While the job is running, you will see how Debugger retrieves activation outputs of the first convolutional layer from each of 100 batches and visualizes them. Also, this will show you how to visualize weights after the job is done.

**Save Tensors Using Debugger Built-in Collections**

You can use built-in collections of tensors using the `CollectionConfig` API and save them using the `DebuggerHookConfig` API. The following example shows how to use the default settings of
Debugger hook configurations to construct a SageMaker TensorFlow estimator. You can also utilize this for MXNet, PyTorch, and XGBoost estimators.

**: Note**

In the following example code, the `s3_output_path` parameter for DebuggerHookConfig is optional. If you do not specify it, Debugger saves the tensors at `s3://<output_path>/debug-output/`, where the `<output_path>` is the default output path of SageMaker training jobs. For example:

```
"s3://sagemaker-us-east-1-111122223333/sagemaker-debugger-training-YYYY-MM-DD-HH-MM-SS-123/debug-output"
```

```python
import sagemaker
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import DebuggerHookConfig, CollectionConfig

# use Debugger CollectionConfig to call built-in collections
collection_configs=[
    CollectionConfig(name="weights"),
    CollectionConfig(name="gradients"),
    CollectionConfig(name="losses"),
    CollectionConfig(name="biases")
]

# configure Debugger hook
# set a target S3 bucket as you want
sagemaker_session=sagemaker.Session()
BUCKET_NAME=sagemaker_session.default_bucket()
LOCATION_IN_BUCKET='debugger-built-in-collections-hook'

hook_config=DebuggerHookConfig(
    s3_output_path='s3://{BUCKET_NAME}/{LOCATION_IN_BUCKET}'.format(BUCKET_NAME=BUCKET_NAME,
                        LOCATION_IN_BUCKET=LOCATION_IN_BUCKET),
    collection_configs=collection_configs
)

# construct a SageMaker TensorFlow estimator
sagemaker_estimator=TensorFlow(  
```
entry_point='directory/to/your_training_script.py', role=sm.get_execution_role(),
base_job_name='debugger-demo-job',
instance_count=1,
instance_type="ml.p3.2xlarge",
framework_version="2.9.0",
py_version="py39",

# debugger-specific hook argument below
degger_hook_config=hook_config
)

degger=degger_sm.fit()

To see a list of Debugger built-in collections, see Debugger Built-in Collections.

Save Tensors Using Debugger Modified Built-in Collections

You can modify the Debugger built-in collections using the CollectionConfig API operation. The following example shows how to tweak the built-in losses collection and construct a SageMaker TensorFlow estimator. You can also use this for MXNet, PyTorch, and XGBoost estimators.

```python
import sagemaker
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import DebuggerHookConfig, CollectionConfig

# use Debugger CollectionConfig to call and modify built-in collections
collection_configs=[
    CollectionConfig(
        name="losses",
        parameters={"save_interval": "50"})
]

# configure Debugger hook
# set a target S3 bucket as you want
sagemaker_session=sagemaker.Session()
BUCKET_NAME=sagemaker_session.default_bucket()
LOCATION_IN_BUCKET='debugger-modified-collections-hook'

hook_config=DebuggerHookConfig(
    s3_output_path='s3://{BUCKET_NAME}/{LOCATION_IN_BUCKET}'.format(BUCKET_NAME=BUCKET_NAME,
        LOCATION_IN_BUCKET=LOCATION_IN_BUCKET),
```
collection_configs=collection_configs
)

# construct a SageMaker TensorFlow estimator
sagemaker_estimator=TensorFlow(
    entry_point='directory/to/your_training_script.py',
    role=sm.get_execution_role(),
    base_job_name='debugger-demo-job',
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="2.9.0",
    py_version="py39",

    # debugger-specific hook argument below
    debugger_hook_config=hook_config
)

sagemaker_estimator.fit()

For a full list of CollectionConfig parameters, see Debugger CollectionConfig API.

Save Tensors Using Debugger Custom Collections

You can also save a reduced number of tensors instead of the full set of tensors (for example, if you want to reduce the amount of data saved in your Amazon S3 bucket). The following example shows how to customize the Debugger hook configuration to specify target tensors that you want to save. You can use this for TensorFlow, MXNet, PyTorch, and XGBoost estimators.

import sagemaker
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import DebuggerHookConfig, CollectionConfig

# use Debugger CollectionConfig to create a custom collection
collection_configs=[
    CollectionConfig(
        name="custom_activations_collection",
        parameters={
            "include_regex": "relu|tanh", # Required
            "reductions": "mean,variance,max,abs_mean,abs_variance,abs_max"
        }
    )
]

# configure Debugger hook

# set a target S3 bucket as you want
sagemaker_session = sagemaker.Session()
BUCKET_NAME = sagemaker_session.default_bucket()
LOCATION_IN_BUCKET = 'debugger-custom-collections-hook'

hook_config = DebuggerHookConfig(
    s3_output_path='s3://{BUCKET_NAME}/{LOCATION_IN_BUCKET}'.format(BUCKET_NAME=BUCKET_NAME,
                                                                       LOCATION_IN_BUCKET=LOCATION_IN_BUCKET),
    collection_configs=collection_configs
)

# construct a SageMaker TensorFlow estimator
sagemaker_estimator = TensorFlow(
    entry_point='directory/to/your_training_script.py',
    role=sm.get_execution_role(),
    base_job_name='debugger-demo-job',
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="2.9.0",
    py_version="py39",

    # debugger-specific hook argument below
    debugger_hook_config=hook_config
)

sagemaker_estimator.fit()

For a full list of CollectionConfig parameters, see Debugger CollectionConfig.

Configure Debugger Built-in Rules

Amazon SageMaker Debugger's built-in rules analyze tensors emitted during the training of a model. SageMaker Debugger offers the Rule API operation that monitors training job progress and errors for the success of training your model. For example, the rules can detect whether gradients are getting too large or too small, whether a model is overfitting or overtraining, and whether a training job does not decrease loss function and improve. To see a full list of available built-in rules, see List of Debugger Built-in Rules.

In the following topics, you'll learn how to use the SageMaker Debugger built-in rules.

Topics
Use Debugger Built-in Rules with the Default Parameter Settings

To specify Debugger built-in rules in an estimator, you need to configure a list object. The following example code shows the basic structure of listing the Debugger built-in rules:

```python
from sagemaker.debugger import Rule, rule_configs

rules=[
    Rule.sagemaker(rule_configs.built_in_rule_name_1()),
    Rule.sagemaker(rule_configs.built_in_rule_name_2()),
    ...
    Rule.sagemaker(rule_configs.built_in_rule_name_n()),
    ...
    # You can also append more profiler rules in the ProfilerRule.sagemaker(rule_configs.*()) format.
]
```

For more information about default parameter values and descriptions of the built-in rule, see List of Debugger Built-in Rules.

To find the SageMaker Debugger API reference, see `sagemaker.debugger.rule_configs` and `sagemaker.debugger.Rule`.

For example, to inspect the overall training performance and progress of your model, construct a SageMaker estimator with the following built-in rule configuration.

```python
from sagemaker.debugger import Rule, rule_configs

rules=[
    Rule.sagemaker(rule_configs.loss_not_decreasing()),
    Rule.sagemaker(rule_configs.overfit()),
    Rule.sagemaker(rule_configs.overtraining()),
    Rule.sagemaker(rule_configs.stalled_training_rule())
]
```

When you start the training job, Debugger collects system resource utilization data every 500 milliseconds and the loss and accuracy values every 500 steps by default. Debugger
analyzes the resource utilization to identify if your model is having bottleneck problems. The loss_not_decreasing, overfit, overtraining, and stalled_training_rule monitors if your model is optimizing the loss function without those training issues. If the rules detect training anomalies, the rule evaluation status changes to IssueFound. You can set up automated actions, such as notifying training issues and stopping training jobs using Amazon CloudWatch Events and AWS Lambda. For more information, see Action on Amazon SageMaker Debugger Rules.

Use Debugger Built-in Rules with Custom Parameter Values

If you want to adjust the built-in rule parameter values and customize tensor collection regex, configure the base_config and rule_parameters parameters for the ProfilerRule.sagemaker and Rule.sagemaker class methods. In case of the Rule.sagemaker class methods, you can also customize tensor collections through the collections_to_save parameter. The instruction of how to use the CollectionConfig class is provided at Configure Tensor Collections Using the CollectionConfig API.

Use the following configuration template for built-in rules to customize parameter values. By changing the rule parameters as you want, you can adjust the sensitivity of the rules to be triggered.

- The base_config argument is where you call the built-in rule methods.
- The rule_parameters argument is to adjust the default key values of the built-in rules listed in List of Debugger Built-in Rules.
- The collections_to_save argument takes in a tensor configuration through the CollectionConfig API, which requires name and parameters arguments.
  - To find available tensor collections for name, see Debugger Built-in Tensor Collections.
  - For a full list of adjustable parameters, see Debugger CollectionConfig API.

For more information about the Debugger rule class, methods, and parameters, see SageMaker Debugger Rule class in the Amazon SageMaker Python SDK.

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs, CollectionConfig
rules=[
    Rule.sagemaker(
        base_config=rule_configs.built_in_rule_name(),
        rule_parameters={
            "key": "value"
    
```
The parameter descriptions and value customization examples are provided for each rule at List of Debugger Built-in Rules.

Example Notebooks and Code Samples to Configure Debugger Rules

In the following sections, notebooks and code samples of how to use Debugger rules to monitor SageMaker training jobs are provided.

Topics

- Debugger Built-in Rules Example Notebooks
- Debugger Built-in Rules Example Code
- Use Debugger Built-in Rules with Parameter Modifications

Debugger Built-in Rules Example Notebooks

The following example notebooks show how to use Debugger built-in rules when running training jobs with Amazon SageMaker:

- Using a SageMaker Debugger built-in rule with TensorFlow
- Using a SageMaker Debugger built-in rule with Managed Spot Training and MXNet
- Using a SageMaker Debugger built-in rule with parameter modifications for a real-time training job analysis with XGBoost

While running the example notebooks in SageMaker Studio, you can find the training job trial created on the Studio Experiment List tab. For example, as shown in the following screenshot, you can find and open a Describe Trial Component window of your current training job. On the Debugger tab, you can check if the Debugger rules, vanishing_gradient() and
loss_not_decreasing(), are monitoring the training session in parallel. For a full instruction of how to find your training job trial components in the Studio UI, see SageMaker Studio - View Experiments, Trials, and Trial Components.

```python
[29]: rules = [  
    Rule.sagemaker(rule_config.sagemaker.vanishing_gradient()),  
    Rule.sagemaker(  
        base_config=sagemaker.loss_not_decreasing(),  
        collections_to_save=[  
            CollectionConfig(  
                name="losses",  
                parameters={  
                    "save_interval": "50",  
                    "train.save_interval": "50",  
                    "eval.save_interval": "10"  
                }  
            )  
        ]  
    )  
]  

estimator = TensorFlow(  
    role=sagemaker.get_execution_role(),  
    base_job_name='sagemaker-demo-mnist-tensorflow',  
    train_instance_count=1,  
    train_instance_type='ml.m4.xlarge',  
    train_volume_size=400,  
    entry_point=entrypoint_script,  
    framework_version='1.15',  
    py_version='py3',  
    train_max_run=3600,  
    script_mode=True,  
    hyperparameters=hyperparameters,  
    ## New parameter  
    rules = rules  
)
```

**Describe Trial Component**

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<th>Metrics</th>
<th>Parameters</th>
<th>Artifacts</th>
<th>AWS Settings</th>
<th>Debugger</th>
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- Status: In Progress
- Last modified: 7 seconds ago
- Rule name: VanishingGradient
- Job ARN: arn:aws:sagemaker:us-east-1...

- Status: In Progress
- Last modified: 7 seconds ago
- Rule name: LossNotDecreasing
- Job ARN: arn:aws:sagemaker:us-east-1...
There are two ways of using the Debugger built-in rules in the SageMaker environment: deploy the built-in rules as it is prepared or adjust their parameters as you want. The following topics show you how to use the built-in rules with example codes.

**Debugger Built-in Rules Example Code**

The following code sample shows how to set the Debugger built-in rules using the `Rule.sagemaker` method. To specify built-in rules that you want to run, use the `rules_configs` API operation to call the built-in rules. To find a full list of Debugger built-in rules and default parameter values, see [List of Debugger Built-in Rules](#).

```python
import sagemaker
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import Rule, CollectionConfig, rule_configs

# call built-in rules that you want to use.
built_in_rules = [
    Rule.sagemaker(rule_configs.vanishing_gradient()),
    Rule.sagemaker(rule_configs.loss_not_decreasing())
]

# construct a SageMaker estimator with the Debugger built-in rules
sagemaker_estimator = TensorFlow(
    entry_point='directory/to/your_training_script.py',
    role=sm.get_execution_role(),
    base_job_name='debugger-built-in-rules-demo',
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="2.9.0",
    py_version="py39",

    # debugger-specific arguments below
    rules=built_in_rules
)

sagemaker_estimator.fit()
```

⚠️ Note

The Debugger built-in rules run in parallel with your training job. The maximum number of built-in rule containers for a training job is 20.
For more information about the Debugger rule class, methods, and parameters, see the SageMaker Debugger Rule class in the Amazon SageMaker Python SDK.

To find an example of how to adjust the Debugger rule parameters, see the following Use Debugger Built-in Rules with Parameter Modifications section.

Use Debugger Built-in Rules with Parameter Modifications

The following code example shows the structure of built-in rules to adjust parameters. In this example, the stalled_training_rule collects the losses tensor collection from a training job at every 50 steps and an evaluation stage at every 10 steps. If the training process starts stalling and not collecting tensor outputs for 120 seconds, the stalled_training_rule stops the training job.

```python
import sagemaker
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import Rule, CollectionConfig, rule_configs

# call the built-in rules and modify the CollectionConfig parameters

base_job_name_prefix= 'smdebug-stalled-demo-' + str(int(time.time()))

built_in_rules_modified=[
    Rule.sagemaker(
        base_config=rule_configs.stalled_training_rule(),
        rule_parameters={
            'threshold': '120',
            'training_job_name_prefix': base_job_name_prefix,
            'stop_training_on_fire' : 'True'
        },
        collections_to_save=[
            CollectionConfig(
                name="losses",
                parameters={
                    'train.save_interval': "50",
                    'eval.save_interval': "10"
                }
            )
        ]
    )
]  

# construct a SageMaker estimator with the modified Debugger built-in rule
```
sagemaker_estimator=TensorFlow(
    entry_point='directory/to/your_training_script.py',
    role=sm.get_execution_role(),
    base_job_name=base_job_name_prefix,
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="2.9.0",
    py_version="py39",

    # debugger-specific arguments below
    rules=built_in_rules_modified
)
sagemaker_estimator.fit()

For an advanced configuration of the Debugger built-in rules using the CreateTrainingJob API, see Configure Debugger Using Amazon SageMaker API.

**Turn Off Debugger**

If you want to completely turn off Debugger, do one of the following:

- Before starting a training job, do the following:

  To stop both monitoring and profiling, include the disable_profiler parameter to your estimator and set it to True.

  > **Warning**
  > If you disable it, you won't be able to view the comprehensive Studio Debugger insights dashboard and the autogenerated profiling report.

  To stop debugging, set the debugger_hook_config parameter to False.

  > **Warning**
  > If you disable it, you won't be able to collect output tensors and cannot debug your model parameters.
For more information about the Debugger-specific parameters, see SageMaker Estimator in the Amazon SageMaker Python SDK.

- While a training job is running, do the following:

To disable both monitoring and profiling while your training job is running, use the following estimator classmethod:

```python
estimator.disable_profiling()
```

To disable framework profiling only and keep system monitoring, use the `update_profiler` method:

```python
estimator.update_profiler(disable_framework_metrics=true)
```

For more information about the estimator extension methods, see the `estimator.disable_profiling` and `estimator.update_profiler` classmethods in the Amazon SageMaker Python SDK documentation.

**Useful SageMaker Estimator Classmethods for Debugger**

The following estimator class methods are useful for accessing your SageMaker training job information and retrieving output paths of training data collected by Debugger. The following methods are executable after you initiate a training job with the `estimator.fit()` method.

- To check the base S3 bucket URI of a SageMaker training job:

  ```python
  estimator.output_path
  ```

- To check the base job name of a SageMaker training job:

  ```python
  estimator.latest_training_job.job_name
  ```

- To see a full `CreateTrainingJob` API operation configuration of a SageMaker training job:
estimator.latest_training_job.describe()

- To check a full list of the Debugger rules while a SageMaker training job is running:
  estimator.latest_training_job.rule_job_summary()

- To check the S3 bucket URI where the model parameter data (output tensors) are saved:
  estimator.latest_job_debugger_artifacts_path()

- To check the S3 bucket URI at where the model performance data (system and framework metrics) are saved:
  estimator.latest_job_profiler_artifacts_path()

- To check the Debugger rule configuration for debugging output tensors:
  estimator.debugger_rule_configs

- To check the list of the Debugger rules for debugging while a SageMaker training job is running:
  estimator.debugger_rules

- To check the Debugger rule configuration for monitoring and profiling system and framework metrics:
  estimator.profiler_rule_configs

- To check the list of the Debugger rules for monitoring and profiling while a SageMaker training job is running:
  estimator.profiler_rules

For more information about the SageMaker estimator class and its methods, see [Estimator API](#) in the [Amazon SageMaker Python SDK](#).
SageMaker Debugger Interactive Report for XGBoost

Receive training reports autogenerated by Debugger. The Debugger reports provide insights into your training jobs and suggest recommendations to improve your model performance.

⚠️ Note
You can download a Debugger reports while your training job is running or after the job has finished. During training, Debugger concurrently updates the report reflecting the current rules’ evaluation status. You can download a complete Debugger report only after the training job has completed.

⚠️ Important
In the report, plots and and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

SageMaker Debugger XGBoost Training Report

For SageMaker XGBoost training jobs, use the Debugger CreateXgboostReport rule to receive a comprehensive training report of the training progress and results. Following this guide, specify the CreateXgboostReport rule while constructing an XGBoost estimator, download the report using the Amazon SageMaker Python SDK or the Amazon S3 console, and gain insights into the training results.

⚠️ Important
In the report, plots and and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

Topics

- [Construct a SageMaker XGBoost Estimator with the Debugger XGBoost Report Rule](#)
- [Download the Debugger XGBoost Training Report](#)
• Debugger XGBoost Training Report Walkthrough

**Construct a SageMaker XGBoost Estimator with the Debugger XGBoost Report Rule**

The [CreateXgboostReport](https://sagemaker.readthedocs.io/en/stable/guide/debugger.html#create-xgboost-report) rule collects the following output tensors from your training job:

- **hyperparameters** – Saves at the first step.
- **metrics** – Saves loss and accuracy every 5 steps.
- **feature_importance** – Saves every 5 steps.
- **predictions** – Saves every 5 steps.
- **labels** – Saves every 5 steps.

The output tensors are saved at a default S3 bucket. For example, `s3://sagemaker-<region>-<12digit_account_id>/<base-job-name>/debug-output/`.

When you construct a SageMaker estimator for an XGBoost training job, specify the rule as shown in the following example code.

**Using the SageMaker generic estimator**

```python
import boto3
import sagemaker
from sagemaker.estimator import Estimator
from sagemaker import image_uris
from sagemaker.debugger import Rule, rule_configs

rules = [Rule.sagemaker(rule_configs.create_xgboost_report())]

region = boto3.Session().region_name
xgboost_container = sagemaker.image_uris.retrieve("xgboost", region, "1.2-1")

estimator = Estimator(
    role=sagemaker.get_execution_role(),
    image_uri=xgboost_container,
    base_job_name="debugger-xgboost-report-demo",
    instance_count=1,
    instance_type="ml.m5.2xlarge",
)
```

Use SageMaker Debugger
```python
# Add the Debugger XGBoost report rule
rules=rules
)
estimator.fit(wait=False)
```

**Download the Debugger XGBoost Training Report**

Download the Debugger XGBoost training report while your training job is running or after the job has finished using the [Amazon SageMaker Python SDK](https://docs.aws.amazon.com/sagemaker/latest/dg/python-sdk.html) and AWS Command Line Interface (CLI).

**Download using the SageMaker Python SDK and AWS CLI**

1. Check the current job's default S3 output base URI.

   ```python
   estimator.output_path
   ```

2. Check the current job name.

   ```python
   estimator.latest_training_job.job_name
   ```

3. The Debugger XGBoost report is stored under `<default-s3-output-base-uri>/`<training-job-name>/rule-output. Configure the rule output path as follows:

   ```python
   rule_output_path = estimator.output_path + ""/
   estimator.latest_training_job.job_name + "/rule-output"
   ```

4. To check if the report is generated, list directories and files recursively under the `rule_output_path` using `aws s3 ls` with the `--recursive` option.

   ```bash
   ! aws s3 ls {rule_output_path} --recursive
   ```

This should return a complete list of files under autogenerated folders that are named `CreateXgboostReport` and `ProfilerReport-1234567890`. The XGBoost training report is stored in the `CreateXgboostReport`, and the profiling report is stored in the `ProfilerReport-1234567890` folder. To learn more about the profiling report generated by default with the XGBoost training job, see [SageMaker Debugger profiling report](https://docs.aws.amazon.com/sagemaker/latest/dg/debugger-profile-xgboost-report.html).
The `xgboost_report.html` is an autogenerated XGBoost training report by Debugger. The `xgboost_report.ipynb` is a Jupyter notebook that's used to aggregate training results into the report. You can download all of the files, browse the HTML report file, and modify the report using the notebook.

5. Download the files recursively using `aws s3 cp`. The following command saves all of the rule output files to the `ProfilerReport-1234567890` folder under the current working directory.

   ```bash
   ! aws s3 cp {rule_output_path} ./ --recursive
   ```

   **Tip**
   If you are using a Jupyter notebook server, run `!pwd` to verify the current working directory.

6. Under the `/CreateXgboostReport` directory, open `xgboost_report.html`. If you are using JupyterLab, choose **Trust HTML** to see the autogenerated Debugger training report.
7. Open the xgboost_report.ipynb file to explore how the report is generated. You can customize and extend the training report using the Jupyter notebook file.

Download using the Amazon S3 console

1. Sign in to the AWS Management Console and open the Amazon S3 console at https://console.aws.amazon.com/s3/.

2. Search for the base S3 bucket. For example, if you haven't specified any base job name, the base S3 bucket name should be in the following format: sagemaker-<region>-111122223333. Look up the base S3 bucket through the **Find bucket by name** field.

3. In the base S3 bucket, look up the training job name by entering your job name prefix in **Find objects by prefix** and then choosing the training job name.
4. In the training job's S3 bucket, choose **rule-output/** subfolder. There must be three subfolders for training data collected by Debugger: **debug-output/**, **profiler-output/**, and **rule-output/**.

5. In the **rule-output/** folder, choose the **CreateXgboostReport/** folder. The folder contains **xbgoost_report.html** (the autogenerated report in html) and **xbgoost_report.ipynb** (a Jupyter notebook with scripts that are used for generating the report).

6. Choose the **xbgoost_report.html** file, choose **Download actions**, and then choose **Download**.
CreateXgboost

Folder overview

Region
US West (Oregon) us-west-2

Download actions
- Download
- Download as

Edit actions
- Rename object
- Edit storage class
- Edit server-side encryption
- Edit metadata

Objects (2)
Objects are the fundamental building blocks of an S3 bucket. They can be files or directories.

Find objects by prefix

<table>
<thead>
<tr>
<th>Name</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>xgboost_report.html</td>
<td>html</td>
</tr>
<tr>
<td>xgboost_report.ipynb</td>
<td>ipynb</td>
</tr>
</tbody>
</table>
7. Open the downloaded `xbgoost_report.html` file in a web browser.

**Debugger XGBoost Training Report Walkthrough**

This section walks you through the Debugger XGBoost training report. The report is automatically aggregated depending on the output tensor regex, recognizing what type of your training job is among binary classification, multiclass classification, and regression.

⚠️ **Important**

In the report, plots and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

**Topics**

- **Distribution of True Labels of the Dataset**
- **Loss versus Step Graph**
- **Feature Importance**
- **Confusion Matrix**
- **Evaluation of the Confusion Matrix**
- **Accuracy Rate of Each Diagonal Element Over Iteration**
- **Receiver Operating Characteristic Curve**
- **Distribution of Residuals at the Last Saved Step**
- **Absolute Validation Error per Label Bin Over Iteration**

**Distribution of True Labels of the Dataset**

This histogram shows the distribution of labeled classes (for classification) or values (for regression) in your original dataset. Skewness in your dataset could contribute to inaccuracies. This visualization is available for the following model types: binary classification, multiclassification, and regression.
Loss versus Step Graph

This is a line chart that shows the progression of loss on training data and validation data throughout training steps. The loss is what you defined in your objective function, such as mean squared error. You can gauge whether the model is overfit or underfit from this plot. This section also provides insights that you can use to determine how to resolve the overfit and underfit problems. This visualization is available for the following model types: binary classification, multiclassification, and regression.
Feature Importance

There are three different types of feature importance visualizations provided: Weight, Gain and Coverage. We provide detailed definitions for each of the three in the report. Feature importance visualizations help you learn what features in your training dataset contributed to the predictions. Feature importance visualizations are available for the following model types: binary classification, multiclassification, and regression.
Confusion Matrix

This visualization is only applicable to binary and multiclass classification models. Accuracy alone might not be sufficient for evaluating the model performance. For some use cases, such as healthcare and fraud detection, it’s also important to know the false positive rate and false
negative rate. A confusion matrix gives you the additional dimensions for evaluating your model performance.
Evaluation of the Confusion Matrix

This section provides you with more insights on the micro, macro, and weighted metrics on precision, recall, and F1-score for your model.

**Overall Accuracy**
- Overall Accuracy: 0.919

**Micro Performance Metrics**
- Performance metrics calculated globally by counting the total true positives, false negatives, and false positives.
  - Micro Precision: 0.919
  - Micro Recall: 0.919
  - Micro F1-score: 0.919

**Macro Performance Metrics**
- Performance metrics calculated for each label, and find their unweighted mean.
  This does not take the class imbalance problem into account.
  - Macro Precision: 0.919
  - Macro Recall: 0.918
  - Macro F1-score: 0.918

**Weighted Performance Metrics**
- Performance metrics calculated for each label and their average weighted by support (the number of true instances for each label).
  This extends the macro option to take the class imbalance into account.
  It might result in an F-score that is not between precision and recall.
  - Weighted Precision: 0.92
  - Weighted Recall: 0.919
  - Weighted F1-score: 0.919

**Classification Report**
The summary of the precision, recall, and F1-score for each class.

<table>
<thead>
<tr>
<th>precision</th>
<th>recall</th>
<th>f1-score</th>
<th>support</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.95</td>
<td>0.97</td>
<td>0.96</td>
</tr>
<tr>
<td>1.0</td>
<td>0.94</td>
<td>0.98</td>
<td>0.96</td>
</tr>
<tr>
<td>2.0</td>
<td>0.93</td>
<td>0.90</td>
<td>0.92</td>
</tr>
<tr>
<td>3.0</td>
<td>0.91</td>
<td>0.89</td>
<td>0.90</td>
</tr>
<tr>
<td>4.0</td>
<td>0.92</td>
<td>0.89</td>
<td>0.90</td>
</tr>
<tr>
<td>5.0</td>
<td>0.94</td>
<td>0.89</td>
<td>0.91</td>
</tr>
<tr>
<td>6.0</td>
<td>0.95</td>
<td>0.94</td>
<td>0.95</td>
</tr>
<tr>
<td>7.0</td>
<td>0.92</td>
<td>0.94</td>
<td>0.93</td>
</tr>
<tr>
<td>8.0</td>
<td>0.87</td>
<td>0.87</td>
<td>0.87</td>
</tr>
<tr>
<td>9.0</td>
<td>0.85</td>
<td>0.90</td>
<td>0.88</td>
</tr>
</tbody>
</table>

- accuracy          0.92          12000
- macro avg         0.92          12000
- weighted avg      0.92          12000

**Accuracy Rate of Each Diagonal Element Over Iteration**

This visualization is only applicable to binary classification and multiclass classification models. This is a line chart that plots the diagonal values in the confusion matrix throughout the training steps for each class. This plot shows you how the accuracy of each class progresses throughout the training steps. You can identify the under-performing classes from this plot.
Receiver Operating Characteristic Curve

This visualization is only applicable to binary classification models. The Receiver Operating Characteristic curve is commonly used to evaluate binary classification model performance. The y-axis of the curve is True Positive Rate (TPF) and x-axis is false positive rate (FPR). The plot also displays the value for the area under the curve (AUC). The higher the AUC value, the more predictive your classifier. You can also use the ROC curve to understand the trade-off between TPR and FPR and identify the optimum classification threshold for your use case. The classification threshold can be adjusted to tune the behavior of the model to reduce more of one or another type of error (FP/FN).
Distribution of Residuals at the Last Saved Step

This visualization is a column chart that shows the residual distributions in the last step Debugger captures. In this visualization, you can check whether the residual distribution is close to normal distribution that’s centered at zero. If the residuals are skewed, your features may not be sufficient for predicting the labels.
**Absolute Validation Error per Label Bin Over Iteration**

This visualization is only applicable to regression models. The actual target values are split into 10 intervals. This visualization shows how validation errors progress for each interval throughout the training steps in line plots. Absolute validation error is the absolute value of difference between prediction and actual during validation. You can identify the underperforming intervals from this visualization.
**Action on Amazon SageMaker Debugger Rules**

Based on the Debugger rule evaluation status, you can set up automated actions such as stopping a training job and sending notifications using Amazon Simple Notification Service (Amazon SNS). You can also create your own actions using Amazon CloudWatch Events and AWS Lambda. To learn how to set up automated actions based on the Debugger rule evaluation status, see the following topics.

**Topics**

- [Debugger Built-in Actions for Rules](#)
- [Create Actions on Rules Using Amazon CloudWatch and AWS Lambda](#)

**Debugger Built-in Actions for Rules**

Use Debugger built-in actions to respond to issues found by Debugger Rule. The Debugger rule_configs class provides tools to configure a list of actions, including automatically stopping training jobs and sending notifications using Amazon Simple Notification Service (Amazon SNS) when the Debugger rules find training issues.
Step 1: Set Up Amazon SNS, Create an SMDebugRules Topic, and Subscribe to the Topic

This section walks you through how to set up an Amazon SNS SMDebugRules topic, subscribe to it, and confirm the subscription to receive notifications from the Debugger rules.

Note
For more information about billing for Amazon SNS, see Amazon SNS pricing and Amazon SNS FAQs.

To create a SMDebugRules topic

2. In the left navigation pane, choose Topics.
3. On the Topics page, choose Create topic.
4. On the Create topic page, in the Details section, do the following:
   a. For Type, choose Standard for topic type.
   b. In Name, enter SMDebugRules.
5. Skip all other optional settings and choose Create topic. If you want to learn more about the optional settings, see Creating an Amazon SNS topic.

To subscribe to the SMDebugRules topic

2. In the left navigation pane, choose Subscriptions.
3. On the Subscriptions page, choose Create subscription.
4. On the Create subscription page, in the Details section, do the following:
   b. For Protocol, choose Email or SMS.
   c. For Endpoint, enter the endpoint value, such as an email address or a phone number that you want to receive notifications.
Note

Make sure you type the correct email address and phone number. Phone numbers must include +, a country code, and phone number, with no special characters or spaces. For example, the phone number +1 (222) 333-4444 is formatted as +12223334444.

5. Skip all other optional settings and choose Create subscription. If you want to learn more about the optional settings, see Subscribing to an Amazon SNS topic.

After you subscribe to the SMDdebugRules topic, you receive the following confirmation message in email or by phone:

**AWS Notification - Subscription Confirmation**

SMDdebugRules <no-reply@sns.amazonaws.com>
To:

You have chosen to subscribe to the topic:

To confirm this subscription, click or visit the link below (If this was in error no action is necessary):
Confirm subscription

Please do not reply directly to this email. If you wish to remove yourself from receiving all future SNS subscription confirmation requests please send an email to sns-opt-out

For more information about Amazon SNS, see Mobile text messaging (SMS) and Email notifications in the Amazon SNS Developer Guide.

**Step 2: Set Up Your IAM Role to Attach Required Policies**

In this step, you add the required policies to your IAM role.

To add the required policies to your IAM role

1. Sign in to the AWS Management Console and open the IAM console at https://console.aws.amazon.com/iam/.
2. In the left navigation pane, choose Policies, and choose Create policy.
3. On the Create policy page, do the following to create a new sns-access policy:
a. Choose the JSON tab.

b. Paste the JSON strings formatted in bold in the following code into the "Statement", replacing the 12-digit AWS account ID with your AWS account ID.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "VisualEditor0",
            "Effect": "Allow",
            "Action": [
                "sns:Publish",
                "sns:CreateTopic",
                "sns:Subscribe"
            ],
            "Resource": "arn:aws:sns::*:111122223333:SMDebugRules"
        }
    ]
}
```

c. At the bottom of the page, choose Review policy.

d. On the Review policy page, for Name, enter sns-access.

e. At the bottom of the page, choose Create policy.

4. Go back to the IAM console, and choose Roles in the left navigation pane.

5. Look up the IAM role that you use for SageMaker model training and choose that IAM role.

6. On the Permissions tab of the Summary page, choose Attach policies.

7. Search for the sns-access policy, select the check box next to the policy, and then choose Attach policy.

For more examples of setting up IAM policies for Amazon SNS, see Example cases for Amazon SNS access control.

**Step 3: Configure Debugger Rules with the Built-in Actions**

After successfully finishing the required settings in the preceding steps, you can configure the Debugger built-in actions for debugging rules as shown in the following example script. You can choose which built-in actions to use while building the actions list object. The rule_configs
is a helper module that provides high-level tools to configure Debugger built-in rules and actions. The following built-in actions are available for Debugger:

- **rule_configs.StopTraining()** – Stops a training job when the Debugger rule finds an issue.
- **rule_configs.Email("abc@abc.com")** – Sends a notification via email when the Debugger rule finds an issue. Use the email address that you used when you set up your SNS topic subscription.
- **rule_configs.SMS("+1234567890")** – Sends a notification via text message when the Debugger rule finds an issue. Use the phone number that you used when you set up your SNS topic subscription.

**Note**

Make sure you type the correct email address and phone number. Phone numbers must include +, a country code, and a phone number, with no special characters or spaces. For example, the phone number +1 (222) 333-4444 is formatted as +12223334444.

You can use all of the built-in actions or a subset of actions by wrapping up using the `rule_configs.ActionList()` method, which takes the built-in actions and configures a list of actions.

**To add all of the three built-in actions to a single rule**

If you want to assign all of the three built-in actions to a single rule, configure a Debugger built-in action list while constructing an estimator. Use the following template to construct the estimator, and Debugger will stop training jobs and send notifications through email and text for any rules that you use to monitor your training job progress.

```python
from sagemaker.debugger import Rule, rule_configs

# Configure an action list object for Debugger rules
actions = rule_configs.ActionList(
    rule_configs.StopTraining(),
    rule_configs.Email("abc@abc.com"),
    rule_configs.SMS("+1234567890")
)

# Configure rules for debugging with the actions parameter
rules = [
```
To create multiple built-in action objects to assign different actions to a single rule

If you want to assign the built-in actions to be triggered at different threshold values of a single rule, you can create multiple built-in action objects as shown in the following script. To avoid a conflict error by running the same rule, you must submit different rule job names (specify different strings for the rules' name attribute) as shown in the following example script template. This example shows how to set up `StalledTrainingRule` to take two different actions: send an email to abc@abc.com when a training job stalls for 60 seconds, and stop the training job if stalling for 120 seconds.

```python
from sagemaker.debugger import Rule, rule_configs
import time

base_job_name_prefix = 'smdebug-stalled-demo-' + str(int(time.time()))

# Configure an action object for StopTraining
action_stop_training = rule_configs.ActionList(
    rule_configs.StopTraining()
)

# Configure an action object for Email
action_email = rule_configs.ActionList(
    rule_configs.Email("abc@abc.com")
)

# Configure a rule with the Email built-in action to trigger if a training job stalls for 60 seconds
stalled_training_job_rule_email = Rule.sagemaker(
    base_config=rule_configs.built_in_rule(), # Required
    rule_parameters={"paramter_key": value }, # Optional
    actions=actions
)

estimator = Estimator(
    ...
    rules = rules
)

estimator.fit(wait=False)
```
base_config=rule_configs.stalled_training_rule(),
rule_parameters={
    "threshold": "60",
    "training_job_name_prefix": base_job_name_prefix
},
actions=action_email
)
stalled_training_job_rule_text.name="StalledTrainingJobRuleEmail"

# Configure a rule with the StopTraining built-in action to trigger if a training job
stalls for 120 seconds
stalled_training_job_rule = Rule.sagemaker(
    base_config=rule_configs.stalled_training_rule(),
    rule_parameters={
        "threshold": "120",
        "training_job_name_prefix": base_job_name_prefix
    },
    actions=action_stop_training
)
stalled_training_job_rule.name="StalledTrainingJobRuleStopTraining"

estimator = Estimator(
    ...
    rules = [stalled_training_job_rule_email, stalled_training_job_rule]
)
estimator.fit(wait=False)

While the training job is running, the Debugger built-in action sends notification emails and text
messages whenever the rule finds issues with your training job. The following screenshot shows an
example of email notification for a training job that has a stalled training job issue.
The following screenshot shows an example text notification that Debugger sends when the rule finds a StalledTraining issue.

Considerations for Using the Debugger Built-in Actions

- To use the Debugger built-in actions, an internet connection is required. This feature is not supported in the network isolation mode provided by Amazon SageMaker or Amazon VPC.
- The built-in actions cannot be used for Profiler rules.
- The built-in actions cannot be used on training jobs with spot training interruptions.
• In email or text notifications, None appears at the end of messages. This does not have any meaning, so you can disregard the text None.

Create Actions on Rules Using Amazon CloudWatch and AWS Lambda

Amazon CloudWatch collects Amazon SageMaker model training job logs and Amazon SageMaker Debugger rule processing job logs. Configure Debugger with Amazon CloudWatch Events and AWS Lambda to take action based on Debugger rule evaluation status.

CloudWatch Logs for Debugger Rules and Training Jobs

To find training job logs and Debugger rule job logs

2. In the left navigation pane under the Log node, choose Log Groups.
3. In the log groups list, do the following:
   • Choose /aws/sagemaker/TrainingJobs for training job logs.
   • Choose /aws/sagemaker/ProcessingJobs for Debugger rule job logs.

You can use the training and Debugger rule job status in the CloudWatch logs to take further actions when there are training issues.

For more information about monitoring training jobs using CloudWatch, see Monitor Amazon SageMaker.

Set Up Debugger for Automated Training Job Termination Using CloudWatch and Lambda

The Debugger rules monitor training job status, and a CloudWatch Events rule watches the Debugger rule training job evaluation status.

Step 1: Create a Lambda Function

To create a Lambda function

1. Open the AWS Lambda console at https://console.aws.amazon.com/lambda/.
2. In the left navigation pane, choose Functions and then choose Create function.
3. On the Create function page, choose Author from scratch option.
4. In the **Basic information** section, enter a **Function name** (for example, `debugger-rule-stop-training-job`).

5. For **Runtime**, choose **Python 3.7**.

6. For **Permissions**, expand the drop down option, and choose **Change default execution role**.

7. For **Execution role**, choose **Use an existing role** and choose the IAM role that you use for training jobs on SageMaker.

   **Note**
   Make sure you use the execution role with `AmazonSageMakerFullAccess` and `AWSLambdaBasicExecutionRole` attached. Otherwise, the Lambda function won't properly react to the Debugger rule status changes of the training job. If you are unsure which execution role is being used, run the following code in a Jupyter notebook cell to retrieve the execution role output:

   ```python
   import sagemaker
   sagemaker.get_execution_role()
   ```

8. At the bottom of the page, choose **Create function**.

The following figure shows an example of the **Create function** page with the input fields and selections completed.
Create function

Choose one of the following options to create your function.

- Author from scratch
  Start with a simple Hello World example.

- Use a blueprint
  Build a Lambda application from sample code and configuration presets for common use cases.

- Container image
  Select a container image to deploy for your function.

- Browse serverless app repository
  Deploy a sample Lambda application from the AWS Serverless Application Repository.

Basic information

Function name
Enter a name that describes the purpose of your function.

debugger-rule-stop-training-job
Use only letters, numbers, hyphens, or underscores with no spaces.

Runtime
Choose the language to use to write your function. Note that the console code editor supports only Node.js, Python, and Ruby.

Python 3.7

Permissions
By default, Lambda will create an execution role with permissions to upload logs to Amazon CloudWatch Logs. You can customize this default role later when adding triggers.

Change default execution role

Execution role
Choose a role that defines the permissions of your function. To create a custom role, go to the IAM console.

- Create a new role with basic Lambda permissions
- Use an existing role
- Create a new role from AWS policy templates

Existing role
Choose an existing role that you've created to be used with this Lambda function. The role must have permission to upload logs to Amazon CloudWatch Logs.

service-role/AmazonSageMaker-ExecutionRole-20200611T110452
View the AmazonSageMaker-ExecutionRole-20200611T110452 role on the IAM console.

Advanced settings

Cancel  Create function
Step 2: Configure the Lambda function

To configure the Lambda function

1. In the **Function code** section of the configuration page, paste the following Python script in the Lambda code editor pane. The `lambda_handler` function monitors the Debugger rule evaluation status collected by CloudWatch and triggers the `StopTrainingJob` API operation. The AWS SDK for Python (Boto3) client for SageMaker provides a high-level method, `stop_training_job`, which triggers the `StopTrainingJob` API operation.

```python
import json
import boto3
import logging

logger = logging.getLogger()
logger.setLevel(logging.INFO)

def lambda_handler(event, context):
    training_job_name = event.get("detail").get("TrainingJobName")
    logging.info(f'Evaluating Debugger rules for training job: {training_job_name}"
    eval_statuses = event.get("detail").get("DebugRuleEvaluationStatuses", None)

    if eval_statuses is None or len(eval_statuses) == 0:
        logging.info("Couldn't find any debug rule statuses, skipping...")
        return {
            'statusCode': 200,
            'body': json.dumps('Nothing to do')
        }

    # should only attempt stopping jobs with InProgress status
    training_job_status = event.get("detail").get("TrainingJobStatus", None)
    if training_job_status != 'InProgress':
        logging.debug(f"Current Training job status({training_job_status}) is not 'InProgress'. Exiting")
        return {
            'statusCode': 200,
            'body': json.dumps('Nothing to do')
        }
    
    client = boto3.client('sagemaker')
    for status in eval_statuses:
        
```
logging.info(status.get("RuleEvaluationStatus") + ', RuleEvaluationStatus=' + str(status))
if status.get("RuleEvaluationStatus") == "IssuesFound":
    secondary_status = event.get("detail").get("SecondaryStatus", None)
    logging.info(
        f'About to stop training job, since evaluation of rule configuration {status.get("RuleConfigurationName")} resulted in "IssuesFound". ' +
        f'\ntraining job "{training_job_name}" status is "{training_job_status}", secondary status is "{secondary_status}" ' +
        f'\nAttempting to stop training job "{training_job_name}"'
    )
    try:
        client.stop_training_job(
            TrainingJobName=training_job_name
        )
    except Exception as e:
        logging.error(
            "Encountered error while trying to "
            "stop training job {}": {}".format(
                training_job_name, str(e)
            )
        )
    raise e
return None

For more information about the Lambda code editor interface, see Creating functions using the AWS Lambda console editor.

2. Skip all other settings and choose **Save** at the top of the configuration page.

### Step 3: Create a CloudWatch Events Rule and Link to the Lambda Function for Debugger

**To create a CloudWatch Events rule and link to the Lambda function for Debugger**

2. In the left navigation pane, choose **Rules** under the **Events** node.
3. Choose **Create rule**.
4. In the **Event Source** section of the **Step 1: Create rule** page, choose **SageMaker** for **Service Name**, and choose **SageMaker Training Job State Change** for **Event Type**. The Event Pattern Preview should look like the following example JSON strings:
5. In the **Targets** section, choose **Add target***, and choose the **debugger-rule-stop-training-job** Lambda function that you created. This step links the CloudWatch Events rule with the Lambda function.

6. Choose **Configure details** and go to the **Step 2: Configure rule details** page.

7. Specify the CloudWatch rule definition name. For example, **debugger-cw-event-rule**.

8. Choose **Create rule** to finish.

9. Go back to the Lambda function configuration page and refresh the page. Confirm that it’s configured correctly in the **Designer** panel. The CloudWatch Events rule should be registered as a trigger for the Lambda function. The configuration design should look like the following example:
Run Example Notebooks to Test Automated Training Job Termination

You can run the following example notebooks, which are prepared for experimenting with stopping a training job using Debugger's built-in rules.

- **Amazon SageMaker Debugger - Reacting to CloudWatch Events from Rules**

  This example notebook runs a training job that has a vanishing gradient issue. The Debugger VanishingGradient built-in rule is used while constructing the SageMaker TensorFlow estimator. When the Debugger rule detects the issue, the training job is terminated.

- **Detect Stalled Training and Invoke Actions Using SageMaker Debugger Rule**

  This example notebook runs a training script with a code line that forces it to sleep for 10 minutes. The Debugger StalledTrainingRule built-in rule invokes issues and stops the training job.
Disable the CloudWatch Events Rule to Stop Using the Automated Training Job Termination

If you want to disable the automated training job termination, you need to disable the CloudWatch Events rule. In the Lambda Designer panel, choose the EventBridge (CloudWatch Events) block linked to the Lambda function. This shows an EventBridge panel below the Designer panel (for example, see the previous screen shot). Select the check box next to EventBridge (CloudWatch Events): debugger-cw-event-rule, and then choose Disable. If you want to use the automated termination functionality later, you can enable the CloudWatch Events rule again.

Visualize Amazon SageMaker Debugger Output Tensors in TensorBoard

⚠️ Important
This page is deprecated in favor of Amazon SageMaker with TensorBoard, which provides a comprehensive TensorBoard experience integrated with SageMaker Training and the access control functionalities of SageMaker domain. To learn more, see Use TensorBoard to debug and analyze training jobs in Amazon SageMaker.

Use SageMaker Debugger to create output tensor files that are compatible with TensorBoard. Load the files to visualize in TensorBoard and analyze your SageMaker training jobs. Debugger automatically generates output tensor files that are compatible with TensorBoard. For any hook configuration you customize for saving output tensors, Debugger has the flexibility to create scalar summaries, distributions, and histograms that you can import to TensorBoard.

Use SageMaker Debugger
You can enable this by passing DebuggerHookConfig and TensorBoardOutputConfig objects to an estimator.

The following procedure explains how to save scalars, weights, and biases as full tensors, histograms, and distributions that can be visualized with TensorBoard. Debugger saves them to the training container's local path (the default path is /opt/ml/output/tensors) and syncs to the Amazon S3 locations passed through the Debugger output configuration objects.

**To save TensorBoard compatible output tensor files using Debugger**

1. Set up a tensorboard_output_config configuration object to save TensorBoard output using the Debugger TensorBoardOutputConfig class. For the s3_output_path parameter, specify the default S3 bucket of the current SageMaker session or a preferred S3 bucket. This example does not add the container_local_output_path parameter; instead, it is set to the default local path /opt/ml/output/tensors.

```python
import sagemaker
from sagemaker.debugger import TensorBoardOutputConfig

bucket = sagemaker.Session().default_bucket()
tensorboard_output_config = TensorBoardOutputConfig(
    s3_output_path='s3://{}'.format(bucket)
)
```


2. Configure the Debugger hook and customize the hook parameter values. For example, the following code configures a Debugger hook to save all scalar outputs every 100 steps in training phases and 10 steps in validation phases, the weights parameters every 500 steps (the default save_interval value for saving tensor collections is 500), and the bias parameters every 10 global steps until the global step reaches 500.

```python
from sagemaker.debugger import CollectionConfig, DebuggerHookConfig

hook_config = DebuggerHookConfig(
    hook_parameters={
        "train.save_interval": "100",
        "eval.save_interval": "10"
    },
    collection_configs=[
```
CollectionConfig("weights"),
CollectionConfig(
    name="biases",
    parameters={
        "save_interval": "10",
        "end_step": "500",
        "save_histogram": "True"
    
    }
),
]
)

For more information about the Debugger configuration APIs, see the Debugger CollectionConfig and DebuggerHookConfig APIs in the Amazon SageMaker Python SDK.

3. Construct a SageMaker estimator with the Debugger parameters passing the configuration objects. The following example template shows how to create a generic SageMaker estimator. You can replace `estimator` and `Estimator` with other SageMaker frameworks' estimator parent classes and estimator classes. Available SageMaker framework estimators for this functionality are TensorFlow, PyTorch, and MXNet.

```python
from sagemaker.estimator import Estimator

estimator = Estimator(
    ...
    # Debugger parameters
degger_hook_config=hook_config,
tensorboard_output_conf=tensorboard_output_config
)
estimator.fit()
```

The `estimator.fit()` method starts a training job, and Debugger writes the output tensor files in real time to the Debugger S3 output path and to the TensorBoard S3 output path. To retrieve the output paths, use the following estimator methods:

- For the Debugger S3 output path, use `estimator.latest_job_debugger_artifacts_path()`.
- For the TensorBoard S3 output path, use `estimator.latest_job_tensorboard_artifacts_path()`.

4. After the training has completed, check the names of saved output tensors:
from smdebug.trials import create_trial
trial = create_trial(estimator.latest_job_debugger_artifacts_path())
trial.tensor_names()

5. Check the TensorBoard output data in Amazon S3:

tensorboard_output_path=estimator.latest_job_tensorboard_artifacts_path()
print(tensorboard_output_path)
!aws s3 ls {tensorboard_output_path}/

6. Download the TensorBoard output data to your notebook instance. For example, the following
AWS CLI command downloads the TensorBoard files to /logs/fit under the current working
directory of your notebook instance.

!aws s3 cp --recursive {tensorboard_output_path} ./logs/fit

7. Compress the file directory to a TAR file to download to your local machine.

!tar -cf logs.tar logs

8. Download and extract the Tensorboard TAR file to a directory on your device, launch a Jupyter
notebook server, open a new notebook, and run the TensorBoard app.

!tar -xf logs.tar
%load_ext tensorboard
%tensorboard --logdir logs/fit

List of Debugger Built-in Rules

Use the Debugger built-in rules provided by Amazon SageMaker Debugger and analyze metrics and
tensors collected while training your models. The Debugger built-in rules monitor various common
conditions that are critical for the success of a training job. You can call the built-in rules using
Amazon SageMaker Python SDK or the low-level SageMaker API operations. There’s no additional
cost for using the built-in rules. For more information about billing, see the Amazon SageMaker
Pricing page.
Note
The maximum numbers of built-in rules that you can attach to a training job is 20. SageMaker Debugger fully manages the built-in rules and analyzes your training job synchronously.

Important
To use the new Debugger features, you need to upgrade the SageMaker Python SDK and the SMDebug client library. In your iPython kernel, Jupyter notebook, or JupyterLab environment, run the following code to install the latest versions of the libraries and restart the kernel.

```python
import sys
import IPython
!{sys.executable} -m pip install -U sagemaker smdebug
IPython.Application.instance().kernel.do_shutdown(True)
```

Debugger Rule

The following rules are the Debugger built-in rules that are callable using the `Rule.sagemaker` classmethod.

Debugger built-in rules for generating training reports

<table>
<thead>
<tr>
<th>Scope of Validity</th>
<th>Built-in Rules</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training Report for SageMaker XGboost training job</td>
<td>• <a href="#">create_xgboost_report</a></td>
</tr>
</tbody>
</table>

Debugger built-in rules for debugging model training data (output tensors)

<table>
<thead>
<tr>
<th>Scope of Validity</th>
<th>Built-in Rules</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deep learning frameworks (TensorFlow, MXNet, and PyTorch)</td>
<td>• <a href="#">dead_relu</a></td>
</tr>
<tr>
<td></td>
<td>• <a href="#">exploding_tensor</a></td>
</tr>
</tbody>
</table>

Use SageMaker Debugger
<table>
<thead>
<tr>
<th>Scope of Validity</th>
<th>Built-in Rules</th>
</tr>
</thead>
<tbody>
<tr>
<td>Built-in Rules</td>
<td></td>
</tr>
<tr>
<td>• poor_weight_initialization</td>
<td></td>
</tr>
<tr>
<td>• saturated_activation</td>
<td></td>
</tr>
<tr>
<td>• vanishing_gradient</td>
<td></td>
</tr>
<tr>
<td>• weight_update_ratio</td>
<td></td>
</tr>
<tr>
<td>Deep learning frameworks (TensorFlow, MXNet, and PyTorch) and the XGBoost algorithm</td>
<td></td>
</tr>
<tr>
<td>• all_zero</td>
<td></td>
</tr>
<tr>
<td>• class_imbalance</td>
<td></td>
</tr>
<tr>
<td>• loss_not_decreasing</td>
<td></td>
</tr>
<tr>
<td>• overfit</td>
<td></td>
</tr>
<tr>
<td>• overtraining</td>
<td></td>
</tr>
<tr>
<td>• similar_across_runs</td>
<td></td>
</tr>
<tr>
<td>• stalled_training_rule</td>
<td></td>
</tr>
<tr>
<td>• tensor_variance</td>
<td></td>
</tr>
<tr>
<td>• unchanged_tensor</td>
<td></td>
</tr>
<tr>
<td>Deep learning applications</td>
<td></td>
</tr>
<tr>
<td>• check_input_images</td>
<td></td>
</tr>
<tr>
<td>• nlp_sequence_ratio</td>
<td></td>
</tr>
<tr>
<td>XGBoost algorithm</td>
<td></td>
</tr>
<tr>
<td>• confusion</td>
<td></td>
</tr>
<tr>
<td>• feature_importance_overweight</td>
<td></td>
</tr>
<tr>
<td>• tree_depth</td>
<td></td>
</tr>
</tbody>
</table>

To use the built-in rules with default parameter values – use the following configuration format:

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs

rules = [
    Rule.sagemaker(rule_configs.built_in_rule_name_1()),
    Rule.sagemaker(rule_configs.built_in_rule_name_2()),
    ...
    Rule.sagemaker(rule_configs.built_in_rule_name_n())
]
```
To use the built-in rules with customizing the parameter values – use the following configuration format:

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs

rules = [
    Rule.sagemaker(
        base_config=rule_configs.built_in_rule_name(),
        rule_parameters={
            "key": "value"
        },
        collections_to_save=[
            CollectionConfig(
                name="tensor_collection_name",
                parameters={
                    "key": "value"
                },
            )
        ]
    )
]
```

To find available keys for the `rule_parameters` parameter, see the parameter description tables.

Sample rule configuration codes are provided for each built-in rule below the parameter description tables.

- For a full instruction and examples of using the Debugger built-in rules, see Debugger Built-in Rules Example Code.
- For a full instruction on using the built-in rules with the low-level SageMaker API operations, see Configure Debugger Using Amazon SageMaker API.

**CreateXgboostReport**

The CreateXgboostReport rule collects output tensors from an XGBoost training job and autogenerates a comprehensive training report. You can download a comprehensive profiling report while a training job is running or after the training job is complete, and check progress of training or the final result of the training job. The CreateXgboostReport rule collects the following output tensors by default:

- `hyperparameters` – Saves at the first step
• metrics – Saves loss and accuracy every 5 steps
• feature_importance – Saves every 5 steps
• predictions – Saves every 5 steps
• labels – Saves every 5 steps

Parameter Descriptions for the CreateXgboostReport Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
</tbody>
</table>

**DeadRelu**

This rule detects when the percentage of rectified linear unit (ReLU) activation functions in a trial are considered dead because their activation activity has dropped below a threshold. If the percent of inactive ReLUs in a layer is greater than the threshold_layer value of inactive ReLUs, the rule returns True.

Parameter Descriptions for the DeadRelu Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>tensor_regex</strong></td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong>                                                                                                                                                                                                                                                                ---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;.*relu_output&quot;</td>
</tr>
<tr>
<td><strong>threshold_inactivity</strong></td>
<td>Defines a level of activity below which a ReLU is considered to be dead. A ReLU might be active in the beginning of a trial and then slowly die during the training process. If the ReLU is active less than the threshold_inactivity, it is considered to be dead.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: 1.0 (in percentage)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>threshold_layer</td>
<td>Returns <code>True</code> if the percentage of inactive ReLUs in a layer is greater than <code>threshold_layer</code>. Returns <code>False</code> if the percentage of inactive ReLUs in a layer is less than <code>threshold_layer</code>.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: 50.0 (in percentage)</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.dead_relu(),
        rule_parameters={
            "tensor_regex": ".*relu_output|.*ReLU_output",
            "threshold_inactivity": "1.0",
            "threshold_layer": "50.0"
        },
        collections_to_save=[
            CollectionConfig(
                name="custom_relu_collection",
                parameters={
                    "include_regex": ".*relu_output|.*ReLU_output",
                    "save_interval": "500"
                }
            )
        ]
    )
]
```

For an example of how to configure and deploy a built-in rule, see [Configure Debugger Built-in Rules](#).
ExplodingTensor

This rule detects whether the tensors emitted during training have non-finite values, either infinite or NaN (not a number). If a non-finite value is detected, the rule returns True.

Parameter Descriptions for the ExplodingTensor Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
</tbody>
</table>
### Parameter Name

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>only_nan</td>
<td>True to monitor the base_trial tensors only for NaN values and not for infinity. False to treat both NaN and infinity as exploding values and to monitor for both.</td>
</tr>
</tbody>
</table>

**Valid values:** String

**Default value:** None

**Optional**

**Default value:** False

---

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.exploding_tensor(),
        rule_parameters={
            "tensor_regex": ".*gradient",
            "only_nan": "False"
        },
        collections_to_save=[
            CollectionConfig(
                name="gradients",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```

For an example of how to configure and deploy a built-in rule, see [Configure Debugger Built-in Rules](#).

---

⚠️ **Note**

This rule is not available for the XGBoost algorithm.
**PoorWeightInitialization**

This rule detects if your model parameters have been poorly initialized.

Good initialization breaks the symmetry of the weights and gradients in a neural network and maintains commensurate activation variances across layers. Otherwise, the neural network doesn't learn effectively. Initializers like Xavier aim to keep variance constant across activations, which is especially relevant for training very deep neural nets. Too small an initialization can lead to vanishing gradients. Too large an initialization can lead to exploding gradients. This rule checks the variance of activation inputs across layers, the distribution of gradients, and the loss convergence for the initial steps to determine if a neural network has been poorly initialized.

**Parameter Descriptions for the PoorWeightInitialization Rule**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required. Valid values: String</td>
</tr>
<tr>
<td>activation_inputs_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched. Optional. Valid values: String. Default value: &quot;.*relu_input&quot;</td>
</tr>
<tr>
<td>threshold</td>
<td>If the ratio between minimum and maximum variance of weights per layer exceeds the threshold at a step, the rule returns True.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>distribution_range</td>
<td>If the minimum difference between 5th and 95th percentiles of the gradient distribution is less than the <code>distribution_range</code>, the rule returns True.</td>
</tr>
<tr>
<td>patience</td>
<td>The number of steps to wait until the loss is considered to be no longer decreasing.</td>
</tr>
<tr>
<td>steps</td>
<td>The number of steps this rule analyzes. You typically need to check only the first few iterations.</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.poor_weight_initialization()
    ),
```
rule_parameters={
    "activation_inputs_regex": ".*relu_input|.*ReLU_input",
    "threshold": "10.0",
    "distribution_range": "0.001",
    "patience": "5",
    "steps": "10"
},
collections_to_save=[
    CollectionConfig(
        name="custom_relu_collection",
        parameters={
            "include_regex": ".*relu_input|.*ReLU_input",
            "save_interval": "500"
        }
    )
]

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

⚠️ Note
This rule is not available for the XGBoost algorithm.

**SaturatedActivation**

This rule detects if the tanh and sigmoid activation layers are becoming saturated. An activation layer is saturated when the input of the layer is close to the maximum or minimum of the activation function. The minimum and maximum of the tanh and sigmoid activation functions are defined by their respective min_threshold and max_thresholds values. If the activity of a node drops below the threshold_inactivity percentage, it is considered saturated. If more than a threshold_layer percent of the nodes are saturated, the rule returns True.
# Parameter Descriptions for the SaturatedActivation Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
<th>Required</th>
<th>Valid values</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
<td>Required</td>
<td>String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects.</td>
<td>Optional</td>
<td>List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td></td>
<td>default</td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td></td>
<td>value</td>
<td>None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched.</td>
<td>Optional</td>
<td>String</td>
</tr>
<tr>
<td></td>
<td></td>
<td>default</td>
<td>&quot;.*tanh_input</td>
</tr>
<tr>
<td>threshold_tanh_min</td>
<td>The minimum and maximum thresholds that define the extremes of the input for a tanh activation function, defined as: (min_threshold, max_threshold). The default</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Use SageMaker Debugger
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>values are determined based on a vanishing gradient threshold of 0.0000001.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: -9.4999</td>
</tr>
<tr>
<td>threshold_tanh_max</td>
<td>The minimum and maximum thresholds that define the extremes of the input for a tanh activation function, defined as: (min_threshold, max_threshold) . The default values are determined based on a vanishing gradient threshold of 0.0000001.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: 9.4999</td>
</tr>
<tr>
<td>threshold_sigmoid_min</td>
<td>The minimum and maximum thresholds that define the extremes of the input for a sigmoid activation function, defined as: (min_threshold, max_threshold) . The default values are determined based on a vanishing gradient threshold of 0.0000001.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: -23</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>threshold_sigmoid_max</td>
<td>The minimum and maximum thresholds that define the extremes of the input for a sigmoid activation function, defined as: ((\text{min_threshold}, \text{max_threshold})). The default values are determined based on a vanishing gradient threshold of 0.0000001.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: 16.99999</td>
</tr>
<tr>
<td>threshold_inactivity</td>
<td>The percentage of inactivity below which the activation layer is considered to be saturated. The activation might be active in the beginning of a trial and then slowly become less active during the training process.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: 1.0</td>
</tr>
<tr>
<td>threshold_layer</td>
<td>Returns True if the number of saturated activations in a layer is greater than the threshold_layer percentage.</td>
</tr>
<tr>
<td></td>
<td>Returns False if the number of saturated activations in a layer is less than the threshold_layer percentage.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default values: 50.0</td>
</tr>
</tbody>
</table>
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.saturated_activation(),
        rule_parameters={
            "tensor_regex": ".*tanh_input|.*sigmoid_input",
            "threshold_tanh_min": "-9.4999",
            "threshold_tanh_max": "9.4999",
            "threshold_sigmoid_min": "-23",
            "threshold_sigmoid_max": "16.9999",
            "threshold_inactivity": "1.0",
            "threshold_layer": "50.0"
        },
        collections_to_save=[
            CollectionConfig(
                name="custom_activations_collection",
                parameters=
                    {
                        "include_regex": ".*tanh_input|.*sigmoid_input",
                        "save_interval": "500"
                    }
            )
        ]
    )
]

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

⚠️ Note
This rule is not available for the XGBoost algorithm.

VanishingGradient

This rule detects if the gradients in a trial become extremely small or drop to a zero magnitude. If the mean of the absolute values of the gradients drops below a specified threshold, the rule returns True.
### Parameters Descriptions for the VanishingGradient Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>threshold</td>
<td>The value at which the gradient is determined to be vanishing.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.0000001</td>
</tr>
</tbody>
</table>

For an example of how to configure and deploy a built-in rule, see [Configure Debugger Built-in Rules](#).

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.vanishing_gradient(),
        rule_parameters={
            "threshold": "0.0000001"
        },
        collections_to_save=[
            CollectionConfig(
                name="gradients",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```
Note
This rule is not available for the XGBoost algorithm.

**WeightUpdateRatio**

This rule keeps track of the ratio of updates to weights during training and detects if that ratio gets too large or too small. If the ratio of updates to weights is larger than the `large_threshold` value or if this ratio is smaller than `small_threshold`, the rule returns `True`.

Conditions for training are best when the updates are commensurate to gradients. Excessively large updates can push the weights away from optimal values, and very small updates result in very slow convergence. This rule requires weights to be available for two training steps, and `train.save_interval` needs to be set equal to `num_steps`.

**Parameter Descriptions for the WeightUpdateRatio Rule**

<table>
<thead>
<tr>
<th>Parameter Name,</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>base_trial</strong></td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required Valid values: String</td>
</tr>
<tr>
<td><strong>num_steps</strong></td>
<td>The number of steps across which the rule checks to determine if the tensor has changed. The number of steps across which you want to compare the weight ratios. If you pass no value, the rule runs by default against the current step and the immediately previous saved step. If you override the default by passing a value for this parameter, the comparison is done between weights at step $s$ and at a step $\geq s - num_steps$.</td>
</tr>
<tr>
<td>Parameter Name,</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>optional</td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td><strong>large_threshold</strong></td>
<td>The maximum value that the ratio of updates to weight can take before the rule returns True.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 10.0</td>
</tr>
<tr>
<td><strong>small_threshold</strong></td>
<td>The minimum value that the ratio of updates to weight can take, below which the rule returns True.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.00000001</td>
</tr>
<tr>
<td><strong>epsilon</strong></td>
<td>A small constant used to ensure that Debugger does not divide by zero when computing the ratio updates to weigh.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.000000001</td>
</tr>
</tbody>
</table>

```
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.weight_update_ratio(),
    )
```
rule_parameters={
    "num_steps": "100",
    "large_threshold": "10.0",
    "small_threshold": "0.00000001",
    "epsilon": "0.00000001"
},
collections_to_save=[
    CollectionConfig(
        name="weights",
        parameters={
            "train.save_interval": "100"
        }
    )
]

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Note
This rule is not available for the XGBoost algorithm.

AllZero
This rule detects if all or a specified percentage of the tensor values are zero.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm. You must specify either the collection_names or tensor_regex parameter. If both the parameters are specified, the rule inspects the union of tensors from both sets.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.
## Parameters Descriptions for the AllZero Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required</td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched. Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>threshold</td>
<td>Specifies the percentage of values in the tensor that needs to be zero for this rule to be invoked.</td>
</tr>
</tbody>
</table>
### ClassImbalance

This rule measures sampling imbalances between classes and throws errors if the imbalance exceeds a threshold or if too many mispredictions for underrepresented classes occur as a result of the imbalance.

Classification models require well-balanced classes in the training dataset or a proper weighting/sampling of classes during training. The rule performs the following checks:

- It counts the occurrences per class. If the ratio of number of samples between smallest and largest class is larger than the `threshold_imbalance`, an error is thrown.
- It checks the prediction accuracy per class. If resampling or weighting has not been correctly applied, then the model can reach high accuracy for the class with many training samples, but...
low accuracy for the classes with few training samples. If a fraction of mispredictions for a certain class is above `threshold_misprediction`, an error is thrown.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see [Configure Debugger Built-in Rules](#).

### Parameter Descriptions for the ClassImbalance Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>base_trial</code></td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required Valid values: String</td>
</tr>
<tr>
<td><code>threshold_imbalance</code></td>
<td>The acceptable imbalance between the number of samples in the smallest class and in the largest class. Exceeding this threshold value throws an error. Optional Valid values: Float Default value: 10</td>
</tr>
<tr>
<td><code>threshold_misprediction</code></td>
<td>A limit on the fraction of mispredictions allowed for each class. Exceeding this threshold throws an error. The underrepresented classes are most at risk of crossing this threshold. Optional Valid values: Float</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td><strong>samples</strong></td>
<td>The number of labels that have to be processed before an imbalance is evaluated. The rule might not be triggered until it has seen sufficient samples across several steps. The more classes that your dataset contains, the larger this sample number should be.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 500 (assuming a dataset like MNIST with 10 classes)</td>
</tr>
<tr>
<td><strong>argmax</strong></td>
<td>If True, \texttt{np.argmax} is applied to the prediction tensor. Required when you have a vector of probabilities for each class. It is used to determine which class has the highest probability.</td>
</tr>
<tr>
<td></td>
<td><strong>Conditional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Boolean</td>
</tr>
<tr>
<td></td>
<td>Default value: False</td>
</tr>
<tr>
<td><strong>labels_regex</strong></td>
<td>The name of the tensor that contains the labels.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;.*labels&quot;</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>predictions_regex</td>
<td>The name of the tensor that contains the predictions.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;.*predictions&quot;</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [  
    Rule.sagemaker(  
        base_config=rule_configs.class_imbalance(),  
        rule_parameters={  
            "threshold_imbalance": "10",  
            "threshold_misprediction": "0.7",  
            "samples": "500",  
            "argmax": "False",  
            "labels_regex": ".*labels",  
            "predictions_regex": ".*predictions"
        },  
        collections_to_save=[  
            CollectionConfig(  
                name="custom_output_collection",  
                parameters={  
                    "include_regex": ".*labels|.*predictions",  
                    "save_interval": "500"
                }
            )  
        ]
    )
]
```

**LossNotDecreasing**

This rule detects when the loss is not decreasing in value at an adequate rate. These losses must be scalars.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm. You must specify either the
collection_names or tensor_regex parameter. If both the parameters are specified, the rule inspects the union of tensors from both sets.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the LossNotDecreasing Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns that is used to restrict this comparison to specific scalar-valued tensors.</td>
</tr>
<tr>
<td></td>
<td>The rule inspects only the tensors that match the regex patterns specified in the list.</td>
</tr>
<tr>
<td></td>
<td>If no patterns are passed, the rule compares all tensors gathered in the trials by default.</td>
</tr>
<tr>
<td></td>
<td>Only scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>use_losses_collection</td>
<td>If set to True, looks for losses in the collection named &quot;losses&quot; when the collection is present.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Boolean</td>
</tr>
<tr>
<td></td>
<td>Default value: True</td>
</tr>
<tr>
<td>num_steps</td>
<td>The minimum number of steps after which the rule checks if the loss has decreased. Rule evaluation happens every num_steps. The rule compares the loss for this step with the loss at a step which is at least num_steps behind the current step. For example, suppose that the loss is being saved every three steps, but num_steps is set to 10. At step 21, loss for step 21 is compared with loss for step 9. The next step at which loss is checked is step 33, because ten steps after step 21 is step 31, and at step 31 and step 32 loss is not saved.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| **diff_percent**            | The minimum percentage difference by which the loss should decrease between `num_steps`.
|                             | Optional                                                                                             |
|                             | Valid values: \( 0.0 < \text{float} < 100 \)                                                          |
|                             | Default value: 0.1 (in percentage)                                                                     |
| **increase_threshold_percent** | The maximum threshold percent that loss is allowed to increase in case loss has been increasing       |
|                             | Optional                                                                                             |
|                             | Valid values: \( 0 < \text{float} < 100 \)                                                            |
|                             | Default value: 5 (in percentage)                                                                      |
| **mode**                    | The name of the Debugger mode to query tensor values for rule checking. If this is not passed, the   |
|                             | rule checks in order by default for the `mode.EVAL`, `mode.TRAIN`, and then `mode.GLOBAL`            |
|                             | Optional                                                                                             |
|                             | Valid values: String (EVAL, TRAIN, or GLOBAL)                                                         |
|                             | Default value: GLOBAL                                                                                 |

```
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.loss_not_decreasing(),
        rule_parameters={
            "tensor_regex": ".*",
            "use_losses_collection": "True",
        }
    )
]```
Overfit

This rule detects if your model is being overfit to the training data by comparing the validation and training losses.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Note

A standard way to prevent overfitting is to regularize your model.

Parameter Descriptions for the Overfit Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><code>parameter_name</code></td>
<td>Valid values: String</td>
</tr>
</tbody>
</table>
| `tensor_regex`     | A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched. **Optional**  
  Valid values: List of strings or a comma-separated string  
  Default value: None |
| `start_step`       | The step from which to start comparing the validation and training loss. **Optional**  
  Valid values: Integer  
  Default value: 0 |
| `patience`         | The number of steps for which the ratio_threshold is allowed to exceed the value set before the model is considered to be overfit. **Optional**  
  Valid values: Integer  
  Default value: 1 |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ratio_threshold</td>
<td>The maximum ratio of the difference between the mean validation loss and mean training loss to the mean training loss. If this threshold is exceeded for a patience number of steps, the model is being overfit and the rule returns True. <strong>Optional</strong> Valid values: Float Default value: 0.1</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.overfit(),
        rule_parameters={
            "tensor_regex": ".*",
            "start_step": "0",
            "patience": "1",
            "ratio_threshold": "0.1"
        },
        collections_to_save=[
            CollectionConfig(
                name="losses",
                parameters={
                    "train.save_interval": "100",
                    "eval.save_interval": "10"
                }
            )
        ]
    )
]
```

**Overtraining**

This rule detects if a model is being overtrained. After a number of training iterations on a well-behaved model (both training and validation loss decrease), the model approaches to a minimum
of the loss function and does not improve anymore. If the model continues training it can happen that validation loss starts increasing, because the model starts overfitting. This rule sets up thresholds and conditions to determine if the model is not improving, and prevents overfitting problems due to overtraining.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

**Note**

Overtraining can be avoided by early stopping. For information on early stopping, see Stop Training Jobs Early. For an example that shows how to use spot training with Debugger, see Enable Spot Training with Amazon SageMaker Debugger.

### Parameter Descriptions for the Overtraining Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td>patience_train</td>
<td>The number of steps to wait before the training loss is considered to not be improving anymore.</td>
</tr>
<tr>
<td></td>
<td>Required</td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>patience_validation</td>
<td>The number of steps to wait before the validation loss is considered to not to be improving anymore.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>delta</td>
<td>The minimum threshold by how much the error should improve before it is considered as a new optimum.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.01</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.overtraining(),
        rule_parameters={
            "patience_train": "5",
            "patience_validation": "10",
            "delta": "0.01"
        },
        collections_to_save=[
            CollectionConfig(
                name="losses",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```
**SimilarAcrossRuns**

This rule compares tensors gathered from a base trial with tensors from another trial.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see [Configure Debugger Built-in Rules](#).

**Parameter Descriptions for the SimilarAcrossRuns Rule**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>other_trials</td>
<td>A completed training job name whose tensors you want to compare to those tensors gathered from the current base_trial.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td></td>
<td>The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.similar_across_runs(),
        rule_parameters={
            "other_trials": "<specify-another-job-name>",
            "collection_names": "losses",
            "tensor_regex": ".*"
        },
        collections_to_save=[
            CollectionConfig(
                name="losses",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```

**StalledTrainingRule**

StalledTrainingRule detects if there is no progress made on training job, and stops the training job if the rule fires. This rule requires tensors to be periodically saved in a time interval defined by its threshold parameter. This rule keeps on monitoring for new tensors, and if no new tensor has been emitted for threshold interval rule gets fired.
# Parameter Descriptions for the StalledTrainingRule Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required Valid values: String</td>
</tr>
<tr>
<td>threshold</td>
<td>A threshold that defines by how much time in seconds the rule waits for a tensor output until it fires a stalled training issue. Default value is 1800 seconds. Optional Valid values: Integer Default value: 1800</td>
</tr>
<tr>
<td>stop_training_on_fire</td>
<td>If set to True, watches if the base training job outputs tensors in &quot;threshold &quot; seconds. Optional Valid values: Boolean Default value: False</td>
</tr>
<tr>
<td>training_job_name_prefix</td>
<td>The prefix of base training job name. If stop_training_on_fire is true, the rule searches for SageMaker training jobs with this prefix in the same account. If there is an inactivity found, the rule takes a StopTrainingJob action. Note if there are multiple jobs found with same prefix, the rule skips termination. It is important that the prefix is set unique per each training job.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>built_in_rules</td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.stalled_training_rule(),
        rule_parameters={
            "threshold": "1800",
            "stop_training_on_fire": "True",
            "training_job_name_prefix": "<specify-training-base-job-name>"
        },
        collections_to_save=[
            CollectionConfig(
                name="losses",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```

**TensorVariance**

This rule detects if you have tensors with very high or low variances. Very high or low variances in a tensor could lead to neuron saturation, which reduces the learning ability of the neural network. Very high variance in tensors can also eventually lead to exploding tensors. Use this rule to detect such issues early.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm. You must specify either the `collection_names` or `tensor_regex` parameter. If both the parameters are specified, the rule inspects the union of tensors from both sets.

For an example of how to configure and deploy a built-in rule, see [Configure Debugger Built-in Rules](#).
## Parameter Descriptions for the TensorVariance Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>max_threshold</td>
<td>The threshold for the upper bound of tensor variance.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>min_threshold</td>
<td>The threshold for the lower bound of tensor variance.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.tensor_variance(),
        rule_parameters={
            "collection_names": "weights",
            "max_threshold": "10",
            "min_threshold": "0.00001",
        },
        collections_to_save=[
            CollectionConfig(
                name="weights",
                parameters={
                    "save_interval": "500"
                } |
            ]
        ]
    )
]
```

**UnchangedTensor**

This rule detects whether a tensor is no longer changing across steps.

This rule runs the `numpy.allclose` method to check if the tensor isn't changing.

This rule can be applied either to one of the supported deep learning frameworks (TensorFlow, MXNet, and PyTorch) or to the XGBoost algorithm. You must specify either the
collection_names or tensor_regex parameter. If both the parameters are specified, the rule inspects the union of tensors from both sets.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the UnchangedTensor Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>collection_names</td>
<td>The list of collection names whose tensors the rule inspects.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match</td>
</tr>
<tr>
<td></td>
<td>the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only</td>
</tr>
<tr>
<td></td>
<td>scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
</tr>
<tr>
<td>num_steps</td>
<td>The number of steps across which the rule checks to determine if the tensor has changed. This checks the last <code>num_steps</code> that are available. They don't need to be consecutive. If <code>num_steps</code> is 2, at step <code>s</code> it doesn't necessarily check for <code>s-1</code> and <code>s</code>. If <code>s-1</code> isn't available, it checks the last available step along with <code>s</code>. In that case, it checks the last available step with the current step. <strong>Optional</strong> Valid values: Integer Default value: 3</td>
</tr>
<tr>
<td>rtol</td>
<td>The relative tolerance parameter to be passed to the <code>numpy.allclose</code> method. <strong>Optional</strong> Valid values: Float Default value: 1e-05</td>
</tr>
<tr>
<td>atol</td>
<td>The absolute tolerance parameter to be passed to the <code>numpy.allclose</code> method. <strong>Optional</strong> Valid values: Float Default value: 1e-08</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>equal_nan</td>
<td>Whether to compare NaNs as equal. If True, NaNs in input array a are considered equal to NaNs in input array b in the output array. This parameter is passed to the <code>numpy.allclose</code> method.</td>
</tr>
</tbody>
</table>

**Optional**

Valid values: Boolean

Default value: False

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.unchanged_tensor(),
        rule_parameters={
            "collection_names": "losses",
            "tensor_regex": "",
            "num_steps": "3",
            "rtol": "1e-05",
            "atol": "1e-08",
            "equal_nan": "False"
        },
        collections_to_save=[
            CollectionConfig(
                name="losses",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```
CheckInputImages

This rule checks if input images have been correctly normalized. Specifically, it detects if the mean of the sample data differs by more than a threshold value from zero. Many computer vision models require that input data has a zero mean and unit variance.

This rule is applicable to deep learning applications.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the CheckInputImages Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>threshold_mean</td>
<td>A threshold that defines by how much mean of the input data can differ from 0.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.2</td>
</tr>
<tr>
<td>threshold_samples</td>
<td>The number of images that have to be sampled before an error can be thrown. If the value is too low, the estimation of the dataset mean will be inaccurate.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 500</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| regex          | The name of the input data tensor.  
**Optional**  
Valid values: String  
Default value: ".*hybridsequential0_input_0" (the name of the input tensor for Apache MXNet models using HybridSequential) |
| channel        | The position of the color channel in the input tensor shape array.  
**Optional**  
Valid values: Integer  
Default value: 1 (for example, MXNet expects input data in the form of (batch_size, channel, height, width)) |

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.check_input_images(),
        rule_parameters={
            "threshold_mean": "0.2",
            "threshold_samples": "500",
            "regex": ".*hybridsequential0_input_0",
            "channel": "1"
        },
        collections_to_save=[
            CollectionConfig(
                name="custom_inputs_collection",
                parameters={
                    "include_regex": ".*hybridsequential0_input_0",
                    "save_interval": "500"
                }
            )
        ]
    )
]```
NLPSequenceRatio

This rule calculates the ratio of specific tokens given the rest of the input sequence that is useful for optimizing performance. For example, you can calculate the percentage of padding end-of-sentence (EOS) tokens in your input sequence. If the number of EOS tokens is too high, an alternate bucketing strategy should be performed. You also can calculate the percentage of unknown tokens in your input sequence. If the number of unknown words is too high, an alternate vocabulary could be used.

This rule is applicable to deep learning applications.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the NLPSequenceRatio Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>A list of regex patterns used to restrict this comparison to specific scalar-valued tensors. The rule inspects only the tensors that match the regex patterns specified in the list. If no patterns are passed, the rule compares all tensors gathered in the trials by default. Only scalar-valued tensors can be matched.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Valid values: List of strings or a comma-separated string</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;.*embedding0_input_0&quot; (assuming an embedding as the initial layer of the network)</td>
</tr>
<tr>
<td>token_values</td>
<td>A string of a list of the numerical values of the tokens. For example, &quot;3, 0&quot;.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Comma-separated string of numerical values</td>
</tr>
<tr>
<td></td>
<td>Default value: 0</td>
</tr>
<tr>
<td>token_thresholds_percent</td>
<td>A string of a list of thresholds (in percentages) that correspond to each of the token values. For example,&quot;50.0, 50.0&quot;.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Comma-separated string of floats</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;50&quot;</td>
</tr>
</tbody>
</table>

```
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.nlp_sequence_ratio(),
        rule_parameters={
            "tensor_regex": ".*embedding0_input_0",
            "token_values": "0",
            "token_thresholds_percent": "50"
        },
        collections_to_save=[
            CollectionConfig(
                name="custom_inputs_collection",
                parameters={
```
Confusion

This rule evaluates the goodness of a confusion matrix for a classification problem.

It creates a matrix of size category_no*category_no and populates it with data coming from (labels, predictions) pairs. For each (labels, predictions) pair, the count in confusion[labels][predictions] is incremented by 1. When the matrix is fully populated, the ratio of data on-diagonal values and off-diagonal values are evaluated as follows:

- For elements on the diagonal: confusion[i][i]/sum_j(confusion[j][j])>=min_diag
- For elements off the diagonal: confusion[j][i])/sum_j(confusion[j][i])<=max_off_diag

This rule can be applied to the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the Confusion Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td>category_no</td>
<td>The number of categories.</td>
</tr>
</tbody>
</table>

Required

Valid values: String

Optional
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Valid values: Integer ≥2</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;None&quot;</td>
</tr>
<tr>
<td>labels</td>
<td>The labels tensor collection or an 1-d vector of true labels.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;labels&quot;</td>
</tr>
<tr>
<td>predictions</td>
<td>The predictions tensor collection or an 1-d vector of estimated labels.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;predictions&quot;</td>
</tr>
<tr>
<td>labels_collection</td>
<td>The rule inspects the tensors in this collection for labels.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;labels&quot;</td>
</tr>
<tr>
<td>predictions_collection</td>
<td>The rule inspects the tensors in this collection for predictions.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;predictions&quot;</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>min_diag</td>
<td>The minimum threshold for the ratio of data on the diagonal.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: 0≤float≤1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.9</td>
</tr>
<tr>
<td>max_off_diag</td>
<td>The maximum threshold for the ratio of data off the diagonal.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: 0≤float≤1</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.1</td>
</tr>
</tbody>
</table>

```
built_in_rules = [  
    Rule.sagemaker(  
        base_config=rule_configs.confusion(),  
        rule_parameters={  
            "category_no": "10",  
            "labels": "labels",  
            "predictions": "predictions",  
            "labels_collection": "labels",  
            "predictions_collection": "predictions",  
            "min_diag": "0.9",  
            "max_off_diag": "0.1"  
        },  
        collections_to_save=[  
            CollectionConfig(  
                name="labels",  
                parameters={  
                    "save_interval": "500"  
                }  
            ),  
            CollectionConfig(  
                name="predictions",  
            )  
        ]  
    )  
]```
parameters={
    "include_regex": "500"
}
]
]
]

Note
This rule infers default values for the optional parameters if their values aren't specified.

FeatureImportanceOverweight

This rule accumulates the weights of the n largest feature importance values per step and ensures that they do not exceed the threshold. For example, you can set the threshold for the top 3 features to not hold more than 80 percent of the total weights of the model.

This rule is valid only for the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the FeatureImportanceOverweight Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required Valid values: String</td>
</tr>
<tr>
<td>threshold</td>
<td>Defines the threshold for the proportion of the cumulative sum of the n largest features. The number n is defined by the nfeatures parameter. Optional</td>
</tr>
</tbody>
</table>

Use SageMaker Debugger
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Valid values: Float</td>
</tr>
<tr>
<td></td>
<td>Default value: 0.8</td>
</tr>
<tr>
<td>nfeatures</td>
<td>The number of largest features.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 3</td>
</tr>
<tr>
<td>tensor_regex</td>
<td>Regular expression (regex) of tensor names the rule to analyze.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td></td>
<td>Default value: &quot;.*feature_importance/weight&quot;</td>
</tr>
</tbody>
</table>

```python
built_in_rules = [
    Rule.sagemaker(
        base_config=rule_configs.feature_importance_overweight(),
        rule_parameters=
        {
            "threshold": "0.8",
            "nfeatures": "3",
            "tensor_regex": ".*feature_importance/weight"
        },
        collections_to_save=[
            CollectionConfig(
                name="feature_importance",
                parameters={
                    "save_interval": "500"
                }
            )
        ]
    )
]
```
TreeDepth

This rule measures the depth of trees in an XGBoost model. XGBoost rejects splits if they do not improve loss. This regularizes the training. As a result, the tree might not grow as deep as defined by the depth parameter.

This rule is valid only for the XGBoost algorithm.

For an example of how to configure and deploy a built-in rule, see Configure Debugger Built-in Rules.

Parameter Descriptions for the TreeDepth Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. Required Valid values: String</td>
</tr>
<tr>
<td>depth</td>
<td>The depth of the tree. The depth of the tree is obtained by computing the base 2 logarithm of the largest node ID. Optional Valid values: Float Default value: 4</td>
</tr>
</tbody>
</table>

Use SageMaker Debugger
collections_to_save=[
    CollectionConfig(
        name="tree",
        parameters={
            "save_interval": "500"
        }
    )
]
]

Create Debugger Custom Rules for Training Job Analysis

You can create custom rules to monitor your training job using the Debugger Rule APIs and the open source smdebug Python library that provide tools to build your own rule containers.

Topics
- Prerequisites for Creating Debugger Custom Rules
- Use the Debugger Client Library smdebug to Create a Custom Rule Python Script
- Use the Debugger APIs to Run Your Own Custom Rules

Prerequisites for Creating Debugger Custom Rules

To create Debugger custom rules, you need the following prerequisites.

- SageMaker Debugger Rule.custom API
- The open source smdebug Python library
- Your own custom rule python script
- Amazon SageMaker Debugger Registry URLs for Custom Rule Evaluators

Use the Debugger Client Library smdebug to Create a Custom Rule Python Script

The smdebug Rule API provides an interface to set up your own custom rules. The following python script is a sample of how to construct a custom rule, CustomGradientRule. This tutorial custom rule watches if the gradients are getting too large and set the default threshold as 10. The custom rule takes a base trial created by a SageMaker estimator when it initiates training job.

```python
from smdebug.rules.rule import Rule
```
class CustomGradientRule(Rule):
    def __init__(self, base_trial, threshold=10.0):
        super().__init__(base_trial)
        self.threshold = float(threshold)

    def invoke_at_step(self, step):
        for tname in self.base_trial.tensor_names(collection="gradients"):
            t = self.base_trial.tensor(tname)
            abs_mean = t.reduction_value(step, "mean", abs=True)
            if abs_mean > self.threshold:
                return True
        return False

You can add multiple custom rule classes as many as you want in the same python script and deploy them to any training job trials by constructing custom rule objects in the following section.

Use the Debugger APIs to Run Your Own Custom Rules

The following code sample shows how to configure a custom rule with the Amazon SageMaker Python SDK. This example assumes that the custom rule script you created in the previous step is located at 'path/to/my_custom_rule.py'.

```python
from sagemaker.debugger import Rule, CollectionConfig

custom_rule = Rule.custom(
    name='MyCustomRule',
    image_uri='759209512951.dkr.ecr.us-west-2.amazonaws.com/sagemaker-debugger-rule-evaluator:latest',
    instance_type='ml.t3.medium',
    source='path/to/my_custom_rule.py',
    rule_to_invoke='CustomGradientRule',
    collections_to_save=[CollectionConfig("gradients")],
    rule_parameters={"threshold": "20.0"}
)
```

The following list explains the Debugger Rule.custom API arguments.

- **name (str)**: Specify a custom rule name as you want.
- **image_uri (str)**: This is the image of the container that has the logic of understanding your custom rule. It sources and evaluates the specified tensor collections you save in the training job.
You can find the list of open source SageMaker rule evaluator images from [Amazon SageMaker Debugger Registry URLs for Custom Rule Evaluators](https://docs.aws.amazon.com/sagemaker/latest/dg/debugger-registry-rule-evaluators.html).

- **instance_type (str)**: You need to specify an instance to build a rule docker container. This spins up the instance in parallel with a training container.

- **source (str)**: This is the local path or the Amazon S3 URI to your custom rule script.

- **rule_to_invoke (str)**: This specifies the particular Rule class implementation in your custom rule script. SageMaker supports only one rule to be evaluated at a time in a rule job.

- **collections_to_save (str)**: This specifies which tensor collections you will save for the rule to run.

- **rule_parameters (dictionary)**: This accepts parameter inputs in a dictionary format. You can adjust the parameters that you configured in the custom rule script.

After you set up the `custom_rule` object, you can use it for building a SageMaker estimator for any training jobs. Specify the `entry_point` to your training script. You do not need to make any change of your training script.

```python
from sagemaker.tensorflow import TensorFlow

estimator = TensorFlow(
    role=sagemaker.get_execution_role(),
    base_job_name='smdebug-custom-rule-demo-tf-keras',
    entry_point='path/to/your_training_script.py'
    train_instance_type='ml.p2.xlarge'
    ...

    # debugger-specific arguments below
    rules = [custom_rule]
)

estimator.fit()
```

For more variations and advanced examples of using Debugger custom rules, see the following example notebooks.

- [Monitor your training job with Amazon SageMaker Debugger custom rules](https://docs.aws.amazon.com/sagemaker/latest/dg/debugger-custom-rule-examples.html)
- [PyTorch iterative model pruning of ResNet and AlexNet](https://docs.aws.amazon.com/sagemaker/latest/dg/tensorflow-pruning.html)
Use Debugger with Custom Training Containers

Amazon SageMaker Debugger is available for any deep learning models that you bring to Amazon SageMaker. The AWS CLI, SageMaker Estimator API, and the Debugger APIs enable you to use any Docker base images to build and customize containers to train your models. To use Debugger with customized containers, you need to make a minimal change to your training script to implement the Debugger hook callback and retrieve tensors from training jobs.

You need the following resources to build a customized container with Debugger.

- Amazon SageMaker Python SDK
- The SMDebug open source client library
- A Docker base image of your choice
- Your training script with a Debugger hook registered – For more information about registering a Debugger hook to your training script, see Register Debugger Hook to Your Training Script.

For an end-to-end example of using Debugger with a custom training container, see the following example notebook.

- Build a Custom Training Container and Debug Training Jobs with Debugger

Tip

This custom container with Debugger guide is an extension of the Adapting your own training container guide which walks you thorough how to build and push your custom training container to Amazon ECR.

Prepare to Build a Custom Training Container

To build a docker container, the basic structure of files should look like the following:

```bash
### debugger_custom_container_test_notebook.ipynb # a notebook to run python snippet codes
```
Register Debugger Hook to Your Training Script

To debug your model training, you need to add a Debugger hook to your training script.

**Note**

This step is required to collect model parameters (output tensors) for debugging your model training. If you only want to monitor and profile, you can skip this hook registration step and exclude the `debugger_hook_config` parameter when constructing an estimator.

The following example code shows the structure of a training script using the Keras ResNet50 model and how to pass the Debugger hook as a Keras callback for debugging. To find a complete training script, see [TensorFlow training script with SageMaker Debugger hook](#).

```python
# An example of training script (your-training-script.py)
import tensorflow.compat.v2 as tf
from tensorflow.keras.applications.resnet50 import ResNet50
import smdebug.tensorflow as smd

def train(batch_size, epoch, model, hook):
    ...
    model.fit(X_train, Y_train,
        batch_size=batch_size,
        epochs=epoch,
        validation_data=(X_valid, Y_valid),
        shuffle=True,
        callbacks=[hook])

def main():
    parser=argparse.ArgumentParser(description="Train resnet50 cifar10")
```
# hyperparameter settings
parser.add_argument(...)

args = parser.parse_args()

model=ResNet50(weights=None, input_shape=(32,32,3), classes=10)

# Add the following line to register the Debugger hook for Keras.
hook=smd.KerasHook.create_from_json_file()

# Start the training.
train(args.batch_size, args.epoch, model, hook)

if __name__ == "__main__":
    main()

For more information about registering the Debugger hook for the supported frameworks and algorithm, see the following links in the SMDebug client library:

- [SMDebug TensorFlow hook](#)
- [SMDebug PyTorch hook](#)
- [SMDebug MXNet hook](#)
- [SMDebug XGBoost hook](#)

In the following example notebooks’ training scripts, you can find more examples about how to add the Debugger hooks to training scripts and collect output tensors in detail:

- [Debugger in script mode with the TensorFlow 2.1 framework](#)

To see the difference between using Debugger in a Deep Learning Container and in script mode, open this notebook and put it and [the previous Debugger in a Deep Learning Container TensorFlow v2.1 notebook example](#) side by side.

In script mode, the hook configuration part is removed from the script in which you set the estimator. Instead, the Debugger hook feature is merged into the training script, [TensorFlow Keras ResNet training script in script mode](#). The training script imports the smdebug library in the required TensorFlow Keras environment to communicate with the TensorFlow ResNet50 algorithm. It also manually implements the smdebug hook functionality by adding the
callbacks=[hook] argument inside the train function (in line 49), and by adding the manual hook configuration (in line 89) provided through SageMaker Python SDK.

This script mode example runs the training job in the TF 2.1 framework for direct comparison with the zero script change in the TF 2.1 example. The benefit of setting up Debugger in script mode is the flexibility to choose framework versions not covered by AWS Deep Learning Containers.

• **Using Amazon SageMaker Debugger in a PyTorch Container in Script Mode**

This notebook enables Debugger in script mode in PyTorch v1.3.1 framework. PyTorch v1.3.1 is supported by SageMaker containers, and this example shows details of how to modify a training script.

The SageMaker PyTorch estimator is already in script mode by default. In the notebook, the line to activate `script_mode` is not included in the estimator configuration.

This notebook shows detailed steps to change the original PyTorch training script to a modified version to enable Debugger. Additionally, this example shows how you can use Debugger built-in rules to detect training issues such as the vanishing gradients problem, and the Debugger trial features to call and analyze the saved tensors.

**Create and Configure a Dockerfile**

Open your SageMaker JupyterLab and create a new folder, `debugger_custom_container_test_folder` in this example, to save your training script and Dockerfile. The following code example is a Dockerfile that includes essential docker build commend. Paste the following code into the Dockerfile text file and save it. Upload your training script to the same folder.

```bash
# Specify a docker base image
FROM tensorflow/tensorflow:2.2.0rc2-gpu-py3
RUN /usr/bin/python3 -m pip install --upgrade pip
RUN pip install --upgrade protobuf

# Install required packages to enable the SageMaker Python SDK and the smdebug library
RUN pip install sagemaker-training
RUN pip install smdebug

CMD ["bin/bash"]
```
If you want to use a pre-built AWS Deep Learning Container image, see Available AWS Deep Learning Containers Images.

### Build and Push the Custom Training Container to Amazon ECR

Create a test notebook, debugger_custom_container_test_notebook.ipynb, and run the following code in the notebook cell. This will access the debugger_byoc_test_docker directory, build the docker with the specified algorithm_name, and push the docker container to your Amazon ECR.

```python
import boto3

account_id = boto3.client('sts').get_caller_identity().get('Account')
ecr_repository = 'sagemaker-debugger-mnist-byoc-tf2'
tag = ':latest'

region = boto3.session.Session().region_name

uri_suffix = 'amazonaws.com'
if region in ['cn-north-1', 'cn-northwest-1']:
    uri_suffix = 'amazonaws.com.cn'
byoc_image_uri = '{}.dkr.ecr.{}.{}/{}'.format(account_id, region, uri_suffix, ecr_repository + tag)

!docker build -t $ecr_repository docker
!$(aws ecr get-login --region $region --registry-ids $account_id --no-include-email)
!aws ecr create-repository --repository-name $ecr_repository
!docker tag {ecr_repository + tag} $byoc_image_uri
!docker push $byoc_image_uri
```

#### Tip

If you use one of the AWS Deep Learning Container base images, run the following code to log in to Amazon ECR and access to the Deep Learning Container image repository.

```bash
! aws ecr get-login-password --region {region} | docker login --username AWS --password-stdin 763104351884.dkr.ecr.us-east-1.amazonaws.com
```
Run and Debug Training Jobs Using the Custom Training Container

After you build and push your docker container to Amazon ECR, configure a SageMaker estimator with your training script and the Debugger-specific parameters. After you execute the estimator.fit(), Debugger will collect output tensors, monitor them, and detect training issues. Using the saved tensors, you can further analyze the training job by using the smedebug core features and tools. Configuring a workflow of Debugger rule monitoring process with Amazon CloudWatch Events and AWS Lambda, you can automate a stopping training job process whenever the Debugger rules spots training issues.

```python
import sagemaker
from sagemaker.estimator import Estimator
from sagemaker.debugger import Rule, DebuggerHookConfig, CollectionConfig, rule_configs

profiler_config = ProfilerConfig(...)
double_hook_config = DebuggerHookConfig(...)

rules = [
    Rule.sagemaker(rule_configs.built_in_rule()),
    ProfilerRule.sagemaker(rule_configs.BuiltInRule())
]

estimator = Estimator(
    image_uri=byoc_image_uri,
    entry_point="./debugger_custom_container_test_folder/your-training-script.py",
    role=sagemaker.get_execution_role(),
    base_job_name='debugger-custom-container-test',
    instance_count=1,
    instance_type='ml.p3.2xlarge',

    # Debugger-specific parameters
    profiler_config=profiler_config,
    debugger_hook_config=debugger_hook_config,
    rules=rules
)

# start training
estimator.fit()
```

Configure Debugger Using Amazon SageMaker API

The preceding topics focus on using Debugger through Amazon SageMaker Python SDK, which is a wrapper around AWS SDK for Python (Boto3) and SageMaker API operations. This offers a high-
level experience of accessing the Amazon SageMaker API operations. In case you need to manually configure the SageMaker API operations using AWS Boto3 or AWS Command Line Interface (CLI) for other SDKs, such as Java, Go, and C++, this section covers how to configure the following low-level API operations.

Topics

- **JSON (AWS CLI)**
- **AWS Boto3**

### JSON (AWS CLI)

Amazon SageMaker Debugger built-in rules can be configured for a training job using the `DebugHookConfig`, `DebugRuleConfiguration`, `ProfilerConfig`, and `ProfilerRuleConfiguration` objects through the SageMaker `CreateTrainingJob` API operation. You need to specify the right image URI in the `RuleEvaluatorImage` parameter, and the following examples walk you through how to set up the JSON strings to request `CreateTrainingJob`.

The following code shows a complete JSON template to run a training job with required settings and Debugger configurations. Save the template as a JSON file in your working directory and run the training job using AWS CLI. For example, save the following code as `debugger-training-job-cli.json`.

```json
{
    "TrainingJobName": "debugger-aws-cli-test",
    "RoleArn": "arn:aws:iam::111122223333:role/service-role/AmazonSageMaker-ExecutionRole-YYYYMMDDT123456",
    "AlgorithmSpecification": {
        // Specify a training Docker container image URI (Deep Learning Container or your own training container) to TrainingImage.
    }
}
```

**Note**

Ensure that you use the correct Docker container images. To find AWS Deep Learning Container images, see [Available Deep Learning Containers Images](#). To find a complete list of available Docker images for using the Debugger rules, see [Use Debugger Docker Images for Built-in or Custom Rules](#).
"TrainingInputMode": "File",
    "EnableSageMakerMetricsTimeSeries": false
},
    "HyperParameters": {
        "sagemaker_program": "entry_point/tf-hvd-train.py",
        "sagemaker_submit_directory": "s3://sagemaker-us-west-2-111122223333/debugger-boto3-profiling-test/source.tar.gz"
    },
    "OutputDataConfig": {
        "S3OutputPath": "s3://sagemaker-us-west-2-111122223333/debugger-aws-cli-test/output"
    },
    "DebugHookConfig": {
        "S3OutputPath": "s3://sagemaker-us-west-2-111122223333/debugger-aws-cli-test/debug-output",
        "CollectionConfigurations": [
            {
                "CollectionName": "losses",
                "CollectionParameters": {
                    "train.save_interval": "50"
                }
            }
        ],
        "DebugRuleConfigurations": [
            {
                "RuleConfigurationName": "LossNotDecreasing",
                "RuleParameters": {"rule_to_invoke": "LossNotDecreasing"}
            }
        ],
        "ProfilerConfig": {
            "S3OutputPath": "s3://sagemaker-us-west-2-111122223333/debugger-aws-cli-test/profiler-output",
            "ProfilingIntervalInMilliseconds": 500,
            "ProfilingParameters": {
                "DataloaderProfilingConfig": "{"StartStep": 5, "NumSteps": 3, "MetricsRegex": "\.*\"},",
                "DetailedProfilingConfig": "{"StartStep": 5, "NumSteps": 3 },",
                "PythonProfilingConfig": "{"StartStep": 5, "NumSteps": 3, "ProfilerName ": "cprofile", "cProfileTimer": "total_time" }",
                "LocalPath": "/opt/ml/output/profiler/"}
After saving the JSON file, run the following command in your terminal. (Use ! at the beginning of the line if you use a Jupyter notebook.)

```bash
aws sagemaker create-training-job --cli-input-json file://debugger-training-job-cli.json
```

**To configure a Debugger rule for debugging model parameters**

The following code samples show how to configure a built-in VanishingGradient rule using this SageMaker API.

**To enable Debugger to collect output tensors**

Specify the Debugger hook configuration as follows:

```json
"DebugHookConfig": {
  "S3OutputPath": "s3://<default-bucket>/<training-job-name>/debug-output",
  "CollectionConfigurations": [
    {
      "CollectionName": "gradients",
      "CollectionParameters": {
        "save_interval": "500"
      }
    }
  ]
}
```
This will make the training job save the tensor collection, gradients, every save_interval of 500 steps. To find available CollectionName values, see Debugger Built-in Collections in the SMDebug client library documentation. To find available CollectionParameters parameter keys and values, see the sagemaker.debugger.CollectionConfig class in the SageMaker Python SDK documentation.

**To enable Debugger rules for debugging the output tensors**

The following DebugRuleConfigurations API example shows how to run the built-in VanishingGradient rule on the saved gradients collection.

```
"DebugRuleConfigurations": [
  {
    "RuleConfigurationName": "VanishingGradient",
    "RuleEvaluatorImage": "503895931360.dkr.ecr.us-east-1.amazonaws.com/sagemaker-debugger-rules:latest",
    "RuleParameters": {
      "rule_to_invoke": "VanishingGradient",
      "threshold": "20.0"
    }
  }
]
```

With a configuration like the one in this sample, Debugger starts a rule evaluation job for your training job using the VanishingGradient rule on the collection of gradients tensor. To find a complete list of available Docker images for using the Debugger rules, see Use Debugger Docker Images for Built-in or Custom Rules. To find the key-value pairs for RuleParameters, see List of Debugger Built-in Rules.

**To configure a Debugger built-in rule for profiling system and framework metrics**

The following example code shows how to specify the ProfilerConfig API operation to enable collecting system and framework metrics.

**To enable Debugger profiling to collect system and framework metrics**
Target Step

```
"ProfilerConfig": {
   // Optional. Path to an S3 bucket to save profiling outputs
   "S3OutputPath": "s3://<default-bucket>/<training-job-name>/profiler-output",
   // Available values for ProfilingIntervalInMilliseconds: 100, 200, 500, 1000 (1 second), 5000 (5 seconds), and 60000 (1 minute) milliseconds.
   "ProfilingIntervalInMilliseconds": 500,
   "ProfilingParameters": {
      "DataloaderProfilingConfig": "{ "StartStep": 5, "NumSteps": 3, "MetricsRegex": ".*" }
   , "DetailedProfilingConfig": "{ "StartStep": 5, "NumSteps": 3 }",
   // For PythonProfilingConfig,
   // available ProfilerName options: cProfile, Pyinstrument
   // available cProfileTimer options only when using cProfile: cpu, off_cpu, total_time
   "PythonProfilingConfig": "{ "StartTimeInSecSinceEpoch": 12345567789, "DurationInSeconds": 10, "MetricsRegex": ".*" }
   }
}
```

Target Time Duration

```
"ProfilerConfig": {
   // Optional. Path to an S3 bucket to save profiling outputs
   "S3OutputPath": "s3://<default-bucket>/<training-job-name>/profiler-output",
   // Available values for ProfilingIntervalInMilliseconds: 100, 200, 500, 1000 (1 second), 5000 (5 seconds), and 60000 (1 minute) milliseconds.
   "ProfilingIntervalInMilliseconds": 500,
   "ProfilingParameters": {
      "DataloaderProfilingConfig": "{ "StartTimeInSecSinceEpoch": 12345567789, "DurationInSeconds": 10, "MetricsRegex": ".*" }
   , "DetailedProfilingConfig": "{ "StartTimeInSecSinceEpoch": 12345567789, "DurationInSeconds": 10 }",
      // For PythonProfilingConfig,
      // available ProfilerName options: cProfile, Pyinstrument
      // available cProfileTimer options only when using cProfile: cpu, off_cpu, total_time
      "PythonProfilingConfig": "{ "StartTimeInSecSinceEpoch": 12345567789, "DurationInSeconds": 10, "ProfilerName": "cProfile", "cProfileTimer": "total_time" }
   }
```

Use SageMaker Debugger
To enable Debugger rules for profiling the metrics

The following example code shows how to configure the ProfilerReport rule.

```json
"ProfilerRuleConfigurations": [  {
      "RuleConfigurationName": "ProfilerReport",
      "RuleEvaluatorImage": "89574180848.dkr.ecr.us-west-2.amazonaws.com/sagemaker-
       debugger-rules:latest",
      "RuleParameters": {
        "rule_to_invoke": "ProfilerReport",
        "CPUBottleneck_cpu_threshold": "90",
        "IOBottleneck_threshold": "90"
      }
    }
]
```

To find a complete list of available Docker images for using the Debugger rules, see Use Debugger Docker Images for Built-in or Custom Rules. To find the key-value pairs for RuleParameters, see List of Debugger Built-in Rules.

**Update Debugger Profiling Configuration Using the UpdateTrainingJob API Operation**

Debugger profiling configuration can be updated while your training job is running by using the UpdateTrainingJob API operation. Configure new ProfilerConfig and ProfilerRuleConfiguration objects, and specify the training job name to the TrainingJobName parameter.
"RuleConfigurationName": "string",
"RuleEvaluatorImage": "string",
"RuleParameters": {
    "string": "string"
}
],
"TrainingJobName": "your-training-job-name-YYYY-MM-DD-HH-MM-SS-SSS"
}

Add Debugger Custom Rule Configuration to the CreateTrainingJob API Operation

A custom rule can be configured for a training job using the DebugHookConfig and DebugRuleConfiguration objects in the CreateTrainingJob API operation. The following code sample shows how to configure a custom ImproperActivation rule written with the smdebug library using this SageMaker API operation. This example assumes that you've written the custom rule in custom_rules.py file and uploaded it to an Amazon S3 bucket. The example provides pre-built Docker images that you can use to run your custom rules. These are listed at Amazon SageMaker Debugger Registry URLs for Custom Rule Evaluators. You specify the URL registry address for the pre-built Docker image in the RuleEvaluatorImage parameter.

"DebugHookConfig": {
    "S3OutputPath": "s3://<default-bucket>/<training-job-name>/debug-output",
    "CollectionConfigurations": [
        {
            "CollectionName": "relu_activations",
            "CollectionParameters": {
                "include_regex": "relu",
                "save_interval": "500",
                "end_step": "5000"
            }
        }
    ],
    "DebugRulesConfigurations": [
        {
            "RuleConfigurationName": "improper_activation_job",
            "InstanceType": "ml.c4.xlarge",
            "VolumeSizeInGB": 400,
            "RuleParameters": {
                "string": "string"
            }
        }
    ]
}
To find a complete list of available Docker images for using the Debugger rules, see Use Debugger Docker Images for Built-in or Custom Rules. To find the key-value pairs for RuleParameters, see List of Debugger Built-in Rules.

AWS Boto3

Amazon SageMaker Debugger built-in rules can be configured for a training job using the create_training_job() function of the AWS Boto3 SageMaker client. You need to specify the right image URI in the RuleEvaluatorImage parameter, and the following examples walk you through how to set up the request body for the create_training_job() function.

The following code shows a complete example of how to configure Debugger for the create_training_job() request body and start a training job in us-west-2, assuming that a training script entry_point/train.py is prepared using TensorFlow. To find an end-to-end example notebook, see Profiling TensorFlow Multi GPU Multi Node Training Job with Amazon SageMaker Debugger (Boto3).

```python
import sagemaker, boto3
import datetime, tarfile

# Start setting up a SageMaker session and a Boto3 SageMaker client
session = sagemaker.Session()
region = session.boto_region_name
bucket = session.default_bucket()
```

Note

Ensure that you use the correct Docker container images. To find available AWS Deep Learning Container images, see Available Deep Learning Containers Images. To find a complete list of available Docker images for using the Debugger rules, see Use Debugger Docker Images for Built-in or Custom Rules.
# Upload a training script to a default Amazon S3 bucket of the current SageMaker session

```python
source = 'source.tar.gz'
project = 'debugger-boto3-test'

tar = tarfile.open(source, 'w:gz')
tar.add('entry_point/train.py') # Specify the directory and name of your training script
tar.close()

s3 = boto3.client('s3')
s3.upload_file(source, bucket, project+'/'+source)

# Set up a Boto3 session client for SageMaker
sm = boto3.Session(region_name=region).client("sagemaker")

# Start a training job
sm.create_training_job(
    TrainingJobName='debugger-boto3-')+datetime.datetime.now().strftime('%Y-%m-%d-%H-%M-%S'),
    HyperParameters={
        'sagemaker_submit_directory': 's3://'+bucket+'/'+project+'/'+source,
        'sagemaker_program': '/entry_point/train.py' # training script file location and name under the sagemaker_submit_directory
    },
    AlgorithmSpecification={
        # Specify a training Docker container image URI (Deep Learning Container or your own training container) to TrainingImage.
        'TrainingInputMode': 'File',
        'EnableSageMakerMetricsTimeSeries': False
    },
    RoleArn='arn:aws:iam::111122223333:role/service-role/AmazonSageMaker-ExecutionRole-20201014T161125',
    OutputDataConfig={'S3OutputPath': 's3://'+bucket+'/'+project+'/output'},
    ResourceConfig={
        'InstanceType': 'ml.p3.8xlarge',
        'InstanceCount': 1,
        'VolumeSizeInGB': 30
    },
    StoppingCondition={
        'MaxRuntimeInSeconds': 86400
    },
)
DebugHookConfig={
    'S3OutputPath': 's3://'+bucket+'/'+project+'/debug-output',
    'CollectionConfigurations': [
        {
            'CollectionName': 'losses',
            'CollectionParameters': {
                'train.save_interval': '500',
                'eval.save_interval': '50'
            }
        }
    ],
    'DebugRuleConfigurations':[{
        'RuleConfigurationName': 'LossNotDecreasing',
        'RuleParameters': {'rule_to_invoke': 'LossNotDecreasing'}
    }],
    'ProfilerConfig={
        'S3OutputPath': 's3://'+bucket+'/'+project+'/profiler-output',
        'ProfilingIntervalInMilliseconds': 500,
        'ProfilingParameters': {
            'DataloaderProfilingConfig': '{"StartStep": 5, "NumSteps": 3, "MetricsRegex": ".*"}',
            'DetailedProfilingConfig': '{"StartStep": 5, "NumSteps": 3, }',
            'PythonProfilingConfig': '{"StartStep": 5, "NumSteps": 3, "ProfilerName": "cprofile", "cProfileTimer": "total_time"}',
            'LocalPath': '/opt/ml/output/profiler/' # Optional. Local path for profiling outputs
        },
        'ProfilerRuleConfigurations':[
            {
                'RuleConfigurationName': 'ProfilerReport',
                'RuleParameters': {'rule_to_invoke': 'ProfilerReport'}
            }
        ]
    }
}
To configure a Debugger rule for debugging model parameters

The following code samples show how to configure a built-in VanishingGradient rule using this SageMaker API.

To enable Debugger to collect output tensors

Specify the Debugger hook configuration as follows:

```
DebugHookConfig={
    'S3OutputPath': 's3://<default-bucket>/<training-job-name>/debug-output',
    'CollectionConfigurations': [
        {
            'CollectionName': 'gradients',
            'CollectionParameters': {
                'train.save_interval': '500',
                'eval.save_interval': '50'
            }
        }
    ]
}
```

This will make the training job save a tensor collection, gradients, every save_interval of 500 steps. To find available CollectionName values, see Debugger Built-in Collections in the SMDebug client library documentation. To find available CollectionParameters parameter keys and values, see the sagemaker.debugger.CollectionConfig class in the SageMaker Python SDK documentation.

To enable Debugger rules for debugging the output tensors

The following DebugRuleConfigurations API example shows how to run the built-in VanishingGradient rule on the saved gradients collection.

```
DebugRuleConfigurations=[
    {
        'RuleConfigurationName': 'VanishingGradient',
        'RuleParameters': {
            'rule_to_invoke': 'VanishingGradient',
            'threshold': '20.0'
        }
    }
]
```
With a configuration like the one in this sample, Debugger starts a rule evaluation job for your training job using the VanishingGradient rule on the collection of gradients tensor. To find a complete list of available Docker images for using the Debugger rules, see Use Debugger Docker Images for Built-in or Custom Rules. To find the key-value pairs for RuleParameters, see List of Debugger Built-in Rules.

To configure a Debugger built-in rule for profiling system and framework metrics

The following example code shows how to specify the ProfilerConfig API operation to enable collecting system and framework metrics.

To enable Debugger profiling to collect system and framework metrics

Target Step

```json
ProfilerConfig={
    'S3OutputPath': 's3://<default-bucket>/<training-job-name>/profiler-output', # Optional. Path to an S3 bucket to save profiling outputs
    'ProfilingIntervalInMilliseconds': 500,
    'ProfilingParameters': {
        'DataloaderProfilingConfig': '{
            "StartStep": 5,
            "NumSteps": 3,
            "MetricsRegex": ".*"
        }
    },
    'DetailedProfilingConfig': '{
        "StartStep": 5,
        "NumSteps": 3
    }
},
'PythonProfilingConfig': '{
        "StartStep": 5,
        "NumSteps": 3,
        "ProfilerName": "cprofile", # Available options: cprofile, pyinstrument
        "cProfileTimer": "total_time" # Include only when using cprofile.
    }',
    'LocalPath': '/opt/ml/output/profiler/' # Optional. Local path for profiling outputs
}
```

Use SageMaker Debugger
Target Time Duration

ProfilerConfig=
   
   'S3OutputPath': 's3://<default-bucket>/<training-job-name>/profiler-output', #
   Optional. Path to an S3 bucket to save profiling outputs
   # Available values for ProfilingIntervalInMilliseconds: 100, 200, 500, 1000 (1
   second), 5000 (5 seconds), and 60000 (1 minute) milliseconds.
   'ProfilingIntervalInMilliseconds': 500,
   'ProfilingParameters': {
       'DataloaderProfilingConfig': '{
           "StartTimeInSecSinceEpoch": 12345567789,
           "DurationInSeconds": 10,
           "MetricsRegex": ".*"
       }',
       'DetailedProfilingConfig': '{
           "StartTimeInSecSinceEpoch": 12345567789,
           "DurationInSeconds": 10
       }',
       'PythonProfilingConfig': '{
           "StartTimeInSecSinceEpoch": 12345567789,
           "DurationInSeconds": 10,
           "ProfilerName": "cprofile", # Available options: cprofile, pynstrument
           "cProfileTimer": "total_time" # Include only when using cprofile.
           Available options: cpu, off_cpu, total_time
       }',
       'LocalPath': '/opt/ml/output/profiler/' # Optional. Local path for profiling
   }
}

To enable Debugger rules for profiling the metrics

The following example code shows how to configure the ProfilerReport rule.

ProfilerRuleConfigurations=[
   {
       'RuleConfigurationName': 'ProfilerReport',
       'RuleEvaluatorImage': '895741380848.dkr.ecr.us-west-2.amazonaws.com/sagemaker-
                             debugger-rules:latest',
   }]

'RuleParameters': {
    'rule_to_invoke': 'ProfilerReport',
    'CPUBottleneck_cpu_threshold': '90',
    'IOBottleneck_threshold': '90'
}

To find a complete list of available Docker images for using the Debugger rules, see [Use Debugger Docker Images for Built-in or Custom Rules](#). To find the key-value pairs for RuleParameters, see [List of Debugger Built-in Rules](#).

### Update Debugger Profiling Configuration Using the UpdateTrainingJob API Operation

Debugger profiling configuration can be updated while your training job is running by using the `update_training_job()` function of the AWS Boto3 SageMaker client. Configure new `ProfilerConfig` and `ProfilerRuleConfiguration` objects, and specify the training job name to the `TrainingJobName` parameter.

```python
ProfilerConfig={
    'DisableProfiler': boolean,
    'ProfilingIntervalInMilliseconds': number,
    'ProfilingParameters': {
        'string': 'string'
    }
},
ProfilerRuleConfigurations=[
    {
        'RuleConfigurationName': 'string',
        'RuleEvaluatorImage': 'string',
        'RuleParameters': {
            'string': 'string'
        }
    }
],
TrainingJobName='your-training-job-name-YYYY-MM-DD-HH-MM-SS-SSS'
```

### Add Debugger Custom Rule Configuration to the CreateTrainingJob API Operation

A custom rule can be configured for a training job using the `DebugHookConfig` and `DebugRuleConfiguration` objects using the AWS Boto3 SageMaker client's `create_training_job()` function. The following code sample shows how to configure a
custom ImproperActivation rule written with the smdebug library using this SageMaker API operation. This example assumes that you've written the custom rule in custom_rules.py file and uploaded it to an Amazon S3 bucket. The example provides pre-built Docker images that you can use to run your custom rules. These are listed at Amazon SageMaker Debugger Registry URLs for Custom Rule Evaluators. You specify the URL registry address for the pre-built Docker image in the RuleEvaluatorImage parameter.

```json
DebugHookConfig={
    'S3OutputPath': 's3://<default-bucket>/<training-job-name>/debug-output',
    'CollectionConfigurations': [
        {
            'CollectionName': 'relu_activations',
            'CollectionParameters': {
                'include_regex': 'relu',
                'save_interval': '500',
                'end_step': '5000'
            }
        }
    ],
    DebugRulesConfigurations=[
        {
            'RuleConfigurationName': 'improper_activation_job',
            'InstanceType': 'ml.c4.xlarge',
            'VolumeSizeInGB': 400,
            'RuleParameters': {
                'source_s3_uri': 's3://bucket/custom_rules.py',
                'rule_to_invoke': 'ImproperActivation',
                'collection_names': 'relu_activations'
            }
        }
    ]
}
```

To find a complete list of available Docker images for using the Debugger rules, see Use Debugger Docker Images for Built-in or Custom Rules. To find the key-value pairs for RuleParameters, see List of Debugger Built-in Rules.

**Best Practices for Amazon SageMaker Debugger**

Use the following guidelines when you run training jobs with Debugger.
Choose a Machine Learning Framework

You can choose a machine learning framework and use SageMaker pre-built training containers or your own containers. Use Debugger to detect training and performance issues, and analyze training progress of your training job in SageMaker. SageMaker provides you options to use pre-built containers that are prepared for a number of machine learning framework environments to train your model on Amazon EC2. Any training job can be adapted to run in AWS Deep Learning Containers, SageMaker training containers, and custom containers.

Use Studio Debugger Insights Dashboard

With Studio Debugger insights dashboard, you are in control of your training jobs. Use the Studio Debugger dashboards to keep your model performance on Amazon EC2 instances in control and optimized. For any SageMaker training jobs running on Amazon EC2 instance, Debugger monitors resource utilization and basic model output data (loss and accuracy values). Through the Studio Debugger dashboards, gain insights into your training jobs and improve your model training performance. To learn more, see Amazon SageMaker Debugger UI in Amazon SageMaker Studio Classic Experiments.

Download Debugger Reports and Gain More Insights

You can view aggregated results and gain insights in Debugger reports. Debugger aggregates training and profiling results collected from the built-in rule analysis into a report per training job.
You can find more detailed information about your training results through the Debugger reports. To learn more, see SageMaker Debugger interactive report.

**Capture Data from Your Training Job and Save Data to Amazon S3**

You can use a Debugger hook to save output tensors. After you choose a container and a framework that fit your training script, use a Debugger hook to configure which tensors to save and to which directory to save them, such as an Amazon S3 bucket. A Debugger hook helps you to build the configuration and to keep it in your account to use in subsequent analyses, where it is secured for use with the most privacy-sensitive applications. To learn more, see Configure SageMaker Debugger to Save Tensors.

**Analyze the Data with a Fleet of Debugger Built-in Rules**

You can use Debugger built-in rules to inspect tensors in parallel with a training job. To analyze the training performance data, Debugger provides built-in rules that watch for abnormal training process behaviors. For example, a Debugger rule detects issues when the training process suffers from system bottleneck issues or training issues, such as vanishing gradients, exploding tensors, overfitting, or overtraining. If necessary, you can also build customized rules by creating a rule definition with your own criteria to define a training issue. To learn more about the Debugger rules, see Configure Debugger Built-in Rules for detailed instructions of using the Amazon SageMaker Python SDK. For a full list of the Debugger built-in rules, see List of Debugger Built-in Rules. If you want to create a custom rule, see Create Debugger Custom Rules for Training Job Analysis.

**Take Actions Based on the Built-in Rule Status**

You can use Debugger with Amazon CloudWatch Events and AWS Lambda. You can automate actions based on the rule status, such as stopping training jobs early and setting up notifications through email or text. When the Debugger rules detect problems and triggers an "IssuesFound" evaluation status, CloudWatch Events detects the rule status changes and invokes the Lambda function to take actions. To configure automated actions to your training issues, see Create Actions on Rules Using Amazon CloudWatch and AWS Lambda.

**Dive Deep into the Data Using the SMDebug Client Library**

You can use the SMDebug tools to access and analyze training data collected by Debugger. The TrainingJob and create_trial classes load the metrics and tensors saved by Debugger. These classes provide extended class methods to analyze the data in real time or after the training has finished. The SMDebug library also provides visualization tools: merge timelines of framework
metrics to aggregate different profiling, line charts and heatmap to track the system utilization, and histograms to find step duration outliers. To learn more about the SMDebug library tools, see Analyze data using the Debugger Python client library.

Monitor and Analyze Training Job Metrics

Amazon CloudWatch supports high-resolution custom metrics, and its finest resolution is 1 second. However, the finer the resolution, the shorter the lifespan of the CloudWatch metrics. For the 1-second frequency resolution, the CloudWatch metrics are available for 3 hours. For more information about the resolution and the lifespan of the CloudWatch metrics, see GetMetricStatistics in the Amazon CloudWatch API Reference.

If you want to profile your training job with a finer resolution down to 100-millisecond (0.1 second) granularity and store the training metrics indefinitely in Amazon S3 for custom analysis at any time, consider using Amazon SageMaker Debugger. SageMaker Debugger provides built-in rules to automatically detect common training issues; it detects hardware resource utilization issues (such as CPU, GPU, and I/O bottlenecks) and non-converging model issues (such as overfit, vanishing gradients, and exploding tensors).

SageMaker Debugger also provides visualizations through Studio Classic and its profiling report. Unlike CloudWatch metrics, which accumulates resource utilization rates of CPU and GPU cores and averages those out across multiple instances, Debugger tracks the utilization rate of each core. This enables you to identify unbalanced usage of hardware resources as you scale up to larger compute clusters. To explore the Debugger visualizations, see SageMaker Debugger Insights Dashboard Walkthrough, Debugger Profiling Report Walkthrough, and Analyze Data Using the SMDebug Client Library.

Monitoring System Utilization and Detect Bottlenecks

With Amazon SageMaker Debugger monitoring, you can measure hardware system resource utilization of Amazon EC2 instances. Monitoring is available for any SageMaker training job constructed with the SageMaker framework estimators (TensorFlow, PyTorch, and MXNet) and the generic SageMaker estimator (SageMaker built-in algorithms and your own custom containers). Debugger built-in rules for monitoring detect system bottleneck issues and notify you when they detect the bottleneck issues.

To learn how to enable Debugger system monitoring, see Configure an estimator with parameters for basic profiling using the Amazon SageMaker Debugger Python modules and then Configure settings for basic profiling of system resource utilization.
For a full list of available built-in rules for monitoring, see [Debugger built-in rules for profiling](#) hardware system resource utilization (system metrics).

### Profiling Framework Operations

With Amazon SageMaker Debugger profiling you can profile deep learning frameworks operations. You can profile your model training with the SageMaker TensorFlow training containers, the SageMaker PyTorch framework containers, and your own training containers. Using the profiling feature of Debugger, you can drill down into the Python operators and functions that are executed to perform the training job. Debugger supports detailed profiling, Python profiling, data loader profiling, and Horovod distributed training profiling. You can merge the profiled timelines to correlate with the system bottlenecks. Debugger built-in rules for profiling watch framework operation related issues, including excessive training initialization time due to data downloading before training starts and step duration outliers in training loops.

To learn how to configure Debugger for framework profiling, see [Configure an estimator with parameters for basic profiling using the Amazon SageMaker Debugger Python modules](#) and then [Configure for framework profiling](#).

For a complete list of available built-in rules for profiling, see [Debugger built-in rules for profiling framework metrics](#).

### Debugging Model Output Tensors

Debugging is available for deep learning frameworks using AWS Deep Learning Containers and the SageMaker training containers. For fully supported framework versions (see the versions at [Supported Frameworks and Algorithms](#)), Debugger automatically registers hooks to collect output tensors, and you can directly run your training script. For the versions with one asterisk sign, you need to manually register the hooks to collect tensors. Debugger provides preconfigured tensor collections with generalized names that you can utilize across the different frameworks.

If you want to customize output tensor configuration, you can also use the CollectionConfig and DebuggerHookConfig API operations and the [Amazon SageMaker Python SDK](#) to configure your own tensor collections. Debugger built-in rules for debugging analyze the output tensors and identifies model optimization problems that blocks your model from minimizing the loss function. For example, the rules identify overfitting, overtraining, loss not decreasing, exploding tensors, and vanishing gradients.

To learn how to configure Debugger for debugging output tensors, see [Step 2: Launch and Debug Training Jobs Using SageMaker Python SDK](#) and then [Configure SageMaker Debugger to Save Tensors](#).
For a full list of available built-in rules for debugging, see Debugger built-in rules for debugging model training data (output tensors).

Amazon SageMaker Debugger Advanced Topics and Reference Documentation

The following sections contain advanced topics, reference documentation for the API operations, exceptions, and known limitations for Debugger.

Topics

- Amazon SageMaker Debugger API Operations
- Use Debugger Docker Images for Built-in or Custom Rules
- Amazon SageMaker Debugger Exceptions
- Considerations for Amazon SageMaker Debugger
- Amazon SageMaker Debugger Usage Statistics

Amazon SageMaker Debugger API Operations

Amazon SageMaker Debugger has API operations in several locations that are used to implement its monitoring and analysis of model training.

Amazon SageMaker Debugger also provides the open source sagemaker-debugger Python SDK that is used to configure built-in rules, define custom rules, and register hooks to collect output tensor data from training jobs.

The Amazon SageMaker Python SDK is a high-level SDK focused on machine learning experimentation. The SDK can be used to deploy built-in or custom rules defined with the SMDebug Python library to monitor and analyze these tensors using SageMaker estimators.

Debugger has added operations and types to the Amazon SageMaker API that enable the platform to use Debugger when training a model and to manage the configuration of inputs and outputs.

- CreateTrainingJob and UpdateTrainingJob use the following Debugger APIs to configure tensor collections, rules, rule images, and profiling options:
  - CollectionConfiguration
  - DebugHookConfig
  - DebugRuleConfiguration
The rule configuration API operations use the SageMaker Processing functionality when analyzing a model training. For more information about SageMaker Processing, see [Process data](#).

**Use Debugger Docker Images for Built-in or Custom Rules**

Amazon SageMaker provides two sets of Docker images for rules: one set for evaluating rules provided by SageMaker (built-in rules) and one set for evaluating custom rules provided in Python source files.

If you use the [Amazon SageMaker Python SDK](#), you can simply use SageMaker high-level Debugger API operations with SageMaker Estimator API operations, without having to manually retrieve the Debugger Docker images and configure the `ConfigureTrainingJob` API.

If you are not using the SageMaker Python SDK, you have to retrieve a relevant pre-built container base image for the Debugger rules. Amazon SageMaker Debugger provides pre-built Docker images for built-in and custom rules, and the images are stored in Amazon Elastic Container Registry (Amazon ECR). To pull an image from an Amazon ECR repository (or to push an image to one), use the full name registry URL of the image using the `CreateTrainingJob` API. SageMaker uses the following URL patterns for the Debugger rule container image registry address.

```bash
<account_id>.dkr.ecr.<Region>.amazonaws.com/<ECR repository name>:<tag>
```

For the account ID in each AWS Region, Amazon ECR repository name, and tag value, see the following topics.

**Amazon SageMaker Python SDK**

[Amazon SageMaker Python SDK](#)
Topics

- Amazon SageMaker Debugger Registry URLs for Built-in Rule Evaluators
- Amazon SageMaker Debugger Registry URLs for Custom Rule Evaluators

Amazon SageMaker Debugger Registry URLs for Built-in Rule Evaluators

Use the following values for the components of the registry URLs for the images that provide built-in rules for Amazon SageMaker Debugger. For account IDs, see the following table.

ECR Repository Name: sagemaker-debugger-rules

Tag: latest

Example of a full registry URL:

904829902805.dkr.ecr.ap-south-1.amazonaws.com/sagemaker-debugger-rules:latest

Account IDs for Built-in Rules Container Images by AWS Region

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<td>us-gov-west-1</td>
<td>515509971035</td>
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</table>

**Amazon SageMaker Debugger Registry URLs for Custom Rule Evaluators**

Use the following values for the components of the registry URL for the images that provide custom rule evaluators for Amazon SageMaker Debugger. For account IDs, see the following table.

**ECR Repository Name:** sagemaker-debugger-rule-evaluator

**Tag:** latest

**Example of a full registry URL:**

552407032007.dkr.ecr.us-east-1.amazonaws.com/sagemaker-debugger-rule-evaluator:latest
# Account IDs for Custom Rules Container Images by AWS Region

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</tr>
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<td>759209512951</td>
</tr>
<tr>
<td>us-gov-west-1</td>
<td>515361955729</td>
</tr>
</tbody>
</table>

**Amazon SageMaker Debugger Exceptions**

Amazon SageMaker Debugger is designed to be aware that tensors required to execute a rule might not be available at every step. As a result, it raises a few exceptions, which enable you to control what happens when a tensor is missing. These exceptions are available in the `smdebug.exceptions` module. You can import them as follows:

```python
from smdebug.exceptions import *
```

The following exceptions are available:

- **TensorUnavailableForStep** – The tensor requested is not available for the step. This might mean that this step might not be saved at all by the hook, or that this step might have saved some tensors but the requested tensor is not part of them. Note that when you see this exception, it means that this tensor can never become available for this step in the future. If the tensor has reductions saved for the step, it notifies you they can be queried.

- **TensorUnavailable** – This tensor is not being saved or has not been saved by the `smdebug` API. This means that this tensor is never seen for any step in `smdebug`.

- **StepUnavailable** – The step was not saved and Debugger has no data from the step.

- **StepNotYetAvailable** – The step has not yet been seen by `smdebug`. It might be available in the future if the training is still going on. Debugger automatically loads new data as it becomes available.

- **NoMoreData** – Raised when the training ends. Once you see this, you know that there are no more steps and no more tensors to be saved.

- **IndexReaderException** – The index reader is not valid.

- **InvalidWorker** – A worker was invoked that was not valid.
• RuleEvaluationConditionMet – Evaluation of the rule at the step resulted in the condition being met.

• InsufficientInformationForRuleInvocation – Insufficient information was provided to invoke the rule.

Considerations for Amazon SageMaker Debugger

Consider the following when using Amazon SageMaker Debugger.

Considerations for Distributed Training

The following list shows the scope of validity and considerations for using Debugger on training jobs with deep learning frameworks and various distributed training options.

• Horovod

Scope of validity of using Debugger for training jobs with Horovod

<table>
<thead>
<tr>
<th>Deep Learning Framework</th>
<th>Apache MXNet</th>
<th>TensorFlow 1.x</th>
<th>TensorFlow 2.x</th>
<th>TensorFlow 2.x with Keras</th>
<th>PyTorch</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monitorin system bottlenecks</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Profiling framework operations</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Debugging model output tensors</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

• SageMaker distributed data parallel
Scope of validity of using Debugger for training jobs with SageMaker distributed data parallel

<table>
<thead>
<tr>
<th>Deep Learning Framework</th>
<th>TensorFlow 2.x</th>
<th>TensorFlow 2.x with Keras</th>
<th>PyTorch</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monitoring system bottlenecks</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Profiling framework operations</td>
<td>No*</td>
<td>No**</td>
<td>Yes</td>
</tr>
<tr>
<td>Debugging model output tensors</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

* Debugger does not support framework profiling for TensorFlow 2.x.

** SageMaker distributed data parallel does not support TensorFlow 2.x with Keras implementation.

- **SageMaker distributed model parallel** – Debugger does not support SageMaker distributed model parallel training.

- **Distributed training with SageMaker checkpoints** – Debugger is not available for training jobs when both the distributed training option and SageMaker checkpoints are enabled. You might see an error that looks like the following:

```
SMDebug Does Not Currently Support Distributed Training Jobs With Checkpointing Enabled
```

To use Debugger for training jobs with distributed training options, you need to disable SageMaker checkpointing and add manual checkpointing functions to your training script. For more information about using Debugger with distributed training options and checkpoints, see [Using SageMaker distributed data parallel with Amazon SageMaker Debugger and checkpoints](#) and [Saving Checkpoints](#).

- **Parameter Server** – Debugger does not support parameter server-based distributed training.

- Profiling distributed training framework operations, such as the `AllReduced` operation of SageMaker distributed data parallel and [Horovod operations](#), is not available.
Considerations for Monitoring System Bottlenecks and Profiling Framework Operations

- For AWS TensorFlow, data loader metrics cannot be collected using the default `local_path` setting of the `FrameworkProfile` class. The path has to be manually configured and end in "/". For example:

  ```python
  FrameworkProfile(local_path="/opt/ml/output/profiler/"
  ```

- For AWS TensorFlow, the data loader profiling configuration cannot be updated while a training job is running.

- For AWS TensorFlow, a `NoneType` error might occur when you use analysis tools and notebook examples with TensorFlow 2.3 training jobs and the detailed profiling option.

- Python profiling and detailed profiling are only supported for Keras API.

- To access the deep profiling feature for TensorFlow and PyTorch, currently you must specify the latest AWS deep learning container images with CUDA 11. For example, you must specify the specific image URI in the TensorFlow and PyTorch estimator as follows:

  - For TensorFlow

    ```python
    image_uri = f"763104351884.dkr.ecr.{region}.amazonaws.com/tensorflow-training:2.3.1-gpu-py37-cu110-ubuntu18.04"
    ```

  - For PyTorch

    ```python
    image_uri = f"763104351884.dkr.ecr.{region}.amazonaws.com/pytorch-training:1.6.0-gpu-py36-cu110-ubuntu18.04"
    ```

Considerations for Debugging Model Output Tensors

- Avoid using functional API operations. Debugger cannot collect model output tensors from PyTorch and MXNet training scripts composed of functional API operations.

- Debugger cannot collect model output tensors from the `torch.nn.functional` API operations. When you write a PyTorch training script, it is recommended to use the `torch.nn` modules instead.

- Debugger cannot collect model output tensors from MXNet functional objects in hybrid blocks. For example, the ReLu activation (`F.relu`) outputs cannot be collected from the following example of `mxnet.gluon.HybridBlock` with `F` in the `hybrid_forward` function.
import mxnet as mx
from mxnet.gluon import HybridBlock, nn

class Model(HybridBlock):
    def __init__(self, **kwargs):
        super(Model, self).__init__(**kwargs)
        # use name_scope to give child Blocks appropriate names.
        with self.name_scope():
            self.dense0 = nn.Dense(20)
            self.dense1 = nn.Dense(20)

    def hybrid_forward(self, F, x):
        x = F.relu(self.dense0(x))
        return F.relu(self.dense1(x))

model = Model()
model.initialize(ctx=mx.cpu(0))
model.hybridize()
model(mx.nd.zeros((10, 10), ctx=mx.cpu(0)))

Amazon SageMaker Debugger Usage Statistics

Consider the following when using autogenerated reports by Amazon SageMaker Debugger.

Debugger Profiling Report Usage

For all SageMaker training jobs, Amazon SageMaker Debugger runs the ProfilerReport rule and autogenerates a SageMaker Debugger profiling report. The ProfilerReport rule provides a Jupyter notebook file (profiler-report.ipynb) that generates a corresponding HTML file (profiler-report.html).

Debugger collects profiling report usage statistics by including code in the Jupyter notebook that collects the unique ProfilerReport rule's processing job ARN if the user opens the final profiler-report.html file.

Debugger only collects information about whether a user opens the final HTML report. It DOES NOT collect any information from training jobs, training data, training scripts, processing jobs, logs, or the content of the profiling report itself.

You can opt out of the collection of usage statistics using either of the following options.
(Recommended) Option 1: Opt Out before Running a Training Job

To opt out, you need to add the following Debugger ProfilerReport rule configuration to your training job request.

SageMaker Python SDK

```python
estimator=sagemaker.estimator.Estimator(
    ...
    rules=ProfilerRule.sagemaker(
        base_config=rule_configs.ProfilerReport()
        rule_parameters={"opt_out_telemetry": "True"}
    )
)
```

AWS CLI

```
"ProfilerRuleConfigurations": [
    {
        "RuleConfigurationName": "ProfilerReport-1234567890",
        "RuleParameters": {
            "rule_to_invoke": "ProfilerReport",
            "opt_out_telemetry": "True"
        }
    }
]
```

AWS SDK for Python (Boto3)

```python
ProfilerRuleConfigurations=[
    {
        'RuleConfigurationName': 'ProfilerReport-1234567890',
        'RuleParameters': {
            'rule_to_invoke': 'ProfilerReport',
            'opt_out_telemetry': 'True'
        }
    }
]`
Option 2: Opt Out after a Training Job Has Completed

To opt out after training has completed, you need to modify the profiler-report.ipynb file.

**Note**

HTML reports autogenerated without Option 1 already added to your training job request still report the usage statistics even after you opt out using Option 2.

1. Follow the instructions on downloading the Debugger profiling report files in the [Download the SageMaker Debugger profiling report](#) page.
2. In the /ProfilerReport-1234567890/profiler-output directory, open profiler-report.ipynb.
3. Add `opt_out=True` to the `setup_profiler_report()` function in the fifth code cell as shown in the following example code:

   ```python
   setup_profiler_report(processing_job_arn, opt_out=True)
   ```

4. Run the code cell to finish opting out.

### Access a training container through AWS Systems Manager for remote debugging

You can securely connect to SageMaker training containers through AWS Systems Manager (SSM). This gives you a shell-level access to debug training jobs that are running within the container. You can also log commands and responses that are streamed to Amazon CloudWatch. If you use your own Amazon Virtual Private Cloud (VPC) to train a model, you can use AWS PrivateLink to set up a VPC endpoint for SSM and connect to containers privately through SSM.

You can connect to [SageMaker Framework Containers](#) or connect to your own training container set up with the SageMaker Training environment.
Set up IAM permissions

To enable SSM in your SageMaker training container, you need to set up an IAM role for the container. For you or users in your AWS account to access the training containers through SSM, you need to set up IAM users with permissions to use SSM.

IAM role

For a SageMaker training container to start with the SSM agent, provide an IAM role with SSM permissions.

To enable remote debugging for your training job, SageMaker needs to start the SSM agent in the training container when the training job starts. To allow the SSM agent to communicate with the SSM service, add the following policy to the IAM role that you use to run your training job.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "ssmmessages:CreateControlChannel",
            "ssmmessages:CreateDataChannel",
            "ssmmessages:OpenControlChannel",
            "ssmmessages:OpenDataChannel"
         ],
         "Resource": "*"
      }
   ]
}
```

IAM user

Add the following policy to provide an IAM user with SSM session permissions to connect to an SSM target. In this case, the SSM target is a SageMaker training container.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
```

Access a training container through SSM for remote debugging
You can restrict IAM users to connect only to containers for specific training jobs by adding the Condition key, as shown in the following policy sample.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "ssm:StartSession",
        "ssm:TerminateSession"
      ],
      "Resource": [
        "*
      ],
      "Condition": {
        "StringLike": {
          "ssm:resourceTag/aws:ssmmessages:target-id": [
            "sagemaker-training-job:*"
          ]
        }
      }
    }
  ]
}
```

You can also explicitly use the `sagemaker:EnableRemoteDebug` condition key to restrict remote debugging. The following is an example policy for IAM users to restrict remote debugging.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "DenyRemoteDebugInTrainingJob",
      "Effect": "Deny",
      "Action": [
        "sagemaker:EnableRemoteDebug"
      ],
      "Resource": [
        "*
      ],
      "Condition": {
        "StringLike": {
          "ssm:resourceTag/aws:ssmmessages:target-id": [
            "sagemaker-training-job:*"
          ]
        }
      }
    }
  ]
}
```
How to enable remote debugging for a SageMaker training job

In this section, learn how to enable remote debugging when starting or updating a training job in Amazon SageMaker.

SageMaker Python SDK

Using the estimator class in the SageMaker Python SDK, you can turn remote debugging on or off using the enable_remote_debug parameter or the enable_remote_debug() and disable_remote_debug() methods.

To enable remote debugging when you create a training job

To enable remote debugging when you create a new training job, set the enable_remote_debug parameter to True. The default value is False, so if you don’t set this parameter at all, or you explicitly set it to False, remote debugging functionality is disabled.

```python
import sagemaker

session = sagemaker.Session()

estimator = sagemaker.estimator.Estimator(
    ...,  
    sagemaker_session=session,

```
To enable remote debugging by updating a training job

Using the following estimator class methods, you can enable or disable remote debugging while a training job is running when the SecondaryStatus of the job is Downloading or Training.

```
# Enable RemoteDebug
estimator.enable_remote_debug()

# Disable RemoteDebug
estimator.disable_remote_debug()
```

AWS SDK for Python (Boto3)

To enable remote debugging when you create a training job

To enable remote debugging when you create a new training job, set the value for the EnableRemoteDebug key to True in the RemoteDebugConfig parameter.

```
import boto3

sm = boto3.Session(region_name=region).client("sagemaker")

# Start a training job
sm.create_training_job(
    ..., 
    TrainingJobName=job_name, 
    AlgorithmSpecification={
        // Specify a training Docker container image URI 
        // (Deep Learning Container or your own training container) to TrainingImage. 
        "TrainingImage": ":<your_image_uri>" ,
```
"TrainingInputMode": "File",
},
RoleArn=iam_role_arn,
OutputDataConfig=output_path,
ResourceConfig={
  "InstanceType": "ml.m5.xlarge",
  "InstanceCount": 1,
  "VolumeSizeInGB": 30
},
StoppingCondition={
  "MaxRuntimeInSeconds": 86400
},
RemoteDebugConfig={
  "EnableRemoteDebug": True
}
)

To enable remote debugging by updating a training job

Using the update_training_job API, you can enable or disable remote debugging while a training job is running when the SecondaryStatus of the job is Downloading or Training.

```python
# Update a training job
sm.update_training_job(
    TrainingJobName=job_name,
    RemoteDebugConfig={
        "EnableRemoteDebug": True  # True | False
    }
)
```

AWS Command Line Interface (CLI)

To enable remote debugging when you create a training job

Prepare a CreateTrainingJob request file in JSON format, as follows.

```json
// train-with-remote-debug.json
{
    "TrainingJobName": job_name,
    "RoleArn": iam_role_arn,
    "AlgorithmSpecification": {
        // Specify a training Docker container image URI (Deep Learning Container or your own training container) to TrainingImage.
```
"TrainingImage": "<your_image_uri>",
   "TrainingInputMode": "File"
 },
 "OutputDataConfig": {
   "S3OutputPath": output_path
 },
 "ResourceConfig": {
   "InstanceType": "ml.m5.xlarge",
   "InstanceCount": 1,
   "VolumeSizeInGB": 30
 },
 "StoppingCondition": {
   "MaxRuntimeInSeconds": 86400
 },
 "RemoteDebugConfig": {
   "EnableRemoteDebug": True
 }
}

After saving the JSON file, run the following command in the terminal where you submit the training job. The following example command assumes that the JSON file is named `train-with-remote-debug.json`. If you run it from a Jupyter notebook, add an exclamation point (!) to the beginning of the line.

```
aws sagemaker create-training-job \
   --cli-input-json file://train-with-remote-debug.json
```

**To enable remote debugging by updating a training job**

Prepare an UpdateTrainingJob request file in JSON format, as follows.

```
// update-training-job-with-remote-debug-config.json
{
   "TrainingJobName": job_name,
   "RemoteDebugConfig": {
      "EnableRemoteDebug": True
   }
}
```

After saving the JSON file, run the following command in the terminal where you submit the training job. The following example command assumes that the JSON file is named `train-
with-remote-debug.json. If you run it from a Jupyter notebook, add an exclamation point (!) to the beginning of the line.

```bash
aws sagemaker update-training-job \
--cli-input-json file://update-training-job-with-remote-debug-config.json
```

**Access your training container**

You can access a training container when the SecondaryStatus of the corresponding training job is Training. The following code examples demonstrate how to check the status of your training job using the DescribeTrainingJob API, how to check the training job logs in CloudWatch, and how to log in to the training container.

**To check the status of a training job**

**SageMaker Python SDK**

To check the SecondaryStatus of a training job, run the following SageMaker Python SDK code.

```python
import sagemaker

session = sagemaker.Session()

# Describe the job status
training_job_info = session.describe_training_job(job_name)
print(training_job_info)
```

**AWS SDK for Python (Boto3)**

To check the SecondaryStatus of a training job, run the following SDK for Python (Boto3) code.

```python
import boto3

session = boto3.session.Session()
region = session.region_name
sm = boto3.Session(region_name=region).client("sagemaker")

# Describe the job status
```

Access a training container through SSM for remote debugging
To check the SecondaryStatus of a training job, run the following AWS CLI command for SageMaker.

```bash
aws sagemaker describe-training-job --training-job-name job_name
```

**To find the host name of a training container**

To connect to the training container through SSM, use this format for the target ID: `sagemaker-training-job:<training-job-name>_algo-<n>`, where `algo-<n>` is the name of the container host. If your job is running on a single instance, the host is always `algo-1`. If you run a distributed training job on multiple instances, SageMaker creates an equal number of hosts and log streams. For example, if you use 4 instances, SageMaker creates `algo-1`, `algo-2`, `algo-3`, and `algo-4`. You must determine which log stream you want to debug, and its host number. To access log streams that are associated with a training job, do the following.

2. In the left navigation pane, choose **Training**, then choose **Training jobs**.
3. From the **Training jobs** list, choose the training job that you want to debug. The training job details page opens.
4. In the **Monitor** section, choose **View logs**. The related training job log stream list opens in the CloudWatch console.
5. Log stream names appear in `<training-job-name>/algo-<n>-<time-stamp>` format, with `algo-<n>` representing the host name.

To learn more about how SageMaker manages configuration information for multi-instance distributed training, see [Distributed Training Configuration](https://docs.aws.amazon.com/sagemaker/latest/dg/distributed-training-parameters.html).

**To access the training container**

Use the following command in terminal to start the SSM session (`aws ssm start-session`) and connect to the training container.
For example, if the training job name is `training-job-test-remote-debug` and the host name is `algo-1`, the target ID becomes `sagemaker-training-job:training-job-test-remote-debug_algo-1`. If the output of this command is similar to `Starting session with SessionId:xxxxx`, the connection is successful.

### SSM access with AWS PrivateLink

If your training containers run within an Amazon Virtual Private Cloud that is not connected to the public internet, you can use AWS PrivateLink to enable SSM. AWS PrivateLink restricts all network traffic between your endpoint instances, SSM, and Amazon EC2 to the Amazon network. For more information on how to setup SSM access with AWS PrivateLink, see [Set up an Amazon VPC endpoint for Session Manager](#).

### Log SSM session commands and results

After following the instructions at [Create a Session Manager preferences document (command line)](#), you can create SSM documents that define your preferences for SSM sessions. You can use SSM documents to configure session options, including data encryption, session duration, and logging. For example, you can specify whether to store session log data in an Amazon Simple Storage Service (Amazon S3) bucket or in an Amazon CloudWatch Logs group. You can create documents that define general preferences for all sessions for an AWS account and AWS Region, or documents that define preferences for individual sessions.

### Troubleshooting issues by checking error logs from SSM

Amazon SageMaker uploads errors from the SSM agent to your CloudWatch Logs in the `/aws/sagemaker/TrainingJobs` log group. SSM agent log streams are named in this format: `<job-name>/algo-<n>-<timestamp>/ssm`. For example, if you create a two-node training job named `training-job-test-remote-debug`, the training job log `training-job-test-remote-debug/algo-<n>-<timestamp>` and multiple SSM agent error logs `training-job-test-remote-debug/algo-<n>-<timestamp>/ssm` are uploaded to your CloudWatch Logs. In this example, you can review the `*/ssm` log streams to troubleshoot SSM issues.
Considerations

Consider the following when using SageMaker remote debugging.

- Remote debugging isn't supported for SageMaker algorithm containers or containers from SageMaker on AWS Marketplace.
- You can't start an SSM session for containers that have network isolation enabled because the isolation prevents outbound network calls.

Release notes for debugging capabilities of Amazon SageMaker

See the following release notes to track the latest updates for debugging capabilities of Amazon SageMaker.

December 21, 2023

New features

Released a remote debugging functionality, a new debugging capability of SageMaker that gives you a shell-level access to training containers. With this release, you can debug training jobs by logging into the job containers running on SageMaker ML instances. To learn more, see the section called “Access a training container through SSM for remote debugging”.

September 7, 2023

New features

Added a new utility module sagemaker.interactive_apps.tensorboard.TensorBoardApp that provides a function called get_app_url(). The get_app_url() function generates unsigned or presigned URLs to open the TensorBoard application in any environment in SageMaker or Amazon EC2. This is to provide a unified experience for both Studio Classic and non-Studio Classic users. For the Studio Classic environment, you can open TensorBoard by running the get_app_url() function as it is, or you can also specify a job name to start tracking as the TensorBoard application opens. For non-Studio Classic environments, you can open TensorBoard by providing your domain information to the utility function. With this functionality, regardless of where or how you run training code and launch training jobs, you can directly access TensorBoard.
by running the get_app_url function in your Jupyter notebook or terminal. This functionality is available in the SageMaker Python SDK v2.184.0 and later. For more information, see the section called “How to access TensorBoard on SageMaker”.

April 4, 2023

New features

Released SageMaker with TensorBoard, a capability that hosts TensorBoard on SageMaker. TensorBoard is available as an application through SageMaker domain, and the SageMaker Training platform supports TensorBoard output data collection to S3 and loading them automatically to the hosted TensorBoard on SageMaker. With this capability, you can run training jobs set up with TensorBoard summary writers in SageMaker, save the TensorBoard output files in Amazon S3, open the TensorBoard application directly from the SageMaker console, and load the output files using SageMaker Data Manager plugin implemented to the hosted TensorBoard interface. You don't need to install TensorBoard manually and host locally on the SageMaker IDEs or local machine. To learn more, see the section called “Use TensorBoard”.

March 16, 2023

Deprecation notes

SageMaker Debugger deprecates the framework profiling feature starting from TensorFlow 2.11 and PyTorch 2.0. You can still use the feature in the previous versions of the frameworks and SDKs as follows.

- SageMaker Python SDK <= v2.130.0
- PyTorch >= v1.6.0, < v2.0
- TensorFlow >= v2.3.1, < v2.11

With the deprecation, SageMaker Debugger also discontinues support for the following three ProfilerRules for framework profiling.

- MaxInitializationTime
- OverallFrameworkMetrics
- StepOutlier
February 21, 2023

Other changes

- The XGBoost report tab has been removed from the SageMaker Debugger's profiler dashboard. You can still access the XGBoost report by downloading it as a Jupyter notebook or a HTML file. For more information, see SageMaker Debugger XGBoost Training Report.

- Starting from this release, the built-in profiler rules are not activated by default. To use the SageMaker Debugger profiler rules to detect certain computational problems, you need to add the rules when you configure a SageMaker training job launcher.

December 1, 2020

Amazon SageMaker Debugger launched deep profiling features at re:Invent 2020.

December 3, 2019

Amazon SageMaker Debugger initially launched at re:Invent 2019.

Profile and optimize computational performance

When training state-of-the-art deep learning models that rapidly grow in size, scaling the training job of such models to a large GPU cluster and identifying computational performance issues from billions and trillions of operations and communications in every iteration of the gradient descent process become a challenge.

SageMaker provides profiling tools to visualize and diagnose such complex computation issues arising from running training jobs on AWS cloud computing resources. There are two profiling options that SageMaker offers: Amazon SageMaker Profiler and a resource utilization monitor in Amazon SageMaker Studio Classic. See the following introductions of the two functionalities to gain quick insights and learn which one to use depending on your needs.

Amazon SageMaker Profiler

Amazon SageMaker Profiler is a profiling capability of SageMaker with which you can deep dive into compute resources provisioned while training deep learning models, and gain visibility into operation-level details. SageMaker Profiler provides Python modules for adding annotations throughout PyTorch or TensorFlow training scripts and activating SageMaker Profiler. You can access the modules through the SageMaker Python SDK and AWS Deep Learning Containers.
With SageMaker Profiler, you can track all activities on CPUs and GPUs, such as CPU and GPU utilizations, kernel runs on GPUs, kernel launches on CPUs, sync operations, memory operations across CPUs and GPUs, latencies between kernel launches and corresponding runs, and data transfer between CPUs and GPUs.

SageMaker Profiler also offers a user interface (UI) that visualizes the profile, a statistical summary of profiled events, and the timeline of a training job for tracking and understanding the time relationship of the events between GPUs and CPUs.

To learn more about SageMaker Profiler, see the section called “Use SageMaker Profiler”.

**Monitoring AWS compute resources in Amazon SageMaker Studio Classic**

SageMaker also provides a user interface in Studio Classic for monitoring resource utilization at high level, but with more granularity compared to the default utilization metrics collected from SageMaker to CloudWatch.

For any training job you run in SageMaker using the SageMaker Python SDK, SageMaker starts profiling basic resource utilization metrics, such as CPU utilization, GPU utilization, GPU memory utilization, network, and I/O wait time. It collects these resource utilization metrics every 500 milliseconds.

Compared to Amazon CloudWatch metrics, which collect metrics at intervals of 1 second, the monitoring functionality of SageMaker provides finer granularity into the resource utilization metrics down to 100-millisecond (0.1 second) intervals, so you can dive deep into the metrics at the level of an operation or a step.

To access the dashboard for monitoring the resource utilization metrics of a training job, see the SageMaker Debugger UI in SageMaker Studio Experiments.

**Topics**

- Use Amazon SageMaker Profiler to profile activities on AWS compute resources
- Monitor AWS compute resource utilization in Amazon SageMaker Studio Classic
- Release notes for profiling capabilities of Amazon SageMaker
Use Amazon SageMaker Profiler to profile activities on AWS compute resources

Amazon SageMaker Profiler is currently in preview release and available at no cost in supported AWS Regions. The generally available version of Amazon SageMaker Profiler (if any) may include features and pricing that are different than those offered in preview.

Amazon SageMaker Profiler is a capability of Amazon SageMaker that provides a detailed view into the AWS compute resources provisioned during training deep learning models on SageMaker. It focuses on profiling the CPU and GPU usage, kernel runs on GPUs, kernel launches on CPUs, sync operations, memory operations across CPUs and GPUs, latencies between kernel launches and corresponding runs, and data transfer between CPUs and GPUs. SageMaker Profiler also offers a user interface (UI) that visualizes the profile, a statistical summary of profiled events, and the timeline of a training job for tracking and understanding the time relationship of the events between GPUs and CPUs.

**Note**

SageMaker Profiler supports PyTorch and TensorFlow and is available in [AWS Deep Learning Containers for SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/deep-learning-container-ecr.html). To learn more, see the section called “Supported framework images, AWS Regions, and instance types”.

**For data scientists**

Training deep learning models on a large compute cluster often has computational optimization problems, such as bottlenecks, kernel launch latencies, memory limit, and low resource utilization.

To identify such computational performance issues, you need to profile deeper into the compute resources to understand which kernels introduce latencies and which operations cause bottlenecks. Data scientists can take the benefit from using the SageMaker Profiler UI for visualizing the detailed profile of training jobs. The UI provides a dashboard furnished with summary charts and a timeline interface to track every event on the compute resources. Data scientists can also add custom annotations to track certain parts of the training job using the SageMaker Profiler Python modules.

**For administrators**
Through the Profiler landing page in the SageMaker console or SageMaker domain, you can manage the Profiler application users if you are an administrator of an AWS account or SageMaker domain. Each domain user can access their own Profiler application given the granted permissions. As a SageMaker domain administrator and domain user, you can create and delete the Profiler application given the permission level you have.

**Supported framework images, AWS Regions, and instance types**

This feature supports the following machine learning frameworks and AWS Regions.

ℹ️ **Note**

To use this feature, make sure that you have at least version 2.180.0 of the SageMaker Python SDK installed.

**SageMaker framework images pre-installed with SageMaker Profiler**

SageMaker Profiler is pre-installed in the following AWS Deep Learning Containers for SageMaker.

**PyTorch images**

<table>
<thead>
<tr>
<th>PyTorch versions</th>
<th>AWS DLC image URI</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.2.0</td>
<td><code>763104351884.dkr.ecr.&lt;region&gt;.amazonaws.com/pytorch-training:2.2.0-gpu-py310-cu121-ubuntu20.04-sagemaker</code></td>
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<tr>
<td></td>
<td><code>763104351884.dkr.ecr.&lt;region&gt;.amazonaws.com/pytorch-training:2.0.1-gpu-py310-cu121-ubuntu20.04-sagemaker</code></td>
</tr>
</tbody>
</table>
## PyTorch versions

<table>
<thead>
<tr>
<th>PyTorch versions</th>
<th>AWS DLC image URI</th>
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</thead>
</table>

## TensorFlow images

<table>
<thead>
<tr>
<th>TensorFlow versions</th>
<th>AWS DLC image URI</th>
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<tr>
<td>2.11.0</td>
<td>763104351884 .dkr.ecr.&lt;region&gt;.amazonaws.com/tensorflow-training:2.11.0-gpu-py39-cu112-ubuntu20.04-sagemaker</td>
</tr>
</tbody>
</table>

### Important

Distribution and maintenance of the framework containers in the preceding tables are under the [Framework Support Policy](#) managed by the AWS Deep Learning Containers service. We highly recommend you to upgrade to the currently supported framework versions, if you are using prior framework versions that are no longer supported.

### Note

If you want to use SageMaker Profiler for other framework images or your own Docker images, you can install SageMaker Profiler using the SageMaker Profiler Python package binary files provided in the following section.
SageMaker Profiler Python package binary files

If you want to configure your own Docker container, use SageMaker Profiler in other pre-built containers for PyTorch and TensorFlow, or install the SageMaker Profiler Python package locally, use one the following binary files. Depending on the Python and CUDA versions in your environment, choose one of the following.

**PyTorch**

- Python3.8, CUDA 11.3: [https://smppy.s3.amazonaws.com/pytorch/cu113/smprof-0.3.334-cp38-cp38-linux_x86_64.whl](https://smppy.s3.amazonaws.com/pytorch/cu113/smprof-0.3.334-cp38-cp38-linux_x86_64.whl)
- Python3.9, CUDA 11.7: [https://smppy.s3.amazonaws.com/pytorch/cu117/smprof-0.3.334-cp39-cp39-linux_x86_64.whl](https://smppy.s3.amazonaws.com/pytorch/cu117/smprof-0.3.334-cp39-cp39-linux_x86_64.whl)
- Python3.10, CUDA 11.8: [https://smppy.s3.amazonaws.com/pytorch/cu118/smprof-0.3.334-cp310-cp310-linux_x86_64.whl](https://smppy.s3.amazonaws.com/pytorch/cu118/smprof-0.3.334-cp310-cp310-linux_x86_64.whl)
- Python3.10, CUDA 12.1: [https://smppy.s3.amazonaws.com/pytorch/cu121/smprof-0.3.334-cp310-cp310-linux_x86_64.whl](https://smppy.s3.amazonaws.com/pytorch/cu121/smprof-0.3.334-cp310-cp310-linux_x86_64.whl)

**TensorFlow**

- Python3.9, CUDA 11.2: [https://smppy.s3.amazonaws.com/tensorflow/cu112/smprof-0.3.334-cp39-cp39-linux_x86_64.whl](https://smppy.s3.amazonaws.com/tensorflow/cu112/smprof-0.3.334-cp39-cp39-linux_x86_64.whl)
- Python3.10, CUDA 11.8: [https://smppy.s3.amazonaws.com/tensorflow/cu118/smprof-0.3.334-cp310-cp310-linux_x86_64.whl](https://smppy.s3.amazonaws.com/tensorflow/cu118/smprof-0.3.334-cp310-cp310-linux_x86_64.whl)

For more information about how to install SageMaker Profiler using the binary files, see the section called “(Optional) Install the SageMaker Profiler Python package”.

**Supported AWS Regions**

SageMaker Profiler is available in the following AWS Regions.

- US East (N. Virginia) (us-east-1)
- US East (Ohio) (us-east-2)
- US West (Oregon) (us-west-2)
- Europe (Frankfurt) (eu-central-1)
- Europe (Ireland) (eu-west-1)
Supported instance types

SageMaker Profiler supports profiling of training jobs on the following instance types.

CPU and GPU profiling

- ml.g4dn.12xlarge
- ml.g5.24xlarge
- ml.g5.48xlarge
- ml.p3dn.24xlarge
- ml.p4de.24xlarge
- ml.p4d.24xlarge
- ml.p5.48xlarge

GPU profiling only

- ml.g5.2xlarge
- ml.g5.4xlarge
- ml.g5.8xlarge
- ml.g5.16.xlarge

Prerequisites

The following list shows the prerequisites to start using SageMaker Profiler.

- A SageMaker domain set up with Amazon VPC in your AWS account.

  For instructions on setting up a domain, see [Onboard to Amazon SageMaker domain using quick setup](#). You also need to add domain user profiles for individual users to access the Profiler UI application. For more information, see [Add and remove SageMaker domain user profiles](#).

- The following list is the minimum set of permissions for using the Profiler UI application.
  - sagemaker:CreateApp
  - sagemaker:DeleteApp
  - sagemaker:DescribeTrainingJob
  - sagemaker:Search
• s3:GetObject
• s3:ListBucket

Prepare and run a training job with SageMaker Profiler

Setting up to running a training job with the SageMaker Profiler consists of two steps: adapting the training script and configuring the SageMaker training job launcher.

Topics
• Step 1: Adapt your training script using the SageMaker Profiler Python modules
• Step 2: Create a SageMaker framework estimator and activate SageMaker Profiler
• (Optional) Install the SageMaker Profiler Python package

Step 1: Adapt your training script using the SageMaker Profiler Python modules

To start capturing kernel runs on GPUs while the training job is running, modify your training script using the SageMaker Profiler Python modules. Import the library and add the start_profiling() and stop_profiling() methods to define the beginning and the end of profiling. You can also use optional custom annotations to add markers in the training script to visualize hardware activities during particular operations in each step.

Note that the annotators extract operations from GPUs. For profiling operations in CPUs, you don’t need to add any additional annotations. CPU profiling is also activated when you specify the profiling configuration, which you’ll practice in the section called “Step 2: Create a SageMaker framework estimator and activate SageMaker Profiler”.

⚠️ Note
Profiling an entire training job is not the most efficient use of resources. We recommend profiling at most 300 steps of a training job.

⚠️ Important
The release on December 14, 2023 involves a breaking change. The SageMaker Profiler Python package name is changed from smppy to smprof. This is effective in the SageMaker Framework Containers for TensorFlow v2.12 and later.
If you use one of the previous versions of the SageMaker Framework Containers such as TensorFlow v2.11.0, the SageMaker Profiler Python package is still available as smppy. If you are uncertain about which version or the package name you should use, replace the import statement of the SageMaker Profiler package with the following code snippet.

```python
try:
    import smprof
except ImportError:
    # backward-compatibility for TF 2.11 and PT 1.13.1 images
    import smppy as smprof
```

**Approach 1. Use the context manager `smprof.annotate` to annotate full functions**

You can wrap full functions with the `smprof.annotate()` context manager. This wrapper is recommended if you want to profile by functions instead of code lines. The following example script shows how to implement the context manager to wrap the training loop and full functions in each iteration.

```python
import smprof

SMProf = smprof.SMProfiler.instance()
config = smprof.Config()
config.profiler = {
    "EnableCuda": "1",
}
SMProf.configure(config)
SMProf.start_profiling()

for epoch in range(args.epochs):
    if world_size > 1:
        sampler.set_epoch(epoch)
    tstart = time.perf_counter()
    for i, data in enumerate(trainloader, 0):
        with smprof.annotate("step_"+str(i)):
            inputs, labels = data
            inputs = inputs.to("cuda", non_blocking=True)
            labels = labels.to("cuda", non_blocking=True)

            optimizer.zero_grad()
```
Approach 2. Use `smprof.annotation_begin()` and `smprof.annotation_end()` to annotate specific code line in functions

You can also define annotations to profile specific code lines. You can set the exact starting point and end point of profiling at the level of individual code lines, not by the functions. For example, in the following script, the `step_annotator` is defined at the beginning of each iteration and ends at the end of the iteration. Meanwhile, other detailed annotators for each operations are defined and wrap around the target operations throughout each iteration.

```python
import smprof

SMProf = smprof.SMProfiler.instance()
cfg = smprof.Config()
cfg.profiler = {
    "EnableCuda": "1",
}
SMProf.configure(cfg)
SMProf.start_profiling()

for epoch in range(args.epochs):
    if world_size > 1:
        sampler.set_epoch(epoch)
    tstart = time.perf_counter()
    for i, data in enumerate(trainloader, 0):
        step_annotator = smprof.annotation_begin("step_" + str(i))
        inputs, labels = data
        inputs = inputs.to("cuda", non_blocking=True)
        labels = labels.to("cuda", non_blocking=True)
        optimizer.zero_grad()
        forward_annotator = smprof.annotation_begin("Forward")
```
outputs = net(inputs)
smprof.annotation_end(forward_annotator)

loss_annotator = smprof.annotation_begin("Loss")
loss = criterion(outputs, labels)
smprof.annotation_end(loss_annotator)

backward_annotator = smprof.annotation_begin("Backward")
loss.backward()
smprof.annotation_end(backward_annotator)

optimizer_annotator = smprof.annotation_begin("Optimizer")
optimizer.step()
smprof.annotation_end(optimizer_annotator)

smprof.annotation_end(step_annotator)
SMProf.stop_profiling()

After annotating and setting up the profiler initiation modules, save the script to submit using a SageMaker training job launcher in the following Step 2. The sample launcher assumes that the training script is named train_with_profiler_demo.py.

**Step 2: Create a SageMaker framework estimator and activate SageMaker Profiler**

The following procedure shows how to prepare a SageMaker framework estimator for training using the SageMaker Python SDK.

1. Set up a profiler_config object using the ProfilerConfig and Profiler modules as follows.

```python
from sagemaker import ProfilerConfig, Profiler
profiler_config = ProfilerConfig(
    profile_params = Profiler(cpu_profiling_duration=3600)
)
```

The following is the description of the Profiler module and its argument.

- **Profiler**: The module for activating SageMaker Profiler with the training job.
- **cpu_profiling_duration (int)**: Specify the time duration in seconds for profiling on CPUs. Default is 3600 seconds.
2. Create a SageMaker framework estimator with the `profiler_config` object created in the previous step. The following code shows an example of creating a PyTorch estimator. If you want to create a TensorFlow estimator, import `sagemaker.tensorflow.TensorFlow` instead, and specify one of the TensorFlow versions supported by SageMaker Profiler. For more information about supported frameworks and instance types, see the section called “SageMaker framework images pre-installed with SageMaker Profiler”.

```python
import sagemaker
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    framework_version="2.0.0",
    role=sagemaker.get_execution_role(),
    entry_point="train_with_profiler_demo.py",  # your training job entry point
    source_dir=source_dir,  # source directory for your training script
    output_path=output_path,
    base_job_name="sagemaker-profiler-demo",
    hyperparameters=hyperparameters,  # if any
    instance_count=1,  # Recommended to test with < 8
    instance_type=ml.p4d.24xlarge,
    profiler_config=profiler_config
)
```

3. Start the training job by running the `fit` method. With `wait=False`, you can silence the training job logs and let it run in the background.

```python
estimator.fit(wait=False)
```

While running the training job or after the job has completed, you can go to the next topic at the section called “Open the SageMaker Profiler UI application” and start exploring and visualizing the saved profiles.

If you want to directly access the profile data saved in the Amazon S3 bucket, use the following script to retrieve the S3 URI.

```python
import os
# This is an ad-hoc function to get the S3 URI
# to where the profile output data is saved
def get_detailed_profiler_output_uri(estimator):
    config_name = None
```
for processing in estimator.profiler_rule_configs:
    params = processing.get("RuleParameters", dict())
    rule = config_name = params.get("rule_to_invoke", "")
    if rule == "DetailedProfilerProcessing":
        config_name = processing.get("RuleConfigurationName")
        break
    return os.path.join(
        estimator.output_path,
        estimator.latest_training_job.name,
        "rule-output",
        config_name,
    )

print(
    f"Profiler output S3 bucket: ",
    get_detailed_profiler_output_uri(estimator)
)

(Optional) Install the SageMaker Profiler Python package

To use SageMaker Profiler on PyTorch or TensorFlow framework images not listed in the section called “SageMaker framework images pre-installed with SageMaker Profiler”, or on your own custom Docker container for training, you can install SageMaker Profiler by using one of the section called “SageMaker Profiler Python package binary files”.

Option 1: Install the SageMaker Profiler package while launching a training job

If you want to use SageMaker Profiler for training jobs using PyTorch or TensorFlow images not listed in the section called “SageMaker framework images pre-installed with SageMaker Profiler”, create a requirements.txt file and locate it under the path you specify to the source_dir parameter of the SageMaker framework estimator in Step 2. For more information about setting up a requirements.txt file in general, see Using third-party libraries in the SageMaker Python SDK documentation. In the requirements.txt file, add one of the S3 bucket paths for the section called “SageMaker Profiler Python package binary files”.

# requirements.txt
https://smppy.s3.amazonaws.com/tensorflow/cu112/smprof-0.3.332-cp39-cp39-linux_x86_64.whl

Option 2: Install the SageMaker Profiler package in your custom Docker containers
If you use a custom Docker container for training, add one of the the section called “SageMaker Profiler Python package binary files” to your Dockerfile.

```bash
# Install the smprof package version compatible with your CUDA version
RUN pip install https://smppy.s3.amazonaws.com/tensorflow/cu112/smprof-0.3.332-cp39-cp39-linux_x86_64.whl
```

For guidance on running a custom Docker container for training on SageMaker in general, see Adapting your own training container.

**Open the SageMaker Profiler UI application**

You can access the SageMaker Profiler UI application through the following options.

**Topics**

- **Option 1: Launch the SageMaker Profiler UI from the domain details page**
- **Option 2: Launch the SageMaker Profiler UI application from the SageMaker Profiler landing page in the SageMaker console**
- **Option 3: Use the application launcher function in the SageMaker Python SDK**

**Option 1: Launch the SageMaker Profiler UI from the domain details page**

If you have access to the SageMaker console, you can take this option.

**Navigate to the domain details page**

The following procedure shows how to navigate to the domain details page.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the left navigation pane, choose domains.
3. From the list of domains, select the domain in which you want to launch the SageMaker Profiler application.

**Launch the SageMaker Profiler UI application**

The following procedure shows how to launch the SageMaker Profiler application that is scoped to a user profile.

1. On the domain details page, choose the **User profiles** tab.
2. Identify the user profile for which you want to launch the SageMaker Profiler UI application.
3. Choose **Launch** for the selected user profile, and choose **Profiler**.

**Option 2: Launch the SageMaker Profiler UI application from the SageMaker Profiler landing page in the SageMaker console**

The following procedure describes how to launch the SageMaker Profiler UI application from the SageMaker Profiler landing page in the SageMaker console. If you have access to the SageMaker console, you can take this option.

2. On the left navigation pane, choose **Profiler**.
3. Under **Get started**, select the domain in which you want to launch the Studio Classic application.
   - If your user profile only belongs to one domain, you do not see the option for selecting a domain.
4. Select the user profile for which you want to launch the SageMaker Profiler UI application. If there is no user profile in the domain, choose **Create user profile**. For more information about creating a new user profile, see [Add and Remove User Profiles](#).
5. Choose **Open Profiler**.

**Option 3: Use the application launcher function in the SageMaker Python SDK**

If you are a SageMaker domain user and have access only to SageMaker Studio, you can access the SageMaker Profiler UI application through SageMaker Studio Classic by running the `sagemaker.interactive_apps.detail_profiler_app.DetailProfilerApp` function.

Note that SageMaker Studio Classic is the previous Studio UI experience before re:Invent 2023, and is migrated as an application into a newly designed Studio UI at re:Invent 2023. The SageMaker Profiler UI application is available at SageMaker domain level, and thus requires your domain ID and user profile name. Currently, the `DetailedProfilerApp` function only works within the SageMaker Studio Classic application; the function properly takes in the domain and user profile information from SageMaker Studio Classic.

For domain, domain users, and Studio created before re:Invent 2023, Studio Classic would be the default experience unless you have updated it following the instructions at [Migrating from Amazon SageMaker Studio Classic](#). If this is your case, there's no further action needed, and you can directly launch the SageMaker Profiler UI application by running the `DetailProfilerApp` function.
If you created a new domain and Studio after re:Invent 2023, launch the Studio Classic application within the Studio UI and then run the `DetailProfilerApp` function to launch the SageMaker Profiler UI application.

Note that the `DetailProfilerApp` function doesn’t work in other SageMaker machine learning IDEs, such as the SageMaker Studio JupyterLab application, the SageMaker Studio Code Editor application, and SageMaker Notebook instances. If you run the `DetailProfilerApp` function in those IDEs, it returns a URL to the Profiler landing page in the SageMaker console, instead of a direct link to open the Profiler UI application.

**Explore the profile output data visualized in the SageMaker Profiler UI**

This section walks through the SageMaker Profiler UI and provides tips for how to use and gain insights from it.

**Load profile**

When you open the SageMaker Profiler UI, the Load profile page opens up. To load and generate the Dashboard and Timeline, go through the following procedure.

**To load the profile of a training job**

1. From the List of training jobs section, use the check box to choose the training job for which you want to load the profile.
2. Choose Load. The job name should appear in the Loaded profile section at the top.
3. Choose the radio button on the left of the Job name to generate the Dashboard and Timeline. Note that when you choose the radio button, the UI automatically opens the Dashboard. Note also that if you generate the visualizations while the job status and loading status still appear to be in progress, the SageMaker Profiler UI generates Dashboard plots and a Timeline up to the most recent profile data collected from the ongoing training job or the partially loaded profile data.

**Tip**

You can load and visualize one profile at a time. To load another profile, you must first unload the previously loaded profile. To unload a profile, use the trash bin icon on the right end of the profile in the Loaded profile section.
After you finish loading and selecting the training job, the UI opens the **Dashboard page** furnished with the following panels by default.

- **GPU active time** – This pie chart shows the percentage of GPU active time versus GPU idle time. You can check if your GPUs are more active than idle throughout the entire training job. GPU active time is based on the profile data points with a utilization rate greater than 0%, whereas GPU idle time is the profiled data points with 0% utilization.

- **GPU utilization over time** – This timeline graph shows the average GPU utilization rate over time per node, aggregating all of the nodes in a single chart. You can check if the GPUs have an unbalanced workload, under-utilization issues, bottlenecks, or idle issues during certain time intervals. To track the utilization rate at the individual GPU level and related kernel runs, use the [section called “Timeline interface”](#). Note that the GPU activity collection starts from where you added the profiler starter function `SMProf.start_profiling()` in your training script, and stops at `SMProf.stop_profiling()`.
- **CPU active time** – This pie chart shows the percentage of CPU active time versus CPU idle time. You can check if your CPUs are more active than idle throughout the entire training job. CPU active time is based on the profiled data points with a utilization rate greater than 0%, whereas CPU idle time is the profiled data points with 0% utilization.

- **CPU utilization over time** – This timeline graph shows the average CPU utilization rate over time per node, aggregating all of the nodes in a single chart. You can check if the CPUs are bottlenecked or underutilized during certain time intervals. To track the utilization rate of the CPUs aligned with the individual GPU utilization and kernel runs, use the the section called “Timeline interface”. Note that the utilization metrics start from the start from the job initialization.

- **Time spent by all GPU kernels** – This pie chart shows all GPU kernels operated throughout the training job. It shows the top 15 GPU kernels by default as individual sectors and all other kernels in one sector. Hover over the sectors to see more detailed information. The value shows the total time of the GPU kernels operated in seconds, and the percentage is based on the entire time of the profile.

- **Time spent by top 15 GPU kernels** – This pie chart shows all GPU kernels operated throughout the training job. It shows the top 15 GPU kernels as individual sectors. Hover over the sectors to see more detailed information. The value shows the total time of the GPU kernels operated in seconds, and the percentage is based on the entire time of the profile.

- **Launch counts of all GPU kernels** – This pie chart shows the number of counts for every GPU kernel launched throughout the training job. It shows the top 15 GPU kernels as individual sectors and all other kernels in one sector. Hover over the sectors to see more detailed information. The value shows the total count of the launched GPU kernels, and the percentage is based on the entire count of all kernels.

- **Launch counts of top 15 GPU kernels** – This pie chart shows the number of counts of every GPU kernel launched throughout the training job. It shows the top 15 GPU kernels. Hover over the sectors to see more detailed information. The value shows the total count of the launched GPU kernels, and the percentage is based on the entire count of all kernels.

- **Step time distribution** – This histogram shows the distribution of step durations on GPUs. This plot is generated only after you add the step annotator in your training script.

- **Kernel precision distribution** – This pie chart shows the percentage of time spent on running kernels in different data types such as FP32, FP16, INT32, and INT8.

- **GPU activity distribution** – This pie chart shows the percentage of time spent on GPU activities, such as running kernels, memory (memcpy and memset), and synchronization (sync).
• **GPU memory operations distribution** – This pie chart shows the percentage of time spent on GPU memory operations. This visualizes the memcpy activities and helps identify if your training job is spending excessive time on certain memory operations.

• **Create a new histogram** – Create a new diagram of a custom metric you annotated manually during the section called “Step 1: Adapt your training script using the SageMaker Profiler Python modules”. When adding a custom annotation to a new histogram, select or type the name of the annotation you added in the training script. For example, in the demo training script in Step 1, step, Forward, Backward, Optimize, and Loss are the custom annotations. While creating a new histogram, these annotation names should appear in the drop-down menu for metric selection. If you choose Backward, the UI adds the histogram of the time spent on backward passes throughout the profiled time to the Dashboard. This type of histogram is useful for checking if there are outliers taking abnormally longer time and causing bottleneck problems.

The following screenshots show the GPU and CPU active time ratio and the average GPU and CPU utilization rate with respect to time per compute node.

The following screenshot shows an example of pie charts for comparing how many times the GPU kernels are launched and measuring the time spent on running them. In the **Time spent by all GPU kernels** and **Launch counts of all GPU kernels** panels, you can also specify an integer to the input
field for $k$ to adjust the number of legend to show in the plots. For example, if you specify 10, the plots show the top 10 most run and launched kernels respectively.

The following screenshot shows an example of step time duration histogram, and pie charts for the kernel precision distribution, GPU activity distribution, and GPU memory operation distribution.
Timeline interface

To gain a detailed view into the compute resources at the level of operations and kernels scheduled on the CPUs and run on the GPUs, use the Timeline interface.

You can zoom in and out and pan left or right in the timeline interface using your mouse, the [w, a, s, d] keys, or the four arrow keys on the keyboard.

Tip

For more tips on the keyboard shortcuts to interact with the Timeline interface, choose Keyboard shortcuts in the left pane.

The timeline tracks are organized in a tree structure, giving you information from the host level to the device level. For example, if you run N instances with eight GPUs in each, the timeline structure of each instance would be as follows.

- algo-i_node – This is what SageMaker tags to assign jobs to provisioned instances. The digit i_node is randomly assigned. For example, if you use 4 instances, this section expands from algo-1 to algo-4.

- CPU – In this section, you can check the average CPU utilization rate and performance counters.

- GPUs – In this section, you can check the average GPU utilization rate, individual GPU utilization rate, and kernels.

- SUM Utilization – The average GPU utilization rates per instance.

- HOST-0 PID-123 – A unique name assigned to each process track. The acronym PID is the process ID, and the number appended to it is the process ID number that's recorded during data capture from the process. This section shows the following information from the process.

  - GPU-i_num_gpu utilization – The utilization rate of the i_num_gpu-th GPU over time.
  - GPU-i_num_gpu device – The kernel runs on the i_num_gpu-th GPU device.
  - stream i_cuda_stream – CUDA streams showing kernel runs on the GPU device. To learn more about CUDA streams, see the slides in PDF at CUDA C/C++ Streams and Concurrency provided by NVIDIA.
  - GPU-i_num_gpu host – The kernel launches on the i_num_gpu-th GPU host.
The following several screenshots show the **Timeline** of the profile of a training job run on `ml.p4d.24xlarge` instances, which are equipped with 8 NVIDIA A100 Tensor Core GPUs in each.

The following is a zoomed-out view of the profile, printing a dozen of steps including an intermittent data loader between `step_232` and `step_233` for fetching the next data batch.

For each CPU, you can track the CPU utilization and performance counters, such as "clk_unhalted_ref.tsc" and "itlb_misses.miss_causes_a_walk", which are indicative of instructions run on the CPU.

For each GPU, you can see a host timeline and a device timeline. Kernel launches are on the host timeline and kernel runs are on the device timeline. You can also see annotations (such as forward, backward, and optimize) if you have added in training script in the GPU host timeline.

In the timeline view, you can also track kernel launch-and-run pairs. This helps you understand how a kernel launch scheduled on a host (CPU) is run on the corresponding GPU device.

**Tip**

Press the f key to zoom into the selected kernel.
The following screenshot is a zoomed-in view into step_233 and step_234 from the previous screenshot. The timeline interval selected in the following screenshot is the AllReduce operation, an essential communication and synchronization step in distributed training, run on the GPU-0 device. In the screenshot, note that the kernel launch in the GPU-0 host connects to the kernel run in the GPU-0 device stream 1, indicated with the arrow in cyan color.

Also two information tabs appear in the bottom pane of the UI when you select a timeline interval, as shown in the previous screenshot. The **Current Selection** tab shows the details of the selected kernel and the connected kernel launch from the host. The connection direction is always from host (CPU) to device (GPU) since each GPU kernel is always called from a CPU. The **Connections** tab shows the chosen kernel launch and run pair. You can select either of them to move it to the center of the **Timeline** view.

The following screenshot zooms in further into the AllReduce operation launch and run pair.
In **Information**, you can access information about the loaded training job, such as the instance type, Amazon Resource Names (ARNs) of compute resources provisioned for the job, node names, and hyperparameters.

**Settings**

The SageMaker Profiler UI application instance is configured to shut down after 2 hours of idle time by default. In **Settings**, use the following settings to adjust the auto shutdown timer.

- **Enable app auto shutdown** – Choose and set to **Enabled** to let the application automatically shut down after the specified number of hours of idle time. To turn off the auto-shutdown functionality, choose **Disabled**.

- **Auto shutdown threshold in hours** – If you choose **Enabled** for **Enable app auto shutdown**, you can set the threshold time in hours for the application to shut down automatically. This is set to 2 by default.

**Frequently asked questions about using SageMaker Profiler**

Use the following frequently asked questions to find answers about using SageMaker Profiler.
Q. I’m getting an error message, ModuleNotFoundError: No module named 'smppy'

Since December 2023, the name of the SageMaker Profiler Python package has changed from smppy to smprof to resolve a duplicate package name issue; smppy is already used by an open source package.

Therefore, if you have been using smppy since before December 2023 and experiencing this ModuleNotFoundError issue, it might be due to the outdated package name in your training script while having the latested smprof package installed or using one of the latest the section called “SageMaker framework images pre-installed with SageMaker Profiler”. In this case, make sure that you replace all mentions of smppy with smprof throughout your training script.

While updating the SageMaker Profiler Python package name in your training scripts, to avoid confusion around which version of the package name you should use, consider using a conditional import statement as shown in the following code snippet.

```python
try:
    import smprof
except ImportError:
    # backward-compatibility for TF 2.11 and PT 1.13.1 images
    import smppy as smprof
```

Also note that if you have been using smppy while upgrading to the latest PyTorch or TensorFlow versions, make sure that you install the latest smprof package by following instructions at the section called “(Optional) Install the SageMaker Profiler Python package”.

Q. I’m getting an error message, ModuleNotFoundError: No module named 'smprof'

First, make sure that you use one of the officially supported SageMaker Framework Containers. If you don’t use one of those, you can install the smprof package by following instructions at the section called “(Optional) Install the SageMaker Profiler Python package”.

Q. I’m not able to import ProfilerConfig

If you are unable to import ProfilerConfig in your job launcher script using the SageMaker Python SDK, your local environment or the Jupyter kernel might have a significantly outdated version of the SageMaker Python SDK. Make sure that you upgrade the SDK to the latest version.

```
$ pip install --upgrade sagemaker
```
Q. I'm getting an error message, aborted: core dumped when importing smprof into my training script

In an earlier version of smprof, this issue occurs with PyTorch 2.0+ and PyTorch Lightning. To resolve this issue, also install the latest smprof package by following instructions at the section called “(Optional) Install the SageMaker Profiler Python package”.

Q. I cannot find the SageMaker Profiler UI from SageMaker Studio. How can I find it?

If you have access to the SageMaker console, choose one of the following options.

- the section called “Option 1: Launch the SageMaker Profiler UI from the domain details page”
- the section called “Option 2: Launch the SageMaker Profiler UI application from the SageMaker Profiler landing page in the SageMaker console”

If you are a domain user and don’t have access to the SageMaker console, you can access the application through SageMaker Studio Classic. If this is your case, choose the following option.

- the section called “Option 3: Use the application launcher function in the SageMaker Python SDK”

Considerations

Consider the following when using SageMaker Profiler.

- SageMaker Profiler is not compatible with SageMaker managed warm pools.

Monitor AWS compute resource utilization in Amazon SageMaker Studio Classic

To track compute resource utilization of your training job, use the monitoring tools offered by Amazon SageMaker Debugger.

For any training job you run in SageMaker using the SageMaker Python SDK, Debugger collects basic resource utilization metrics, such as CPU utilization, GPU utilization, GPU memory utilization, network, and I/O wait time every 500 milliseconds. To see the dashboard of the resource utilization metrics of your training job, simply use the SageMaker Debugger UI in SageMaker Studio Experiments.
Deep learning operations and steps might operate in intervals of milliseconds. Compared to Amazon CloudWatch metrics, which collect metrics at intervals of 1 second, Debugger provides finer granularity into the resource utilization metrics down to 100-millisecond (0.1 second) intervals so you can dive deep into the metrics at the level of an operation or a step.

If you want to change the metric collection time interval, you can add a parameter for profiling configuration to your training job launcher. For example, if you're using the SageMaker Python SDK, you need to pass the `profiler_config` parameter when you create an estimator object. To learn how to adjust the resource utilization metric collection interval, see the section called “Code template for configuring a SageMaker estimator object with the SageMaker Debugger Python modules in the SageMaker Python SDK” and then the section called “Configure settings for basic profiling of system resource utilization”.

Additionally, you can add issue detecting tools called built-in profiling rules provided by SageMaker Debugger. The built-in profiling rules run analysis against the resource utilization metrics and detect computational performance issues. For more information, see the section called “Configure built-in profiler rules”. You can receive rule analysis results through the SageMaker Debugger UI in SageMaker Studio Experiments or the SageMaker Debugger Profiling Report. You can also create custom profiling rules using the SageMaker Python SDK.

To learn more about monitoring functionalities provided by SageMaker Debugger, see the following topics.

Topics

- Configure an estimator with parameters for basic profiling using the Amazon SageMaker Debugger Python modules
- Configure built-in profiler rules managed by Amazon SageMaker Debugger
- List of Debugger built-in profiler rules
- Amazon SageMaker Debugger UI in Amazon SageMaker Studio Classic Experiments
- SageMaker Debugger interactive report
- Analyze data using the Debugger Python client library

Configure an estimator with parameters for basic profiling using the Amazon SageMaker Debugger Python modules

By default, SageMaker Debugger basic profiling is on by default and monitors resource utilization metrics, such as CPU utilization, GPU utilization, GPU memory utilization, Network, and I/O

Monitor AWS compute resource utilization in SageMaker Studio Classic
wait time, of all SageMaker training jobs submitted using the Amazon SageMaker Python SDK. SageMaker Debugger collects these resource utilization metrics every 500 milliseconds. You don't need to make any additional changes in your code, training script, or the job launcher for tracking basic resource utilization. If you want to access the resource utilization metrics dashboard of your training job in SageMaker Studio, you can jump onto the Amazon SageMaker Debugger UI in Amazon SageMaker Studio Classic Experiments.

If you want to change the metric collection interval for basic profiling, you can specify Debugger-specific parameters while creating a SageMaker training job launcher using the SageMaker Python SDK, AWS SDK for Python (Boto3), or AWS Command Line Interface (CLI). In this guide, we focus on how to change profiling options using the Amazon SageMaker Python SDK.

If you want to activate the rules that detect system resource utilization problems automatically, you can add the rules parameter in the estimator object for activating the rules.

⚠️ **Important**

To use the latest SageMaker Debugger features, you need to upgrade the SageMaker Python SDK and the SMDebug client library. In your iPython kernel, Jupyter Notebook, or JupyterLab environment, run the following code to install the latest versions of the libraries and restart the kernel.

```python
import sys
import IPython
!{sys.executable} -m pip install -U sagemaker smdebug
IPython.Application.instance().kernel.do_shutdown(True)
```

**Code template for configuring a SageMaker estimator object with the SageMaker Debugger Python modules in the SageMaker Python SDK**

To adjust the basic profiling configuration (profiler_config) or add the profiler rules (rules), choose one of the tabs to get the template for setting up a SageMaker estimator. In the subsequent pages, you can find more information about how to configure the two parameters.
Note

The following code examples are not directly executable. Proceed to the next sections to learn how to configure each parameter.

PyTorch

```python
# An example of constructing a SageMaker PyTorch estimator
import boto3
import sagemaker
from sagemaker.pytorch import PyTorch
from sagemaker.debugger import ProfilerConfig, ProfilerRule, rule_configs

session=boto3.session.Session()
region=session.region_name

profiler_config=ProfilerConfig(...)
rules=[
    ProfilerRule.sagemaker(rule_configs.BuiltInRule())
]

estimator=PyTorch(
    entry_point="directory/to/your_training_script.py",
    role=sagemaker.get_execution_role(),
    base_job_name="debugger-profiling-demo",
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="1.12.0",
    py_version="py37",

    # SageMaker Debugger parameters
    profiler_config=profiler_config,
    rules=rules
)

estimator.fit(wait=False)
```

TensorFlow

```python
# An example of constructing a SageMaker TensorFlow estimator
import boto3
```
import sagemaker
from sagemaker.tensorflow import TensorFlow
from sagemaker.debugger import ProfilerConfig, ProfilerRule, rule_configs

session=boto3.session.Session()
region=session.region_name

profiler_config=ProfilerConfig(...)
rules=[
    ProfilerRule.sagemaker(rule_configs.BuiltInRule())
]

estimator=TensorFlow(
    entry_point="directory/to/your_training_script.py",
    role=sagemaker.get_execution_role(),
    base_job_name="debugger-profiling-demo",
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="2.8.0",
    py_version="py37",

    # SageMaker Debugger parameters
    profiler_config=profiler_config,
    rules=rules
)

estimator.fit(wait=False)

MXNet

# An example of constructing a SageMaker MXNet estimator
import sagemaker
from sagemaker.mxnet import MXNet
from sagemaker.debugger import ProfilerConfig, ProfilerRule, rule_configs

profiler_config=ProfilerConfig(...)
rules=[
    ProfilerRule.sagemaker(rule_configs.BuiltInRule())
]

estimator=MXNet(
    entry_point="directory/to/your_training_script.py",
    role=sagemaker.get_execution_role(),
base_job_name="debugger-profiling-demo",
instance_count=1,
instance_type="ml.p3.2xlarge",
framework_version="1.7.0",
py_version="py37",

# SageMaker Debugger parameters
profiler_config=profiler_config,
rules=rules
)

estimator.fit(wait=False)

Note

For MXNet, when configuring the profiler_config parameter, you can only configure for system monitoring. Profiling framework metrics is not supported for MXNet.

XGBoost

# An example of constructing a SageMaker XGBoost estimator
import sagemaker
from sagemaker.xgboost.estimator import XGBoost
from sagemaker.debugger import ProfilerConfig, ProfilerRule, rule_configs

profiler_config=ProfilerConfig(...)
rules=[
    ProfilerRule.sagemaker(rule_configs.BuiltInRule())
]

estimator=XGBoost(
    entry_point="directory/to/your_training_script.py",
    role=sagemaker.get_execution_role(),
    base_job_name="debugger-profiling-demo",
    instance_count=1,
    instance_type="ml.p3.2xlarge",
    framework_version="1.5-1",

    # Debugger-specific parameters
    profiler_config=profiler_config,
rules=rules
)
estimator.fit(wait=False)

## Note

For XGBoost, when configuring the `profiler_config` parameter, you can only configure for system monitoring. Profiling framework metrics is not supported for XGBoost.

Generic estimator

```python
# An example of constructing a SageMaker generic estimator using the XGBoost algorithm base image
import boto3
import sagemaker
from sagemaker.estimator import Estimator
from sagemaker import image_uris
from sagemaker.debugger import ProfilerConfig, DebuggerHookConfig, Rule, ProfilerRule, rule_configs

profiler_config = ProfilerConfig(...)  
rules = [ProfilerRule.sagemaker(rule_configs.BuiltInRule())]

region = boto3.Session().region_name
xgboost_container = sagemaker.image_uris.retrieve("xgboost", region, "1.5-1")

estimator = Estimator(
    role=sagemaker.get_execution_role(),
    image_uri=xgboost_container,
    base_job_name="debugger-demo",
    instance_count=1,
    instance_type="ml.m5.2xlarge",

    # Debugger-specific parameters
    profiler_config=profiler_config,
    rules=rules
    )
```
estimator.fit(wait=False)

The following provides brief descriptions of the parameters.

- **profiler_config** – Configure Debugger to collect system metrics and framework metrics from your training job and save into your secured S3 bucket URI or local machine. You can set how frequently or loosely collect the system metrics. To learn how to configure the `profiler_config` parameter, see [Configure settings for basic profiling of system resource utilization](#) and [Configure for framework profiling](#).

- **rules** – Configure this parameter to activate SageMaker Debugger built-in rules that you want to run in parallel. Make sure that your training job has access to this S3 bucket. The rules runs on processing containers and automatically analyze your training job to find computational and operational performance issues. The `ProfilerReport` rule is the most integrated rule that runs all built-in profiling rules and saves the profiling results as a report into your secured S3 bucket. To learn how to configure the `rules` parameter, see [Configure built-in profiler rules managed by Amazon SageMaker Debugger](#).

**Note**

Debugger securely saves output data in subfolders of your default S3 bucket. For example, the format of the default S3 bucket URI is `s3://sagemaker-<region>-<12digit_account_id>/<base-job-name>/<debugger-subfolders>/`. There are three subfolders created by Debugger: debug-output, profiler-output, and rule-output. You can also retrieve the default S3 bucket URIs using the `SageMaker estimator` classmethods.

See the following topics to find out how to configure the Debugger-specific parameters in detail.

**Topics**

- [Configure settings for basic profiling of system resource utilization](#)
- [Configure for framework profiling](#)
- [Updating Debugger system monitoring and framework profiling configuration while a training job is running](#)
- [Turn off Debugger](#)
Configure settings for basic profiling of system resource utilization

To adjust the time interval for collecting the utilization metrics, use the `ProfilerConfig` API operation to create a parameter object while constructing a SageMaker framework or generic estimator depending on your preference.

**Note**

By default, for all SageMaker training jobs, Debugger collects resource utilization metrics from Amazon EC2 instances every 500 milliseconds for system monitoring, without any Debugger-specific parameters specified in SageMaker estimators. Debugger saves the system metrics in the default S3 bucket. The format of the default S3 bucket URI is `s3://sagemaker-<region>-<12digit_account_id>/<training-job-name>/profiler-output/`.

The following code example shows how to set up the `profiler_config` parameter with a system monitoring time interval of 1000 milliseconds.

```python
from sagemaker.debugger import ProfilerConfig

profiler_config=ProfilerConfig(
    system_monitor_interval_millis=1000
)
```

- `system_monitor_interval_millis (int)` – Specify the monitoring intervals in milliseconds to record system metrics. Available values are 100, 200, 500, 1000 (1 second), 5000 (5 seconds), and 60000 (1 minute) milliseconds. The default value is 500 milliseconds.

To see the progress of system monitoring, see [Open the Amazon SageMaker Debugger Insights dashboard](#).

Configure for framework profiling

**Warning**

In favor of [Amazon SageMaker Profiler](#), SageMaker Debugger deprecates the framework profiling feature starting from TensorFlow 2.11 and PyTorch 2.0. You can still use the feature in the previous versions of the frameworks and SDKs as follows.
To enable Debugger framework profiling, configure the `framework_profile_params` parameter when you construct an estimator. Debugger framework profiling collects framework metrics, such as data from initialization stage, data loader processes, Python operators of deep learning frameworks and training scripts, detailed profiling within and between steps, with cProfile or Pyinstrument options. Using the `FrameworkProfile` class, you can configure custom framework profiling options.

**Note**

Before getting started with Debugger framework profiling, verify that the framework used to build your model is supported by Debugger for framework profiling. For more information, see [Supported Frameworks and Algorithms](#).

Debugger saves the framework metrics in a default S3 bucket. The format of the default S3 bucket URI is `s3://sagemaker-<region>-<12digit_account_id>/<training-job-name>/profiler-output/`.

### Start a training job with the default framework profiling

The following example code is the simplest `profiler_config` parameter setting to start the default system monitoring and the default framework profiling. The `FrameworkProfile` class in the following example code initiates the default framework profiling when a training job starts. Debugger framework profiling includes the following options: detailed profiling, data loader profiling, and Python profiling.

```python
from sagemaker.debugger import ProfilerConfig, FrameworkProfile

profiler_config=ProfilerConfig(
    framework_profile_params=FrameworkProfile()
)
```
With this profiler_config parameter configuration, Debugger calls the default settings of monitoring and profiling. Debugger monitors system metrics every 500 milliseconds; profiles the fifth step with the detailed profiling option; the seventh step with the data loader profiling option; and the ninth, tenth, and eleventh steps with the Python profiling option.

To find available profiling configuration options, the default parameter settings, and examples of how to configure them, see Start a training job with the default system monitoring and customized framework profiling with different profiling options and SageMaker Debugger APIs – FrameworkProfile in the Amazon SageMaker Python SDK.

If you want to change the system monitoring interval and enable the default framework profiling, you can specify the system_monitor_interval_millis parameter explicitly with the framework_profile_params parameter. For example, to monitor every 1000 milliseconds and enable the default framework profiling, use the following example code.

```python
from sagemaker.debugger import ProfilerConfig, FrameworkProfile

profiler_config=ProfilerConfig(
    system_monitor_interval_millis=1000,
    framework_profile_params=FrameworkProfile()
)
```

For more information about the FrameworkProfile class, see SageMaker Debugger APIs – FrameworkProfile in the Amazon SageMaker Python SDK.

**Start a training job with the default system monitoring and customized framework profiling for target steps or a target time range**

If you want to specify target steps or target time intervals to profile your training job, you need to specify parameters for the FrameworkProfile class. The following code examples show how to specify the target ranges for profiling along with system monitoring.

- **For a target step range**

  With the following example configuration, Debugger monitors the entire training job every 500 milliseconds (the default monitoring) and profiles a target step range from step 5 to step 15 (for 10 steps).

  ```python
  from sagemaker.debugger import ProfilerConfig, FrameworkProfile
  
  profiler_config=ProfilerConfig(
      system_monitor_interval_millis=1000,
      framework_profile_params=FrameworkProfile()
  )
  ```
With the following example configuration, Debugger monitors the entire training job every 1000 milliseconds and profiles a target step range from step 5 to step 15 (for 10 steps).

```python
from sagemaker.debugger import ProfilerConfig, FrameworkProfile
profiler_config=ProfilerConfig(
    system_monitor_interval_millis=1000,
    framework_profile_params=FrameworkProfile(start_step=5, num_steps=10))
```

- **For a target time range**

With the following example configuration, Debugger monitors the entire training job every 500 milliseconds (the default monitoring) and profiles a target time range from the current Unix time for 600 seconds.

```python
import time
from sagemaker.debugger import ProfilerConfig, FrameworkProfile
profiler_config=ProfilerConfig(
    framework_profile_params=FrameworkProfile(start_unix_time=int(time.time()),
    duration=600))
```

With the following example configuration, Debugger monitors the entire training job every 1000 milliseconds and profiles a target time range from the current Unix time for 600 seconds.

```python
import time
from sagemaker.debugger import ProfilerConfig, FrameworkProfile
profiler_config=ProfilerConfig(
    system_monitor_interval_millis=1000,
    framework_profile_params=FrameworkProfile(start_unix_time=int(time.time()),
    duration=600))
```

The framework profiling is performed for all of the profiling options at the target step or time range.
To find more information about available profiling options, see SageMaker Debugger APIs – FrameworkProfile in the Amazon SageMaker Python SDK.

The next section shows you how to script the available profiling options.

Start a training job with the default system monitoring and customized framework profiling with different profiling options

You can use the following profiling configuration classes to manage the framework profiling options:

- **DetailedProfilingConfig** – Specify a target step or time range to profile framework operations using the native framework profilers (TensorFlow profiler and PyTorch profiler). For example, if using TensorFlow, the Debugger hooks enable the TensorFlow profiler to collect TensorFlow-specific framework metrics. Detailed profiling enables you to profile all framework operators at a pre-step (before the first step), within steps, and between steps of a training job.

  **Note**
  Detailed profiling might significantly increase GPU memory consumption. We do not recommend enabling detailed profiling for more than a couple of steps.

- **DataloaderProfilingConfig** – Specify a target step or time range to profile deep learning framework data loader processes. Debugger collects every data loader event of the frameworks.

  **Note**
  Data loader profiling might lower the training performance while collecting information from data loaders. We don't recommend enabling data loader profiling for more than a couple of steps.
  Debugger is preconfigured to annotate data loader processes only for the AWS deep learning containers. Debugger cannot profile data loader processes from any other custom or external training containers.

- **PythonProfilingConfig** – Specify a target step or time range to profile Python functions. You can also choose between two Python profilers: cProfile and Pyinstrument.
• **cProfile** – The standard Python profiler. cProfile collects information for every Python operator called during training. With cProfile, Debugger saves cumulative time and annotation for each function call, providing complete detail about Python functions. In deep learning, for example, the most frequently called functions might be the convolutional filters and backward pass operators, and cProfile profiles every single of them. For the cProfile option, you can further select a timer option: total time, CPU time, and off-CPU time. While you can profile every function call executing on processors (both CPU and GPU) in CPU time, you can also identify I/O or network bottlenecks with the off-CPU time option. The default is total time, and Debugger profiles both CPU and off-CPU time. With cProfile, you are able to drill down to every single functions when analyzing the profile data.

• **Pyinstrument** – Pyinstrument is a low-overhead Python profiler that works based on sampling. With the Pyinstrument option, Debugger samples profiling events every millisecond. Because Pyinstrument measures elapsed wall-clock time instead of CPU time, the Pyinstrument option can be a better choice over the cProfile option for reducing profiling noise (filtering out irrelevant function calls that are cumulatively fast) and capturing operators that are actually compute intensive (cumulatively slow) for training your model. With Pyinstrument, you are able to see a tree of function calls and better understand the structure and root cause of the slowness.

**Note**

Enabling Python profiling might slow down the overall training time. cProfile profiles the most frequently called Python operators at every call, so the processing time on profiling increases with respect to the number of calls. For Pyinstrument, the cumulative profiling time increases with respect to time because of its sampling mechanism.

The following example configuration shows the full structure when you use the different profiling options with specified values.

```python
import time
from sagemaker.debugger import (ProfilerConfig,
                              FrameworkProfile,
                              DetailedProfilingConfig,
                              DataloaderProfilingConfig,
                              PythonProfilingConfig,
                              PythonProfiler, cProfileTimer)
```
import profiler_config=ProfilerConfig(
    system_monitor_interval_millis=500,
    framework_profile_params=FrameworkProfile(
        detailed_profiling_config=DetailedProfilingConfig(
            start_step=5,
            num_steps=1
        ),
        dataloader_profiling_config=DataloaderProfilingConfig(
            start_step=7,
            num_steps=1
        ),
        python_profiling_config=PythonProfilingConfig(
            start_step=9,
            num_steps=1,
            python_profiler=PythonProfiler.CPROFILE,
            cprofile_timer=cProfileTimer.TOTAL_TIME
        )
    )
)

For more information about available profiling options, see DetailedProfilingConfig, DataloaderProfilingConfig, and PythonProfilingConfig in the Amazon SageMaker Python SDK.

Updating Debugger system monitoring and framework profiling configuration while a training job is running

If you want to activate or update the Debugger monitoring configuration for a training job that is currently running, use the following SageMaker estimator extension methods:

- To activate Debugger system monitoring for a running training job and receive a Debugger profiling report, use the following:

  ```python
  estimator.enable_default_profiling()
  ```

  When you use the enable_default_profiling method, Debugger initiates the default system monitoring and the ProfileReport built-in rule, which generates a comprehensive profiling report at the end of the training job. This method can be called only if the current training job is running without both Debugger monitoring and profiling.

  For more information, see estimator.enable_default_profiling in the Amazon SageMaker Python SDK.
To update system monitoring configuration, use the following:

```python
estimator.update_profiler(
    system_monitor_interval_millis=500
)
```

For more information, see `estimator.update_profiler` in the Amazon SageMaker Python SDK.

### Turn off Debugger

If you want to completely turn off Debugger, do one of the following:

- Before starting a training job, do the following:
  
  To turn off profiling, include the `disable_profiler` parameter to your estimator and set it to True.

  ![Warning]
  
  If you disable it, you won't be able to view the comprehensive Studio Debugger insights dashboard and the autogenerated profiling report.

  
  To turn off debugging, set the `debugger_hook_config` parameter to False.

  ![Warning]
  
  If you disable it, you won't be able to collect output tensors and cannot debug your model parameters.

```python
estimator=Estimator(
    ...
    disable_profiler=True
    debugger_hook_config=False
)
```

For more information about the Debugger-specific parameters, see SageMaker Estimator in the Amazon SageMaker Python SDK.
While a training job is running, do the following:

To disable both monitoring and profiling while your training job is running, use the following estimator classmethod:

```python
estimator.disable_profiling()
```

To disable framework profiling only and keep system monitoring, use the `update_profiler` method:

```python
estimator.update_profiler(disable_framework_metrics=true)
```

For more information about the estimator extension methods, see the `estimator.disable_profiling` and `estimator.update_profiler` classmethods in the Amazon SageMaker Python SDK documentation.

### Configure built-in profiler rules managed by Amazon SageMaker Debugger

The Amazon SageMaker Debugger built-in profiler rules analyze system metrics and framework operations collected during the training of a model. Debugger offers the `ProfilerRule` API operation that helps configure the rules to monitor training compute resources and operations and to detect anomalies. For example, the profiling rules can help you detect whether there are computational problems such as CPU bottlenecks, excessive I/O wait time, imbalanced workload across GPU workers, and compute resource underutilization. To see a full list of available built-in profiling rules, see [List of Debugger built-in profiler rules](#).

#### Note

The built-in rules are provided through Amazon SageMaker processing containers and fully managed by SageMaker Debugger at no additional cost. For more information about billing, see the [Amazon SageMaker Pricing](#) page.

In the following topics, learn how to use the Debugger built-in rules.

### Topics

- Use SageMaker Debugger built-in profiler rules with their default parameter settings
Use Debugger built-in profiler rules with custom parameter values

Use SageMaker Debugger built-in profiler rules with their default parameter settings

To add SageMaker Debugger built-in rules in your estimator, you need to configure a `rules` list object. The following example code shows the basic structure of listing the SageMaker Debugger built-in rules.

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs

rules=[
    ProfilerRule.sagemaker(rule_configs.BuiltInProfilerRuleName_1()),
    ProfilerRule.sagemaker(rule_configs.BuiltInProfilerRuleName_2()),
    ...
    ProfilerRule.sagemaker(rule_configs.BuiltInProfilerRuleName_n()),
    ...
] # You can also append more debugging rules in the Rule.sagemaker(rule_configs.*()) format.

estimator=Estimator(
    ...
    rules=rules
)
```

For a complete list of available built-in rules, see [List of Debugger built-in profiler rules](#).

To use the profiling rules and inspect the computational performance and progress of your training job, add the `ProfilerReport` rule of SageMaker Debugger. This rule activates all built-in rules under the `ProfilerRule` `ProfilerRule` family. Furthermore, this rule generates an aggregated profiling report. For more information, see [Profiling Report Generated Using SageMaker Debugger](#). You can use the following code to add the profiling report rule to your training estimator.

```python
from sagemaker.debugger import Rule, rule_configs

rules=[
    ProfilerRule.sagemaker(rule_configs.ProfilerReport())
]
```

When you start the training job with the `ProfilerReport` rule, Debugger collects resource utilization data every 500 milliseconds. Debugger analyzes the resource utilization to identify
if your model is having bottleneck problems. If the rules detect training anomalies, the rule evaluation status changes to IssueFound. You can set up automated actions, such as notifying training issues and stopping training jobs using Amazon CloudWatch Events and AWS Lambda. For more information, see Action on Amazon SageMaker Debugger Rules.

Use Debugger built-in profiler rules with custom parameter values

If you want to adjust the built-in rule parameter values and customize tensor collection regex, configure the base_config and rule_parameters parameters for the ProfilerRule.sagemaker and Rule.sagemaker class methods. In case of the Rule.sagemaker class methods, you can also customize tensor collections through the collections_to_save parameter. For instruction on how to use the CollectionConfig class, see Configure Tensor Collections Using the CollectionConfig API.

Use the following configuration template for built-in rules to customize parameter values. By changing the rule parameters as you want, you can adjust the sensitivity of the rules to be initiated.

- The base_config argument is where you call the built-in rule methods.
- The rule_parameters argument is to adjust the default key values of the built-in rules listed in List of Debugger built-in profiler rules.

For more information about the Debugger rule class, methods, and parameters, see SageMaker Debugger Rule class in the Amazon SageMaker Python SDK.

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs, CollectionConfig

rules=[
    ProfilerRule.sagemaker(
        base_config=rule_configs.BuiltInProfilerRuleName(),
        rule_parameters={
            "key": "value"
        }
    )
]
```

The parameter descriptions and value customization examples are provided for each rule at List of Debugger built-in profiler rules.

For a low-level JSON configuration of the Debugger built-in rules using the CreateTrainingJob API, see Configure Debugger Using Amazon SageMaker API.
List of Debugger built-in profiler rules

Use the Debugger built-in profiler rules provided by Amazon SageMaker Debugger and analyze metrics collected while training your models. The Debugger built-in rules monitor various common conditions that are critical for the success of running a performant training job. You can call the built-in profiler rules using Amazon SageMaker Python SDK or the low-level SageMaker API operations. There's no additional cost for using the built-in rules. For more information about billing, see the Amazon SageMaker Pricing page.

**Note**

The maximum numbers of built-in profiler rules that you can attach to a training job is 20. SageMaker Debugger fully manages the built-in rules and analyzes your training job synchronously.

**Important**

To use the new Debugger features, you need to upgrade the SageMaker Python SDK and the SMDebug client library. In your iPython kernel, Jupyter notebook, or JupyterLab environment, run the following code to install the latest versions of the libraries and restart the kernel.

```python
import sys
import IPython
!{sys.executable} -m pip install -U sagemaker smdebug
IPython.Application.instance().kernel.do_shutdown(True)
```

**Profiler rules**

The following rules are the Debugger built-in rules that are callable using the ProfilerRule.sagemaker classmethod.
**Debugger built-in rule for generating the profiling report**

<table>
<thead>
<tr>
<th>Scope of Validity</th>
<th>Built-in Rules</th>
</tr>
</thead>
<tbody>
<tr>
<td>Profiling Report for any SageMaker training job</td>
<td>• <a href="#">ProfilerReport</a></td>
</tr>
</tbody>
</table>

**Debugger built-in rules for profiling hardware system resource utilization (system metrics)**

<table>
<thead>
<tr>
<th>Scope of Validity</th>
<th>Built-in Rules</th>
</tr>
</thead>
</table>
| Generic system monitoring rules for any SageMaker training job | • [BatchSize](#)  
• [CPUBottleneck](#)  
• [GPUMemoryIncrease](#)  
• [IOBottleneck](#)  
• [LoadBalancing](#)  
• [LowGPUUtilization](#)  
• [OverallSystemUsage](#) |

**Debugger built-in rules for profiling framework metrics**

<table>
<thead>
<tr>
<th>Scope of Validity</th>
<th>Built-in Rules</th>
</tr>
</thead>
</table>
| Profiling rules for deep learning frameworks (TensorFlow and PyTorch) | • [MaxInitializationTime](#)  
• [OverallFrameworkMetrics](#)  
• [StepOutlier](#) |

**Warning**

In favor of [Amazon SageMaker Profiler](#), SageMaker Debugger deprecates the framework profiling feature starting from TensorFlow 2.11 and PyTorch 2.0. You can still use the feature in the previous versions of the frameworks and SDKs as follows.

- SageMaker Python SDK <= v2.130.0
- PyTorch >= v1.6.0, < v2.0
• TensorFlow >= v2.3.1, < v2.11

See also March 16, 2023.

To use the built-in rules with default parameter values – use the following configuration format:

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs

rules = [
    ProfilerRule.sagemaker(rule_configs.BuiltInRuleName_1()),
    ProfilerRule.sagemaker(rule_configs.BuiltInRuleName_2()),
    ...
    ProfilerRule.sagemaker(rule_configs.BuiltInRuleName_n())
]
```

To use the built-in rules with customizing the parameter values – use the following configuration format:

```python
from sagemaker.debugger import Rule, ProfilerRule, rule_configs

rules = [
    ProfilerRule.sagemaker(
        base_config=rule_configs.BuiltInRuleName(),
        rule_parameters={
            "key": "value"
        }
    )
]
```

To find available keys for the rule_parameters parameter, see the parameter description tables.

Sample rule configuration codes are provided for each built-in rule below the parameter description tables.

• For a full instruction and examples of using the Debugger built-in rules, see Debugger Built-in Rules Example Code.

• For a full instruction on using the built-in rules with the low-level SageMaker API operations, see Configure Debugger Using Amazon SageMaker API.
ProfilerReport

The ProfilerReport rule invokes all of the built-in rules for monitoring and profiling. It creates a profiling report and updates when the individual rules are triggered. You can download a comprehensive profiling report while a training job is running or after the training job is complete. You can adjust the rule parameter values to customize sensitivity of the built-in monitoring and profiling rules. The following example code shows the basic format to adjust the built-in rule parameters through the ProfilerReport rule.

```python
rules=[
    ProfilerRule.sagemaker(
        rule_configs.ProfilerReport(
            <BuiltInRuleName>_<parameter_name> = value
        )
    )
]
```

If you trigger this ProfilerReport rule without any customized parameter as shown in the following example code, then the ProfilerReport rule triggers all of the built-in rules for monitoring and profiling with their default parameter values.

```python
rules=[ProfilerRule.sagemaker(rule_configs.ProfilerReport())]
```

The following example code shows how to specify and adjust the CPUBottleneck rule's cpu_threshold parameter and the IOBottleneck rule's threshold parameter.

```python
rules=[
    ProfilerRule.sagemaker(
        rule_configs.ProfilerReport(
            CPUBottleneck_cpu_threshold = 90,
            IOBottleneck_threshold = 90
        )
    )
]
```

To explore what's in the profiler report, see SageMaker Debugger Profiling Report. Also, because this rule activates all of the profiling rules, you can also check the rule analysis status using the SageMaker Debugger UI in SageMaker Studio Experiments.
## Parameter Descriptions for the OverallSystemUsage Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>&lt;BuiltInRuleName&gt;_&lt;parameter_name&gt;</td>
<td>Customizable parameter to adjust thresholds of other built-in monitoring and profiling rules.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Default value: None</td>
</tr>
</tbody>
</table>

### BatchSize

The BatchSize rule helps detect if GPU is underutilized due to a small batch size. To detect this issue, this rule monitors the average CPU utilization, GPU utilization, and GPU memory utilization. If utilization on CPU, GPU, and GPU memory is low on average, it may indicate that the training job can either run on a smaller instance type or can run with a bigger batch size. This analysis does not work for frameworks that heavily overallocate memory. However, increasing the batch size can lead to processing or data loading bottlenecks because more data preprocessing time is required in each iteration.

## Parameter Descriptions for the BatchSize Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>cpu_threshold_p95</td>
<td>Defines the threshold for 95th quantile of CPU utilization in percentage.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 70 (in percentage)</td>
</tr>
<tr>
<td>gpu_threshold_p95</td>
<td>Defines the threshold for 95th quantile of GPU utilization in percentage.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 70 (in percentage)</td>
</tr>
<tr>
<td>gpu_memory_threshold_p95</td>
<td>Defines the threshold for 95th quantile of GPU memory utilization in percentage.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 70 (in percentage)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| patience      | Defines the number of data points to skip until the rule starts evaluation. The first several steps of training jobs usually show high volume of data processes, so keep the rule patient and prevent it from being invoked too soon with a given number of profiling data that you specify with this parameter. **Optional**
|               | Valid values: Integer
|               | Default values: 100 |
| window        | Window size for computing quantiles. **Optional**
|               | Valid values: Integer
|               | Default values: 500 |
| scan_interval_us | Time interval that timeline files are scanned. **Optional**
|               | Valid values: Integer
|               | Default values: 60000000 (in microseconds) |

**CPUBottleneck**

The CPUBottleneck rule helps detect if GPU is underutilized due to CPU bottlenecks. Rule returns True if number of CPU bottlenecks exceeds a predefined threshold.
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>threshold</td>
<td>Defines the threshold for proportion of bottlenecked time to the total training time. If the proportion exceeds the percentage specified to the threshold parameter, the rule switches the rule status to True.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 50 (in percentage)</td>
</tr>
<tr>
<td>gpu_threshold</td>
<td>A threshold that defines low GPU utilization.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10 (in percentage)</td>
</tr>
<tr>
<td>cpu_threshold</td>
<td>A threshold that defines high CPU utilization.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 90 (in percentage)</td>
</tr>
<tr>
<td>patience</td>
<td>Defines the number of data points to skip until the rule starts evaluation. The first</td>
</tr>
</tbody>
</table>
### Parameter Name

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| several steps of training jobs usually show high volume of data processes, so keep the rule patient and prevent it from being invoked too soon with a given number of profiling data that you specify with this parameter. | Optional  
| Valid values: Integer  
| Default values: 100 | scan_interval_us  
| Time interval with which timeline files are scanned. | Optional  
| Valid values: Integer  
| Default values: 60000000 (in microseconds) | GPUMemoryIncrease  
| The GPUMemoryIncrease rule helps detect a large increase in memory usage on GPUs. | Parameter Descriptions for the GPUMemoryIncrease Rule  
| base_trial       | The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.  
<p>| Required | increase       | Defines the threshold for absolute memory increase. |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>patience</td>
<td>Defines the number of data points to skip until the rule starts evaluation. The first several steps of training jobs usually show high volume of data processes, so keep the rule patient and prevent it from being invoked too soon with a given number of profiling data that you specify with this parameter.</td>
</tr>
<tr>
<td>window</td>
<td>Window size for computing quantiles.</td>
</tr>
<tr>
<td>scan_interval_us</td>
<td>Time interval that timeline files are scanned.</td>
</tr>
</tbody>
</table>
**IOBottleneck**

This rule helps to detect if GPU is underutilized due to data IO bottlenecks. Rule returns True if number of IO bottlenecks exceeds a predefined threshold.

### Parameter Descriptions for the IOBottleneck Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>threshold</td>
<td>Defines the threshold when Rule to return True.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 50 (in percentage)</td>
</tr>
<tr>
<td>gpu_threshold</td>
<td>A threshold that defines when GPU is considered underutilized.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 70 (in percentage)</td>
</tr>
<tr>
<td>io_threshold</td>
<td>A threshold that defines high IO wait time.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 50 (in percentage)</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| patience       | Defines the number of data points to skip until the rule starts evaluation. The first several steps of training jobs usually show high volume of data processes, so keep the rule patient and prevent it from being invoked too soon with a given number of profiling data that you specify with this parameter. **Optional**
|                | Valid values: Integer |
|                | Default values: 1000 |
| scan_interval_us | Time interval that timeline files are scanned. **Optional**
|                | Valid values: Integer |
|                | Default values: 60000000 (in microseconds) |

**LoadBalancing**

The LoadBalancing rule helps detect issues in workload balancing among multiple GPUs.

**Parameter Descriptions for the LoadBalancing Rule**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| base_trial     | The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger. **Required**
<p>|                | Valid values: String |
| threshold      | Defines the workload percentage. |</p>
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>patience</td>
<td>Defines the number of data points to skip until the rule starts evaluation. The first several steps of training jobs usually show high volume of data processes, so keep the rule patient and prevent it from being invoked too soon with a given number of profiling data that you specify with this parameter.</td>
</tr>
<tr>
<td>scan_interval_us</td>
<td>Time interval that timeline files are scanned.</td>
</tr>
</tbody>
</table>

**LowGPUUtilization**

The LowGPUUtilization rule helps detect if GPU utilization is low or suffers from fluctuations. This is checked for each GPU on each worker. Rule returns True if 95th quantile is below threshold_p95 which indicates underutilization. Rule returns true if 95th quantile is above threshold_p95 and 5th quantile is below threshold_p5 which indicates fluctuations.
### Parameter Descriptions for the LowGPUUtilization Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| `base_trial`   | The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.  
*Required*  
Valid values: String |
| `threshold_p95` | A threshold for 95th quantile below which GPU is considered to be underutilized.  
*Optional*  
Valid values: Integer  
Default value: 70 (in percentage) |
| `threshold_p5`  | A threshold for 5th quantile. Default is 10 percent.  
*Optional*  
Valid values: Integer  
Default values: 10 (in percentage) |
| `patience`     | Defines the number of data points to skip until the rule starts evaluation. The first several steps of training jobs usually show high volume of data processes, so keep the rule patient and prevent it from being invoked too soon with a given number of profiling data that you specify with this parameter.  
*Optional*  
Valid values: Integer |
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>window</td>
<td>Window size for computing quantiles.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 500</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>scan_interval_us</td>
<td>Time interval that timeline files are scanned.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 60000000 (in microseconds)</td>
</tr>
</tbody>
</table>

**OverallSystemUsage**

The OverallSystemUsage rule measures overall system usage per worker node. The rule currently only aggregates values per node and computes their percentiles.

**Parameter Descriptions for the OverallSystemUsage Rule**

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>scan_interval_us</td>
<td>Time interval to scan timeline files.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
</tbody>
</table>
MaxInitializationTime

The MaxInitializationTime rule helps detect if the training initialization is taking too much time. The rule waits until the first step is available.

Parameter Descriptions for the MaxInitializationTime Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
</table>
| base_trial           | The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.  
Required                                                                 |
|                      | Valid values: String                                                        |
| threshold            | Defines the threshold in minutes to wait for the first step to become available.  
Optional                                                                 |
|                      | Valid values: Integer                                                       |
|                      | Default value: 20 (in minutes)                                              |
| scan_interval_us     | Time interval with which timeline files are scanned.  
Optional                                                                 |
|                      | Valid values: Integer                                                       |
|                      | Default values: 60000000 (in microseconds)                                  |
OverallFrameworkMetrics

The OverallFrameworkMetrics rule summarizes the time spent on framework metrics, such as forward and backward pass, and data loading.

Parameter Descriptions for the OverallFrameworkMetrics Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>scan_interval_us</td>
<td>Time interval to scan timeline files.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 60000000 (in microseconds)</td>
</tr>
</tbody>
</table>

StepOutlier

The StepOutlier rule helps detect outliers in step durations. This rule returns True if there are outliers with step durations larger than stddev sigmas of the entire step durations in a time range.

Parameter Descriptions for the StepOutlier Rule

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>base_trial</td>
<td>The base trial training job name. This parameter is automatically set to the current training job by Amazon SageMaker Debugger.</td>
</tr>
<tr>
<td></td>
<td><strong>Required</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: String</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------</td>
</tr>
<tr>
<td>stddev</td>
<td>Defines a factor by which to multiply the standard deviation. For example, the rule is invoked by default when a step duration is larger or smaller than 5 times the standard deviation.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5 (in minutes)</td>
</tr>
<tr>
<td>mode</td>
<td>Mode under which steps have been saved and on which Rule should run on. Per default rule will run on steps from EVAL and TRAIN phase</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 5 (in minutes)</td>
</tr>
<tr>
<td>n_outliers</td>
<td>How many outliers to ignore before rule returns True</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default value: 10</td>
</tr>
<tr>
<td>scan_interval_us</td>
<td>Time interval with which timeline files are scanned.</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
</tr>
<tr>
<td></td>
<td>Valid values: Integer</td>
</tr>
<tr>
<td></td>
<td>Default values: 60000000 (in microseconds)</td>
</tr>
</tbody>
</table>
Amazon SageMaker Debugger UI in Amazon SageMaker Studio Classic Experiments

Use the Amazon SageMaker Debugger Insights dashboard in Amazon SageMaker Studio Classic Experiments to analyze your model performance and system bottlenecks while running training jobs on Amazon Elastic Compute Cloud (Amazon EC2) instances. Gain insights into your training jobs and improve your model training performance and accuracy with the Debugger dashboards. By default, Debugger monitors system metrics (CPU, GPU, GPU memory, network, and data I/O) every 500 milliseconds and basic output tensors (loss and accuracy) every 500 iterations for training jobs. You can also further customize Debugger configuration parameter values and adjust the saving intervals through the Studio Classic UI or using the Amazon SageMaker Python SDK.

Important
If you're using an existing Studio Classic app, delete the app and restart to use the latest Studio Classic features. For instructions on how to restart and update your Studio Classic environment, see Update Amazon SageMaker Studio Classic.

Topics

- Open the Amazon SageMaker Debugger Insights dashboard
- Amazon SageMaker Debugger Insights dashboard controller
- Explore the Amazon SageMaker Debugger Insights dashboard
- Shut down the Amazon SageMaker Debugger Insights instance

Open the Amazon SageMaker Debugger Insights dashboard

In the SageMaker Debugger Insights dashboard in Studio Classic, you can see the compute resource utilization, resource utilization, and system bottleneck information of your training job that runs on Amazon EC2 instances in real time and after trainings.

Note
The SageMaker Debugger Insights dashboard runs a Studio Classic application on an ml.m5.4xlarge instance to process and render the visualizations. Each SageMaker Debugger Insights tab runs one Studio Classic kernel session. Multiple kernel sessions for
multiple SageMaker Debugger Insights tabs run on the single instance. When you close a SageMaker Debugger Insights tab, the corresponding kernel session is also closed. The Studio Classic application remains active and accrues charges for the ml.m5.4xlarge instance usage. For information about pricing, see the Amazon SageMaker Pricing page.

⚠️ Important

When you are done using the SageMaker Debugger Insights dashboard, you must shut down the ml.m5.4xlarge instance to avoid accruing charges. For instructions on how to shut down the instance, see Shut down the Amazon SageMaker Debugger Insights instance.

To open the SageMaker Debugger Insights dashboard

1. On the Studio Classic Home page, choose Experiments in the left navigation pane.
2. Search your training job in the Experiments page. If your training job is set up with an Experiments run, the job should appear in the Experiments tab; if you didn't set up an Experiments run, the job should appear in the Unassigned runs tab.
3. Choose (click) the link of the training job name to see the job details.
4. Under the OVERVIEW menu, choose Debugger. This should show the following two sections.
   - In the Debugger rules section, you can browse the status of the Debugger built-in rules associated with the training job.
   - In the Debugger insights section, you can find links to open SageMaker Debugger Insights on the dashboard.
5. In the SageMaker Debugger Insights section, choose the link of the training job name to open the SageMaker Debugger Insights dashboard. This opens a Debug [your-training-job-name] window. In this window, Debugger provides an overview of the computational performance of your training job on Amazon EC2 instances and helps you identify issues in compute resource utilization.

You can also download an aggregated profiling report by adding the built-in ProfilerReport rule of SageMaker Debugger. For more information, see Configure Built-in Profiler Rules and Profiling Report Generated Using SageMaker Debugger.
Amazon SageMaker Debugger Insights dashboard controller

There are different components of the Debugger controller for monitoring and profiling. In this guide, you learn about the Debugger controller components.

**Note**

The SageMaker Debugger Insights dashboard runs a Studio Classic app on an `ml.m5.4xlarge` instance to process and render the visualizations. Each SageMaker Debugger Insights tab runs one Studio Classic kernel session. Multiple kernel sessions for multiple SageMaker Debugger Insights tabs run on the single instance. When you close a SageMaker Debugger Insights tab, the corresponding kernel session is also closed. The Studio Classic app remains active and accrues charges for the `ml.m5.4xlarge` instance usage. For information about pricing, see the [Amazon SageMaker Pricing](#) page.

**Important**

When you are done using the SageMaker Debugger Insights dashboard, shut down the `ml.m5.4xlarge` instance to avoid accruing charges. For instructions on how to shut down the instance, see [Shut down the Amazon SageMaker Debugger Insights instance](#).

SageMaker Debugger Insights controller UI

Using the Debugger controller located at the upper-left corner of the Insights dashboard, you can refresh the dashboard, configure or update Debugger settings for monitoring system metrics, stop a training job, and download a Debugger profiling report.
• If you want to manually refresh the dashboard, choose the refresh button (the round arrow at the upper-left corner) as shown in the preceding screenshot.

• The **Monitoring** toggle button is on by default for any SageMaker training job initiated using the SageMaker Python SDK. If not activated, you can use the toggle button to start monitoring. During monitoring, Debugger only collects resource utilization metrics to detect computational problems such as CPU bottlenecks and GPU underutilization. For a complete list of resource utilization problems that Debugger monitors, see [Debugger built-in rules for profiling hardware system resource utilization (system metrics)](https://docs.aws.amazon.com/sagemaker/latest/dg/debugger-system-metrics.html).

• The **Configure monitoring** button opens a pop-up window that you can use to set or update the data collection frequency and the S3 path to save the data.

---

**Configure Debugger monitoring**

S3 bucket URI for Debugger output data
Set up the S3 bucket URI to save the Debugger monitoring and profiling output data.

Note: The S3 bucket URI must be in the same AWS region where your training job is running. AWS Region does not allow cross-region requests.

**S3 bucket URI**

```
s3://sagemaker-us-east-2-11112222333
```

**Collect monitoring data every**

- 500ms
- 100ms
- 200ms
- 500ms
- 1s
- 5s
- 1min
You can specify values for the following fields.

- **S3 bucket URI**: Specify the base S3 bucket URI.

- **Collect monitoring data every**: Select a time interval to collect system metrics. You can choose one of the monitoring intervals from the dropdown list. Available intervals are 100 milliseconds, 200 milliseconds, 500 milliseconds (default), 1 second, 5 seconds, and 1 minute.

  ⚠️ **Note**
  
  If you choose one of the lower time intervals, you increase the granularity of resource utilization metrics, so you can capture spikes and anomalies with a higher time resolution. However, higher the resolution, larger the size of system metrics to process. This might introduce additional overhead and impact the overall training and processing time.

- Using the **Stop training** button, you can stop the training job when you find anomalies in resource utilization.

- Using the **Download report** button, you can download an aggregated profiling report by using the built-in **ProfilerReport** rule of SageMaker Debugger. The button is activated when you add the built-in **ProfilerReport** rule to the estimator. For more information, see [Configure Built-in Profiler Rules](#) and [Profiling Report Generated Using SageMaker Debugger](#).

---

**Explore the Amazon SageMaker Debugger Insights dashboard**

When you initiate a SageMaker training job, SageMaker Debugger starts monitoring the resource utilization of the Amazon EC2 instances by default. You can track the system utilization rates, statistics overview, and built-in rule analysis through the Insights dashboard. This guide walks you through the content of the SageMaker Debugger Insights dashboard under the following tabs: **System Metrics** and **Rules**.

⚠️ **Note**

The SageMaker Debugger Insights dashboard runs a Studio Classic application on an **ml.m5.4xlarge** instance to process and render the visualizations. Each SageMaker Debugger Insights tab runs one Studio Classic kernel session. Multiple kernel sessions for multiple SageMaker Debugger Insights tabs run on the single instance. When you close a SageMaker Debugger Insights tab, the corresponding kernel session is also closed. The
Studio Classic application remains active and accrues charges for the \( \text{ml.m5.4xlarge} \) instance usage. For information about pricing, see the Amazon SageMaker Pricing page.

⚠️ **Important**

When you are done using the SageMaker Debugger Insights dashboard, shut down the \( \text{ml.m5.4xlarge} \) instance to avoid accruing charges. For instructions on how to shut down the instance, see Shut down the Amazon SageMaker Debugger Insights instance.

⚠️ **Important**

In the reports, plots and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

**Topics**

- **System metrics**
- **Rules**

**System metrics**

In the **System Metrics** tab, you can use the summary table and timeseries plots to understand resource utilization.

**Resource utilization summary**

This summary table shows the statistics of compute resource utilization metrics of all nodes (denoted as algo-\( n \)). The resource utilization metrics include the total CPU utilization, the total GPU utilization, the total CPU memory utilization, the total GPU memory utilization, the total I/O wait time, and the total network in bytes. The table shows the minimum and the maximum values, and p99, p90, and p50 percentiles.
Resource utilization time series plots

Use the time series graphs to see more details of resource utilization and identify at what time interval each instance shows any undesired utilization rate, such as low GPU utilization and CPU bottlenecks that can cause a waste of the expensive instance.

The time series graph controller UI

The following screenshot shows the UI controller for adjusting the time series graphs.

- **algo-1**: Use this dropdown menu to choose the node that you want to look into.
- **Zoom In**: Use this button to zoom in the time series graphs and view shorter time intervals.
- **Zoom Out**: Use this button to zoom out the time series graphs and view wider time intervals.
- **Pan Left**: Move the time series graphs to an earlier time interval.
- **Pan Right**: Move the time series graphs to a later time interval.
- **Fix Timeframe**: Use this check box to fix or bring back the time series graphs to show the whole view from the first data point to the last data point.

**CPU utilization and I/O wait time**

The first two graphs show CPU utilization and I/O wait time over time. By default, the graphs show the average of CPU utilization rate and I/O wait time spent on the CPU cores. You can select one or more CPU cores by selecting the labels to graph them on single chart and compare utilization across cores. You can drag and zoom in and out to have a closer look at specific time intervals.

**GPU utilization and GPU memory utilization**

The following graphs show GPU utilization and GPU memory utilization over time. By default, the graphs show the mean utilization rate over time. You can select the GPU core labels to see the utilization rate of each core. Taking the mean of utilization rate over the total number of GPU cores shows the mean utilization of the entire hardware system resource. By looking at the mean utilization rate, you can check the overall system resource usage of an Amazon EC2 instance. The
following figure shows an example training job on an ml.p3.16xlarge instance with 8 GPU cores. You can monitor if the training job is well distributed, fully utilizing all GPUs.

![GPU utilization over time](image)

**Overall system utilization over time**

The following heatmap shows an example of the entire system utilization of an ml.p3.16xlarge instance over time, projected onto the two-dimensional plot. Every CPU and GPU core is listed in the vertical axis, and the utilization is recorded over time with a color scheme, where the bright colors represent low utilization and the darker colors represent high utilization. See the labeled color bar on the right side of the plot to find out which color level corresponds to which utilization rate.
Use the Rules tab to find a summary of the profiling rule analysis on your training job. If the profiling rule is activated with the training job, the text appears highlighted with the solid white text. Inactive rules are dimmed in gray text. To activate these rules, follow instructions at the section called “Configure built-in profiler rules”.

Rules
### Insights

The following list shows a summary of Debugger rule analysis on your training job. Expand the following rule items to find suggestions and additional details, such as the number of times each rule triggered, the rule parameters, and the default threshold values to evaluate your training job performance.

Showing 8 suggestions

- **BatchSize - Issue Found**
- **LowGPUtilization - Issue Found**
  - Check for bottlenecks, minimize blocking calls, change distributed training strategy, increase batch-size.
  - Number of times the rule triggered: 14
  - Number of violations: 14
  - Number of datapoints: 1797
  - Rule parameters:
    - threshold_p95: 70%
    - threshold_p5: 10%
    - window: 500
    - patience: 1000
  - For more information, see the [LowGPUtilization](#) rule description.

- **CPUBottleneck - No Issue Found**
- **IOBottleneck - No Issue Found**
- **GPUMemoryIncrease - No Issue Found**
- **StepOutlier - No Issue Found**
- **MaxInitializationTime - No Issue Found**
- **LoadBalancing - No Issue Found**

---

**Shut down the Amazon SageMaker Debugger Insights instance**

When you are not using the SageMaker Debugger Insights dashboard, you should shut down the app instance to avoid incurring additional fees.

**To shut down the SageMaker Debugger Insights app instance in Studio Classic**
1. In Studio Classic, select the Running Instances and Kernels icon ( ).

2. Under the RUNNING APPS list, look for the sagemaker-debugger-1.0 app. Select the shutdown icon ( ) next to the app. The SageMaker Debugger Insights dashboards run on an ml.m5.4xlarge instance. This instance also disappears from the RUNNING INSTANCES when you shut down the sagemaker-debugger-1.0 app.

SageMaker Debugger interactive report

Receive profiling reports autogenerated by Debugger. The Debugger report provide insights into your training jobs and suggest recommendations to improve your model performance. The following screenshot shows a collage of the Debugger profiling report. To learn more, see SageMaker Debugger profiling report.

Note

You can download a Debugger reports while your training job is running or after the job has finished. During training, Debugger concurrently updates the report reflecting the
current rules' evaluation status. You can download a complete Debugger report only after the training job has completed.

Important

In the reports, plots and and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

SageMaker Debugger profiling report

For any SageMaker training jobs, the SageMaker Debugger ProfilerReport rule invokes all of the monitoring and profiling rules and aggregates the rule analysis into a comprehensive report.

Following this guide, download the report using the Amazon SageMaker Python SDK or the S3 console, and learn what you can interpret from the profiling results.
Important

In the report, plots and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

Download the SageMaker Debugger profiling report

Download the SageMaker Debugger profiling report while your training job is running or after the job has finished using the Amazon SageMaker Python SDK and AWS Command Line Interface (CLI).

Note

To get the profiling report generated by SageMaker Debugger, you must use the built-in ProfilerReport rule offered by SageMaker Debugger. To activate the rule with your training job, see Configure Built-in Profiler Rules.

Tip

You can also download the report with a single click in the SageMaker Studio Debugger insights dashboard. This doesn't require any additional scripting to download the report. To find out how to download the report from Studio, see Open the Amazon SageMaker Debugger Insights dashboard.

Download using SageMaker Python SDK and AWS CLI

1. Check the current job’s default S3 output base URI.

   ```python
   estimator.output_path
   ```

2. Check the current job name.

   ```python
   estimator.latest_training_job.job_name
   ```

3. The Debugger profiling report is stored under `<default-s3-output-base-uri>/ <training-job-name>/rule-output`. Configure the rule output path as follows:
rule_output_path = estimator.output_path + estimator.latest_training_job.job_name + "/rule-output"

4. To check if the report is generated, list directories and files recursively under the rule_output_path using `aws s3 ls` with the `--recursive` option.

```bash
! aws s3 ls {rule_output_path} --recursive
```

This should return a complete list of files under an autogenerated folder that’s named as ProfilerReport-1234567890. The folder name is a combination of strings: ProfilerReport and a unique 10-digit tag based on the Unix timestamp when the ProfilerReport rule is initiated.

The `profiler-report.html` is an autogenerated profiling report by Debugger. The remaining files are the built-in rule analysis components stored in JSON and a Jupyter notebook that are used to aggregate them into the report.

5. Download the files recursively using `aws s3 cp`. The following command saves all of the rule output files to the ProfilerReport-1234567890 folder under the current working directory.

```bash
! aws s3 cp {rule_output_path} ./ --recursive
```

Tip

If using a Jupyter notebook server, run `!pwd` to double check the current working directory.

6. Under the /ProfilerReport-1234567890/profiler-output directory, open `profiler-report.html`. If using JupyterLab, choose Trust HTML to see the autogenerated Debugger profiling report.
7. Open the `profiler-report.ipynb` file to explore how the report is generated. You can also customize and extend the profiling report using the Jupyter notebook file.

Download using Amazon S3 Console

1. Sign in to the AWS Management Console and open the Amazon S3 console at https://console.aws.amazon.com/s3/.

2. Search for the base S3 bucket. For example, if you haven't specified any base job name, the base S3 bucket name should be in the following format: `sagemaker-<region>-111122223333`. Look up the base S3 bucket through the *Find bucket by name* field.

3. In the base S3 bucket, look up the training job name by specifying your job name prefix into the *Find objects by prefix* input field. Choose the training job name.
4. In the training job's S3 bucket, there must be three subfolders for training data collected by Debugger: `debug-output/`, `profiler-output/`, and `rule-output/`. Choose `rule-output/`.

5. In the `rule-output/` folder, choose `ProfilerReport-1234567890`, and choose `profiler-output/` folder. The `profiler-output/` folder contains `profiler-report.html` (the autogenerated profiling report in html), `profiler-report.ipynb` (a Jupyter notebook with scripts that are used for generating the report), and a `profiler-report/` folder (contains rule analysis JSON files that are used as components of the report).

6. Select the `profiler-report.html` file, choose `Actions`, and `Download`.
Monitor AWS compute resource utilization in SageMaker Studio Classic

- Open
- Calculate total size
- Copy
- Move
- Initiate restore
- Query with S3 Select
- **Download actions**
  - Download
  - Download as
- Edit actions
  - Rename object
  - Edit storage class
  - Edit server-side encryption
- Edit metadata

Folder overview

Region
US East (Ohio) us-east-2

Objects (3)
Objects are the fundamental unit of data in Amazon S3.

<table>
<thead>
<tr>
<th>Name</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>profiler-report.html</td>
<td>html</td>
</tr>
<tr>
<td>profiler-report.ipynb</td>
<td>ipynb</td>
</tr>
<tr>
<td>profiler-reports/</td>
<td>Folder</td>
</tr>
</tbody>
</table>
7. Open the downloaded `profiler-report.html` file in a web browser.

**Note**

If you started your training job without configuring the Debugger-specific parameters, Debugger generates the report based only on the system monitoring rules because the Debugger parameters are not configured to save framework metrics. To enable framework metrics profiling and receive an extended Debugger profiling report, configure the `profiler_config` parameter when constructing or updating SageMaker estimators. To learn how to configure the `profiler_config` parameter before starting a training job, see [Configure for framework profiling](#).

To update the current training job and enable framework metrics profiling, see [Update Debugger Framework Profiling Configuration](#).

**Debugger profiling report walkthrough**

This section walks you through the Debugger profiling report section by section. The profiling report is generated based on the built-in rules for monitoring and profiling. The report shows result plots only for the rules that found issues.

**Important**

In the report, plots and and recommendations are provided for informational purposes and are not definitive. You are responsible for making your own independent assessment of the information.

**Topics**

- Training job summary
- System usage statistics
- Framework metrics summary
- Rules summary
- Analyzing the training loop – step durations
- GPU utilization analysis
Training job summary

At the beginning of the report, Debugger provides a summary of your training job. In this section, you can overview the time durations and timestamps at different training phases.

The following table gives a summary about the training job. The table includes information about when the training job started and ended, how much time initialization, training loop and finalization took. Your training job started on 11/29/2020 at 23:12:42 and ran for 737 seconds.

<table>
<thead>
<tr>
<th>#</th>
<th>Job Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Start time 23:12:42 11/29/2020</td>
</tr>
<tr>
<td>1</td>
<td>End time 23:24:59 11/29/2020</td>
</tr>
<tr>
<td>2</td>
<td>Job duration 737 seconds</td>
</tr>
<tr>
<td>3</td>
<td>Training loop start 23:17:31 11/29/2020</td>
</tr>
<tr>
<td>4</td>
<td>Training loop end 23:24:59 11/29/2020</td>
</tr>
<tr>
<td>5</td>
<td>Training loop duration 448 seconds</td>
</tr>
<tr>
<td>6</td>
<td>Initialization time 288 seconds</td>
</tr>
<tr>
<td>7</td>
<td>Finalization time 0 seconds</td>
</tr>
<tr>
<td>8</td>
<td>Initialization 39 %</td>
</tr>
<tr>
<td>9</td>
<td>Training loop 60 %</td>
</tr>
<tr>
<td>10</td>
<td>Finalization 0 %</td>
</tr>
</tbody>
</table>

The summary table contains the following information:

- **start_time** – The exact time when the training job started.
- **end_time** – The exact time when the training job finished.
- **job_duration_in_seconds** – The total training time from the **start_time** to the **end_time**.
- **training_loop_start** – The exact time when the first step of the first epoch has started.
- **training_loop_end** – The exact time when the last step of the last epoch has finished.
- **training_loop_duration_in_seconds** – The total time between the training loop start time and the training loop end time.
• **initialization_in_seconds** – Time spent on initializing the training job. The initialization phase covers the period from the `start_time` to the `training_loop_start` time. The initialization time is spent on compiling the training script, starting the training script, creating and initializing the model, initiating EC2 instances, and downloading training data.

• **finalization_in_seconds** – Time spent on finalizing the training job, such as finishing the model training, updating the model artifacts, and closing the EC2 instances. The finalization phase covers the period from the `training_loop_end` time to the `end_time`.

• **initialization (%)** – The percentage of time spent on **initialization** over the total **job_duration_in_seconds**.

• **training loop (%)** – The percentage of time spent on **training loop** over the total **job_duration_in_seconds**.

• **finalization (%)** – The percentage of time spent on **finalization** over the total **job_duration_in_seconds**.

**System usage statistics**

In this section, you can see an overview of system utilization statistics.

**System usage statistics**

The 95th percentile of the total GPU utilization on node algo-2 is 74%. GPUs on node algo-2 are well utilized.

The following table shows usage statistics per worker node such as total CPU and GPU utilization, total CPU and memory footprint. The table also include total IO wait time and total sent/received bytes. The table shows min and max values as well as p99, p90 and p50 percentiles.

<table>
<thead>
<tr>
<th>node</th>
<th>metric</th>
<th>unit</th>
<th>max</th>
<th>p99</th>
<th>p95</th>
<th>p50</th>
<th>min</th>
</tr>
</thead>
<tbody>
<tr>
<td>algo-1</td>
<td>Network</td>
<td>bytes</td>
<td>218817581.57</td>
<td>168.02</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-1</td>
<td>I/O</td>
<td>percentage</td>
<td>13.2653125</td>
<td>5.928312500000000</td>
<td>0.1955937499999999</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-1</td>
<td>GPU memory</td>
<td>percentage</td>
<td>32.25</td>
<td>26.25</td>
<td>21</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-1</td>
<td>CPU memory</td>
<td>percentage</td>
<td>75</td>
<td>74.5</td>
<td>74.25</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-1</td>
<td>CPU</td>
<td>percentage</td>
<td>5.05</td>
<td>5.01</td>
<td>4.98</td>
<td>2.17</td>
<td>0.55</td>
</tr>
<tr>
<td>algo-1</td>
<td>CPU</td>
<td>percentage</td>
<td>32.955625</td>
<td>22.629131250000000</td>
<td>17.034</td>
<td>3.702499999999999</td>
<td>0</td>
</tr>
<tr>
<td>algo-2</td>
<td>Network</td>
<td>bytes</td>
<td>4135.24</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-2</td>
<td>I/O</td>
<td>percentage</td>
<td>20.1875</td>
<td>8.155250000000000</td>
<td>1.747812499999999</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-2</td>
<td>GPU memory</td>
<td>percentage</td>
<td>38</td>
<td>31.75</td>
<td>21.75</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-2</td>
<td>GPU</td>
<td>percentage</td>
<td>75</td>
<td>74.5</td>
<td>74.25</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>algo-2</td>
<td>CPU memory</td>
<td>percentage</td>
<td>5.05</td>
<td>5.02</td>
<td>4.99</td>
<td>2.17</td>
<td>0.55</td>
</tr>
<tr>
<td>algo-2</td>
<td>CPU</td>
<td>percentage</td>
<td>35.00437499999999</td>
<td>25.699968750000000</td>
<td>18.3342998875</td>
<td>3.77828125</td>
<td>0</td>
</tr>
</tbody>
</table>

The Debugger profiling report includes the following information:
- **node** – Lists the name of nodes. If using distributed training on multi nodes (multiple EC2 instances), the node names are in format of algo-n.

- **metric** – The system metrics collected by Debugger: CPU, GPU, CPU memory, GPU memory, I/O, and Network metrics.

- **unit** – The unit of the system metrics.

- **max** – The maximum value of each system metric.

- **p99** – The 99th percentile of each system utilization.

- **p95** – The 95th percentile of each system utilization.

- **p50** – The 50th percentile (median) of each system utilization.

- **min** – The minimum value of each system metric.

**Framework metrics summary**

In this section, the following pie charts show the breakdown of framework operations on CPUs and GPUs.
Each of the pie charts analyzes the collected framework metrics in various aspects as follows:

- **Ratio between TRAIN/EVAL phase and others** – Shows the ratio between time durations spent on different training phases.
- **Ratio between forward and backward pass** – Shows the ratio between time durations spent on forward and backward pass in the training loop.
- **Ratio between CPU/GPU operators** – Shows the ratio between time spent on operators running on CPU or GPU, such as convolutional operators.
- **General metrics recorded in framework** – Shows the ratio between time spent on major framework metrics, such as data loading, forward and backward pass.

**Overview: CPU Operators**

This section provides information of the CPU operators in detail. The table shows the percentage of the time and the absolute cumulative time spent on the most frequently called CPU operators.
Overview: GPU operators

This section provides information of the GPU operators in detail. The table shows the percentage of the time and the absolute cumulative time spent on the most frequently called GPU operators.

<table>
<thead>
<tr>
<th>#</th>
<th>Percentage</th>
<th>Cumulative time</th>
<th>GPU operator</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>31.17</td>
<td>8013464</td>
<td>CudnnConvolutionBackward</td>
</tr>
<tr>
<td>1</td>
<td>27.41</td>
<td>578830</td>
<td>cudnn_convolution_backward</td>
</tr>
<tr>
<td>2</td>
<td>12.8</td>
<td>2430857</td>
<td>add_</td>
</tr>
<tr>
<td>3</td>
<td>11.84</td>
<td>2284879</td>
<td>torch.autograd._accumulateGrad</td>
</tr>
<tr>
<td>4</td>
<td>4.91</td>
<td>948134</td>
<td>CudnnBatchNormBackward</td>
</tr>
<tr>
<td>5</td>
<td>4.44</td>
<td>737918</td>
<td>add</td>
</tr>
<tr>
<td>6</td>
<td>3.18</td>
<td>164127</td>
<td>mul_</td>
</tr>
<tr>
<td>7</td>
<td>2.45</td>
<td>473492</td>
<td>conv2d</td>
</tr>
<tr>
<td>8</td>
<td>2.28</td>
<td>440157</td>
<td>convolution</td>
</tr>
</tbody>
</table>

Overview: GPU operators

The following table shows a list of operators that your training job run on GPU. The most expensive operator on GPU was "CudnnConvolutionBackward" with 34%.

<table>
<thead>
<tr>
<th>#</th>
<th>Percentage</th>
<th>Cumulative time</th>
<th>GPU operator</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>34.46</td>
<td>13866569</td>
<td>CudnnConvolutionBackward</td>
</tr>
<tr>
<td>1</td>
<td>34.44</td>
<td>13867210</td>
<td>cudnn_convolution_backward</td>
</tr>
<tr>
<td>2</td>
<td>8.16</td>
<td>2462529</td>
<td>conv2d</td>
</tr>
<tr>
<td>3</td>
<td>6.13</td>
<td>2473099</td>
<td>convolution</td>
</tr>
<tr>
<td>4</td>
<td>6.11</td>
<td>2463505</td>
<td>_<em>convolution</em></td>
</tr>
<tr>
<td>5</td>
<td>5.05</td>
<td>2444423</td>
<td>cudnn_convolution</td>
</tr>
<tr>
<td>6</td>
<td>3.34</td>
<td>1348774</td>
<td>CudnnBatchNormBackward</td>
</tr>
<tr>
<td>7</td>
<td>3.3</td>
<td>1330055</td>
<td>cudnn_batch_norm_backward</td>
</tr>
</tbody>
</table>

Rules summary

In this section, Debugger aggregates all of the rule evaluation results, analysis, rule descriptions, and suggestions.
Analyzing the training loop – step durations

In this section, you can find a detailed statistics of step durations on each GPU core of each node. Debugger evaluates mean, maximum, p99, p95, p50, and minimum values of step durations, and evaluate step outliers. The following histogram shows the step durations captured on different worker nodes and GPUs. You can enable or disable the histogram of each worker by choosing the legends on the right side. You can check if there is a particular GPU that's causing step duration outliers.
GPU utilization analysis

This section shows the detailed statistics about GPU core utilization based on LowGPUUtilization rule. It also summarizes the GPU utilization statistics, mean, p95, and p5 to determine if the training job is underutilizing GPUs.

Batch size

This section shows the detailed statistics of total CPU utilization, individual GPU utilizations, and GPU memory footprints. The BatchSize rule determines if you need to change the batch size to better utilize the GPUs. You can check whether the batch size is too small resulting in underutilization or too large causing overutilization and out of memory issues. In the plot, the boxes show the p25 and p75 percentile ranges (filled with dark purple and bright yellow respectively) from the median (p50), and the error bars show the 5th percentile for the lower bound and 95th percentile for the upper bound.
In this section, you can drill down into the CPU bottlenecks that the CPUBottleneck rule detected from your training job. The rule checks if the CPU utilization is above `cpu_threshold` (90% by default) and also if the GPU utilization is below `gpu_threshold` (10% by default).

**CPU bottlenecks**

In this section, you can drill down into the CPU bottlenecks that the CPUBottleneck rule detected from your training job. The rule checks if the CPU utilization is above `cpu_threshold` (90% by default) and also if the GPU utilization is below `gpu_threshold` (10% by default).
The pie charts show the following information:

- **Low GPU usage caused by CPU bottlenecks** – Shows the ratio of data points between the ones with GPU utilization above and below the threshold and the ones that matches the CPU bottleneck criteria.

- **Ratio between TRAIN/EVAL phase and others** – Shows the ratio between time durations spent on different training phases.

- **Ratio between forward and backward pass** – Shows the ratio between time durations spent on forward and backward pass in the training loop.

- **Ratio between CPU/GPU operators** – Shows the ratio between time durations spent on GPUs and CPUs by Python operators, such as data loader processes and forward and backward pass operators.

- **General metrics recorded in framework** – Shows major framework metrics and the ratio between time durations spent on the metrics.

**I/O bottlenecks**

In this section, you can find a summary of I/O bottlenecks. The rule evaluates the I/O wait time and GPU utilization rates and monitors if the time spent on the I/O requests exceeds a threshold percent of the total training time. It might indicate I/O bottlenecks where GPUs are waiting for data to arrive from storage.
Load balancing in multi-GPU training

In this section, you can identify workload balancing issue across GPUs.

![Workloads on node algo-1](image)

GPU memory analysis

In this section, you can analyze the GPU memory utilization collected by the GPUMemoryIncrease rule. In the plot, the boxes show the p25 and p75 percentile ranges (filled with dark purple and bright yellow respectively) from the median (p50), and the error bars show the 5th percentile for the lower bound and 95th percentile for the upper bound.

![Node algo-1](image)
Analyze data using the Debugger Python client library

While your training job is running or after it has completed, you can access the training data collected by Debugger using the Amazon SageMaker Python SDK and the SMDebug client library. The Debugger Python client library provides analysis and visualization tools that enable you to drill down into your training job data.

To install the library and use its analysis tools (in a JupyterLab notebook or an iPython kernel)

```
! pip install -U smdebug
```

The following topics walk you through how to use the Debugger Python tools to visualize and analyze the training data collected by Debugger.

Analyze system and framework metrics

- Access the profile data
- Plot the system metrics and framework metrics data
- Access the profiling data using the pandas data parsing tool
- Access the Python profiling stats data
- Merge timelines of multiple profile trace files
- Profiling data loaders

Access the profile data

The SMDebug TrainingJob class reads data from the S3 bucket where the system and framework metrics are saved.

To set up a TrainingJob object and retrieve profiling event files of a training job

```python
from smdebug.profiler.analysis.notebook_utils.training_job import TrainingJob
tj = TrainingJob(training_job_name, region)
```

Tip

You need to specify the training_job_name and region parameters to log to a training job. There are two ways to specify the training job information:
• Use the SageMaker Python SDK while the estimator is still attached to the training job.

```python
import sagemaker
training_job_name=estimator.latest_training_job.job_name
region=sagemaker.Session().boto_region_name
```

• Pass strings directly.

```python
training_job_name="your-training-job-name-YYYY-MM-DD-HH-MM-SS-SSS"
region="us-west-2"
```

**Note**

By default, SageMaker Debugger collects system metrics to monitor hardware resource utilization and system bottlenecks. Running the following functions, you might receive error messages regarding unavailability of framework metrics. To retrieve framework profiling data and gain insights into framework operations, you must enable framework profiling.

• If you use SageMaker Python SDK to manipulate your training job request, pass the `framework_profile_params` to the `profiler_config` argument of your estimator. To learn more, see [Configure SageMaker Debugger Framework Profiling](#).

• If you use Studio Classic, turn on profiling using the Profiling toggle button in the Debugger insights dashboard. To learn more, see [SageMaker Debugger Insights Dashboard Controller](#).

To retrieve a description of the training job description and the S3 bucket URI where the metric data are saved

```python
j.describe_training_job()
j.get_config_and_profiler_s3_output_path()
```

To check if the system and framework metrics are available from the S3 URI

```python
j.wait_for_sys_profiling_data_to_be_available()
```
To create system and framework reader objects after the metric data become available

```python
system_metrics_reader = tj.get_systems_metrics_reader()
framework_metrics_reader = tj.get_framework_metrics_reader()
```

To refresh and retrieve the latest training event files

The reader objects have an extended method, `refresh_event_file_list()`, to retrieve the latest training event files.

```python
system_metrics_reader.refresh_event_file_list()
framework_metrics_reader.refresh_event_file_list()
```

Plot the system metrics and framework metrics data

You can use the system and algorithm metrics objects for the following visualization classes to plot timeline graphs and histograms.

**Note**

To visualize the data with narrowed-down metrics in the following visualization object plot methods, specify `select_dimensions` and `select_events` parameters. For example, if you specify `select_dimensions=['GPU']`, the plot methods filter the metrics that include the "GPU" keyword. If you specify `select_events=['total']`, the plot methods filter the metrics that include the "total" event tags at the end of the metric names. If you enable these parameters and give the keyword strings, the visualization classes return the charts with filtered metrics.

- The `MetricsHistogram` class

```python
from smdebug.profiler.analysis.notebook_utils.metrics_histogram import MetricsHistogram

metrics_histogram = MetricsHistogram(system_metrics_reader)
metrics_histogram.plot(
    starttime=0,
```
Monitor AWS compute resource utilization in SageMaker Studio Classic

```python
datetime = system_metrics_reader.get_timestamp_of_latest_available_file(),
select_dimensions=['CPU', 'GPU', 'I/O'], # optional
select_events=['total'] # optional
)

• The StepTimelineChart class

```python
from smdebug.profiler.analysis.notebook_utils.step_timeline_chart import StepTimelineChart

view_step_timeline_chart = StepTimelineChart(framework_metrics_reader)

• The StepHistogram class

```python
from smdebug.profiler.analysis.notebook_utils.step_histogram import StepHistogram

step_histogram = StepHistogram(framework_metrics_reader)
step_histogram.plot(
    start_time=step_histogram.last_timestamp - 5 * 1000 * 1000,
    end_time=step_histogram.last_timestamp,
    show_workers=True
)

• The TimelineCharts class

```python
from smdebug.profiler.analysis.notebook_utils.timeline_charts import TimelineCharts

view_timeline_charts = TimelineCharts(
    system_metrics_reader,
    framework_metrics_reader,
    select_dimensions=['CPU', 'GPU', 'I/O'], # optional
    select_events=['total'] # optional
)

view_timeline_charts.plot_detailed_profiler_data([700,710])

• The Heatmap class

```python
from smdebug.profiler.analysis.notebook_utils.heatmap import Heatmap

view_heatmap = Heatmap(
    system_metrics_reader,
    framework_metrics_reader,
)```
Access the profiling data using the pandas data parsing tool

The following PandasFrame class provides tools to convert the collected profiling data to Pandas data frame.

```python
from smdebug.profiler.analysis.utils.profiler_data_to_pandas import PandasFrame
```

The PandasFrame class takes the `tj` object's S3 bucket output path, and its methods `get_all_system_metrics()` and `get_all_framework_metrics()` return system metrics and framework metrics in the Pandas data format.

```python
pf = PandasFrame(tj.profiler_s3_output_path)
system_metrics_df = pf.get_all_system_metrics()
framework_metrics_df = pf.get_all_framework_metrics(
    selected_framework_metrics=[
        'Step:ModeKeys.TRAIN',
        'Step:ModeKeys.GLOBAL'
    ]
)
```

Access the Python profiling stats data

The Python profiling provides framework metrics related to Python functions and operators in your training scripts and the SageMaker deep learning frameworks.

Training Modes and Phases for Python Profiling

To profile specific intervals during training to partition statistics for each of these intervals, Debugger provides tools to set modes and phases.

For training modes, use the following `PythonProfileModes` class:

```python
from smdebug.profiler.python_profile_utils import PythonProfileModes
```

This class provides the following options:
• **PythonProfileModes.TRAIN** – Use if you want to profile the target steps in the training phase. This mode option available only for TensorFlow.

• **PythonProfileModes.EVAL** – Use if you want to profile the target steps in the evaluation phase. This mode option available only for TensorFlow.

• **PythonProfileModes.PREDICT** – Use if you want to profile the target steps in the prediction phase. This mode option available only for TensorFlow.

• **PythonProfileModes.GLOBAL** – Use if you want to profile the target steps in the global phase, which includes the previous three phases. This mode option available only for PyTorch.

• **PythonProfileModes.PRE_STEP_ZERO** – Use if you want to profile the target steps in the initialization stage before the first training step of the first epoch starts. This phase includes the initial job submission, uploading the training scripts to EC2 instances, preparing the EC2 instances, and downloading input data. This mode option available for both TensorFlow and PyTorch.

• **PythonProfileModes.POST_HOOK_CLOSE** – Use if you want to profile the target steps in the finalization stage after the training job has done and the Debugger hook is closed. This phase includes profiling data while the training jobs are finalized and completed. This mode option available for both TensorFlow and PyTorch.

For training phases, use the following StepPhase class:

```python
from smdebug.profiler.analysis.utils.python_profile_analysis_utils import StepPhase
```

This class provides the following options:

• **StepPhase.START** – Use to specify the start point of the initialization phase.

• **StepPhase.STEP_START** – Use to specify the start step of the training phase.

• **StepPhase.FORWARD_PASS_END** – Use to specify the steps where the forward pass ends. This option is available only for PyTorch.

• **StepPhase.STEP_END** – Use to specify the end steps in the training phase. This option is available only for TensorFlow.

• **StepPhase.END** – Use to specify the ending point of the finalization (post-hook-close) phase. If the callback hook is not closed, the finalization phase profiling does not occur.

**Python Profiling Analysis Tools**

Monitor AWS compute resource utilization in SageMaker Studio Classic
Debugger supports the Python profiling with two profiling tools:

- cProfile – The standard python profiler. cProfile collects framework metrics on CPU time for every function called when profiling was enabled.
- Pyinstrument – This is a low overhead Python profiler sampling profiling events every milliseconds.

To learn more about the Python profiling options and what's collected, see [Start a training job with the default system monitoring and customized framework profiling with different profiling options](#).

The following methods of the `PythonProfileAnalysis`, `cProfileAnalysis`, `PyinstrumentAnalysis` classes are provided to fetch and analyze the Python profiling data. Each function loads the latest data from the default S3 URI.

```python
from smdebug.profiler.analysis.python_profile_analysis import PythonProfileAnalysis, cProfileAnalysis, PyinstrumentAnalysis
```

To set Python profiling objects for analysis, use the `cProfileAnalysis` or `PyinstrumentAnalysis` classes as shown in the following example code. It shows how to set a `cProfileAnalysis` object, and if you want to use `PyinstrumentAnalysis`, replace the class name.

```python
python_analysis = cProfileAnalysis(
    local_profile_dir=tf_python_stats_dir,
    s3_path=tj.profiler_s3_output_path
)
```

The following methods are available for the `cProfileAnalysis` and `PyinstrumentAnalysis` classes to fetch the Python profiling stats data:

- `python_analysis.fetch_python_profile_stats_by_time(start_time_since_epoch_in_secs, end_time_since_epoch_in_secs)` – Takes in a start time and end time, and returns the function stats of step stats whose start or end times overlap with the provided interval.
- `python_analysis.fetch_python_profile_stats_by_step(start_step, end_step, mode, start_phase, end_phase)` – Takes in a start step and end step and returns the function stats of all step stats whose profiled step satisfies `start_step <= step < end_step`. 
• **start_step** and **end_step** (str) – Specify the start step and end step to fetch the Python profiling stats data.

• **mode** (str) – Specify the mode of training job using the `PythonProfileModes` enumerator class. The default is `PythonProfileModes.TRAIN`. Available options are provided in the [Training Modes and Phases for Python Profiling](#) section.

• **start_phase** (str) – Specify the start phase in the target step(s) using the `StepPhase` enumerator class. This parameter enables profiling between different phases of training. The default is `StepPhase.STEP_START`. Available options are provided in the [Training Modes and Phases for Python Profiling](#) section.

• **end_phase** (str) – Specify the end phase in the target step(s) using the `StepPhase` enumerator class. This parameter sets up the end phase of training. Available options are as same as the ones for the `start_phase` parameter. The default is `StepPhase.STEP_END`. Available options are provided in the [Training Modes and Phases for Python Profiling](#) section.

• `python_analysis.fetch_profile_stats_between_modes(start_mode, end_mode)` – Fetches stats from the Python profiling between the start and end modes.

• `python_analysis.fetch_pre_step_zero_profile_stats()` – Fetches the stats from the Python profiling until step 0.

• `python_analysis.fetch_post_hook_close_profile_stats()` – Fetches stats from the Python profiling after the hook is closed.

• `python_analysis.list_profile_stats()` – Returns a DataFrame of the Python profiling stats. Each row holds the metadata for each instance of profiling and the corresponding stats file (one per step).

• `python_analysis.list_available_node_ids()` – Returns a list the available node IDs for the Python profiling stats.

The `cProfileAnalysis` class specific methods:

• `fetch_profile_stats_by_training_phase()` – Fetches and aggregates the Python profiling stats for every possible combination of start and end modes. For example, if a training and validation phases are done while detailed profiling is enabled, the combinations are (PRE_STEP_ZERO, TRAIN), (TRAIN, TRAIN), (TRAIN, EVAL), (EVAL, EVAL), and (EVAL, POST_HOOK_CLOSE). All stats files within each of these combinations are aggregated.
- `fetch_profile_stats_by_job_phase()` – Fetches and aggregates the Python profiling stats by job phase. The job phases are `initialization` (profiling until step 0), `training_loop` (training and validation), and `finalization` (profiling after the hook is closed).

**Merge timelines of multiple profile trace files**

The SMDebug client library provide profiling analysis and visualization tools for merging timelines of system metrics, framework metrics, and Python profiling data collected by Debugger.

***Tip***

Before proceeding, you need to set a `TrainingJob` object that will be utilized throughout the examples in this page. For more information about setting up a `TrainingJob` object, see [Access the profile data](#).

The `MergedTimeline` class provides tools to integrate and correlate different profiling information in a single timeline. After Debugger captures profiling data and annotations from different phases of a training job, JSON files of trace events are saved in a default `tracefolder` directory.

- For annotations in the Python layers, the trace files are saved in `*pythontimeline.json`.
- For annotations in the TensorFlow C++ layers, the trace files are saved in `*model_timeline.json`.
- Tensorflow profiler saves events in a `*trace.json.gz` file.

***Tip***

If you want to list all of the JSON trace files, use the following AWS CLI command:

```
! aws s3 ls {tj.profiler_s3_output_path} --recursive | grep '\.json$'
```

As shown in the following animated screenshot, putting and aligning the trace events captured from the different profiling sources in a single plot can provide an overview of the entire events occurring in different phases of the training job.
Tip

To interact with the merged timeline on the tracing app using a keyboard, use the W key for zooming in, the A key for shifting to the left, the S key for zooming out, and the D key for shifting to the right.

The multiple event trace JSON files can be merged into one trace event JSON file using the following MergedTimeline API operation and class method from the smdebug.profiler.analysis.utils.merge_timelines module.

```python
from smdebug.profiler.analysis.utils.merge_timelines import MergedTimeline

combined_timeline = MergedTimeline(path, file_suffix_filter, output_directory)
combined_timeline.merge_timeline(start, end, unit)
```

The MergedTimeline API operation passes the following parameters:
• **path (str)** – Specify a root folder ("/profiler-output") that contains system and framework profiling trace files. You can locate the profiler-output using the SageMaker estimator classmethod or the TrainingJob object. For example, `estimator.latest_job_profiler_artifacts_path()` or `tj.profiler_s3_output_path`.

• **file_suffix_filter (list)** – Specify a list of file suffix filters to merge timelines. Available suffix filters are ["model_timeline.json", "pythontimeline.json", "trace.json.gz"]. If this parameter is not manually specified, all of the trace files are merged by default.

• **output_directory (str)** – Specify a path to save the merged timeline JSON file. The default is to the directory specified for the path parameter.

The `merge_timeline()` classmethod passes the following parameters to execute the merging process:

• **start (int)** – Specify start time (in microseconds and in Unix time format) or start step to merge timelines.

• **end (int)** – Specify end time (in microseconds and in Unix time format) or end step to merge timelines.

• **unit (str)** – Choose between "time" and "step". The default is "time".

Using the following example codes, execute the `merge_timeline()` method and download the merged JSON file.

• Merge timeline with the "time" unit option. The following example code merges all available trace files between the Unix start time (the absolute zero Unix time) and the current Unix time, which means that you can merge the timelines for the entire training duration.

```python
import time
from smdebug.profiler.analysis.utils.merge_timelines import MergedTimeline
from smdebug.profiler.profiler_constants import CONVERT_TO_MICROSECS

combined_timeline = MergedTimeline(tj.profiler_s3_output_path, output_directory="./")
combined_timeline.merge_timeline(0, int(time.time() * CONVERT_TO_MICROSECS))
```

• Merge timeline with the "step" unit option. The following example code merges all available timelines between step 3 and step 9.

```python
combined_timeline = MergedTimeline(tj.profiler_s3_output_path, output_directory="./")
combined_timeline.merge_timeline(0, 9)
```
from smdebug.profiler.analysis.utils.merge_timelines import MergedTimeline

combined_timeline = MergedTimeline(tj.profiler_s3_output_path, output_directory="./")
combined_timeline.merge_timeline(3, 9, unit="step")

Open the Chrome tracing app at chrome://tracing on a Chrome browser, and open the JSON file. You can explore the output to plot the merged timeline.

Profiling data loaders

In PyTorch, data loader iterators, such as `SingleProcessingDataLoaderIter` and `MultiProcessingDataLoaderIter`, are initiated at the beginning of every iteration over a dataset. During the initialization phase, PyTorch turns on worker processes depending on the configured number of workers, establishes data queue to fetch data and `pin_memory` threads.

To use the PyTorch data loader profiling analysis tool, import the following `PT_dataloader_analysis` class:

```python
from smdebug.profiler.analysis.utils.pytorch_dataloader_analysis import PT_dataloader_analysis
```

Pass the profiling data retrieved as a Pandas frame data object in the [Access the profiling data using the pandas data parsing tool](#) section:

```python
pt_analysis = PT_dataloader_analysis(pf)
```

The following functions are available for the `pt_analysis` object:

The SMDebug S3SystemMetricsReader class reads the system metrics from the S3 bucket specified to the `s3_trial_path` parameter.

- `pt_analysis.analyze_dataloaderIter_initialization()`

  The analysis outputs the median and maximum duration for these initializations. If there are outliers, (i.e duration is greater than 2 * median), the function prints the start and end times for those durations. These can be used to inspect system metrics during those time intervals.

  The following list shows what analysis is available from this class method:

  - Which type of data loader iterators were initialized.
• The number of workers per iterator.
• Inspect whether the iterator was initialized with or without pin_memory.
• Number of times the iterators were initialized during training.

```
pt_analysis.analyze_dataloaderWorkers()
```

The following list shows what analysis is available from this class method:
• The number of worker processes that were spun off during the entire training.
• Median and maximum duration for the worker processes.
• Start and end time for the worker processes that are outliers.

```
pt_analysis.analyze_dataloader_getnext()
```

The following list shows what analysis is available from this class method:
• Number of GetNext calls made during the training.
• Median and maximum duration in microseconds for GetNext calls.
• Start time, End time, duration and worker id for the outlier GetNext call duration.

```
pt_analysis.analyze_batchtime(start_timestamp, end_timestamp,
select_events=[".*"], select_dimensions=[".*"])
```

Debugger collects the start and end times of all the GetNext calls. You can find the amount of
time spent by the training script on one batch of data. Within the specified time window, you
can identify the calls that are not directly contributing to the training. These calls can be from
the following operations: computing the accuracy, adding the losses for debugging or logging
purposes, and printing the debugging information. Operations like these can be compute
intensive or time consuming. We can identify such operations by correlating the Python profiler,
system metrics, and framework metrics.

The following list shows what analysis is available from this class method:
• Profile time spent on each data batch, BatchTime_in_seconds, by finding the difference
  between start times of current and subsequent GetNext calls.
• Find the outliers in BatchTime_in_seconds and start and end time for those outliers.
• Obtain the system and framework metrics during those BatchTime_in_seconds
timestamps. This indicates where the time was spent.

```
pt_analysis.plot_the_window()
```

Plots a timeline charts between a start timestamp and the end timestamp.
Release notes for profiling capabilities of Amazon SageMaker

See the following release notes to track the latest updates for profiling capabilities of Amazon SageMaker.

March 21, 2024

Currency updates

SageMaker Profiler has added support for PyTorch v2.2.0, v2.1.0, and v2.0.1.

AWS Deep Learning Containers pre-installed with SageMaker Profiler

SageMaker Profiler is packaged in the following AWS Deep Learning Containers.

- SageMaker Framework Container for PyTorch v2.2.0
- SageMaker Framework Container for PyTorch v2.1.0
- SageMaker Framework Container for PyTorch v2.0.1

December 14, 2023

Currency updates

SageMaker Profiler has added support for TensorFlow v2.13.0.

Breaking changes

This release involves a breaking change. The SageMaker Profiler Python package name is changed from smppy to smprof. If you have been using the previous version of the package while you have started using the latest SageMaker Framework Containers for TensorFlow listed in the following section, make sure that you update the package name from smppy to smprof in the import statement in your training script.

AWS Deep Learning Containers pre-installed with SageMaker Profiler

SageMaker Profiler is packaged in the following AWS Deep Learning Containers.

- SageMaker Framework Container for TensorFlow v2.13.0
- SageMaker Framework Container for TensorFlow v2.12.0
If you use the previous versions of the framework containers such as TensorFlow v2.11.0, the SageMaker Profiler Python package is still available as smppy. If you are uncertain which version or the package name you should use, replace the import statement of the SageMaker Profiler package with the following code snippet.

```python
try:
    import smprof
except ImportError:
    # backward-compatibility for TF 2.11 and PT 1.13.1 images
    import smppy as smprof
```

**August 24, 2023**

**New features**

Released Amazon SageMaker Profiler, a profiling and visualization capability of SageMaker to deep dive into compute resources provisioned while training deep learning models and gain visibility into operation-level details. SageMaker Profiler provides Python modules (smppy) for adding annotations throughout PyTorch or TensorFlow training scripts and activating SageMaker Profiler. You can access the modules through the SageMaker Python SDK and AWS Deep Learning Containers. For any jobs run with the SageMaker Profiler Python modules, you can load the profile data in the SageMaker Profiler UI application that provides a summary dashboard and a detailed timeline. To learn more, see [Use Amazon SageMaker Profiler to profile activities on AWS compute resources](https://docs.aws.amazon.com/sagemaker/latest/dg/profiler.html).

This release of the SageMaker Profiler Python package is integrated into the following SageMaker Framework Containers for PyTorch and TensorFlow.

- PyTorch v2.0.0
- PyTorch v1.13.1
- TensorFlow v2.12.0
- TensorFlow v2.11.0

**Distributed training in Amazon SageMaker**

SageMaker provides distributed training libraries and supports various distributed training options for deep learning tasks such as computer vision (CV) and natural language processing (NLP). With
SageMaker's distributed training libraries, you can run highly scalable and cost-effective custom data parallel and model parallel deep learning training jobs. You can also use other distributed training frameworks and packages such as PyTorch DistributedDataParallel (DDP), torchrun, MPI (mpirun), and parameter server. Throughout the documentation, instructions and examples focus on how to set up the distributed training options for deep learning tasks using the SageMaker Python SDK.

Tip
To learn best practices for distributed computing of machine learning (ML) training and processing jobs in general, see Distributed computing with SageMaker best practices.

Before you get started

SageMaker Training supports distributed training on a single instance as well as multiple instances, so you can run any size of training at scale. We recommend you to use the framework estimator classes such as PyTorch and TensorFlow in the SageMaker Python SDK, which are the training job launchers with various distributed training options. When you create an estimator object, the object sets up distributed training infrastructure, runs the CreateTrainingJob API in the backend, finds the Region where your current session is running, and pulls one of the pre-built AWS deep learning container prepackaged with a number of libraries including deep learning frameworks, distributed training frameworks, and the EFA driver. If you want to mount an FSx file system to the training instances, you need to pass your VPC subnet and security group ID to the estimator. Before running your distributed training job in SageMaker, read the following general guidance on the basic infrastructure setup.

Availability zones and network backplane

When using multiple instances (also called nodes), it's important to understand the network that connects the instances, how they read the training data, and how they share information between themselves. For example, when you run a distributed data-parallel training job, a number of factors, such as communication between the nodes of a compute cluster for running the AllReduce operation and data transfer between the nodes and data storage in Amazon Simple Storage Service or Amazon FSx for Lustre, play a crucial role to achieve an optimal use of compute resources and a faster training speed. To reduce communication overhead, make sure that you configure instances, VPC subnet, and data storage in the same AWS Region and Availability Zone.
GPU instances with faster network and high-throughput storage

You can technically use any instances for distributed training. For cases where you need to run multi-node distributed training jobs for training large models, such as large language models (LLMs) and diffusion models, which require faster inter-node commutation, we recommend **EFA-enabled GPU instances supported by SageMaker**. Especially, to achieve the most performant distributed training job in SageMaker, we recommend **P4d and P4de instances equipped with NVIDIA A100 GPUs**. These are also equipped with high-throughput low-latency local instance storage and faster intra-node network. For data storage, we recommend **Amazon FSx for Lustre** that provides high throughput for storing training datasets and model checkpoints.

Get started with distributed training in Amazon SageMaker

If you're already familiar with distributed training, choose one of the following options that matches your preferred strategy or framework to get started. If you want to learn about distributed training in general, see the section called “Basic distributed training concepts”.

The SageMaker distributed training libraries are optimized for the SageMaker training environment, help adapt your distributed training jobs to SageMaker, and improve training speed and throughput. The libraries offer both data parallel and model parallel training strategies. They combine software and hardware technologies to improve inter-GPU and inter-node communications, and extend SageMaker's training capabilities with built-in options that require minimal code changes to your training scripts.

Use the SageMaker distributed data parallelism (SMDDP) library

The SMDDP library improves communication between nodes with implementations of AllReduce and AllGather collective communication operations that are optimized for AWS network infrastructure and Amazon SageMaker ML instance topology. You can use the SMDDP library as the backend of PyTorch-based distributed training packages: PyTorch distributed data parallel (DDP), PyTorch fully sharded data parallelism (FSDP), DeepSpeed, and Megatron-DeepSpeed. The following code example shows how to set a PyTorch estimator for launching a distributed training job on two ml.p4d.24xlarge instances.

```python
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    ...,
    instance_count=2,
    instance_type="ml.p4d.24xlarge",
)```
# Activate distributed training with SMDDP

distribution={ "pytorchddp": { "enabled": True } }  # mpirun, activates SMDDP AllReduce OR AllGather

distribution={ "torch_distributed": { "enabled": True } }  # torchrun, activates SMDDP AllGather

distribution={ "smdistributed": { "dataparallel": { "enabled": True } } }  # mpirun, activates SMDDP AllReduce OR AllGather

To learn how to prepare your training script and launch a distributed data-parallel training job on SageMaker, see the section called “SageMaker distributed data parallelism library”.

**Use the SageMaker model parallelism library (SMP)**

SageMaker provides the SMP library and supports various distributed training techniques, such as sharded data parallelism, pipelining, tensor parallelism, optimizer state sharding, and more. To learn more about what the SMP library offers, see the section called “Core Features”.

To use SageMaker's model parallelism library, configure the distribution parameter of the SageMaker framework estimators. Supported framework estimators are PyTorch and TensorFlow. The following code example shows how to construct a framework estimator for distributed training with the model parallelism library on two ml.p4d.24xlarge instances.

```python
from sagemaker.framework import Framework

distribution={
    "smdistributed": {
        "modelparallel": {
            "enabled":True,
            "parameters": {
                ...  # enter parameter key-value pairs here
            }
        },
    },
    "mpi": {
        "enabled":True,
        ...  # enter parameter key-value pairs here
    }
}

estimator = Framework(
    ...
    instance_count=2,
)```
To learn how to adapt your training script, configure distribution parameters in the estimator class, and launch a distributed training job, see SageMaker's model parallelism library (see also Distributed Training APIs in the SageMaker Python SDK documentation).

**Use open source distributed training frameworks**

SageMaker also supports the following options to operate mpirun and torchrun in the backend.

- To use PyTorch DistributedDataParallel (DDP) in SageMaker with the mpirun backend, add distribution={"pytorchddp": {"enabled": True}} to your PyTorch estimator. For more information, see also PyTorch Distributed Training and SageMaker PyTorch Estimator's distribution argument in the SageMaker Python SDK documentation.

Note

This option is available for PyTorch 1.12.0 and later.

```python
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    ...
    instance_count=2,
    instance_type="ml.p4d.24xlarge",
    distribution={"pytorchddp": {"enabled": True}}  # runs mpirun in the backend
)
```

- SageMaker supports the PyTorch torchrun launcher for distributed training on GPU-based Amazon EC2 instances, such as P3 and P4, as well as Trn1 powered by the AWS Trainium device.

To use PyTorch DistributedDataParallel (DDP) in SageMaker with the torchrun backend, add distribution={"torch_distributed": {"enabled": True}} to the PyTorch estimator.

Note

This option is available for PyTorch 1.13.0 and later.
The following code snippet shows an example of constructing a SageMaker PyTorch estimator to run distributed training on two `ml.p4d.24xlarge` instances with the `torch_distributed` distribution option.

```python
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    ...,  
    instance_count=2,  
    instance_type="ml.p4d.24xlarge",  
    distribution={"torch_distributed": {"enabled": True}}  # runs torchrun in the backend
)
```

For more information, see Distributed PyTorch Training and SageMaker PyTorch Estimator's distribution argument in the SageMaker Python SDK documentation.

**Notes for distributed training on Trn1**

A Trn1 instance consists of up to 16 Trainium devices, and each Trainium device consists of two NeuronCores. For specs of the AWS Trainium devices, see Trainium Architecture in the AWS Neuron Documentation.

To train on the Trainium-powered instances, you only need to specify the Trn1 instance code, `ml.trn1.*`, in string to the `instance_type` argument of the SageMaker PyTorch estimator class. To find available Trn1 instance types, see AWS Trn1 Architecture in the AWS Neuron documentation.

⚠️ **Note**

SageMaker Training on Amazon EC2 Trn1 instances is currently available only for the PyTorch framework in the AWS Deep Learning Containers for PyTorch Neuron starting v1.11.0. To find a complete list of supported versions of PyTorch Neuron, see Neuron Containers in the AWS Deep Learning Containers GitHub repository.

When you launch a training job on Trn1 instances using the SageMaker Python SDK, SageMaker automatically picks up and runs the right container from Neuron Containers provided by AWS.
Deep Learning Containers. The Neuron Containers are prepackaged with training environment settings and dependencies for easier adaptation of your training job to the SageMaker Training platform and Amazon EC2 Trn1 instances.

Note
To run your PyTorch training job on Trn1 instances with SageMaker, you should modify your training script to initialize process groups with the xla backend and use PyTorch/XLA. To support the XLA adoption process, the AWS Neuron SDK provides PyTorch Neuron that uses XLA to make conversion of PyTorch operations to Trainium instructions. To learn how to modify your training script, see Developer Guide for Training with PyTorch Neuron (torch-neuronx) in the AWS Neuron Documentation.

For more information, see Distributed Training with PyTorch Neuron on Trn1 instances and SageMaker PyTorch Estimator's distribution argument in the SageMaker Python SDK documentation.

- To use MPI in SageMaker, add distribution={"mpi": {"enabled": True}} to your estimator. The MPI distribution option is available for the following frameworks: MXNet, PyTorch, and TensorFlow.
- To use a parameter server in SageMaker, add distribution={"parameter_server": {"enabled": True}} to your estimator. The parameter server option is available for the following frameworks: MXNet, PyTorch, and TensorFlow.

Tip
For more information about using the MPI and parameter server options per framework, use the following links to the SageMaker Python SDK documentation.

- MXNet Distributed Training and SageMaker MXNet Estimator's distribution argument
- PyTorch Distributed Training and SageMaker PyTorch Estimator's distribution argument
Basic distributed training concepts

SageMaker's distributed training libraries use the following distributed training terms and features.

Datasets and Batches

- **Training Dataset**: All of the data you use to train the model.
- **Global batch size**: The number of records selected from the training dataset in each iteration to send to the GPUs in the cluster. This is the number of records over which the gradient is computed at each iteration. If data parallelism is used, it is equal to the total number of model replicas multiplied by the per-replica batch size: \(\text{global batch size} = (\text{the number of model replicas}) \times (\text{per-replica batch size})\). A single batch of global batch size is often referred to as the *mini-batch* in machine learning literature.
- **Per-replica batch size**: When data parallelism is used, this is the number of records sent to each model replica. Each model replica performs a forward and backward pass with this batch to calculate weight updates. The resulting weight updates are synchronized (averaged) across all replicas before the next set of per-replica batches are processed.
- **Micro-batch**: A subset of the mini-batch or, if hybrid model and data parallelism is used, it is a subset of the per-replica sized batch. When you use SageMaker's distributed model parallelism library, each micro-batch is fed into the training pipeline one-by-one and follows an execution schedule defined by the library's runtime.

Training

- **Epoch**: One training cycle through the entire dataset. It is common to have multiple iterations per an epoch. The number of epochs you use in training is unique on your model and use case.
- **Iteration**: A single forward and backward pass performed using a global batch sized batch (a mini-batch) of training data. The number of iterations performed during training is determined by the global batch size and the number of epochs used for training. For example, if a dataset includes 5,000 samples, and you use a global batch size of 500, it will take 10 iterations to complete a single epoch.
- **Learning rate**: A variable that influences the amount that weights are changed in response to the calculated error of the model. The learning rate plays an important role in the model's ability to converge as well as the speed and optimality of convergence.

Instances and GPUs
• **Instances**: An AWS machine learning compute instance. These are also referred to as *nodes*.

• **Cluster size**: When using SageMaker's distributed training library, this is the number of instances multiplied by the number of GPUs in each instance. For example, if you use two ml.p3.8xlarge instances in a training job, which have 4 GPUs each, the cluster size is 8. While increasing cluster size can lead to faster training times, communication between instances must be optimized; Otherwise, communication between the nodes can add overhead and lead to slower training times. The SageMaker distributed training library is designed to optimize communication between Amazon EC2 ML compute instances, leading to higher device utilization and faster training times.

**Distributed Training Solutions**

• **Data parallelism**: A strategy in distributed training where a training dataset is split up across multiple GPUs in a compute cluster, which consists of multiple Amazon EC2 ML Instances. Each GPU contains a *replica* of the model, receives different batches of training data, performs a forward and backward pass, and shares weight updates with the other nodes for synchronization before moving on to the next batch and ultimately another epoch.

• **Model parallelism**: A strategy in distributed training where the model partitioned across multiple GPUs in a compute cluster, which consists of multiple Amazon EC2 ML Instances. The model might be complex and have a large number of hidden layers and weights, making it unable to fit in the memory of a single instance. Each GPU carries a subset of the model, through which the data flows and the transformations are shared and compiled. The efficiency of model parallelism, in terms of GPU utilization and training time, is heavily dependent on how the model is partitioned and the execution schedule used to perform forward and backward passes.

• **Pipeline Execution Schedule (Pipelining)**: The pipeline execution schedule determines the order in which computations (micro-batches) are made and data is processed across devices during model training. Pipelining is a technique to achieve true parallelization in model parallelism and overcome the performance loss due to sequential computation by having the GPUs compute simultaneously on different data samples. To learn more, see Pipeline Execution Schedule.

**Advanced concepts**

Machine Learning (ML) practitioners commonly face two scaling challenges when training models: *scaling model size* and *scaling training data*. While model size and complexity can result in better accuracy, there is a limit to the model size you can fit into a single CPU or GPU. Furthermore, scaling model size may result in more computations and longer training times.
Not all models handle training data scaling equally well because they need to ingest all the training data \textit{in memory} for training. They only scale vertically, and to bigger and bigger instance types. In most cases, scaling training data results in longer training times.

Deep Learning (DL) is a specific family of ML algorithms consisting of several layers of artificial neural networks. The most common training method is with mini-batch Stochastic Gradient Descent (SGD). In mini-batch SGD, the model is trained by conducting small iterative changes of its coefficients in the direction that reduces its error. Those iterations are conducted on equally sized subsamples of the training dataset called \textit{mini-batches}. For each mini-batch, the model is run in each record of the mini-batch, its error measured and the gradient of the error estimated. Then the average gradient is measured across all the records of the mini-batch and provides an update direction for each model coefficient. One full pass over the training dataset is called an \textit{epoch}. Model trainings commonly consist of dozens to hundreds of epochs. Mini-batch SGD has several benefits: First, its iterative design makes training time theoretically linear of dataset size. Second, in a given mini-batch each record is processed individually by the model without need for inter-record communication other than the final gradient average. The processing of a mini-batch is consequently particularly suitable for parallelization and distribution.

Parallelizing SGD training by distributing the records of a mini-batch over different computing devices is called \textit{data parallel distributed training}, and is the most commonly used DL distribution paradigm. Data parallel training is a relevant distribution strategy to scale the mini-batch size and process each mini-batch faster. However, data parallel training comes with the extra complexity of having to compute the mini-batch gradient average with gradients coming from all the workers and communicating it to all the workers, a step called \textit{allreduce} that can represent a growing overhead, as the training cluster is scaled, and that can also drastically penalize training time if improperly implemented or implemented over improper hardware subtracts.

Data parallel SGD still requires developers to be able to fit at least the model and a single record in a computing device, such as a single CPU or GPU. When training very large models such as large transformers in Natural Language Processing (NLP), or segmentation models over high-resolution images, there may be situations in which this is not feasible. An alternative way to break up the workload is to partition the model over multiple computing devices, an approach called \textit{model-parallel distributed training}.

**Strategies**

Distributed training is usually split by two approaches: data parallel and model parallel. \textit{Data parallel} is the most common approach to distributed training: You have a lot of data, batch it up,
and send blocks of data to multiple CPUs or GPUs (nodes) to be processed by the neural network or ML algorithm, then combine the results. The neural network is the same on each node. A *model parallel* approach is used with large models that won’t fit in a node’s memory in one piece; it breaks up the model and places different parts on different nodes. In this situation, you need to send your batches of data out to each node so that the data is processed on all parts of the model.

The terms *network* and *model* are often used interchangeably: A large model is really a large network with many layers and parameters. Training with a large network produces a large model, and loading the model back onto the network with all your pre-trained parameters and their weights loads a large model into memory. When you break apart a model to split it across nodes, you’re also breaking apart the underlying network. A network consists of layers, and to split up the network, you put layers on different compute devices.

A common pitfall of naively splitting layers across devices is severe GPU under-utilization. Training is inherently sequential in both forward and backward passes, and at a given time, only one GPU can actively compute, while the others wait on the activations to be sent. Modern model parallel libraries solve this problem by using pipeline execution schedules to improve device utilization. However, only the Amazon SageMaker's distributed model parallel library includes automatic model splitting. The two core features of the library, automatic model splitting and pipeline execution scheduling, simplifies the process of implementing model parallelism by making automated decisions that lead to efficient device utilization.

**Train with data parallel and model parallel**

If you are training with a large dataset, start with a data parallel approach. If you run out of memory during training, you may want to switch to a model parallel approach, or try hybrid model and data parallelism. You can also try the following to improve performance with data parallel:

- Change your model's hyperparameters.
- Reduce the batch size.
- Keep reducing the batch size until it fits. If you reduce batch size to 1, and still run out of memory, then you should try model-parallel training.

Try gradient compression (FP16, INT8):

- On NVIDIA TensorCore-equipped hardware, using *mixed precision training* creates both speed-up and memory consumption reduction.
- SageMaker's distributed data parallelism library supports Automatic Mixed Precision (AMP) out of the box. No extra action is needed to enable AMP other than the framework-level
modifications to your training script. If gradients are in FP16, the SageMaker data parallelism library runs its AllReduce operation in FP16. For more information about implementing AMP APIs to your training script, see the following resources:

- **Frameworks - PyTorch** in the *NVIDIA Deep Learning Performance documentation*
- **Frameworks - TensorFlow** in the *NVIDIA Deep Learning Performance documentation*
- **Automatic Mixed Precision for Deep Learning** in the *NVIDIA Developer Docs*
- **Introducing native PyTorch automatic mixed precision for faster training on NVIDIA GPUs** in the *PyTorch Blog*
- **TensorFlow mixed precision APIs** in the *TensorFlow documentation*

Try reducing the input size:

- Reduce the NLP sequence length if you increase the sequence link, need to adjust the batch size down, or adjust the GPUs up to spread the batch.
- Reduce image resolution.

Check if you use batch normalization, since this can impact convergence. When you use distributed training, your batch is split across GPUs and the effect of a much lower batch size can be a higher error rate thereby disrupting the model from converging. For example, if you prototyped your network on a single GPU with a batch size of 64, then scaled up to using four p3dn.24xlarge, you now have 32 GPUs and your per-GPU batch size drops from 64 to 2. This will likely break the convergence you saw with a single node.

Start with model-parallel training when:

- Your model does not fit on a single device.
- Due to your model size, you’re facing limitations in choosing larger batch sizes, such as if your model weights take up most of your GPU memory and you are forced to choose a smaller, suboptimal batch size.

To learn more about the SageMaker distributed libraries, see the following:

- **Run distributed training with the SageMaker distributed data parallelism library**
- **(Archived) SageMaker model parallelism library v1.x**
Optimize distributed training

Customize hyperparameters for your use case and your data to get the best scaling efficiency. In the following discussion, we highlight some of the most impactful training variables and provide references to state-of-the-art implementations so you can learn more about your options. Also, we recommend that you refer to your preferred framework’s distributed training documentation.

- Apache MXNet distributed training
- PyTorch distributed training
- TensorFlow distributed training

Batch Size

SageMaker distributed toolkits generally allow you to train on bigger batches. For example, if a model fits within a single device but can only be trained with a small batch size, using either model-parallel training or data parallel training enables you to experiment with larger batch sizes.

Be aware that batch size directly influences model accuracy by controlling the amount of noise in the model update at each iteration. Increasing batch size reduces the amount of noise in the gradient estimation, which can be beneficial when increasing from very small batches sizes, but can result in degraded model accuracy as the batch size increases to large values.

Tip

Adjust your hyperparameters to ensure that your model trains to a satisfying convergence as you increase its batch size.

A number of techniques have been developed to maintain good model convergence when batch is increased.

Mini-batch size

In SGD, the mini-batch size quantifies the amount of noise present in the gradient estimation. A small mini-batch results in a very noisy mini-batch gradient, which is not representative of the true gradient over the dataset. A large mini-batch results in a mini-batch gradient close to the true gradient over the dataset and potentially not noisy enough—likely to stay locked in irrelevant minima.
To learn more about these techniques, see the following papers:

- **Accurate, Large Minibatch SGD: Training ImageNet in 1 Hour**, Goya et al.
- **PowerAI DDL**, Cho et al.
- **Scale Out for Large Minibatch SGD: Residual Network Training on ImageNet-1K with Improved Accuracy and Reduced Time to Train**, Codreanu et al.
- **ImageNet Training in Minutes**, You et al.
- **Large Batch Training of Convolutional Networks**, You et al.
- **Large Batch Optimization for Deep Learning: Training BERT in 76 Minutes**, You et al.
- **Accelerated Large Batch Optimization of BERT Pretraining in 54 minutes**, Zheng et al.
- **Deep Gradient Compression**, Lin et al.

**Scenarios**

The following sections cover scenarios in which you may want to scale up training, and how you can do so using AWS resources.

**Scaling from a Single GPU to Many GPUs**

The amount of data or the size of the model used in machine learning can create situations in which the time to train a model is longer that you are willing to wait. Sometimes, the training doesn’t work at all because the model or the training data is too large. One solution is to increase the number of GPUs you use for training. On an instance with multiple GPUs, like a p3.16xlarge that has eight GPUs, the data and processing is split across the eight GPUs. When you use distributed training libraries, this can result in a near-linear speedup in the time it takes to train your model. It takes slightly over 1/8 the time it would have taken on p3.2xlarge with one GPU.

<table>
<thead>
<tr>
<th>Instance type</th>
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<tr>
<td>p3dn.24xlarge</td>
<td>8</td>
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</table>
Note

The ml instance types used by SageMaker training have the same number of GPUs as the corresponding p3 instance types. For example, ml.p3.8xlarge has the same number of GPUs as p3.8xlarge - 4.

Scaling from a single instance to multiple instances

If you want to scale your training even further, you can use more instances. However, you should choose a larger instance type before you add more instances. Review the previous table to see how many GPUs are in each p3 instance type.

If you have made the jump from a single GPU on a p3.2xlarge to four GPUs on a p3.8xlarge, but decide that you require more processing power, you may see better performance and incur lower costs if you choose a p3.16xlarge before trying to increase instance count. Depending on the libraries you use, when you keep your training on a single instance, performance is better and costs are lower than a scenario where you use multiple instances.

When you are ready to scale the number of instances, you can do this with SageMaker Python SDK estimator function by setting your instance_count. For example, you can set instance_type = p3.16xlarge and instance_count = 2. Instead of the eight GPUs on a single p3.16xlarge, you have 16 GPUs across two identical instances. The following chart shows scaling and throughput starting with eight GPUs on a single instance and increasing to 64 instances for a total of 256 GPUs.
Custom training scripts

While SageMaker makes it simple to deploy and scale the number of instances and GPUs, depending on your framework of choice, managing the data and results can be very challenging, which is why external supporting libraries are often used. This most basic form of distributed training requires modification of your training script to manage the data distribution.
SageMaker also supports Horovod and implementations of distributed training native to each major deep learning framework. If you choose to use examples from these frameworks, you can follow SageMaker's container guide for Deep Learning Containers, and various example notebooks that demonstrate implementations.

Run distributed training with the SageMaker distributed data parallelism library

The SageMaker distributed data parallelism (SMDDP) library extends SageMaker training capabilities on deep learning models with near-linear scaling efficiency by providing implementations of collective communication operations optimized for AWS infrastructure.

When training large machine learning (ML) models, such as large language models (LLM) and diffusion models, on a huge training dataset, ML practitioners use clusters of accelerators and distributed training techniques to reduce the time to train or resolve memory constraints for models that cannot fit in each GPU memory. ML practitioners often start with multiple accelerators on a single instance and then scale to clusters of instances as their workload requirements increase. As the cluster size increases, so does the communication overhead between multiple nodes, which leads to drop in overall computational performance.

To address such overhead and memory problems, the SMDDP library offers the following.

- The SMDDP library optimizes training jobs for AWS network infrastructure and Amazon SageMaker ML instance topology.
- The SMDDP library improves communication between nodes with implementations of AllReduce and AllGather collective communication operations that are optimized for AWS infrastructure.

To learn more about the details of the SMDDP library offerings, proceed to the section called “Introduction to the SMDDP library”.

For more information about training with the model-parallel strategy offered by SageMaker, see also (Archived) SageMaker model parallelism library v1.x.

Topics

- Introduction to the SageMaker distributed data parallelism library
- Supported frameworks, AWS Regions, and instances types
Introduction to the SageMaker distributed data parallelism library

The SageMaker distributed data parallelism (SMDDP) library is a collective communication library that improves compute performance of distributed data parallel training. The SMDDP library addresses communications overhead of the key collective communication operations by offering the following.

1. The library offers AllReduce optimized for AWS. AllReduce is a key operation used for synchronizing gradients across GPUs at the end of each training iteration during distributed data training.

2. The library offers AllGather optimized for AWS. AllGather is another key operation used in sharded data parallel training, which is a memory-efficient data parallelism technique offered by popular libraries such as the SageMaker model parallelism (SMP) library, DeepSpeed Zero Redundancy Optimizer (ZeRO), and PyTorch Fully Sharded Data Parallelism (FSDP).

3. The library performs optimized node-to-node communication by fully utilizing AWS network infrastructure and the Amazon EC2 instance topology.

The SMDDP library can increase training speed by offering performance improvement as you scale your training cluster, with near-linear scaling efficiency.

ℹ️ Note

The SageMaker distributed training libraries are available through the AWS deep learning containers for PyTorch and Hugging Face within the SageMaker Training platform. To use the libraries, you must use the SageMaker Python SDK or the SageMaker APIs through SDK for Python (Boto3) or AWS Command Line Interface. Throughout the documentation, instructions and examples focus on how to use the distributed training libraries with the SageMaker Python SDK.
SMDDP collective communication operations optimized for AWS compute resources and network infrastructure

The SMDDP library provides implementations of the AllReduce and AllGather collective operations that are optimized for AWS compute resources and network infrastructure.

SMDDP AllReduce collective operation

The SMDDP library achieves optimal overlapping of the AllReduce operation with the backward pass, significantly improving GPU utilization. It achieves near-linear scaling efficiency and faster training speed by optimizing kernel operations between CPUs and GPUs. The library performs AllReduce in parallel while GPU is computing gradients without taking away additional GPU cycles, which makes the library to achieve faster training.

- **Leverages CPUs**: The library uses CPUs to AllReduce gradients, offloading this task from the GPUs.
- **Improved GPU usage**: The cluster's GPUs focus on computing gradients, improving their utilization throughout training.

The following is the high-level workflow of the SMDDP AllReduce operation.

1. The library assigns ranks to GPUs (workers).
2. At each iteration, the library divides each global batch by the total number of workers (world size) and assigns small batches (batch shards) to the workers.
   - The size of the global batch is \((\text{number of nodes in a cluster}) \times (\text{number of GPUs per node}) \times (\text{per batch shard})\).
   - A batch shard (small batch) is a subset of dataset assigned to each GPU (worker) per iteration.
3. The library launches a training script on each worker.
4. The library manages copies of model weights and gradients from the workers at the end of every iteration.
5. The library synchronizes model weights and gradients across the workers to aggregate a single trained model.

The following architecture diagram shows an example of how the library sets up data parallelism for a cluster of 3 nodes.
SMDDP `AllGather` collective operation

`AllGather` is a collective operation where each worker starts with an input buffer, and then concatenates or *gathers* the input buffers from all other workers into an output buffer.

**Note**

The SMDDP `AllGather` collective operation is available in `smdistributed-datatparallel>=2.0.1` and AWS Deep Learning Containers (DLC) for PyTorch v2.0.1 and later.

`AllGather` is heavily used in distributed training techniques such as sharded data parallelism where each individual worker holds a fraction of a model, or a sharded layer. The workers call `AllGather` before forward and backward passes to reconstruct the sharded layers. The forward
and backward passes continue onward after the parameters are all gathered. During the backward pass, each worker also calls ReduceScatter to collect (reduce) gradients and break (scatter) them into gradient shards to update the corresponding sharded layer. For more details on the role of these collective operations in sharded data parallelism, see the SMP library's implementation of sharded data parallelism, ZeRO in the DeepSpeed documentation, and the blog about PyTorch Fully Sharded Data Parallelism.

Because collective operations like AllGather are called in every iteration, they are the main contributors to GPU communication overhead. Faster computation of these collective operations directly translates to a shorter training time with no side effects on convergence. To achieve this, the SMDDP library offers AllGather optimized for P4d instances.

SMDDP AllGather uses the following techniques to improve computational performance on P4d instances.

1. It transfers data between instances (inter-node) through the Elastic Fabric Adapter (EFA) network with a mesh topology. EFA is the AWS low-latency and high-throughput network solution. A mesh topology for inter-node network communication is more tailored to the characteristics of EFA and AWS network infrastructure. Compared to the NCCL ring or tree topology that involves multiple packet hops, SMDDP avoids accumulating latency from multiple hops as it only needs one hop. SMDDP implements a network rate control algorithm that balances the workload to each communication peer in a mesh topology and achieves a higher global network throughput.

2. It adopts low-latency GPU memory copy library based on NVIDIA GPUDirect RDMA technology (GDRCopy) to coordinate local NVLink and EFA network traffic. GDRCopy, a low-latency GPU memory copy library offered by NVIDIA, provides low-latency communication between CPU processes and GPU CUDA kernels. With this technology, the SMDDP library is able to pipeline the intra-node and inter-node data movement.

3. It reduces the usage of GPU streaming multiprocessors to increase compute power for running model kernels. P4d and P4de instances are equipped with NVIDIA A100 GPUs, which each have 108 streaming multiprocessors. While NCCL takes up to 24 streaming multiprocessors to run collective operations, SMDDP uses fewer than 9 streaming multiprocessors. Model compute kernels pick up the saved streaming multiprocessors for faster computation.
Supported frameworks, AWS Regions, and instances types

Before using the SageMaker distributed data parallelism (SMDDP) library, check what are the supported ML frameworks and instance types and if there are enough quotas in your AWS account and AWS Region.

Supported frameworks

The following tables show the deep learning frameworks and their versions that SageMaker and SMDDP support. The SMDDP library is available in SageMaker Framework Containers, integrated in Docker containers distributed by the SageMaker model parallelism (SMP) library v2, or downloadable as a binary file.

Note

To check the latest updates and release notes of the SMDDP library, see the section called “Release notes”.

Topics

- PyTorch
- PyTorch Lightning
- Hugging Face Transformers
- TensorFlow (deprecated)

PyTorch

<table>
<thead>
<tr>
<th>PyTorch version</th>
<th>SMDDP library version</th>
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<th>SMP Docker images pre-installed with SMDDP</th>
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*Note*: To check the latest updates and release notes of the SMDDP library, see the section called “Release notes”.

**Topics**: PyTorch, PyTorch Lightning, Hugging Face Transformers, TensorFlow (deprecated)

**PyTorch**

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**SageMaker distributed data parallelism library**
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<tr>
<td>PyTorch version</td>
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**The URLs of the binary files are for installing the SMDDP library in custom containers. For more information, see [Create your own Docker container with the SageMaker distributed data parallel library](https://docs.aws.amazon.com/sagemaker/latest/dg/….)

Note

The SMDDP library is available in AWS Regions where the [SageMaker Framework Containers](https://docs.aws.amazon.com/sagemaker/latest/) and the [SMP Docker images](https://docs.aws.amazon.com/sagemaker/latest/) are in service.

Note

The SMDDP library v1.4.0 and later works as a backend of PyTorch distributed (torch.distributed) data parallelism (torch.parallel.DistributedDataParallel). In accordance with the change, the following [smdistributed APIs](https://docs.aws.amazon.com/sagemaker/latest/) for the PyTorch distributed package have been deprecated.

- smdistributed.dataparallel.torch.distributed is deprecated. Use the [torch.distributed](https://docs.aws.amazon.com/sagemaker/latest/) package instead.
- smdistributed.dataparallel.torch.parallel.DistributedDataParallel is deprecated. Use the [torch.nn.parallel.DistributedDataParallel](https://docs.aws.amazon.com/sagemaker/latest/) API instead.

If you need to use the previous versions of the library (v1.3.0 or before), see the [archived SageMaker distributed data parallelism documentation](https://docs.aws.amazon.com/sagemaker/latest/) in the [SageMaker Python SDK documentation](https://docs.aws.amazon.com/sagemaker/latest/).

**PyTorch Lightning**

The SMDDP library is available for PyTorch Lightning in the following SageMaker Framework Containers for PyTorch and the SMP Docker containers.

**PyTorch Lightning v2**
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** The URLs of the binary files are for installing the SMDDP library in custom containers. For more information, see [Create your own Docker container with the SageMaker distributed data parallel library](#).

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**Note**

PyTorch Lightning and its utility libraries such as Lightning Bolts are not preinstalled in the PyTorch DLCs. When you construct a SageMaker PyTorch estimator and submit a training job request in **Step 2**, you need to provide `requirements.txt` to install `pytorch-lightning` and `lightning-bolts` in the SageMaker PyTorch training container.

```
# requirements.txt
pytorch-lightning
lightning-bolts
```
For more information about specifying the source directory to place the requirements.txt file along with your training script and a job submission, see Using third-party libraries in the Amazon SageMaker Python SDK documentation.

Hugging Face Transformers

The AWS Deep Learning Containers for Hugging Face use the SageMaker Training Containers for PyTorch and TensorFlow as their base images. To look up the Hugging Face Transformers library versions and paired PyTorch and TensorFlow versions, see the latest Hugging Face Containers and the Prior Hugging Face Container Versions.

TensorFlow (deprecated)

⚠️ Important

The SMDDP library discontinued support for TensorFlow and is no longer available in DLCs for TensorFlow later than v2.11.0. The following table lists previous DLCs for TensorFlow with the SMDDP library installed.

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<thead>
<tr>
<th>TensorFlow version</th>
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AWS Regions

The SMDDP library is available in all of the AWS Regions where the AWS Deep Learning Containers for SageMaker and the SMP Docker images are in service.

Supported instance types

The SMDDP library requires one of the following instance types.
### Tip

To properly run distributed training on the EFA-enabled instance types, you should enable traffic between the instances by setting up the security group of your VPC to allow all inbound and outbound traffic to and from the security group itself. To learn how to set up the security group rules, see [Step 1: Prepare an EFA-enabled security group](#) in the [Amazon EC2 User Guide](#).

### Important

The SMDDP library discontinued support for P3 instances. The SMDDP library supports instance types that are equipped with NVIDIA A100 GPUs and EFA.

For specs of the instance types, see the [Accelerated Computing](#) section in the [Amazon EC2 Instance Types page](#). For information about instance pricing, see [Amazon SageMaker Pricing](#).

If you encountered an error message similar to the following, follow the instructions at [Request a service quota increase for SageMaker resources](#).

```
ResourceLimitExceeded: An error occurred (ResourceLimitExceeded) when calling the CreateTrainingJob operation: The account-level service limit 'ml.p3dn.24xlarge for training job usage' is 0 Instances, with current utilization of 0 Instances and a request delta of 1 Instances. Please contact AWS support to request an increase for this limit.
```

### How to run a distributed training job with the SageMaker distributed data parallelism library

The SageMaker distributed data parallelism (SMDDP) library is designed for ease of use and to provide seamless integration with PyTorch.
When training a deep learning model with the SMDDP library on SageMaker, you can focus on writing your training script and model training.

To get started, import the SMDDP library to use its collective operations optimized for AWS. The following topics provide instructions on what to add to your training script depending on which collective operation you want to optimize.

Topics

- **Step 1: Adapt your training script to use the SMDDP collective operations**
- **Step 2: Launch a distributed training job using the SageMaker Python SDK**

**Step 1: Adapt your training script to use the SMDDP collective operations**

The training script examples provided in this section are simplified and highlight only the required changes to enable the SageMaker distributed data parallelism (SMDDP) library in your training script. For end-to-end Jupyter notebook examples that demonstrate how to run a distributed training job with the SMDDP library, see [Amazon SageMaker data parallelism library examples](https://docs.aws.amazon.com/sagemaker/latest/dg/data-parallelism-library.html).

Topics

- **Use the SMDDP library in your PyTorch training script**
- **Use the SMDDP library in your PyTorch Lightning training script**
- **Use the SMDDP library in your TensorFlow training script (deprecated)**

**Use the SMDDP library in your PyTorch training script**

Starting from the SageMaker distributed data parallelism (SMDDP) library v1.4.0, you can use the library as a backend option for the PyTorch distributed package. To use the SMDDP AllReduce and AllGather collective operations, you only need to import the SMDDP library at the beginning of your training script and set SMDDP as the the backend of PyTorch distributed modules during process group initialization. With the single line of backend specification, you can keep all the native PyTorch distributed modules and the entire training script unchanged. The following code snippets show how to use the SMDDP library as the backend of PyTorch-based distributed training packages: PyTorch distributed data parallel (DDP), PyTorch fully sharded data parallelism (FSDP), DeepSpeed, and Megatron-DeepSpeed.

**For PyTorch DDP or FSDP**

Initialize the process group as follows.
import torch.distributed as dist
import smdistributed.dataparallel.torch.torch_smddp

dist.init_process_group(backend="smddp")

Note
(For PyTorch DDP jobs only) The smddp backend currently does not support creating subprocess groups with the torch.distributed.new_group() API. You also cannot use the smddp backend concurrently with other process group backends such as NCCL and Gloo.

For DeepSpeed or Megatron-DeepSpeed

Initialize the process group as follows.

import deepspeed
import smdistributed.dataparallel.torch.torch_smddp
deespeed.init_distributed(dist_backend="smddp")

Note
To use SMDDP AllGather with the mpirun-based launchers (smdistributed and pytorchddp) in the section called “Step 2: Launch a distributed training job”, you also need to set the following environment variable in your training script.

export SMDATAPARALLEL_OPTIMIZE_SDP=true

For general guidance on writing a PyTorch FSDP training script, see Advanced Model Training with Fully Sharded Data Parallel (FSDP) in the PyTorch documentation.

For general guidance on writing a PyTorch DDP training script, see Getting started with distributed data parallel in the PyTorch documentation.

After you have completed adapting your training script, proceed to Step 2: Launch a distributed training job using the SageMaker Python SDK.
Use the SMDDP library in your PyTorch Lightning training script

If you want to bring your PyTorch Lightning training script and run a distributed data parallel training job in SageMaker, you can run the training job with minimal changes in your training script. The necessary changes include the following: import the smdistributed.dataparallel library's PyTorch modules, set up the environment variables for PyTorch Lightning to accept the SageMaker environment variables that are preset by the SageMaker training toolkit, and activate the SMDDP library by setting the process group backend to "smddp". To learn more, walk through the following instructions that break down the steps with code examples.

Note
The PyTorch Lightning support is available in the SageMaker data parallel library v1.5.0 and later.

PyTorch Lightning == v2.1.0 and PyTorch == 2.0.1

1. Import the pytorch_lightning library and the smdistributed.dataparallel.torch modules.

   ```python
   import lightning as pl
   import smdistributed.dataparallel.torch.torch_smddp
   ```

2. Instantiate the LightningEnvironment.

   ```python
   from lightning.fabric.plugins.environments.lightning import LightningEnvironment
   
   env = LightningEnvironment()
   env.world_size = lambda: int(os.environ["WORLD_SIZE"])  
   env.global_rank = lambda: int(os.environ["RANK"])  
   ```

3. For PyTorch DDP – Create an object of the DDPStrategy class with "smddp" for process_group_backend and "gpu" for accelerator, and pass that to the Trainer class.

   ```python
   import lightning as pl
   from lightning.pytorch.strategies import DDPStrategy
   
   ddp = DDPStrategy(  
         cluster_environment=env,  
         process_group_backend="smddp",  
   ```
For PyTorch FSDP – Create an object of the `FSDPStrategy` class (with `wrapping policy` of choice) with "sddp" for `process_group_backend` and "gpu" for `accelerator`, and pass that to the `Trainer` class.

```python
import lightning as pl
from lightning.pytorch.strategies import FSDPStrategy
from functools import partial
from torch.distributed.fsdp.wrap import size_based_auto_wrap_policy

policy = partial(
    size_based_auto_wrap_policy,
    min_num_params=10000
)

fsdp = FSDPStrategy(
    auto_wrap_policy=policy,
    process_group_backend="sddp",
    cluster_environment=env
)

trainer = pl.Trainer(
    max_epochs=200,
    strategy=fsdp,
    devices=num_gpus,
    num_nodes=num_nodes
)
```

After you have completed adapting your training script, proceed to Step 2: Launch a distributed training job using the SageMaker Python SDK.
Note

When you construct a SageMaker PyTorch estimator and submit a training job request in the section called “Step 2: Launch a distributed training job”, you need to provide requirements.txt to install pytorch-lightning and lightning-bolts in the SageMaker PyTorch training container.

```
# requirements.txt
pytorch-lightning
lightning-bolts
```

For more information about specifying the source directory to place the requirements.txt file along with your training script and a job submission, see Using third-party libraries in the Amazon SageMaker Python SDK documentation.

---

Use the SMDDP library in your TensorFlow training script (deprecated)

Important

The SMDDP library discontinued support for TensorFlow and is no longer available in DLCs for TensorFlow later than v2.11.0. To find previous TensorFlow DLCs with the SMDDP library installed, see the section called “Supported frameworks”.

The following steps show you how to modify a TensorFlow training script to utilize SageMaker's distributed data parallel library.

The library APIs are designed to be similar to Horovod APIs. For additional details on each API that the library offers for TensorFlow, see the SageMaker distributed data parallel TensorFlow API documentation.

Note

SageMaker distributed data parallel is adaptable to TensorFlow training scripts composed of tf core modules except tf.keras modules. SageMaker distributed data parallel does not support TensorFlow with Keras implementation.
### Note

The SageMaker distributed data parallelism library supports Automatic Mixed Precision (AMP) out of the box. No extra action is needed to enable AMP other than the framework-level modifications to your training script. If gradients are in FP16, the SageMaker data parallelism library runs its AllReduce operation in FP16. For more information about implementing AMP APIs to your training script, see the following resources:

- **Frameworks - TensorFlow** in the *NVIDIA Deep Learning Performance documentation*
- **Automatic Mixed Precision for Deep Learning** in the *NVIDIA Developer Docs*
- **TensorFlow mixed precision APIs** in the *TensorFlow documentation*

1. Import the library's TensorFlow client and initialize it.

```python
import smdistributed.dataparallel.tensorflow as sdp
sdp.init()
```

2. Pin each GPU to a single `smdistributed.dataparallel` process with `local_rank`—this refers to the relative rank of the process within a given node. The `sdp.tensorflow.local_rank()` API provides you with the local rank of the device. The leader node is rank 0, and the worker nodes are rank 1, 2, 3, and so on. This is invoked in the following code block as `sdp.local_rank()`. `set_memory_growth` is not directly related to SageMaker distributed, but must be set for distributed training with TensorFlow.

```python
gpus = tf.config.experimental.list_physical_devices('GPU')
for gpu in gpus:
    tf.config.experimental.set_memory_growth(gpu, True)
if gpus:
    tf.config.experimental.set_visible_devices(gpus[sdp.local_rank()], 'GPU')
```

3. Scale the learning rate by the number of workers. The `sdp.tensorflow.size()` API provides you the number of workers in the cluster. This is invoked in the following code block as `sdp.size()`.

```python
learning_rate = learning_rate * sdp.size()
```

4. Use the library's `DistributedGradientTape` to optimize AllReduce operations during training. This wraps `tf.GradientTape`. 

---

*SageMaker distributed data parallelism library*
with tf.GradientTape() as tape:
    output = model(input)
    loss_value = loss(label, output)

# SageMaker data parallel: Wrap tf.GradientTape with the library's DistributedGradientTape
tape = sdp.DistributedGradientTape(tape)

5. Broadcast the initial model variables from the leader node (rank 0) to all the worker nodes (ranks 1 through n). This is needed to ensure a consistent initialization across all the worker ranks. Use the sdp.tensorflow.broadcast_variables API after the model and optimizer variables are initialized. This is invoked in the following code block as sdp.broadcast_variables().

sdp.broadcast_variables(model.variables, root_rank=0)
sdp.broadcast_variables(opt.variables(), root_rank=0)

6. Finally, modify your script to save checkpoints only on the leader node. The leader node has a synchronized model. This also avoids worker nodes overwriting the checkpoints and possibly corrupting the checkpoints.

if sdp.rank() == 0:
    checkpoint.save(checkpoint_dir)

The following is an example TensorFlow training script for distributed training with the library.

import tensorflow as tf

# SageMaker data parallel: Import the library TF API
import smdistributed.dataparallel.tensorflow as sdp

# SageMaker data parallel: Initialize the library
sdp.init()

gpus = tf.config.experimental.list_physical_devices('GPU')
for gpu in gpus:
    tf.config.experimental.set_memory_growth(gpu, True)
if gpus:
    # SageMaker data parallel: Pin GPUs to a single library process
tf.config.experimental.set_visible_devices(gpus[sdp.local_rank()], 'GPU')
# Prepare Dataset

dataset = tf.data.Dataset.from_tensor_slices(…)

# Define Model

mnist_model = tf.keras.Sequential(…)

loss = tf.losses.SparseCategoricalCrossentropy()

# SageMaker data parallel: Scale Learning Rate
# LR for 8 node run : 0.000125
# LR for single node run : 0.001

opt = tf.optimizers.Adam(0.000125 * sdp.size())

@tf.function

def training_step(images, labels, first_batch):
    with tf.GradientTape() as tape:
        probs = mnist_model(images, training=True)
        loss_value = loss(labels, probs)

    # SageMaker data parallel: Wrap tf.GradientTape with the library's
    DistributedGradientTape
    tape = sdp.DistributedGradientTape(tape)

    grads = tape.gradient(loss_value, mnist_model.trainable_variables)
    opt.apply_gradients(zip(grads, mnist_model.trainable_variables))

    if first_batch:
        # SageMaker data parallel: Broadcast model and optimizer variables
        sdp.broadcast_variables(mnist_model.variables, root_rank=0)
        sdp.broadcast_variables(opt.variables(), root_rank=0)

    return loss_value

...

# SageMaker data parallel: Save checkpoints only from master node.

if sdp.rank() == 0:
    checkpoint.save(checkpoint_dir)

After you have completed adapting your training script, move on to Step 2: Launch a distributed training job using the SageMaker Python SDK.
Step 2: Launch a distributed training job using the SageMaker Python SDK

To run a distributed training job with your adapted script from the section called “Step 1: Adapt your training script to use the SMDDP collective operations”, use the SageMaker Python SDK's framework or generic estimators by specifying the prepared training script as an entry point script and the distributed training configuration.

This page walks you through how to use the SageMaker Python SDK in two ways.

- If you want to achieve a quick adoption of your distributed training job in SageMaker, configure a SageMaker PyTorch or TensorFlow framework estimator class. The framework estimator picks up your training script and automatically matches the right image URI of the pre-built PyTorch or TensorFlow Deep Learning Containers (DLC), given the value specified to the framework_version parameter.

- If you want to extend one of the pre-built containers or build a custom container to create your own ML environment with SageMaker, use the SageMaker generic Estimator class and specify the image URI of the custom Docker container hosted in your Amazon Elastic Container Registry (Amazon ECR).

Your training datasets should be stored in Amazon S3 or Amazon FSx for Lustre in the AWS Region in which you are launching your training job. If you use Jupyter notebooks, you should have a SageMaker notebook instance or a SageMaker Studio Classic app running in the same AWS Region. For more information about storing your training data, see the SageMaker Python SDK data inputs documentation.

**Tip**

We recommend that you use Amazon FSx for Lustre instead of Amazon S3 to improve training performance. Amazon FSx has higher throughput and lower latency than Amazon S3.

**Tip**

To properly run distributed training on the EFA-enabled instance types, you should enables traffic between the instances by setting up the security group of your VPC to allow all inbound and outbound traffic to and from the security group itself. To learn how to set up SageMaker distributed data parallelism library
the security group rules, see Step 1: Prepare an EFA-enabled security group in the Amazon EC2 User Guide.

Choose one of the following topics for instructions on how to run a distributed training job of your training script. After you launch a training job, you can monitor system utilization and model performance using Use Amazon SageMaker Debugger to debug and improve model performance or Amazon CloudWatch.

While you follow instructions in the following topics to learn more about technical details, we also recommend that you try the Amazon SageMaker data parallelism library examples to get started.

Topics
- Using framework estimators in the SageMaker Python SDK
- Using the SageMaker generic estimator to extend prebuilt containers
- Create your own Docker container with the SageMaker distributed data parallel library

Using framework estimators in the SageMaker Python SDK

You can launch distributed training by adding the distribution argument to the SageMaker framework estimators, PyTorch or TensorFlow. For more details, choose one of the frameworks supported by the SageMaker distributed data parallelism (SMDDP) library from the following selections.

PyTorch

The following launcher options are available for launching PyTorch distributed training.

- pytorchddp – This option runs mpirun and sets up environment variables needed for running PyTorch distributed training on SageMaker. To use this option, pass the following dictionary to the distribution parameter.

```json
{ "pytorchddp": { "enabled": True } }
```

- torch_distributed – This option runs torchrun and sets up environment variables needed for running PyTorch distributed training on SageMaker. To use this option, pass the following dictionary to the distribution parameter.
smdistributed – This option also runs mpirun but with smddprun that sets up environment variables needed for running PyTorch distributed training on SageMaker.

If you chose to replace NCCL AllGather to SMDDP AllGather, you can use all three options. Choose one option that fits with your use case.

If you chose to replace NCCL AllReduce with SMDDP AllReduce, you should choose one of the mpirun-based options: smdistributed or pytorchddp. You can also add additional MPI options as follows.

```json
{ "pytorchddp": {  
  "enabled": True,  
  "custom_mpi_options": "-verbose -x NCCL_DEBUG=VERSION" 
 }
}

{ "smdistributed": { 
  "dataparallel": {  
    "enabled": True,  
    "custom_mpi_options": "-verbose -x NCCL_DEBUG=VERSION" 
  } 
 }
}
```

The following code sample shows the basic structure of a PyTorch estimator with distributed training options.

```python
from sagemaker.pytorch import PyTorch

pt_estimator = PyTorch(  
  base_job_name="training_job_name_prefix",  
  source_dir="subdirectory-to-your-code",  
  entry_point="adapted-training-script.py",  
)```
[Amazon SageMaker Developer Guide]

```python
role="SageMakerRole",
py_version="py310",
framework_version="2.0.1",

# For running a multi-node distributed training job, specify a value greater than 1
# Example: 2,3,4,..8
instance_count=2,

# Instance types supported by the SageMaker data parallel library:
# ml.p4d.24xlarge, ml.p4de.24xlarge
instance_type="ml.p4d.24xlarge",

# Activate distributed training with SMDDP
distribution={ "pytorchddp": { "enabled": True } }  # mpirun, activates SMDDP AllReduce OR AllGather
# distribution={ "torch_distributed": { "enabled": True } }  # torchrun, activates SMDDP AllGather
# distribution={ "smdistributed": { "dataparallel": { "enabled": True } } }  # mpirun, activates SMDDP AllReduce OR AllGather
)

pt_estimator.fit("s3://bucket/path/to/training/data")
```

**Note**

PyTorch Lightning and its utility libraries such as Lightning Bolts are not preinstalled in the SageMaker PyTorch DLCs. Create the following `requirements.txt` file and save in the source directory where you save the training script.

```text
# requirements.txt
pytorch-lightning
lightning-bolts
```

For example, the tree-structured directory should look like the following.

```text
### pytorch_training_launcher_jupyter_notebook.ipynb
### sub-folder-for-your-code
    ### adapted-training-script.py
### requirements.txt
```
For more information about specifying the source directory to place the requirements.txt file along with your training script and a job submission, see Using third-party libraries in the Amazon SageMaker Python SDK documentation.

Considerations for activating SMDDP collective operations and using the right distributed training launcher options

- SMDDP AllReduce and SMDDP AllGather are not mutually compatible at present.
- SMDDP AllReduce is activated by default when using smdistributed or pytorchddp, which are mpirun-based launchers, and NCCL AllGather is used.
- SMDDP AllGather is activated by default when using torch_distributed launcher, and AllReduce falls back to NCCL.
- SMDDP AllGather can also be activated when using the mpirun-based launchers with an additional environment variable set as follows.

```bash
export SMDATAPARALLEL_OPTIMIZE_SDP=true
```

TensorFlow

⚠️ Important

The SMDDP library discontinued support for TensorFlow and is no longer available in DLCs for TensorFlow later than v2.11.0. To find previous TensorFlow DLCs with the SMDDP library installed, see the section called “TensorFlow (deprecated)”.

```python
from sagemaker.tensorflow import TensorFlow

tf_estimator = TensorFlow(
    base_job_name = "training_job_name_prefix",
    entry_point="adapted-training-script.py",
    role="SageMakerRole",
    framework_version="2.11.0",
    py_version="py38",
)```
# For running a multi-node distributed training job, specify a value greater
# than 1
# Example: 2,3,4,..8
instance_count=2,

# Instance types supported by the SageMaker data parallel library:
# ml.p4d.24xlarge, ml.p3dn.24xlarge, and ml.p3.16xlarge
instance_type="ml.p3.16xlarge",

# Training using the SageMaker data parallel distributed training strategy
distribution={ "smdistributed": { "dataparallel": { "enabled": True } } }

)  

tf_estimator.fit("s3://bucket/path/to/training/data")

### Using the SageMaker generic estimator to extend prebuilt containers

You can customize SageMaker prebuilt containers or extend them to handle any additional functional requirements for your algorithm or model that the prebuilt SageMaker Docker image doesn't support. For an example of how you can extend a pre-built container, see [Extend a Prebuilt Container](#).

To extend a prebuilt container or adapt your own container to use the library, you must use one of the images listed in [Supported frameworks](#).

**Note**

From TensorFlow 2.4.1 and PyTorch 1.8.1, SageMaker framework DLCs supports EFA-enabled instance types. We recommend that you use the DLC images that contain TensorFlow 2.4.1 or later and PyTorch 1.8.1 or later.

For example, if you use PyTorch, your Dockerfile should contain a FROM statement similar to the following:

```
# SageMaker PyTorch image
FROM 763104351884.dkr.ecr.<aws-region>.amazonaws.com/pytorch-training:<image-tag>

ENV PATH="/opt/ml/code:${PATH}"
```
You can further customize your own Docker container to work with SageMaker using the SageMaker Training toolkit and the binary file of the SageMaker distributed data parallel library. To learn more, see the instructions in the following section.

### Create your own Docker container with the SageMaker distributed data parallel library

To build your own Docker container for training and use the SageMaker data parallel library, you must include the correct dependencies and the binary files of the SageMaker distributed parallel libraries in your Dockerfile. This section provides instructions on how to create a complete Dockerfile with the minimum set of dependencies for distributed training in SageMaker using the data parallel library.

#### Note

This custom Docker option with the SageMaker data parallel library as a binary is available only for PyTorch.

### To create a Dockerfile with the SageMaker training toolkit and the data parallel library

1. Start with a Docker image from NVIDIA CUDA. Use the cuDNN developer versions that contain CUDA runtime and development tools (headers and libraries) to build from the PyTorch source code.

   ```
   FROM nvidia/cuda:11.3.1-cudnn8-devel-ubuntu20.04
   ```
Tip

The official AWS Deep Learning Container (DLC) images are built from the NVIDIA CUDA base images. If you want to use the prebuilt DLC images as references while following the rest of the instructions, see the AWS Deep Learning Containers for PyTorch Dockerfiles.

2. Add the following arguments to specify versions of PyTorch and other packages. Also, indicate the Amazon S3 bucket paths to the SageMaker data parallel library and other software to use AWS resources, such as the Amazon S3 plug-in.

To use versions of the third party libraries other than the ones provided in the following code example, we recommend you look into the official Dockerfiles of AWS Deep Learning Container for PyTorch to find versions that are tested, compatible, and suitable for your application.

To find URLs for the SMDATAPARALLEL_BINARY argument, see the lookup tables at Supported frameworks.

```
ARG PYTORCH_VERSION=1.10.2
ARG PYTHON_SHORT_VERSION=3.8
ARG EFA_VERSION=1.14.1
ARG SMDATAPARALLEL_BINARY=https://smdataparallel.s3.amazonaws.com/binary/pytorch/
  ${PYTORCH_VERSION}/cu113/2022-02-18/smdistributed_dataparallel-1.4.0-cp38-cp38-
  linux_x86_64.whl
ARG PT_S3_WHL_GPU=https://aws-s3-plugin.s3.us-west-2.amazonaws.com/
  binaries/0.0.1/1c3e69e/awsio-0.0.1-cp38-cp38-manylinux1_x86_64.whl
ARG CONDA_PREFIX="/opt/conda"
ARG BRANCH_OFI=1.1.3-aws
```

3. Set the following environment variables to properly build SageMaker training components and run the data parallel library. You use these variables for the components in the subsequent steps.

```
# Set ENV variables required to build PyTorch
ENV TORCH_CUDA_ARCH_LIST="7.0+PTX 8.0"
ENV TORCH_NVCC_FLAGS="-Xfatbin -compress-all"
ENV NCCL_VERSION=2.10.3

# Add OpenMPI to the path.
ENV PATH /opt/amazon/openmpi/bin:$PATH
```
# Add Conda to path
ENV PATH $CONDA_PREFIX/bin:$PATH

# Set this environment variable for SageMaker to launch SMDDP correctly.
ENV SAGEMAKER_TRAINING_MODULE=sagemaker_pytorch_container.training:main

# Add environment variable for processes to be able to call fork()
ENV RDMAV_FORK_SAFE=1

# Indicate the container type
ENV DLC_CONTAINER_TYPE=training

# Add EFA and SMDDP to LD library path
ENV LD_LIBRARY_PATH="/opt/conda/lib/python${PYTHON_SHORT_VERSION}/site-packages/smdistributed/dataparallel/lib:$LD_LIBRARY_PATH"
ENV LD_LIBRARY_PATH=/opt/amazon/efa/lib/:$LD_LIBRARY_PATH

4. Install or update curl, wget, and git to download and build packages in the subsequent steps.

RUN --mount=type=cache,id=apt-final,target=/var/cache/apt \
    apt-get update && apt-get install -y --no-install-recommends \
    curl \ 
    wget \ 
    git \ 
    && rm -rf /var/lib/apt/lists/*

5. Install Elastic Fabric Adapter (EFA) software for Amazon EC2 network communication.

RUN DEBIAN_FRONTEND=noninteractive apt-get update
RUN mkdir /tmp/efa \ 
    && cd /tmp/efa \ 
    && curl --silent -- -O https://efa-installer.amazonaws.com/aws-efa-installer-\ 
$[EFA_VERSION].tar.gz \ 
    && tar -xf aws-efa-installer-\$[EFA_VERSION].tar.gz \ 
    && cd aws-efa-installer \ 
    && ./efa_installer.sh -y --skip-kmod -g \ 
    && rm -rf /tmp/efa

6. Install Conda to handle package management.

RUN curl -fsSL -v -o ~/miniconda.sh -O https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh && 

7. Get, build, and install PyTorch and its dependencies. We build PyTorch from the source code because we need to have control of the NCCL version to guarantee compatibility with the AWS OFI NCCL plug-in.
   a. Following the steps in the PyTorch official dockerfile, install build dependencies and set up ccache to speed up recompilation.

```bash
RUN DEBIAN_FRONTEND=noninteractive \
    apt-get install -y --no-install-recommends \
    build-essential \ 
    ca-certificates \ 
    ccache \ 
    cmake \ 
    git \ 
    libjpeg-dev \ 
    libpng-dev \ 
    && rm -rf /var/lib/apt/lists/*

# Setup ccache
RUN /usr/sbin/update-ccache-symlinks
RUN mkdir /opt/ccache && ccache --set-config=cache_dir=/opt/ccache

b. Install PyTorch’s common and Linux dependencies.

```bash
# Common dependencies for PyTorch
RUN conda install astunparse numpy ninja pyyaml mkl mkl-include setuptools cmake cffi typing_extensions future six requests dataclasses

# Linux specific dependency for PyTorch
RUN conda install -c pytorch magma-cuda113

c. Clone the PyTorch GitHub repository.

```bash
RUN --mount=type=cache,target=/opt/ccache \
    cd / \
```

SageMaker distributed data parallelism library
d. Install and build a specific NCCL version. To do this, replace the content in the PyTorch's default NCCL folder (/pytorch/third_party/nccl) with the specific NCCL version from the NVIDIA repository. The NCCL version was set in the step 3 of this guide.

```bash
RUN cd /pytorch/third_party/nccl \
    && rm -rf nccl \
    && git clone https://github.com/NVIDIA/nccl.git -b v${NCCL_VERSION}-1 \
    && cd nccl \
    && make -j64 src.build CUDA_HOME=/usr/local/cuda NVCC_GENCODE="-gencode=arch=compute_70,code=sm_70 -gencode=arch=compute_80,code=sm_80" \
    && make pkg.txz.build \
    && tar -xvf build/pkg/txz/nccl_*.txz -C $CONDA_PREFIX --strip-components=1
```

e. Build and install PyTorch. This process usually takes slightly more than 1 hour to complete. It is built using the NCCL version downloaded in a previous step.

```bash
RUN cd /pytorch \
    && CMAKE_PREFIX_PATH="$(dirname $(which conda))/../" \
    python setup.py install \
    && rm -rf /pytorch
```

8. Build and install AWS OFI NCCL plugin. This enables libfabric support for the SageMaker data parallel library.

```bash
RUN DEBIAN_FRONTEND=noninteractive apt-get update \
    && apt-get install -y --no-install-recommends \
    autoconf \n    automake \n    libtool
RUN mkdir /tmp/efa-ofi-nccl \
    && cd /tmp/efa-ofi-nccl \
    && git clone https://github.com/aws/aws-ofi-nccl.git -b v${BRANCH_OFI} \
    && cd aws-ofi-nccl \
    && ./autogen.sh \
    && ./configure --with-libfabric=/opt/amazon/efa \
    --with-mpi=/opt/amazon/openmpi \
    --with-cuda=/usr/local/cuda \
    --with-nccl=$CONDA_PREFIX \
    && make \
```

```bash
RUN pip install --no-cache-dir -U \n    packaging \n    mpi4py==3.0.3
RUN cd /tmp \n    && git clone https://github.com/pytorch/vision.git -b v0.9.1 \n    && cd vision \n    && BUILD_VERSION="0.9.1+cu111" python setup.py install \n    && cd /tmp \n    && rm -rf vision
```

10. Install and configure OpenSSH. OpenSSH is required for MPI to communicate between containers. Allow OpenSSH to talk to containers without asking for confirmation.

```bash
RUN apt-get update \n    && apt-get install -y --allow-downgrades --allow-change-held-packages --no-install-recommends \n    && apt-get install -y --no-install-recommends openssh-client openssh-server \n    && mkdir -p /var/run/sshd \n    && cat /etc/ssh/ssh_config | grep -v StrictHostKeyChecking > /etc/ssh/ssh_config.new \n    && echo "    StrictHostKeyChecking no" >> /etc/ssh/ssh_config.new \n    && mv /etc/ssh/ssh_config.new /etc/ssh/ssh_config \n    && rm -rf /var/lib/apt/lists/*

# Configure OpenSSH so that nodes can communicate with each other
RUN mkdir -p /var/run/sshd && \n    sed 's@session\s*required\s*pam_loginuid.so@session optional pam_loginuid.so@g' -i /etc/pam.d/sshd
RUN rm -rf /root/.ssh/ && \n    mkdir -p /root/.ssh/ && \n    ssh-keygen -q -t rsa -N '' -f /root/.ssh/id_rsa && \n    cp /root/.ssh/id_rsa.pub /root/.ssh/authorized_keys && \n    printf "Host *\n    StrictHostKeyChecking no
    " >> /root/.ssh/config
```

11. Install the PT S3 plug-in to efficiently access datasets in Amazon S3.

```bash
RUN pip install --no-cache-dir -U ${PT_S3_WHL_GPU}
```
12. Install the `libboost` library. This package is needed for networking the asynchronous IO functionality of the SageMaker data parallel library.

```bash
WORKDIR /
RUN wget https://sourceforge.net/projects/boost/files/boost/1.73.0/
    boost_1_73_0.tar.gz/download -O boost_1_73_0.tar.gz \
    && tar -xzf boost_1_73_0.tar.gz \
    && cd boost_1_73_0 \
    && ./bootstrap.sh \
    && ./b2 threading=multi --prefix=${CONDA_PREFIX} --prefix=${CONDA_PREFIX} -j 64 cxxflags=-fPIC cflags=-fPIC install || true \
    && cd .. \
    && rm -rf boost_1_73_0.tar.gz \
    && rm -rf boost_1_73_0 \
    && cd ${CONDA_PREFIX}/include/boost
```

13. Install the following SageMaker tools for PyTorch training.

```bash
WORKDIR /root
RUN pip install --no-cache-dir -U  
    smclarify \
    "sagemaker>=2,<3" \
    sagemaker-experiments==0.* \
    sagemaker-pytorch-training
```

14. Finally, install the SageMaker data parallel binary and the remaining dependencies.

```bash
RUN --mount=type=cache,id=apt-final,target=/var/cache/apt \
    apt-get update && apt-get install -y --no-install-recommends \
    jq \
    libhwloc-dev \
    libnuma1 \
    libnuma-dev \
    libssl1.1 \
    libtool \
    hwloc  \
    && rm -rf /var/lib/apt/lists/*

RUN SMDATAPARALLEL_PT=1 pip install --no-cache-dir ${SMDATAPARALLEL_BINARY}
```
After you finish creating the Dockerfile, see Adapting Your Own Training Container to learn how to build the Docker container, host it in Amazon ECR, and run a training job using the SageMaker Python SDK.

The following example code shows a complete Dockerfile after combining all the previous code blocks.

```bash
# This file creates a docker image with minimum dependencies to run SageMaker data parallel training
FROM nvidia/cuda:11.3.1-cudnn8-devel-ubuntu20.04

# Set appropriate versions and location for components
ARG PYTORCH_VERSION=1.10.2
ARG PYTHON_SHORT_VERSION=3.8
ARG EFA_VERSION=1.14.1
ARG SMDATAPARALLEL_BINARY=https://smdataparallel.s3.amazonaws.com/binary/pytorch/
${PYTORCH_VERSION}/cu113/2022-02-18/smdistributed_dataparallel-1.4.0-cp38-cp38-linux_x86_64.whl
ARG PT_S3_WHL_GPU=https://aws-s3-plugin.s3.us-west-2.amazonaws.com/binaries/0.0.1/1c3e69e/awsio-0.0.1-cp38-cp38-manylinux1_x86_64.whl
ARG CONDA_PREFIX="/opt/conda"
ARG BRANCH_OFI=1.1.3-aws

# Set ENV variables required to build PyTorch
ENV TORCH_CUDA_ARCH_LIST="3.7 5.0 7.0+PTX 8.0"
ENV TORCH_NVCC_FLAGS="-Xfatbin -compress-all"
ENV NCCL_VERSION=2.10.3

# Add OpenMPI to the path.
ENV PATH /opt/amazon/openmpi/bin:$PATH

# Add Conda to path
ENV PATH $CONDA_PREFIX/bin:$PATH

# Set this enviroment variable for SageMaker to launch SMDDP correctly.
ENV SAGEMAKER_TRAINING_MODULE=sagemaker_pytorch_container.training:main

# Add enviroment variable for processes to be able to call fork()
ENV RDMAV_FORK_SAFE=1

# Indicate the container type
ENV DLC_CONTAINER_TYPE=training
```
# Add EFA and SMDDP to LD library path
ENV LD_LIBRARY_PATH="/opt/conda/lib/python${PYTHON_SHORT_VERSION}/site-packages/
 smdistributed/dataparallel/lib:$LD_LIBRARY_PATH"
ENV LD_LIBRARY_PATH=/opt/amazon/efa/lib/:$LD_LIBRARY_PATH

# Install basic dependencies to download and build other dependencies
RUN --mount=type=cache,id=apt-final,target=/var/cache/apt \
    apt-get update && apt-get install -y --no-install-recommends  \
    curl \ 
    wget \ 
    git \ 
    && rm -rf /var/lib/apt/lists/*

# Install EFA.  
# This is required for SMDDP backend communication
RUN DEBIAN_FRONTEND=noninteractive apt-get update
RUN mkdir /tmp/efa \ 
    && cd /tmp/efa \ 
    && curl --silent -O https://efa-installer.amazonaws.com/aws-efa-installer-
    ${EFA_VERSION}.tar.gz \ 
    && tar -xf aws-efa-installer-${EFA_VERSION}.tar.gz \ 
    && cd aws-efa-installer \ 
    && ./efa_installer.sh -y --skip-kmod -g \ 
    && rm -rf /tmp/efa

# Install Conda
RUN curl -fsSL -v -o ~/miniconda.sh -O https://repo.anaconda.com/miniconda/Miniconda3-
    latest-Linux-x86_64.sh && \ 
    chmod +x ~/miniconda.sh && \ 
    ~/miniconda.sh -b -p $CONDA_PREFIX && \ 
    rm ~/miniconda.sh && \ 
    $CONDA_PREFIX/bin/conda install -y python=${PYTHON_SHORT_VERSION} conda-build
    pyyaml numpy ipython && 
    $CONDA_PREFIX/bin/conda clean -ya

# Install PyTorch.
# Start with dependencies listed in official PyTorch dockerfile
# https://github.com/pytorch/pytorch/blob/master/Dockerfile
RUN DEBIAN_FRONTEND=noninteractive \
    apt-get install -y --no-install-recommends \
    build-essential \ 
    ca-certificates \ 
    ccache \
cmake  
  git  
  libjpeg-dev  
  libpng-dev & &  
  rm -rf /var/lib/apt/lists/*

# Setup ccache  
RUN /usr/sbin/update-ccache-symlinks  
RUN mkdir /opt/ccache & & ccache --set-config=cache_dir=/opt/ccache

# Common dependencies for PyTorch  
RUN conda install astunparse numpy ninja pyyaml mkl mkl-include setuptools cmake cffi typing_extensions future six requests dataclasses

# Linux specific dependency for PyTorch  
RUN conda install -c pytorch magma-cuda113

# Clone PyTorch  
RUN --mount=type=cache,target=/opt/ccache \    
    cd / \    
    & & git clone --recursive https://github.com/pytorch/pytorch -b v$PYTORCH_VERSION

# Note that we need to use the same NCCL version for PyTorch and OFI plugin.  
# To enforce that, install NCCL from source before building PT and OFI plugin.

# Install NCCL.  
# Required for building OFI plugin (OFI requires NCCL's header files and library)  
RUN cd /pytorch/third_party/nccl \    
    & & rm -rf nccl \    
    & & git clone https://github.com/NVIDIA/nccl.git -b v${NCCL_VERSION}-1 \    
    & & cd nccl \    
    & & make -j64 src.build CUDA_HOME=/usr/local/cuda NVCC_GENCODE="-genencode=arch=compute_70,code=sm_70 -genencode=arch=compute_80,code=sm_80" \    
    & & make pkg.txz.build \    
    & & tar -xvf build/pkg/txz/nccl_* .txz -C $CONDA_PREFIX --strip-components=1

# Build and install PyTorch.  
RUN cd /pytorch \    
    & & CMAKE_PREFIX_PATH="$(dirname $(which conda))/../" \    
    python setup.py install \    
    & & rm -rf /pytorch

RUN ccache -C

# Build and install OFI plugin. 

SageMaker distributed data parallelism library
It is required to use libfabric.

```
RUN DEBIAN_FRONTEND=noninteractive apt-get update \
    && apt-get install -y --no-install-recommends \
        autoconf \n        automake \n        libtool
RUN mkdir /tmp/efa-ofi-nccl \n    && cd /tmp/efa-ofi-nccl \n    && git clone https://github.com/aws/aws-ofi-nccl.git -b v${BRANCH_OFI} \n    && cd aws-ofi-nccl \n    && ./autogen.sh \n    && ./configure --with-libfabric=/opt/amazon/efa \n        --with-mpi=/opt/amazon/openmpi \n        --with-cuda=/usr/local/cuda \n        --with-nccl=$CONDA_PREFIX 
    && make \n    && make install \n    && rm -rf /tmp/efa-ofi-nccl

# Build and install Torchvision
RUN pip install --no-cache-dir -U \
    packaging \n    mpi4py==3.0.3
RUN cd /tmp \n    && git clone https://github.com/pytorch/vision.git -b v0.9.1 \n    && cd vision \n    && BUILD_VERSION="0.9.1+cu111" python setup.py install \n    && cd /tmp \n    && rm -rf vision

# Install OpenSSH.
# Required for MPI to communicate between containers, allow OpenSSH to talk to
# containers without asking for confirmation
RUN apt-get update \
    && apt-get install -y --allow-downgrades --allow-change-held-packages --no-
    install-recommends \
    && apt-get install -y --no-install-recommends openssh-client openssh-server \
    && mkdir -p /var/run/sshd \n    && cat /etc/ssh/ssh_config | grep -v StrictHostKeyChecking > /etc/ssh/
    ssh_config.new \n    && echo "" StrictHostKeyChecking no >> /etc/ssh/ssh_config.new \n    && mv /etc/ssh/ssh_config.new /etc/ssh/ssh_config \n    && rm -rf /var/lib/apt/lists/*
# Configure OpenSSH so that nodes can communicate with each other
```
RUN mkdir -p /var/run/sshd && 
    sed '@session@s*required@session optional pam_loginuid.so' -i /etc/pam.d/sshd
RUN rm -rf /root/.ssh/ && 
    mkdir -p /root/.ssh/ && 
    ssh-keygen -q -t rsa -N '@2@' -f /root/.ssh/id_rsa && 
    cp /root/.ssh/id_rsa.pub /root/.ssh/authorized_keys && 
    printf "Host * \n StrictHostKeyChecking no\n" >> /root/.ssh/config

# Install PT S3 plugin.
# Required to efficiently access datasets in Amazon S3
RUN pip install --no-cache-dir -U ${PT_S3_WHL_GPU}
RUN mkdir -p /etc/pki/tls/certs && cp /etc/ssl/certs/ca-certificates.crt /etc/pki/tls/certs/ca-bundle.crt

# Install libboost from source.
# This package is needed for smdataparallel functionality (for networking asynchronous IO).
WORKDIR /
RUN wget https://sourceforge.net/projects/boost/files/boost/1.73.0/boost_1_73_0.tar.gz/
download -0 boost_1_73_0.tar.gz 
    && tar -xzf boost_1_73_0.tar.gz 
    && cd boost_1_73_0 
    && ./bootstrap.sh 
    && ./b2 threading=multi --prefix=${CONDA_PREFIX} -j 64 cxxflags=-fPIC cflags=-fPIC install || true 
    && cd .. 
    && rm -rf boost_1_73_0.tar.gz 
    && rm -rf boost_1_73_0 
    && cd ${CONDA_PREFIX}/include/boost

# Install SageMaker PyTorch training.
WORKDIR /root
RUN pip install --no-cache-dir -U 
    smclarify 
    "sagemaker>=2,<3" 
    sagemaker-experiments==0.* 
    sagemaker-pytorch-training

# Install SageMaker data parallel binary (SMDDP)
# Start with dependencies
RUN --mount=type=cache,id=apt-final,target=/var/cache/apt 
    apt-get update && apt-get install -y --no-install-recomends 
    jq \

SageMaker distributed data parallelism library
Amazon SageMaker data parallelism library examples

This page provides Jupyter notebooks that present examples of implementing the SageMaker distributed data parallelism (SMDDP) library to run distributed training jobs on SageMaker.

Blogs and Case Studies

The following blogs discuss case studies about using the SMDDP library.

SMDDP v2 blogs


SMDDP v1 blogs
- How I trained 10TB for Stable Diffusion on SageMaker in Medium (November 29, 2022)
- Run PyTorch Lightning and native PyTorch DDP on Amazon SageMaker Training, featuring Amazon Search, AWS Machine Learning Blog (August 18, 2022)
- Training YOLOv5 on AWS with PyTorch and the SageMaker distributed data parallel library, Medium (May 6, 2022)
- Speed up EfficientNet model training on SageMaker with PyTorch and the SageMaker distributed data parallel library, Medium (March 21, 2022)
- Speed up EfficientNet training on AWS with the SageMaker distributed data parallel library, Towards Data Science (January 12, 2022)
- Hyundai reduces ML model training time for autonomous driving models using Amazon SageMaker, AWS Machine Learning Blog (June 25, 2021)
- Distributed Training: Train BART/T5 for Summarization using Transformers and Amazon SageMaker, the Hugging Face website (April 8, 2021)

Example notebooks

Example notebooks are provided in the SageMaker examples GitHub repository. To download the examples, run the following command to clone the repository and go to training/distributed_training/pytorch/data_parallel.

```
git clone https://github.com/aws/amazon-sagemaker-examples.git
cd amazon-sagemaker-examples/training/distributed_training/pytorch/data_parallel
```

**Note**

Clone and run the example notebooks in the following SageMaker ML IDEs.

- SageMaker JupyterLab (available in Studio created after December 2023)
- SageMaker Code Editor (available in Studio created after December 2023)
- Studio Classic (available as an application in Studio created after December 2023)
- SageMaker Notebook Instances

SMDDP v2 examples

- Train Llama 2 using the SageMaker distributed data parallel library (SMDDP) and DeepSpeed
• **Train Falcon using the SageMaker distributed data parallel library (SMDDP) and PyTorch Fully Sharded Data Parallelism (FSDP)**

**SMDDP v1 examples**

• **CNN with PyTorch and the SageMaker data parallelism library**
• **BERT with PyTorch and the SageMaker data parallelism library**
• **CNN with TensorFlow 2.3.1 and the SageMaker data parallelism library**
• **BERT with TensorFlow 2.3.1 and the SageMaker data parallelism library**
• **HuggingFace Distributed Data Parallel Training in PyTorch on SageMaker - Distributed Question Answering**
• **HuggingFace Distributed Data Parallel Training in PyTorch on SageMaker - Distributed Text Summarization**
• **HuggingFace Distributed Data Parallel Training in TensorFlow on SageMaker**

**Configuration tips for the SageMaker distributed data parallelism library**

Review the following tips before using the SageMaker distributed data parallelism (SMDDP) library. This list includes tips that are applicable across frameworks.

**Topics**

• **Data preprocessing**
• **Single versus multiple nodes**
• **Debug scaling efficiency with Debugger**
• **Batch size**
• **Custom MPI options**
• **Use Amazon FSx and set up an optimal storage and throughput capacity**

**Data preprocessing**

If you preprocess data during training using an external library that utilizes the CPU, you may run into a CPU bottleneck because SageMaker distributed data parallel uses the CPU for AllReduce operations. You may be able to improve training time by moving preprocessing steps to a library that uses GPUs or by completing all preprocessing before training.
Single versus multiple nodes

We recommend that you use this library with multiple nodes. The library can be used with a single-host, multi-device setup (for example, a single ML compute instance with multiple GPUs); however, when you use two or more nodes, the library's AllReduce operation gives you significant performance improvement. Also, on a single host, NVLink already contributes to in-node AllReduce efficiency.

Debug scaling efficiency with Debugger

You can use Amazon SageMaker Debugger to monitor and visualize CPU and GPU utilization and other metrics of interest during training. You can use Debugger built-in rules to monitor computational performance issues, such as CPUBottleneck, LoadBalancing, and LowGPUUtilization. You can specify these rules with Debugger configurations when you define an Amazon SageMaker Python SDK estimator. If you use AWS CLI and AWS SDK for Python (Boto3) for training on SageMaker, you can enable Debugger as shown in Configure SageMaker Debugger Using Amazon SageMaker API.

To see an example using Debugger in a SageMaker training job, you can reference one of the notebook examples in the SageMaker Notebook Examples GitHub repository. To learn more about Debugger, see Amazon SageMaker Debugger.

Batch size

In distributed training, as more nodes are added, batch sizes should increase proportionally. To improve convergence speed as you add more nodes to your training job and increase the global batch size, increase the learning rate.

One way to achieve this is by using a gradual learning rate warmup where the learning rate is ramped up from a small to a large value as the training job progresses. This ramp avoids a sudden increase of the learning rate, allowing healthy convergence at the start of training. For example, you can use a Linear Scaling Rule where each time the mini-batch size is multiplied by k, the learning rate is also multiplied by k. To learn more about this technique, see the research paper, Accurate, Large Minibatch SGD: Training ImageNet in 1 Hour, Sections 2 and 3.

Custom MPI options

The SageMaker distributed data parallel library employs Message Passing Interface (MPI), a popular standard for managing communication between nodes in a high-performance cluster, and uses NVIDIA's NCCL library for GPU-level communication. When you use the data parallel library with
a TensorFlow or Pytorch Estimator, the respective container sets up the MPI environment and executes the `mpirun` command to start jobs on the cluster nodes.

You can set custom MPI operations using the `custom_mpi_options` parameter in the Estimator. Any `mpirun` flags passed in this field are added to the `mpirun` command and executed by SageMaker for training. For example, you may define the distribution parameter of an Estimator using the following to use the `NCCL_DEBUG` variable to print the NCCL version at the start of the program:

```python
distribution = {'smdistributed': {'dataparallel': {'enabled': True, "custom_mpi_options": "-verbose -x NCCL_DEBUG=VERSION"}}}
```

**Use Amazon FSx and set up an optimal storage and throughput capacity**

When training a model on multiple nodes with distributed data parallelism, it is highly recommended to use FSx for Lustre. Amazon FSx is a scalable and high-performance storage service that supports shared file storage with a faster throughput. Using Amazon FSx storage at scale, you can achieve a faster data loading speed across the compute nodes.

Typically, with distributed data parallelism, you would expect that the total training throughput scales near-linearly with the number of GPUs. However, if you use suboptimal Amazon FSx storage, the training performance might slow down due to a low Amazon FSx throughput.

For example, if you use the SCRATCH_2 deployment type of Amazon FSx file system with the minimum 1.2 TiB storage capacity, the I/O throughput capacity is 240 MB/s. Amazon FSx storage works in a way that you can assign physical storage devices, and the more devices assigned, the larger throughput you get. The smallest storage increment for the SRATCH_2 type is 1.2 TiB, and the corresponding throughput gain is 240 MB/s.

Assume that you have a model to train on a 4-node cluster over a 100 GB data set. With a given batch size that’s optimized to the cluster, assume that the model can complete one epoch in about 30 seconds. In this case, the minimum required I/O speed is approximately 3 GB/s (100 GB / 30 s). This is apparently a much higher throughput requirement than 240 MB/s. With such a limited Amazon FSx capacity, scaling your distributed training job up to larger clusters might aggravate I/O bottleneck problems; model training throughput might improve in later epochs as cache builds up, but Amazon FSx throughput can still be a bottleneck.

To alleviate such I/O bottleneck problems, you should increase the Amazon FSx storage size to obtain a higher throughput capacity. Typically, to find an optimal I/O throughput, you may experiment with different Amazon FSx throughput capacities, assigning an equal to or slightly
lower throughput than your estimate, until you find that it is sufficient to resolve the I/O bottleneck problems. In case of the aforementioned example, Amazon FSx storage with 2.4 GB/s throughput and 67 GB RAM cache would be sufficient. If the file system has an optimal throughput, the model training throughput should reach maximum either immediately or after the first epoch as cache has built up.

To learn more about how to increase Amazon FSx storage and deployment types, see the following pages in the *Amazon FSx for Lustre documentation*:

- [How to increase storage capacity](#)
- [Aggregate file system performance](#)

## Amazon SageMaker distributed data parallelism library FAQ

Use the following to find answers to commonly asked questions about the SMDDP library.

**Q: When using the library, how are the allreduce-supporting CPU instances managed? Do I have to create heterogeneous CPU-GPU clusters, or does the SageMaker service create extra C5s for jobs that use the SMDDP library?**

The SMDDP library only supports GPU instances, more specifically, P4d and P4de instances with NVIDIA A100 GPUs and EFA. No additional C5 or CPU instances are launched; if your SageMaker training job is on an 8-node P4d cluster, only 8 `ml.p4d.24xlarge` instances are used. No additional instances are provisioned.

**Q: I have a training job taking 5 days on a single `ml.p3.24xlarge` instance with a set of hyperparameters H1 (learning rate, batch size, optimizer, etc). Is using SageMaker's data parallelism library and a five-time bigger cluster enough to achieve an approximate five-time speedup? Or do I have to revisit its training hyperparameters after activating the SMDDP library?**

The library changes the overall batch size. The new overall batch size is scaled linearly with the number of training instances used. As a result of this, hyperparameters, such as learning rate, have to be changed to ensure convergence.

**Q: Does the SMDDP library support Spot?**

Yes. You can use managed spot training. You specify the path to the checkpoint file in the SageMaker training job. You save and restore checkpoints in their training script as mentioned in the last steps of [the section called “TensorFlow (deprecated)”](#) and [the section called “PyTorch”](#).
Q: Is the SMDDP library relevant in a single-host, multi-device setup?

The library can be used in single-host multi-device training but the library offers performance improvements only in multi-host training.

Q: Where should the training dataset be stored?

The training dataset can be stored in an Amazon S3 bucket or on an Amazon FSx drive. See this document for various supported input file systems for a training job.

Q: When using the SMDDP library, is it mandatory to have training data in FSx for Lustre? Can Amazon EFS and Amazon S3 be used?

We generally recommend you use Amazon FSx because of its lower latency and higher throughput. If you prefer, you can use Amazon EFS or Amazon S3.

Q: Can the library be used with CPU nodes?

No. To find instance types supported by the SMDDP library, see the section called “Supported instance types”.

Q: What frameworks and framework versions are currently supported by the SMDDP library at launch?

The SMDDP library currently supports PyTorch v1.6.0 or later and TensorFlow v2.3.0 or later. It doesn't support TensorFlow 1.x. For more information about which version of the SMDDP library is packaged within AWS deep learning containers, see Release Notes for Deep Learning Containers.

Q: Does the library support AMP?

Yes, the SMDDP library supports Automatic Mixed Precision (AMP) out of the box. No extra action is needed to use AMP other than the framework-level modifications to your training script. If gradients are in FP16, the SageMaker data parallelism library runs its AllReduce operation in FP16. For more information about implementing AMP APIs to your training script, see the following resources:

- Frameworks - PyTorch in the NVIDIA Deep Learning Performance documentation
- Frameworks - TensorFlow in the NVIDIA Deep Learning Performance documentation
- Automatic Mixed Precision for Deep Learning in the NVIDIA Developer Docs
- Introducing native PyTorch automatic mixed precision for faster training on NVIDIA GPUs in the PyTorch Blog
• TensorFlow mixed precision APIs in the TensorFlow documentation

Q: How do I identify if my distributed training job is slowed down due to I/O bottleneck?

With a larger cluster, the training job requires more I/O throughput, and therefore the training throughput might take longer (more epochs) to ramp up to the maximum performance. This indicates that I/O is being bottlenecked and cache is harder to build up as you scale nodes up (higher throughput requirement and more complex network topology). For more information about monitoring the Amazon FSx throughput on CloudWatch, see Monitoring FSx for Lustre in the FSx for Lustre User Guide.

Q: How do I resolve I/O bottlenecks when running a distributed training job with data parallelism?

We highly recommend that you use Amazon FSx as your data channel if you are using Amazon S3. If you are already using Amazon FSx but still having I/O bottleneck problems, you might have set up your Amazon FSx file system with a low I/O throughput and a small storage capacity. For more information about how to estimate and choose the right size of I/O throughput capacity, see Use Amazon FSx and set up an optimal storage and throughput capacity.

Q: (For the library v1.4.0 or later) How do I resolve the Invalid backend error while initializing process group.

If you encounter the error message `ValueError: Invalid backend: 'smddp'` when calling `init_process_group`, this is due to the breaking change in the SMDDP library v1.4.0 and later. You must import the PyTorch client of the library, `smdistributed.dataparallel.torch.torch_smddp`, which registers `smddp` as a backend for PyTorch. To learn more, see the section called “PyTorch”.

Q: (For the SMDDP library v1.4.0 or later) I would like to call the collective primitives of the torch.distributed interface. Which primitives does the smddp backend support?

In v1.4.0, the SMDDP library supports `all_reduce`, `broadcast`, `reduce`, `all_gather`, and `barrier` of the `torch.distributed` interface.

Q: (For the SMDDP library v1.4.0 or later) Does this new API work with other custom DDP classes or libraries like Apex DDP?

The SMDDP library is tested with other third-party distributed data parallel libraries and framework implementations that use the `torch.distributed` modules. Using the SMDDP library
with custom DDP classes works as long as the collective operations used by the custom DDP classes are supported by the SMDDP library. See the preceding question for a list of supported collectives. If you have these use cases and need further support, reach out to the SageMaker team through the AWS Support Center or AWS Developer Forums for Amazon SageMaker.

Q: Does the SMDDP library support the bring-your-own-container (BYOC) option? If so, how do I install the library and run a distributed training job by writing a custom Dockerfile?

If you want to integrate the SMDDP library and its minimum dependencies into your own Docker container, BYOC is the right approach. You can build your own container using the binary file of the library. The recommended process is to write a custom Dockerfile with the library and its dependencies, build the Docker container, host it in Amazon ECR, and use the ECR image URI to launch a training job using the SageMaker generic estimator class. For more instructions on how to prepare a custom Dockerfile for distributed training in SageMaker with the SMDDP library, see Create your own Docker container with the SageMaker distributed data parallel library.

Troubleshooting for distributed training in Amazon SageMaker

If you have problems in running a training job when you use the library, use the following list to try to troubleshoot. If you need further support, reach out to the SageMaker team through AWS Support Center or AWS Developer Forums for Amazon SageMaker.

Topics

• Using SageMaker distributed data parallel with Amazon SageMaker Debugger and checkpoints
• An unexpected prefix attached to model parameter keys
• SageMaker distributed training job stalling during initialization
• SageMaker distributed training job stalling at the end of training
• Observing scaling efficiency degradation due to Amazon FSx throughput bottlenecks
• SageMaker distributed training job with PyTorch returns deprecation warnings

Using SageMaker distributed data parallel with Amazon SageMaker Debugger and checkpoints

To monitor system bottlenecks, profile framework operations, and debug model output tensors for training jobs with SageMaker distributed data parallel, use Amazon SageMaker Debugger.

However, when you use SageMaker Debugger, SageMaker distributed data parallel, and SageMaker checkpoints, you might see an error that looks like the following example.
SMDebug Does Not Currently Support Distributed Training Jobs With Checkpointing Enabled

This is due to an internal error between Debugger and checkpoints, which occurs when you enable SageMaker distributed data parallel.

- If you enable all three features, SageMaker Python SDK automatically turns off Debugger by passing `debugger_hook_config=False`, which is equivalent to the following framework estimator example.

```python
bucket=sagemaker.Session().default_bucket()
base_job_name="sagemaker-checkpoint-test"
checkpoint_in_bucket="checkpoints"

# The S3 URI to store the checkpoints
checkpoint_s3_bucket="s3://{}//{}//{}".format(bucket, base_job_name, 
                             checkpoint_in_bucket)

estimator = TensorFlow(
    ...
    distribution={"smdistributed": {"dataparallel": { "enabled": True }},
    checkpoint_s3_uri=checkpoint_s3_bucket,
    checkpoint_local_path="/opt/ml/checkpoints",
    debugger_hook_config=False
)
```

- If you want to keep using both SageMaker distributed data parallel and SageMaker Debugger, a workaround is manually adding checkpointing functions to your training script instead of specifying the `checkpoint_s3_uri` and `checkpoint_local_path` parameters from the estimator. For more information about setting up manual checkpointing in a training script, see Saving Checkpoints.

An unexpected prefix attached to model parameter keys

For PyTorch distributed training jobs, an unexpected prefix (model for example) might be attached to `state_dict` keys (model parameters). The SageMaker data parallel library does not directly alter or prepend any model parameter names when PyTorch training jobs save model artifacts. The PyTorch's distributed training changes the names in the `state_dict` to go over the network, prepending the prefix. If you encounter any model failure problem due to different parameter names while you are using the SageMaker data parallel library and checkpointing for PyTorch...
training, adapt the following example code to remove the prefix at the step you load checkpoints in your training script.

```python
state_dict = {k.partition('model.')[2]: state_dict[k] for k in state_dict.keys()}
```

This takes each `state_dict` key as a string value, separates the string at the first occurrence of 'model.', and takes the third list item (with index 2) of the partitioned string.

For more information about the prefix issue, see a discussion thread at Prefix parameter names in saved model if trained by multi-GPU? in the PyTorch discussion forum.

For more information about the PyTorch methods for saving and loading models, see Saving & Loading Model Across Devices in the PyTorch documentation.

**SageMaker distributed training job stalling during initialization**

If your SageMaker distributed data parallel training job stalls during initialization when using EFA-enabled instances, this might be due to a misconfiguration in the security group of the VPC subnet that's used for the training job. EFA requires a proper security group configuration to enable traffic between the nodes.

**To configure inbound and outbound rules for the security group**

1. Sign in to the AWS Management Console and open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
2. Choose Security Groups in the left navigation pane.
3. Select the security group that's tied to the VPC subnet you use for training.
4. In the Details section, copy the Security group ID.
6. On the Edit inbound rules page, do the following:
   a. Choose Add rule.
   b. For Type, choose All traffic.
   c. For Source, choose Custom, paste the security group ID into the search box, and select the security group that pops up.
7. Choose Save rules to finish configuring the inbound rule for the security group.
9. Repeat the step 6 and 7 to add the same rule as an outbound rule.
After you complete the preceding steps for configuring the security group with the inbound and outbound rules, re-run the training job and verify if the stalling issue is resolved.

For more information about configuring security groups for VPC and EFA, see Security groups for your VPC and Elastic Fabric Adapter.

**SageMaker distributed training job stalling at the end of training**

One of the root causes of stalling issues at the end of training is a mismatch in the number of batches that are processed per epoch across different ranks. All workers (GPUs) synchronize their local gradients in the backward pass to ensure they all have the same copy of the model at the end of the batch iteration. If the batch sizes are unevenly assigned to different worker groups during the final epoch of training, the training job stalls. For example, while a group of workers (group A) finishes processing all batches and exits the training loop, another group of workers (group B) starts processing another batch and still expects communication from group A to synchronize the gradients. This causes group B to wait for group A, which already completed training and does not have any gradients to synchronize.

Therefore, when setting up your training dataset, it is important that each worker gets the same number of data samples so that each worker goes through the same number of batches while training. Make sure each rank gets the same number of batches to avoid this stalling issue.

**Observing scaling efficiency degradation due to Amazon FSx throughput bottlenecks**

One potential cause of lowered scaling efficiency is the FSx throughput limit. If you observe a sudden drop in scaling efficiency when you switch to a larger training cluster, try using a larger FSx for Lustre file system with a higher throughput limit. For more information, see Aggregate file system performance and Managing storage and throughput capacity in the Amazon FSx for Lustre User Guide.

**SageMaker distributed training job with PyTorch returns deprecation warnings**

Since v1.4.0, the SageMaker distributed data parallelism library works as a backend of PyTorch distributed. Because of the breaking change of using the library with PyTorch, you might encounter a warning message that the smdistributed APIs for the PyTorch distributed package are deprecated. The warning message should be similar to the following:

```
smdistributed.dataparallel.torch.dist is deprecated in the SageMaker distributed data parallel library v1.4.0+
Please use torch.distributed and specify 'smddp' as a backend when initializing process group as follows:
```

SageMaker distributed data parallelism library
torch.distributed.init_process_group(backend='smddp')

For more information, see the library's API documentation at

In v1.4.0 and later, the library only needs to be imported once at the top of your training script and set as the backend during the PyTorch distributed initialization. With the single line of backend specification, you can keep your PyTorch training script unchanged and directly use the PyTorch distributed modules. See Use the SMDDP library in your PyTorch training script to learn about the breaking changes and the new way to use the library with PyTorch.

**SageMaker data parallelism library release notes**

See the following release notes to track the latest updates for the SageMaker distributed data parallelism (SMDDP) library.

**The SageMaker distributed data parallelism library v2.2.0**

*Date: March 4, 2024*

**New features**

- Added support for PyTorch v2.2.0 with CUDA v12.1.

**Integration into Docker containers distributed by the SageMaker model parallelism (SMP) library**

This version of the SMDDP library is migrated to the section called “SMP v2.2.0”.

658645717510.dkr.ecr.<region>.amazonaws.com/smdistributed-modelparallel:2.2.0-gpu-py310-cu121

For Regions where the SMP Docker images are available, see the section called “AWS Regions”.

**Binary file of this release**

You can download or install the library using the following URL.

https://smdataparallel.s3.amazonaws.com/binary/pytorch/2.2.0/cu121/2024-03-04/smdistributed_dataparallel-2.2.0-cp310-cp310-linux_x86_64.whl
The SageMaker distributed data parallelism library v2.1.0

Date: March 1, 2024

New features

• Added support for PyTorch v2.1.0 with CUDA v12.1.

Bug fixes

• Fixed the CPU memory leak issue in SMDDP v2.0.1.

Integration into SageMaker Framework Containers

This version of the SMDDP library passed benchmark testing and is migrated to the following SageMaker Framework Container.

• PyTorch v2.1.0

763104351884.dkr.ecr.<region>.amazonaws.com/pytorch-training:2.1.0-gpu-py310-cu121-ubuntu20.04-sagemaker

Integration into Docker containers distributed by the SageMaker model parallelism (SMP) library

This version of the SMDDP library is migrated to the section called “SMP v2.1.0”.

658645717510.dkr.ecr.<region>.amazonaws.com/smdistributed-modelparallel:2.1.2-gpu-py310-cu121

For Regions where the SMP Docker images are available, see the section called “AWS Regions”.

Binary file of this release

You can download or install the library using the following URL.

https://smdataparallel.s3.amazonaws.com/binary/pytorch/2.1.0/cu121/2024-02-04/smdistributed_dataparallel-2.1.0-cp310-cp310-linux_x86_64.whl
The SageMaker distributed data parallelism library v2.0.1

Date: December 7, 2023

New features

- Added a new SMDDP-implementation of AllGather collective operation optimized for AWS compute resources and network infrastructure. To learn more, see the section called “SMDDP AllGather collective operation”.
- The SMDDP AllGather collective operation is compatible with PyTorch FSDP and DeepSpeed. To learn more, see the section called “PyTorch”.
- Added support for PyTorch v2.0.1

Known issues

- There's a CPU memory leak issue from a gradual CPU memory increase while training with SMDDP AllReduce in DDP mode.

Integration into SageMaker Framework Containers

This version of the SMDDP library passed benchmark testing and is migrated to the following SageMaker Framework Container.

- PyTorch v2.0.1

    763104351884.dkr.ecr.<region>.amazonaws.com/pytorch-training:2.0.1-gpu-py310-cu118-ubuntu20.04-sagemaker

Binary file of this release

You can download or install the library using the following URL.

    https://smdataparallel.s3.amazonaws.com/binary/pytorch/2.0.1/cu118/2023-12-07/smdistributed_dataparallel-2.0.2-cp310-cp310-linux_x86_64.whl

Other changes

- Starting from this release, documentation for the SMDDP library is fully available in this Amazon SageMaker Developer Guide. In favor of the complete developer guide for SMDDP v2 housed in
the Amazon SageMaker Developer Guide, documentation for the additional reference for SMDDP v1.x in the SageMaker Python SDK documentation is no longer supported. If you still need SMP v1.x documentation, see the following snapshot of the documentation at SageMaker Python SDK v2.212.0 documentation.

SageMaker model parallelism library v2

Note
Since the release of the SageMaker model parallelism (SMP) library v2.0.0 on December 19, 2023, this documentation is renewed for the SMP library v2. For previous versions of the SMP library, see the section called “(Archived) SageMaker model parallelism library v1.x”.

The Amazon SageMaker model parallelism library is a capability of SageMaker that enables high performance and optimized large scale training on SageMaker accelerate compute instances. The the section called “Core features of SMP v2” include techniques and optimizations to accelerate and simplify large model training, such as hybrid sharded data parallelism, tensor parallelism, activation checkpointing, and activation offloading. You can use the SMP library to accelerate the training and fine-tuning of large language models (LLMs), large vision models (LVMs), and foundation models (FMs) with hundreds of billions of parameters.

The SageMaker model parallelism library v2 (SMP v2) aligns the library's APIs and methods with open source PyTorch Fully Sharded Data Parallelism (FSDP), which gives you the benefit of SMP performance optimizations with minimal code changes. With SMP v2, you can improve the computational performance of training a state-of-the-art large model on SageMaker by bringing your PyTorch FSDP training scripts to SageMaker.

You can use SMP v2 for the general SageMaker Training jobs and distributed training workloads on the section called “SageMaker HyperPod” clusters.

Topics
• Introduction to model parallelism
• Supported frameworks and AWS Regions
• Get started with the SageMaker model parallelism library v2
• Core features of the SageMaker model parallelism library v2
Introduction to model parallelism

Model parallelism is a distributed training method in which the deep learning (DL) model is partitioned across multiple GPUs and instances. The SageMaker model parallel library v2 (SMP v2) is compatible with the native PyTorch APIs and capabilities. This makes it convenient for you to adapt your PyTorch Fully Sharded Data Parallel (FSDP) training script to the SageMaker Training platform and take advantage of the performance improvement that SMP v2 provides.

This introduction page provides a high-level overview about model parallelism and a description of how it can help overcome issues that arise when training deep learning (DL) models that are typically very large in size. It also provides examples of what the SageMaker model parallel library offers to help manage model parallel strategies and memory consumption.

What is model parallelism?

Increasing the size of deep learning models (layers and parameters) yields better accuracy for complex tasks such as computer vision and natural language processing. However, there is a limit to the maximum model size you can fit in the memory of a single GPU. When training DL models, GPU memory limitations can be bottlenecks in the following ways:

- They limit the size of the model that you can train, because the memory footprint of a model scales proportionally to the number of parameters.
- They limit the per-GPU batch size during training, driving down GPU utilization and training efficiency.

To overcome the limitations associated with training a model on a single GPU, SageMaker provides the model parallel library to help distribute and train DL models efficiently on multiple compute nodes. Furthermore, with the library, you can achieve optimized distributed training using EFA-supported devices, which enhance the performance of inter-node communication with low latency, high throughput, and OS bypass.
Estimate memory requirements before using model parallelism

Before you use the SageMaker model parallel library, consider the following to get a sense of the memory requirements of training large DL models.

For a training job that uses automatic mixed precision such as float16 (FP16) or bfloat16 (BF16) and Adam optimizers, the required GPU memory per parameter is about 20 bytes, which we can break down as follows:

- An FP16 or BF16 parameter ~ 2 bytes
- An FP16 or BF16 gradient ~ 2 bytes
- An FP32 optimizer state ~ 8 bytes based on the Adam optimizers
- An FP32 copy of parameter ~ 4 bytes (needed for the optimizer apply (OA) operation)
- An FP32 copy of gradient ~ 4 bytes (needed for the OA operation)

Even for a relatively small DL model with 10 billion parameters, it can require at least 200GB of memory, which is much larger than the typical GPU memory (for example, NVIDIA A100 with 40GB/80GB memory) available on a single GPU. On top of the memory requirements for model and optimizer states, there are other memory consumers such as activations generated in the forward pass. The memory required can be a lot greater than 200GB.

For distributed training, we recommend that you use Amazon EC2 P4 and P5 instances that have NVIDIA A100 and H100 Tensor Core GPUs respectively. For more details about specifications such as CPU cores, RAM, attached storage volume, and network bandwidth, see the Accelerated Computing section in the Amazon EC2 Instance Types page. For instance types that SMP v2 supports, see the section called “Supported instance types”.

Even with the accelerated computing instances, models with about 10 billion parameters such as Megatron-LM and T5, and even larger models with hundreds of billions of parameters such as GPT-3, cannot fit model replicas in each GPU device.

How the library employs model parallelism and memory saving techniques

The library consists of various types of model parallelism features and memory-saving features such as optimizer state sharding, activation checkpointing, and activation offloading. All these techniques can be combined to efficiently train large models that consist of hundreds of billions of parameters.
Topics

- Sharded data parallelism
- Tensor parallelism
- Activation checkpointing and offloading
- Choosing the right techniques for your model

Sharded data parallelism

_Sharded data parallelism_ is a memory-saving distributed training technique that splits the state of a model (model parameters, gradients, and optimizer states) across GPUs within a data-parallel group.

SageMaker implements sharded data parallelism through FSDP, and extends it to implement the scale aware hybrid sharding strategy discussed in the blog post [Near-linear scaling of gigantic-model training on AWS](https://aws.amazon.com/blogs/machine-learning/)

You can apply sharded data parallelism to your model as a standalone strategy. Furthermore, if you are using the most performant GPU instances equipped with NVIDIA A100 Tensor Core GPUs, `ml.p4d.24xlarge` and `ml.p4de.24xlarge`, you can take the advantage of improved training speed from the AllGather operation offered by the [SageMaker data parallelism (SMDDP) library](https://docs.aws.amazon.com/sagemaker/latest/dg/data-parallelism.html).

To dive deep into sharded data parallelism and learn how to set it up or use a combination of sharded data parallelism with other techniques like tensor parallelism and mixed precision training, see the section called “Hybrid sharded data parallelism”.

Tensor parallelism

_Tensor parallelism_ splits individual layers, or `nn.Modules`, across devices to run in parallel. The following figure shows the simplest example of how the SMP library splits a model with four layers to achieve two-way tensor parallelism ("tensor_parallel_degree": 2). In the following figure, the notations for model parallel group, tensor parallel group, and data parallel group are MP_GROUP, TP_GROUP, and DP_GROUP respectively. The layers of each model replica are bisected and distributed into two GPUs. The library manages communication across the tensor-distributed model replicas.
To dive deep into tensor parallelism and other memory-saving features for PyTorch, and to learn how to set a combination of the core features, see the section called “Tensor parallelism”.

Activation checkpointing and offloading

To save GPU memory, the library supports activation checkpointing to avoid storing internal activations in the GPU memory for user-specified modules during the forward pass. The library recomputes these activations during the backward pass. In addition, with activation offloading, it offloads the stored activations to CPU memory and fetches them back to GPU during the backward pass to further reduce the activation memory footprint. For more information about how to use these features, see the section called “Activation checkpointing” and the section called “Activation offloading”.

Sample model with four layers

Model parallel configuration

Batch 1
Batch 3
Batch 5
Batch 7
Batch 2
Batch 4
Batch 6
Batch 8

MP_GROUP=TP_GROUP

GPU 0
GPU 1
GPU 2
GPU 3
GPU 4
GPU 5
GPU 6
GPU 7
Choosing the right techniques for your model

For more information about choosing the right techniques and configurations, see the section called “Best practices”.

Supported frameworks and AWS Regions

Before using the SageMaker model parallelism library v2 (SMP v2), check the supported frameworks and instance types and determine if there are enough quotas in your AWS account and AWS Region.

Note
To check the latest updates and release notes of the library, see the section called “Release notes”.

Supported frameworks

SMP v2 supports the following deep learning frameworks and available through SMP Docker containers and an SMP Conda channel. When you use the framework estimator classes in the SageMaker Python SDK and specify distribution configuration to use SMP v2, SageMaker automatically picks up the SMP Docker containers. To use SMP v2, we recommend that you always keep the SageMaker Python SDK up to date in your development environment.

PyTorch versions that the SageMaker model parallelism library supports

<table>
<thead>
<tr>
<th>PyTorch version</th>
<th>SageMaker model parallelism library version</th>
<th>SMP Docker image URI</th>
</tr>
</thead>
<tbody>
<tr>
<td>v2.2.0</td>
<td>smdistributed-modelparallel==v2.2.0</td>
<td>658645717510.dkr.ecr.us-west-2.amazonaws.com/smdistributed-modelparallel:2.2.0-gpu-py310-cu121</td>
</tr>
<tr>
<td>v2.1.2</td>
<td>smdistributed-modelparallel==v2.1.0</td>
<td>658645717510.dkr.ecr.us-west-2.amazonaws.com/smdistributed-modelparallel:2.1.0-gpu-cu121</td>
</tr>
</tbody>
</table>
The following S3 bucket is a public Conda channel hosted by the SMP service team. If you want to install the SMP v2 library in an environment such as SageMaker HyperPod clusters, use this Conda channel to properly install the SMP library.


For more information about Conda channels in general, see [Channels](#) in the *Conda documentation*.

⚠️ **Note**

To find previous versions of the SMP library v1.x and pre-packaged DLCs, see the section called “Supported Frameworks” in the *SMP v1 documentation*.

### Use SMP v2 with open source libraries

The SMP v2 library works with other PyTorch-based open source libraries such as PyTorch Lightning, Hugging Face Transformers, and Hugging Face Accelerate, because SMP v2 is compatible with the PyTorch FSDP APIs. If you have further questions on using the SMP library with other third party libraries, contact the SMP service team at sm-model-parallel-feedback@amazon.com.
AWS Regions

SMP v2 is available in the following AWS Regions. If you'd like to use the SMP Docker image URIs or the SMP Conda channel, check the following list and choose the AWS Region matching with yours, and update the image URI or the channel URL accordingly.

- ap-northeast-1
- ap-northeast-2
- ap-northeast-3
- ap-south-1
- ap-southeast-1
- ap-southeast-2
- ca-central-1
- eu-central-1
- eu-north-1
- eu-west-1
- eu-west-2
- eu-west-3
- sa-east-1
- us-east-1
- us-east-2
- us-west-1
- us-west-2

Supported instance types

SMP v2 requires one of the following ML instance types.

<table>
<thead>
<tr>
<th>Instance type</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.p4d.24xlarge</td>
<td></td>
</tr>
<tr>
<td>ml.p4de.24xlarge</td>
<td></td>
</tr>
<tr>
<td>ml.p5.48xlarge</td>
<td></td>
</tr>
</tbody>
</table>
Tip

Starting from SMP v2.2.0 supporting PyTorch v2.2.0 and later, the section called “Mixed precision training with FP8 on P5 instances using Transformer Engine” is available.

For specs of the SageMaker machine learning instance types in general, see the Accelerated Computing section in the Amazon EC2 Instance Types page. For information about instance pricing, see Amazon SageMaker Pricing.

If you encountered an error message similar to the following, follow the instructions at Requesting a quota increase in the AWS Service Quotas User Guide.

```
ResourceLimitExceeded: An error occurred (ResourceLimitExceeded) when calling the CreateTrainingJob operation: The account-level service limit 'ml.p3dn.24xlarge for training job usage' is 0 Instances, with current utilization of 0 Instances and a request delta of 1 Instances. Please contact AWS support to request an increase for this limit.
```

Get started with the SageMaker model parallelism library v2

On this page, you'll learn how to use the SageMaker model parallelism library v2 APIs and get started with running a PyTorch Fully Sharded Data Parallel (FSDP) training job in the SageMaker Training platform or on a SageMaker HyperPod cluster.

There are various scenarios for running a PyTorch training job with SMP v2.

1. For SageMaker training, use one of the pre-built SageMaker Framework Containers for PyTorch v2.0.1 and later, which are pre-packaged with SMP v2.
2. Use the SMP v2 binary file to set up a Conda environment for running a distributed training workload on a SageMaker HyperPod cluster.
3. Extend the pre-built SageMaker Framework Containers for PyTorch v2.0.1 and later to install any additional functional requirements for your use case. To learn how to extend a pre-built container, see Extend a Pre-built Container.
4. You can also bring your own Docker container and manually set up all SageMaker Training environment using the SageMaker Training toolkit and install the SMP v2 binary file. This is the least recommended option due to the complexity of dependencies. To learn how to run your own Docker container, see Adapting Your Own Training Container.
This getting started guide covers the first two scenarios.

Topics

- Step 1: Adapt your PyTorch FSDP training script
- Step 2: Launch a training job

Step 1: Adapt your PyTorch FSDP training script

To activate and configure the SMP v2 library, start with importing and adding the `torch.sagemaker.init()` module at the top of the script. This module takes in the SMP configuration dictionary of the section called “SMP v2 core feature configuration parameters” that you'll prepare in the section called “Step 2: Launch a training job”. Also, for using the various core features offered by SMP v2, you might need to make few more changes to adapt your training script. More detailed instructions on adapting your training script for using the SMP v2 core features are provided at the section called “Core features of SMP v2”.

SageMaker Training

In your training script, add the following two lines of code, which is the minimal requirement to start training with SMP v2. In the section called “Step 2: Launch a training job”, you'll set up an object of the SageMaker PyTorch estimator class with an SMP configuration dictionary through the distribution argument of the estimator class.

```python
import torch.sagemaker as tsm
tsm.init()
```

**Note**

You can also directly pass a configuration dictionary of the section called “SMP v2 core feature configuration parameters” to the `torch.sagemaker.init()` module. However, the parameters passed to the PyTorch estimator in the section called “Step 2: Launch a training job” take priority and override the ones specified to the `torch.sagemaker.init()` module.
SageMaker HyperPod

In your training script, add the following two lines of code. In the section called “Step 2: Launch a training job”, you’ll set up a smp_config.json file for setting up SMP configurations in JSON format, and upload it to a storage or a file system mapped with your SageMaker HyperPod cluster. We recommend that you keep the configuration file under the same directory where you upload your training script.

```python
import torch.sagemaker as tsm
tsm.init("/dir_to_training_files/smp_config.json")
```

Note
You can also directly pass a configuration dictionary of the the section called “SMP v2 core feature configuration parameters” into the torch.sagemaker.init() module.

Step 2: Launch a training job

Learn how to configure SMP distribution options for launching a PyTorch FSDP training job with SMP core features.

SageMaker Training

When you set up a training job launcher object of the PyTorch framework estimator class in the SageMaker Python SDK, configure the section called “SMP v2 core feature configuration parameters” through distribution argument as follows.

Note
The distribution configuration for SMP v2 is integrated in the SageMaker Python SDK starting from v2.200. Make sure that you use the SageMaker Python SDK v2.200 or later.

Note
In SMP v2, you should configure smdistributed with torch_distributed for the distribution argument of the SageMaker PyTorch estimator. With
torch\_distributed, SageMaker runs torchrun, which is the default multi-node job launcher of PyTorch Distributed.

```python
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    # Pass the training script you adapted with SMP from Step 1
    entry_point="your-training-script.py",
    ... # Configure other required and optional parameters
    distribution={
        "torch\_distributed": { "enabled": True },
        "smdistributed": {
            "modelparallel": {
                "enabled": True,
                "parameters": {
                    "hybrid\_shard\_degree": \texttt{Integer},
                    "sm\_activation\_offloading": \texttt{Boolean},
                    "activation\_loading\_horizon": \texttt{Integer},
                    "fsdp\_cache\_flush\_warnings": \texttt{Boolean},
                    "allow\_empty\_shards": \texttt{Boolean},
                    "tensor\_parallel\_degree": \texttt{Integer},
                    "tensor\_parallel\_seed": \texttt{Integer}
                }
            }
        }
    }
)
```

SageMaker HyperPod

Before you start, make sure if the following prerequisites are met.

- An Amazon FSx shared directory mounted (/fsx) to your HyperPod cluster.
- Conda installed in the FSx shared directory. To learn how to install Conda, use the instructions at [Installing on Linux](#) in the Conda User Guide.
- cuda11.8 or cuda12.1 installed on your HyperPod cluster head and compute nodes.

If the prerequisites are all met, proceed to the following instructions on launching a workload with SMP v2 on a HyperPod cluster.
1. Prepare an `smp_config.json` file that contains a dictionary of [the section called “SMP v2 core feature configuration parameters”](#). Make sure that you upload this JSON file to where you store your training script, or the path you specified to the `torch.sagemaker.init()` module in Step 1. If you've already passed the configuration dictionary to the `torch.sagemaker.init()` module in the training script in Step 1, you can skip this step.

```json
// smp_config.json
{
    "hybrid_shard_degree": Integer,
    "sm_activation_offloading": Boolean,
    "activation_loading_horizon": Integer,
    "fsdp_cache_flush_warnings": Boolean,
    "allow_empty_shards": Boolean,
    "tensor_parallel_degree": Integer,
    "tensor_parallel_seed": Integer
}
```

2. Upload the `smp_config.json` file to a directory in your file system. The directory path must match with the path you specified in Step 1. If you’ve already passed the configuration dictionary to the `torch.sagemaker.init()` module in the training script, you can skip this step.

3. On the compute node of your HyperPod instance, start a terminal session with the following command.

```
sudo su -l ubuntu
```

4. Create a Conda environment on the compute node.

```
# Run on compute node
SMP_CUDA_VER=<11.8 or 12.1>

source /fsx/<path to miniconda>/miniconda3/bin/activate

export ENV_PATH=/fsx/<path to miniconda>/miniconda3/envs/<ENV NAME>
conda create -p ${ENV_PATH} python=3.10
conda activate ${ENV_PATH}

# Verify aws-cli is installed: Expect something like "aws-cli/2.15.0"
aws --version
```
# Install aws-cli if not already installed
# https://docs.aws.amazon.com/cli/latest/userguide/getting-started-install.html#cliv2-linux-install

conda install pytorch="2.0.1=sm_py3.10_cuda${SMP_CUDA_VER}*" packaging --override-channels \
    smp-2.0.0-pt-2.0.1/2023-12-11/smp-v2/ \
    -c pytorch -c numba/label/dev \
    -c nvidia -c conda-forg

# Install dependencies of the script as below
python -m pip install packaging transformers==4.31.0 accelerate ninja tensorboard h5py datasets \
    && python -m pip install expecttest hypothesis \
    && python -m pip install "flash-attn>=2.0.4" --no-build-isolation

# Install SMDDP wheel (only run for cuda11.8)
SMDDP_WHL="smdistributed_dataparallel-2.0.2-cp310-cp310-linux_x86_64_whl" \
    && wget -q https://smdataparallel.s3.amazonaws.com/binary/pytorch/2.0.1/
cu118/2023-12-07/${SMDDP_WHL} \
    && pip install --force ${SMDDP_WHL} \
    && rm ${SMDDP_WHL}

# cuDNN installation for TransformerEngine installation for cuda11.8
# Please download from below link, you need to agree to terms
local_installers/11.x/cudnn-linux-x86_64-8.9.5.30_cuda11-archive.tar.xz

tar xf cudnn-linux-x86_64-8.9.5.30_cuda11-archive.tar.xz \
    && rm -rf /usr/local/cuda-$SMP_CUDA_VER/include/cudnn* /usr/local/cuda-$SMP_CUDA_VER/lib/cudnn* \
    && cp ./cudnn-linux-x86_64-8.9.5.30_cuda11-archive/include/* /usr/local/cuda-$SMP_CUDA_VER/include/ \
    && cp ./cudnn-linux-x86_64-8.9.5.30_cuda11-archive/lib/* /usr/local/cuda-$SMP_CUDA_VER/lib/ \
    && rm -rf cudnn-linux-x86_64-8.9.5.30_cuda11-archive.tar.xz \
    && rm -rf cudnn-linux-x86_64-8.9.5.30_cuda11-archive/

# Please download from below link, you need to agree to terms
# https://developer.download.nvidia.com/compute/cudnn/secure/8.9.7/
local_installers/12.x/cudnn-linux-x86_64-8.9.7.29_cuda12-archive.tar.xz

# cuDNN installation for TransformerEngine installation for cuda12.1
# Please download from below link, you need to agree to terms
# https://developer.download.nvidia.com/compute/cudnn/secure/8.9.7/
local_installers/12.x/cudnn-linux-x86_64-8.9.7.29_cuda12-archive.tar.xz

tar xf cudnn-linux-x86_64-8.9.7.29_cuda12-archive.tar.xz \
    && rm -rf /usr/local/cuda-$SMP_CUDA_VER/include/cudnn* /usr/local/cuda-$SMP_CUDA_VER/lib/cudnn* \
    && cp ./cudnn-linux-x86_64-8.9.7.29_cuda12-archive/include/* /usr/local/cuda-$SMP_CUDA_VER/include/ \
    && cp ./cudnn-linux-x86_64-8.9.7.29_cuda12-archive/lib/* /usr/local/cuda-$SMP_CUDA_VER/lib/ \
    && rm -rf cudnn-linux-x86_64-8.9.7.29_cuda12-archive.tar.xz \
    && rm -rf cudnn-linux-x86_64-8.9.7.29_cuda12-archive/
5. Run a test training job.
   a. In the shared file system (/fsx), clone the **Awesome Distributed Training GitHub repository**, and go to the `3.test_cases/11.modelparallel` folder.

```
git clone https://github.com/aws-samples/awsome-distributed-training/
cd awsome-distributed-training/3.test_cases/11.modelparallel
```

   b. Submit a job using `sbatch` as follows.

```
conda activate <ENV_PATH>
sbatch -N 16 conda_launch.sh
```

   If the job submission is successful, the output message of this `sbatch` command should be similar to **Submitted batch job ABCDEF**.

   c. Check the log file in the current directory under `logs/`.

```
tail -f ./logs/fsdp_smp_ABCDEF.out
```
Core features of the SageMaker model parallelism library v2

The Amazon SageMaker model parallelism library v2 (SMP v2) offers distribution strategies and memory-saving techniques, such as sharded data parallelism, tensor parallelism, and checkpointing. The model parallelism strategies and techniques offered by SMP v2 help distribute large models across multiple devices while optimizing training speed and memory consumption. SMP v2 also provides a Python package `torch.sagemaker` to help adapt your training script with few lines of code change.

This guide follows the basic two-step flow introduced in the section called “Get started with SMP v2”. To dive deep into the core features of SMP v2 and how to use them, see the following topics.

Note

These core features are available in SMP v2.0.0 and later and the SageMaker Python SDK v2.200.0 and later, and works for PyTorch v2.0.1 and later. To check the versions of the packages, see the section called “Supported frameworks and AWS Regions”.

Topics

- Hybrid sharded data parallelism
- Compatibility with the SMDDP library optimized for AWS infrastructure
- Mixed precision training
- Delayed parameter initialization
- Activation checkpointing
- Activation offloading
- Tensor parallelism
- Fine-tuning
- FlashAttention
- Save and load checkpoints while using SMP

Hybrid sharded data parallelism

*Sharded data parallelism* is a memory-saving distributed training technique that splits the state of a model (model parameters, gradients, and optimizer states) across devices. This helps you fit
a larger model or increase the batch size using the freed-up GPU memory. The SMP library offers a capability of running sharded data parallelism with PyTorch Fully Sharded Data Parallel (FSDP). PyTorch FSDP by default shards across the whole set of GPUs being used. In SMP v2, the library offers this sharded data parallelism on top of PyTorch FSDP by extending PyTorch hybrid sharding (HYBRID_SHARD), which is one of the sharding strategies provided by PyTorch FSDP: FULL_SHARD, SHARD_GRAD_OP, HYBRID_SHARD, _HYBRID_SHARD_ZERO2. Extending hybrid sharding in this manner helps implement scale-aware-sharding as described in the blog Near-linear scaling of gigantic-model training on AWS for PyTorch FSDP.

The SMP library makes it easy to use HYBRID_SHARD and _HYBRID_SHARD_ZERO2 across any configurable number of GPUs, extending the native PyTorch FSDP that supports sharding across a single node (HYBRID_SHARD) or all GPUs (FULL_SHARD). PyTorch FSDP calls can stay as is, and you only need to add the hybrid_shard_degree argument to the SMP configuration, as shown in the following code example. You don't need to change the value of the sharding_strategy argument in the PyTorch FSDP wrapper around your PyTorch model. You can pass ShardingStrategy.HYBRID_SHARD as the value. Alternatively, the SMP library overrides the strategy in the script and sets it to ShardingStrategy.HYBRID_SHARD if you specify a value equal to or greater than 2 to the hybrid_shard_degree parameter.

The following code snippets show how to add the SMP initialization module torch.sagemaker.init() to your training script and set up the SMP configuration dictionary in JSON format for training job launcher while following the two-step process introduced in the section called “Get started with SMP v2”. You don’t need to make any changes to your PyTorch model or PyTorch FSDP configuration. For more information about the hybrid_shard_degree parameter, see the section called “SMP v2 core feature configuration parameters”.

SMP configuration dictionary

```
{ "hybrid_shard_degree": 16 }
```

In training script

```
import torch.sagemaker as tsm
tsm.init()

# Set up a PyTorch model
model = ...

# Wrap the PyTorch model using the PyTorch FSDP module
```
model = FSDP(
    model,
    ...
)

# Optimizer needs to be created after FSDP wrapper
optimizer = ...

Compatibility with the SMDDP library optimized for AWS infrastructure

You can use the SageMaker model parallelism library v2 (SMP v2) in conjunction with the SageMaker distributed data parallelism (SMDDP) library that offers the AllGather collective communication operation optimized for AWS infrastructure. In distributed training, collective communication operations are designed for synchronizing multiple GPU workers and exchange information between them. AllGather is one of the core collective communication operations typically used in sharded data parallelism. To learn more about the SMDDP AllGather operation, see the section called “SMDDP AllGather collective operation”. Optimizing such collective communication operations would directly contribute to a faster end-to-end training without side effects on convergence.

Note

The SMDDP library supports P4 and P4de instances (see also the section called “Supported frameworks, AWS Regions, and instances types” by the SMDDP library).

The SMDDP library integrates natively with PyTorch through the process group layer. To use the SMDDP library, you only need to add two lines of code to your training script. It supports any training frameworks such as SageMaker Model Parallelism Library, PyTorch FSDP, and DeepSpeed.

To activate SMDDP and use its AllGather operation, you need to add two lines of code to your training script as part of the section called “Step 1: Adapt your PyTorch FSDP training script”. Note that you need to initialize PyTorch Distributed with the SMDDP backend first, and then run the SMP initialization.

import torch.distributed as dist

# Initialize with SMDDP
import smdistributed.dataparallel.torch.torch_smddp
dist.init_process_group(backend="smddp") # Replacing "nccl"

# Initialize with SMP
import torch.sagemaker as tsm
tsm.init()

**SageMaker Framework Containers** for PyTorch (see also the section called “Supported frameworks and AWS Regions” by SMP v2 and the section called “Supported frameworks, AWS Regions, and instances types” by the SMDDP library) are pre-packaged with the SMP binary and the SMDDP binary. To learn more about the SMDDP library, see the section called “SageMaker distributed data parallelism library”.

**Mixed precision training**

The SageMaker model parallelism (SMP) library v2 supports mixed precision training out of the box by integrating with open source frameworks such as PyTorch FSDP and Transformer Engine. To learn more, see the following topics.

**Topics**

- Mixed precision training with FP8 on P5 instances using Transformer Engine
- Mixed precision training with half-precision data types using PyTorch FSDP

**Mixed precision training with FP8 on P5 instances using Transformer Engine**

Starting from the SageMaker model parallelism (SMP) library v2.2.0, the SMP library integrates with Transformer Engine and supports FP8 mixed precision training out of the box, keeping compatibility with PyTorch FSDP MixedPrecision. This means that you can use both PyTorch FSDP for mixed precision training and Transformer Engine for FP8 training. For model layers not supported by Transformer Engine's FP8 training feature, those layers fall back to PyTorch FSDP mixed precision.

**Note**

SMP v2 offers FP8 support for the following Hugging Face Transformer models:

- GPT-NeoX
- Llama 2
Note

This FP8 training on the P5 feature is available in the following combination of libraries of SageMaker and the PyTorch library:

- SMP v2.2.0 and later
- the SageMaker Python SDK v2.212.0 and later
- PyTorch v2.2.0 and later

**FP8** (8-bit floating point precision) is a data type that has emerged as another paradigm to accelerate deep learning training of LLM models. With the release of NVIDIA H100 GPUs supporting FP8 data types, you can benefit from the advantages from the performance improvements on P5 instances equipped with the H100 GPUs, while accelerating distributed training with FP8 mixed precision training.

The FP8 data type further branches down to E4M3 and E5M2 formats. **E4M3** offers a better precision, has a limited dynamic range, and is ideal for the forward pass in model training. **E5M2** has a broader dynamic range, but reduced precision, and is better suited for the backward pass, where precision is less critical and a wider dynamic range becomes beneficial. Hence, we recommend that you use the hybrid FP8 strategy recipe to leverage these characteristics effectively.

For half-precision data types (FP16 and BF16), global loss-scaling techniques such as static loss-scaling or dynamic loss-scaling handle convergence issues that arise from information loss due to rounding gradients in half-precision. However, the dynamic range of FP8 is even narrower, and the global loss scaling techniques are not sufficient. At this point, we need a finer-grained per-tensor scaling technique. **Delayed scaling** is a strategy that selects a scaling factor based on the maximum absolute values observed in a number of tensors form previous iterations. There's a trade-off in this strategy; it uses the full performance benefits of FP8 computation but requires memory for keeping the maximum value history of tensors. To learn more about the delayed scaling strategy in general, see the paper [FP8 Formats for Deep Learning](#).

In practice, using FP8 is helpful in all training scenarios on P5 instances. We strongly recommend enabling FP8 whenever possible for enhancing training performance.

SMP v2 supports Transformer Engine out of the box. Therefore, when running FP8 training with SMP v2 on P5 instances of SageMaker (ml.p5.48xlarge), the only thing you need to do is to
import torch.sagemaker as tsm
import transformer_engine.pytorch as te
from transformer_engine.common.recipe import DelayedScaling, Format

# Enable E4M3 during forward pass, E5M2 during backward pass.
fp8_format = Format.HYBRID

# Create an FP8 recipe.
fp8_recipe = DelayedScaling(fp8_format=fp8_format, amax_history_len=32,
    amax_compute_algo="max")

# Enable FP8 autocasting.
with te.fp8_autocast(enabled=True, fp8_recipe=fp8_recipe,
    fp8_group=tsm.state.world_process_group):
    out = model(inp)

    loss = out.sum()
    loss.backward()

To find a practical example of FP8 training with SMP v2 on P5 instances, see the example notebook at [Accelerate SageMaker PyTorch FSDP Training of Llama-v2 (or GPT-NeoX) with FP8 on P5 instances](#).

**Mixed precision training with half-precision data types using PyTorch FSDP**

SMP v2 supports [PyTorch FSDP MixedPrecision](#) for training jobs on P4 and P5 instances. PyTorch FSDP provides various configurations for mixed precision for both performance improvement and memory reduction.

**Note**

This mixed precision training with the PyTorch FSDP feature is available in the following combination of libraries of SageMaker and the PyTorch library.

- SMP v2.0.0 and later
The standard way to configure a model for mixed precision is to create the model in `float32`, and then allow FSDP to cast the parameters to `float16` or `bfloat16` on the fly by passing a MixedPrecision policy, as shown in the following code snippet. For more information about options to change the `dtype` for parameters, reduction, or buffers for mixed precision in PyTorch, see [PyTorch FSDP MixedPrecision API](https://pytorch.org/docs/stable/fsdp.html#mixed-precision) in the *PyTorch documentation*.

```python
# Native PyTorch API
from torch.distributed.fsdp import MixedPrecision
dtype = torch.bfloat16
mixed_precision_policy = MixedPrecision(
    param_dtype=dtype, reduce_dtype=dtype, buffer_dtype=dtype
)
model = FSDP(model,
             ..., mixed_precision=mixed_precision_policy)
```

Note that certain models (such as the Hugging Face Transformers Llama model) expect buffers as `float32`. To use `float32`, replace `torch.bfloat16` with `torch.float32` in the line defining the `dtype` object.

**Delayed parameter initialization**

Initialization of a large model for training is not always possible with the limited GPU memory. To resolve this problem of insufficient GPU memory, you can initialize the model on CPU memory. However, for larger models with more than 20 or 40 billion parameters, even CPU memory might not be enough. For such case, we recommend that you initialize the model on what PyTorch calls a *meta device*, which allows the creation of tensors without any data attached to them. A tensor on a meta device only needs the shape information, and this allows to create a large model with its parameters on meta devices. Hugging Face *Accelerate* provides the context manager `init_empty_weights` to help create such model on meta devices while initializing the buffers on a regular device. Before training starts, PyTorch FSDP initializes the model parameters. This
delayed parameter initialization feature of SMP v2 delays this creation of model parameters to happen after PyTorch FSDP performs parameter sharding. PyTorch FSDP accepts a parameter initialization function (param_init_fn) when sharding the modules, and it calls param_init_fn for each module. The param_init_fn API takes a module as an argument and initializes all the parameters in it, not including the parameters of any child module. Note that this behavior differs from the native PyTorch v2.0.1 which has a bug causing the parameters to be initialized multiple times.

SMP v2 provides the the section called “torch.sagemaker.delayed_param.DelayedParamIniter” API for applying delayed parameter initialization.

The following code snippets show how to apply the torch.sagemaker.delayed_param.DelayedParamIniter API to your training script.

Assume that you have a PyTorch FSDP training script as follows.

```python
# Creation of model on meta device
from accelerate import init_empty_weights
with init_empty_weights():
    model = create_model()

# Define a param init fn, below is an example for Hugging Face GPTNeoX.
def init_weights(module):
    d = torch.cuda.current_device()
    # Note that below doesn't work if you have buffers in the model
    # buffers will need to reinitialized after this call
    module.to_empty(device=d, recurse=False)
    if isinstance(module, (nn.Linear, Conv1D)):
        module.weight.data.normal_(mean=0.0, std=args.initializer_range)
        if module.bias:
            module.bias.data.zero_()
    elif isinstance(module, nn.Embedding):
        module.weight.data.normal_(mean=0.0, std=args.initializer_range)
        if module.padding_idx:
            module.weight.data[module.padding_idx].zero_()
    elif isinstance(module, nn.LayerNorm):
        module.bias.data.zero_()
        module.weight.data.fill_(1.0)

# Changes to FSDP wrapper.
model = FSDP(model)

```

model,
...,
param_init_fn=init_weights
)

# At this point model is initialized and sharded for sharded data parallelism.

Note that the delayed parameter initialization approach is not model agnostic. To resolve this issue, you need to write an init_weights function as shown in the preceding example to match the initialization in the original model definition, and it should cover all the parameters of the model. To simplify this process of preparing such init_weights function, SMP v2 implements this initialization function for the following models: GPT-2, GPT-J, GPT-NeoX, and Llama from Hugging Face Transformers. The torch.sagemaker.delayed_param.DelayedParamIniter API also works with the SMP tensor parallel implementation, torch.sagemaker.tensor_parallel.transformer.TransformerLMHead model, that you can call after the the section called “torch.sagemaker.transform API call.

Using the torch.sagemaker.delayed_param.DelayedParamIniter API, you can adapt your PyTorch FSDP script as follows. After creating a model with empty weights, register the torch.sagemaker.delayed_param.DelayedParamIniter API to the model, and define an object of it. Pass the object to the param_init_fn of the PyTorch FSDP class.

```python
from torch.sagemaker.delayed_param import DelayedParamIniter
from accelerate import init_empty_weights

with init_empty_weights():
    model = create_model()

delayed_initer = DelayedParamIniter(model)

with delayed_initer.validate_params_and_buffers_inited():
    model = FSDP(
        model,
        ...,
        param_init_fn=delayed_initer.get_param_init_fn()
    )
```

Notes on tied weights
When training models with tied weights, we need to take special care to tie the weights after initializing the weights with delayed parameter initialization. PyTorch FSDP does not have a mechanism to tie the weights after initializing them using `param_init_fn` as above. To address such cases we added API to allow a `post_init_hook_fn`, which can be used to tie the weights. You can pass any function in there which accepts the module as argument, but we also have a predefined `post_param_init_fn` defined in `DelayedParamIniter` which calls `tie_weights` method of the module if it exists. Note that it's safe to always pass in `post_param_init_fn` even if there's no `tie_weights` method for the module.

```python
with delayed_initer.validate_params_and_buffers_inited():
    model = FSDP(
        model,
        ...,
        param_init_fn=delayed_initer.get_param_init_fn(),
        post_param_init_fn=delayed_initer.get_post_param_init_fn()
    )
```

**Activation checkpointing**

Activation checkpointing is a technique to reduce memory usage by clearing activations of certain layers and recomputing them during the backward pass. Effectively, this trades extra computation time for reducing memory usage. If a module is checkpointed, at the end of a forward pass, only the initial inputs to the module and final outputs from the module stay in memory. PyTorch releases any intermediate tensors that are part of the computation inside that module during the forward pass. During the backward pass of the checkpointed modules, PyTorch recomputes these tensors. At this point, the layers beyond this checkpointed module have finished their backward pass, so the peak memory usage with checkpointing becomes lower.

SMP v2 supports the PyTorch activation checkpointing module, `apply_activation_checkpointing`. The following are examples of activation checkpointing of the Hugging Face GPT-NeoX model.

**Checkpointing Transformer layers of the Hugging Face GPT-NeoX model**

```python
from transformers.models.gpt_neox import GPTNeoXLayer
from torch.distributed.algorithms._checkpoint.checkpoint_wrapper import ( apply_activation_checkpointing )

# check_fn receives a module as the arg,
```
Checkpointing every other Transformer layer of the Hugging Face GPT-NeoX model

```python
# check_fn receives a module as arg,
# and it needs to return whether the module is to be checkpointed
# here we define that function based on global variable (transformer_layers)
from transformers.models.gpt_neox import GPTNeoXLayer
from torch.distributed.algorithms._checkpoint.checkpoint_wrapper import 
    apply_activation_checkpointing

transformer_layers = 
    m for m model.modules() if isinstance(m, GPTNeoXLayer)

def is_odd_transformer_layer(module):
    return transformer_layers.index(module) % 2 == 0

apply_activation_checkpointing(model, check_fn=is_odd_transformer_layer)
```

Alternatively, PyTorch also has the `torch.utils.checkpoint` module for checkpointing, which is used by a subset of Hugging Face Transformers models. This module also works with SMP v2. However, it requires you to have access to the model definition for adding the checkpoint wrapper. Therefore, we recommend you to use the `apply_activation_checkpointing` method.

Activation offloading

⚠️ **Important**

In SMP v2.2.0, the activation offloading functionality of the SMP library doesn't work. Use the native PyTorch activation offloading instead.

Typically, the forward pass computes activations at each layer and keeps them in GPU memory until the backward pass for the corresponding layer finishes. Offloading these tensors to CPU
memory after the forward pass and fetching them back to GPU when they are needed for the backward pass of the layer can save substantial GPU memory usage. PyTorch supports offloading activations, but the implementation causes GPUs to be idle while activations are fetched back from CPU during backward pass. This causes a major performance degradation when using activation offloading.

SMP v2 improves this activation offloading, and it pre-fetches activations ahead of time before the activations are needed for the GPU to start backward pass on those activations. This pre-fetching feature helps training progresses be more efficiently run without idle GPUs, which results in offering benefits from lower memory usage without a performance degradation.

You can keep the native PyTorch modules for offloading activations in your training script. The following is an example structure of applying the SMP activation offloading feature in your script. Note that activation offloading is applicable only when used together with the section called “Activation checkpointing”. To learn more about the native PyTorch checkpoint tools for activation offloading, see also the checkpoint_wrapper.py in the PyTorch GitHub repository and Activation Checkpointing in the PyTorch blog Scaling Multi-modal Foundation Models in TorchMultimodal with PyTorch Distributed.

To apply the SMP activation offloading feature on PyTorch activation checkpointing, add the sm_activation_offloading and activation_loading_horizon parameters to the SMP configuration dictionary during the section called “Step 2: Launch a training job”.

The following code snippets show how to add the SMP initialization module torch.sagemaker.init() to your training script and set up the SMP configuration dictionary in JSON format for training job launcher while following the two-step process introduced in the section called “Get started with SMP v2”. You don’t need to make any changes to your PyTorch model or PyTorch FSDP configuration. For more information about the sm_activation_offloading and activation_loading_horizon parameters, see the section called “SMP v2 core feature configuration parameters”.

**SMP configuration**

```json
{
    "activation_loading_horizon": 2,
    "sm_activation_offloading": True
}
```

**In training script**
**Note**

While activating the SMP activation offloading feature, make sure that you also use the PyTorch offload_wrapper function and apply it to the root module. The SMP activation offloading feature uses the root module to determine when forward pass is done to start pre-fetching.

```python
import torch.sagemaker as tsm
tsm.init()

# Native PyTorch module for activation offloading
from torch.distributed.algorithms._checkpoint.checkpoint_wrapper import (  
    apply_activation_checkpointing,
    offload_wrapper,
)

model = FSDP(...)

# Activation offloading requires activation checkpointing.
apply_activation_checkpointing(
    model,
    check_fn=checkpoint_transformer_layers_policy,
)

model = offload_wrapper(model)
```

**Tensor parallelism**

Tensor parallelism is a type of model parallelism in which specific model weights, gradients, and optimizer states are split across devices. In contrast to pipeline parallelism, which keeps individual weights intact but partitions the set of weights, gradients, or optimizer across devices, tensor parallelism shreds individual weights. This typically involves distributed computation of specific operations, modules, or layers of the model.

Tensor parallelism is required in cases in which a single parameter consumes most of the GPU memory (such as large embedding tables with a large vocabulary size or a large softmax layer with a large number of classes). In this case, treating this large tensor or operation as an atomic unit is inefficient and impedes balance of the memory load.
SMP v2 integrates with Transformer Engine for the implementation for tensor parallelism, and runs on top of PyTorch FSDP APIs. You can enable PyTorch FSDP and SMP tensor parallelism simultaneously, and determine the best model parallelism for best performance.

In practice, tensor parallelism is especially helpful in the following scenarios.

- When training with long context lengths as that leads to high activation memory with FSDP alone.
- When training with really large clusters on which the global batch size exceeds desired limits.

**Hugging Face Transformer models compatible with the SMP tensor parallelism**

SMP v2 currently offers tensor parallelism support for the following Hugging Face transformer models.

- GPT-NeoX
- Llama 2

For reference configuration for applying tensor parallelism on these models, see the section called “Configuration tips”.

**Configure tensor parallelism**

For `tensor_parallel_degree`, you select a value for the degree of tensor parallelism. The value must evenly divide the number of GPUs in your cluster. For example, to shard your model while using an instance with 8 GPUs, choose 2, 4, or 8. We recommend that you start with a small number, and gradually increase it until the model fits in the GPU memory.

The following code snippets show how to add the SMP initialization module `torch.sagemaker.init()` to your training script and set up the SMP configuration dictionary in JSON format for training job launcher while following the two-step process introduced in the section called “Get started with SMP v2”. You don’t need to make any changes to your PyTorch model or PyTorch FSDP configuration. For more information about the `tensor_parallel_degree` and `tensor_parallel_seed` parameters, see the section called “SMP v2 core feature configuration parameters”.

**SMP configuration**

```json
{
...
```
In your training script

Initialize with `torch.sagemaker.init()` to activate SMP v2 and wrap your model with the `torch.sagemaker.transform` API.

```python
import torch.sagemaker as tsm
tsm.init()

from transformers import AutoModelForCausalLM
model = AutoModelForCausalLM.from_config(..)
model = tsm.transform(model)
```

Saving and loading Hugging Face Transformer checkpoints

After the SMP library transforms a model, it changes the state dictionary (`state_dict`) of the model. This means that the model becomes incompatible with the original Hugging Face Transformer checkpointing functionalities. To handle this, the SMP library provides APIs to save checkpoints from a transformed model in Hugging Face Transformer representation, and the `torch.sagemaker.transform` API to load a Hugging Face Transformer model checkpoint for fine-tuning.

For more information about saving checkpoints while using the tensor parallelism feature of SMP v2, see the section called “Save and load checkpoints while using SMP”.

For more information about fine-tuning a model applying the tensor parallelism feature of SMP v2, see the section called “Fine-tuning”.

Fine-tuning

Fine-tuning is a process of continuously training pre-trained models to improve performance for specific use cases.

Fine-tuning small models that fit fully on a single GPU, or those that fit 8 copies of model fully on CPUs is straightforward. It requires no special change to regular FSDP training. In the realm of models larger than this, you need to consider using the delayed parameter initialization functionality, which can be tricky.
To address this, the SMP library loads the full model on one of the ranks while the rest of the ranks create models with empty weights on a meta device. Then, PyTorch FSDP initializes the weights on non-zero ranks using the `init_weights` function, and synchronizes the weights on all ranks to the weights on the 0th rank with `sync_module_states` set to True. The following code snippet shows how you should set it up in your training script.

```python
import torch.distributed as dist
from transformers import AutoModelForCausalLM
from accelerate import init_empty_weights

if dist.get_rank() == 0:
    model = AutoModelForCausalLM.from_pretrained(..., low_cpu_mem_usage=True)
else:
    with init_empty_weights():
        model = AutoModelForCausalLM.from_config(AutoConfig.from_pretrained(...))
    delayed_initer = DelayedParamIniter(model)

model = FSDP(
    model,
    ...,
    sync_module_states=True,
    param_init_fn=delayed_initer.get_param_init_fn() if dist.get_rank() > 0 else None
)
```

### Fine-tuning a pre-trained Hugging Face Transformer model with SMP tensor parallelism

This section discusses loading Transformer models for two use cases: fine-tuning small Transformer models and fine-tuning large Transformer models. For smaller models without delayed parameter initialization, wrap the model with the `torch.sagemaker.transform` API before wrapping it with PyTorch FSDP.

```python
import functools
from transformers import AutoModelForCausalLM

model = AutoModelForCausalLM.from_pretrained("meta-llama/Llama-2-7b-hf",
    low_cpu_mem_usage=True)

# Transform model while loading state dictionary from rank 0.
```
For larger models, the preceding approach causes to run out of CPU memory. We recommend that you use delayed parameter initialization to avoid such CPU memory issues. In this case, you can apply the torch.sagemaker.transform API and the torch.sagemaker.delayed_param.DelazedParamIniter API as shown in the following code example.

```python
from transformers import AutoModelForCausalLM
from torch.sagemaker import transform
from torch.sagemaker.delayed_param import DelayedParamIniter

# Create one instance of model without delayed param
# on CPU, on one rank.
if dist.get_rank() == 0:
    model = AutoModelForCausalLM.from_pretrained(...,low_cpu_mem_usage=True)
else:
    with init_empty_weights():
        model = AutoModelForCausalLM.from_config(AutoConfig.from_pretrained(...))

# Transform model while loading state dictionary from rank 0
model = transform(model, load_state_dict_from_rank0=True)

if dist.get_rank() != 0: # For fine-tuning, delayed parameter on non-zero ranks
delayed_initer = DelayedParamIniter(model)
else:
    delayed_initer = None

with (delayed_initer.validate_params_and_buffers_inited() if delayed_initer else nullcontext()):
    # Wrap the model with FSDP
    model = FSDP(model, ...,
```
FlashAttention

SMP v2 supports FlashAttention kernels and makes it easy to apply them to various scenarios for Hugging Face Transformer models. Note that if you use FlashAttention package v2.0 or later, SMP uses FlashAttention v2; however, the Triton flash attention defaults to the flash attention kernel in FlashAttention v1.x, making it exclusively supported in FlashAttention v1.

The module (nn.Module) is a low level API that defines the attention layers of a model. It should be applied right after model creation, from the AutoModelForCausalLM.from_config() API for example, and before the model is being transformed or wrapped with FSDP.

Use FlashAttention kernels for self attention

The following code snippet shows how to use the the section called “torch.sagemaker.nn.attn.FlashSelfAttention” API provided by SMP v2.

```python
def new_attn(self, q, k, v, attention_mask=None, head_mask=None):
    return (
        self.flashmod((q, k, v), causal=True, cast_dtype=torch.bfloat16, layout="b h s d"),
        None,
    )

for layer in model.gpt_neox.layers:
    layer.attention.flash_mod = torch.sagemaker.nn.attn.FlashSelfAttention()
    layer.attention._attn = functools.partial(new_attn, layer.attention)
```

Use FlashAttention kernels for grouped-query attention

SMP v2 also supports FlashAttention kernels for grouped-query attention (GQA) and makes it easy to apply them to various scenarios for Hugging Face Transformer models. Different from original attention architecture, GQA equally partitions query heads into groups, and query heads in the same group share the same key and value heads. Therefore, q and kv heads are passed into forward call separately. Note: The number of q heads needs to be divisible by the number of kv heads.

Example of using FlashGroupedQueryAttention
The following code snippet shows how to use the the section called “torch.sagemaker.nn.attn.FlashGroupedQueryAttention” API provided by SMP v2.

```python
from transformers.models.llama.modeling_llama import LlamaAttention
from torch.sagemaker.nn.attn import FlashGroupedQueryAttention

class LlamaFlashAttention(LlamaAttention):
    def __init__(self, config: LlamaConfig):
        super().__init__(config)

        self.flash_attn = FlashGroupedQueryAttention(
            attention_dropout_prob=0.0,
        )

    query_states = self.q_proj(hidden_states)
    key_states = self.k_proj(hidden_states)
    value_states = self.v_proj(hidden_states)
    ...  
    kv = (key_states, value_states)
    attn_output = self.flash_attn(
        query_states,
        kv,
        attn_mask=attention_mask,
        causal=True,
        layout="b h s d",
    )
    ...  
    attn_output = self.o_proj(attn_output)
    ...  
    return attn_output
```

The SMP library also provides the section called “torch.sagemaker.nn.huggingface.llama_flashattn.LlamaFlashAttention”, which uses the the section called “torch.sagemaker.nn.attn.FlashGroupedQueryAttention” API at low level. Hugging Face Transformers has a similar implementation called
**LlamaFlashAttention2** from v4.36.0. The following code snippet shows how to use the SMP v2 LlamaFlashAttention API or the Transformers LlamaFlashAttention2 API to replace the attention layers of an existing Llama model.

```python
from torch.sagemaker.nn.huggingface.llama_flashattn import LlamaFlashAttention
from transformers.models.llama.modeling_llama import LlamaFlashAttention2

flash_attn_class = LlamaFlashAttention # or flash_attn_class = LlamaFlashAttention2

attn_name = "self_attn"
for layer in model.model.layers:
    prev_layer = getattr(layer, attn_name)
    setattr(layer, attn_name, flash_attn_class(model.config))
```

**Save and load checkpoints while using SMP**

The SMP library supports PyTorch APIs for checkpoints, and provides APIs that help checkpoint properly while using the SMP library.

PyTorch FSDP supports three types of checkpoints: full, sharded and local. These serve different purposes. Full checkpoint should ideally be used only when exporting the model after training finishes, because it’s expensive to generate a full checkpoint. Sharded checkpoint is the recommended approach for saving and loading checkpoints during training. Using sharded checkpoints you can also change the cluster size when resuming training. Local checkpoints are more restrictive. With local checkpoints, you need to resume training with same number of GPUs and currently it’s not supported when using tensor parallelism with SMP. Note that checkpoints by FSDP require writing to a shared network file system, such as FSx.

**Sharded checkpoints**

The following procedure highlights what you need to do to adapt your training script to save and load sharded checkpoints with or without the SMP tensor parallelism feature.

1. Import the SMP `torch.sagemaker` package.

   ```python
   import torch.sagemaker as tsm
   ```

2. Set up auxiliary variables to save and load checkpoints.
   
   a. Set up a coordinator rank for performing communicative collective operations such as AllReduce.
b. Using the `torch.sagemaker.state` enumerations, set up the action rank to determine whether to let the ranks take part in checkpointing. And add an if statement for saving checkpoints depending on the usage of SMP v2 tensor parallelism.

```python
action_rank: bool = global_rank < (tsm.state.hybrid_shard_degree * tsm.state.tp_size)

if tsm.state.tp_size > 1:
    # Tensor parallel groups will have their own sub directories.
    sub_dir = f"tp{tsm.state.tp_size}-{tsm.state.tp_rank}"
else:
    sub_dir = ""
```

3. Keep using the PyTorch FSDP checkpoint APIs as is.

The following code example shows a full PyTorch FSDP training script with the FSDP checkpoint APIs.

```python
import torch.distributed as dist
from torch.distributed.checkpoint.optimizer import (  
    load_sharded_optimizer_state_dict
)
from torch.distributed.fsdp import (  
    FullyShardedDataParallel as FSDP,  
    StateDictType
)
import torch.sagemaker as tsm

sharding_strategy, state_dict_type = ..., ...
global_rank = dist.get_rank()

# 0. Auxiliary variables to save and load checkpoints.

# Used when performing comm collectives such as allreduce.
coordinator_rank: int = min(dist.get_process_group_ranks(model.process_group))

# To determine whether to take part in checkpointing.
action_rank: bool = global_rank < (tsm.state.hybrid_shard_degree * tsm.state.tp_size)
```
if tsm.state.tp_size > 1:
    # Tensor parallel groups will have their own sub directories.
    sub_dir = f"tp{tsm.state.tp_size}-{tsm.state.tp_rank}"
else:
    sub_dir = ""

# 1. Save checkpoints.
with FSDP.state_dict_type(model, StateDictType.SHARDED_STATE_DICT):
    state_dict = {
        "model": model.state_dict(),
        "optimizer": FSDP.optim_state_dict(model, optimizer),
        # Potentially add more customized state dicts.
    }

    # Save from one single replication group.
    if action_rank:
        dist.checkpoint.save_state_dict(
            state_dict=state_dict,
            storage_writer=dist.checkpoint.FileSystemWriter(os.path.join(save_dir, sub_dir)),
            process_group=model.process_group,
            coordinator_rank=coordinator_rank,
        )

# 2. Load checkpoints.
with FSDP.state_dict_type(model, StateDictType.SHARDED_STATE_DICT):
    # 2.1 Load model and everything else except the optimizer.
    state_dict = {
        # All states except optimizer state can be passed here.
        "model": model.state_dict()
    }

    dist.checkpoint.load_state_dict(
        state_dict=state_dict,
        storage_reader=dist.checkpoint.FileSystemReader(os.path.join(load_dir, sub_dir)),
        process_group=model.process_group,
        coordinator_rank=coordinator_rank,
    )
    model.load_state_dict(state_dict["model"])
    # Potentially process more customized and non-optimizer dict states.

    # 2.2 Load optimizer.
    optim_state = load_sharded_optimizer_state_dict(....)
Full model checkpoints

At the end of training, you can save a full checkpoint that combines all shards of a model into a single model checkpoint file. The SMP library fully supports the PyTorch full model checkpoints API, so you don't need to make any changes.

Note that if you use the SMP the section called “Tensor parallelism”, the SMP library transforms the model. When checkpointing the full model in this case, the SMP library translates the model back to the Hugging Face Transformers checkpoint format by default.

In cases where you train with the SMP tensor parallelism and turn off the SMP translation process, you can use the translate_on_save argument of the PyTorch FullStateDictConfig API to switch the SMP auto-translation on or off as needed. For example, if you are focusing on training a model, you don't need to add the translation process which adds overhead. In that case, we recommend you to set translate_on_save=False. Also, if you plan to keep using the SMP translation of the model for further training in future, you can switch it off to save the SMP translation of the model for later use. Translating the model back to the Hugging Face Transformers model checkpoint format is needed when you wrap up the training of your model and use that for inference.

```
from torch.distributed.fsdp import FullyShardedDataParallel as FSDP
from torch.distributed.fsdp import FullStateDictConfig
import torch.sagemaker as tsm

# Save checkpoints.
with FSDP.state_dict_type(
    model,
    StateDictType.FULL_STATE_DICT,
    FullStateDictConfig(
        rank0_only=True, offload_to_cpu=True,
```
# Default value is to translate back to Hugging Face Transformers format,
# when saving full checkpoints for models trained with SMP tensor parallelism.
# translate_on_save=True

state_dict = model.state_dict()
if dist.get_rank() == 0:
    logger.info("Processed state dict to save. Starting write to disk now.")
    os.makedirs(save_dir, exist_ok=True)
    # This name is needed for HF from_pretrained API to work.
    torch.save(state_dict, os.path.join(save_dir, "pytorch_model.bin"))
    hf_model_config.save_pretrained(save_dir)
    dist.barrier()

Note that the option FullStateDictConfig(rank0_only=True, offload_to_cpu=True) is to gather the model on the CPU of the 0th rank device to save memory when training large models.

To load the model back for inference, you do so as shown in the following code example. Note that the class AutoModelForCausalLM might change to other factor builder classes in Hugging Face Transformers, such as AutoModelForSeq2SeqLM, depending on your model. For more information, see Hugging Face Transformers documentation.

```python
from transformers import AutoModelForCausalLM
model = AutoModelForCausalLM.from_pretrained(save_dir)
```

### Amazon SageMaker model parallelism library v2 examples

This page provides a list of blogs and Jupyter notebooks that present practical examples of implementing the SageMaker model parallelism (SMP) library v2 to run distributed training jobs on SageMaker.

**Blogs and Case Studies**

The following blogs discuss case studies about using SMP v2.

- Amazon SageMaker model parallel library now accelerates PyTorch FSDP workloads by up to 20%
PyTorch example notebooks

Example notebooks are provided in the SageMaker examples GitHub repository. To download the examples, run the following command to clone the repository and go to training/distributed_training/pytorch/model_parallel_v2.

Note

Clone and run the example notebooks in the following SageMaker ML IDEs.

- SageMaker JupyterLab (available in Studio created after December 2023)
- SageMaker Code Editor (available in Studio created after December 2023)
- Studio Classic (available as an application in Studio created after December 2023)
- SageMaker Notebook Instances

git clone https://github.com/aws/amazon-sagemaker-examples.git
cd amazon-sagemaker-examples/training/distributed_training/pytorch/model_parallel_v2

SMP v2 example notebooks

- Accelerate training of Llama v2 with SMP v2, PyTorch FSDP, and Transformer Engine by running FP8 training on P5 instances
- Fine-tune Llama v2 with SMP v2 and PyTorch FSDP at large-scale using tensor parallelism, hybrid sharding, and activation offloading
- Train GPT-NeoX with SMP v2 and PyTorch FSDP at large scale
- Fine-tune GPT-NeoX with SMP v2 and PyTorch FSDP at large-scale using tensor parallelism, hybrid sharding, and activation offloading

SageMaker distributed model parallelism best practices

Use the following guidelines when you run a distributed training job with the SageMaker model parallel library v2 (SMP v2).
Setting up the right configuration for distributed training

To estimate and find the best starting point to apply distributed training techniques that SMP v2 provides, review the following list. Each list item discusses the advantage of using the the section called “Core features of SMP v2” along with potential tradeoffs.

Configuration tips

This section provides guidelines on how to decide on the best model configurations for optimal throughput with global batch size requirements.

First, we recommend the following setups regardless of the size of your model.

1. Use the most powerful instance type that you can use.

2. Turn on mixed precision all the time, as it provides substantial benefits for performance and memory reduction. We recommend you to use bfloat16 as it’s more precise than float16.

3. Turn on the SageMaker distributed data parallelism library (instead of using NCCL) whenever it’s applicable, as shown in the section called “Compatibility with the SMDDP library”. One exception is for tensor-parallelism-only use cases (hybrid_shard_degree = 1 and tensor_parallel_degree > 1).

4. If your model has more than about 60 billion parameters, we recommend using the section called “Delayed parameter initialization”. You can also use delayed parameter initialization to speed up the initialization for any model.

5. We recommend you to enable the section called “Activation checkpointing”.

Depending on the size of your model, we recommend that you start with the following guidance.

1. Use sharded data parallelism.

   a. Depending on the batch size you intend to fit in the GPU memory, choose the appropriate sharded data parallel degree. Normally, you should start with the lowest degree to fit your model in the GPU memory while minimizing overhead from network communication. If you see a warning that cache flushes are happening, we recommend that you increase the sharding degree.

   b. Determine world_size based on the maximum local batch size and required global batch size, if any.
c. You can experiment with activation offloading. Depending on scenarios, it can address your memory needs without having to increase the sharding degree, which means less communication.

2. Use sharded data parallelism of PyTorch FSDP and tensor parallelism of SMP v2 simultaneously, as introduced in the section called “Tensor parallelism”.

   a. When training on large clusters, with FSDP alone the global batch size can become too large, causing convergence issues for the model. Typically, most research work keeps the batch size under 4 million tokens. In this case, you can resolve the problem by composing PyTorch FSDP with tensor parallelism of SMP v2 to reduce the batch size.

   For example, if you have 256 nodes and sequence length 4096, even a batch size of 1 per GPU leads to global batch size of 8M tokens. However, when you use tensor parallelism with degree 2 and batch size of 1 per tensor parallel group, this becomes 1/2 batch size per GPU, which translates to 4 million tokens.

   b. When training with long context lengths such as 8k, 16k activation memory can become very high. FSDP doesn't shard activations, and activations can cause GPUs to go out of memory. In such scenarios, you can train efficiently by composing PyTorch FSDP with tensor parallelism of SMP v2.

Reference configurations

The SageMaker model parallelism training team provides the following reference points based on experiments with the Llama 2 model transformed to the SMP transformer model using the section called “torch.sagemaker.transform”, and trained on ml.p4d.24xlarge instance(s) with sequence length 4096 and mixed precision (FP16 or BF16).

<table>
<thead>
<tr>
<th>Model</th>
<th>Model size (the number of model parameters)</th>
<th>The number of instances</th>
<th>Sharded data parallel degree</th>
<th>Tensor parallel degree</th>
<th>Activation checkpointing</th>
<th>Activation offloading</th>
<th>Batch size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Llama 2</td>
<td>7B</td>
<td>1</td>
<td>8</td>
<td>1</td>
<td>TRUE</td>
<td>FALSE</td>
<td>4</td>
</tr>
<tr>
<td>70B</td>
<td>32</td>
<td>256</td>
<td>1</td>
<td>TRUE</td>
<td>FALSE</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

SageMaker model parallelism library v2

3666
### Model

<table>
<thead>
<tr>
<th>Model size (the number of model parameters)</th>
<th>The number of instances</th>
<th>Sharded data parallel degree</th>
<th>Tensor parallel degree</th>
<th>Activation checkpointing</th>
<th>Activation offloading</th>
<th>Batch size</th>
</tr>
</thead>
<tbody>
<tr>
<td>175B</td>
<td>64</td>
<td>128</td>
<td>4</td>
<td>TRUE</td>
<td>TRUE</td>
<td>6</td>
</tr>
</tbody>
</table>

You can extrapolate from the preceding configurations to estimate GPU memory usage for your model configuration. For example, if you increase the sequence length for a 10-billion-parameter model or increase the size of the model to 20 billion, you might want to lower batch size first. If the model still doesn’t fit, try increasing the degree of tensor parallelism.

### Monitoring and logging a training job using the SageMaker console and Amazon CloudWatch

To monitor system-level metrics such as CPU memory utilization, GPU memory utilization, and GPU utilization, use visualization provided through the SageMaker console.

1. In the left navigation pane, choose **Training**.
2. Choose **Training jobs**.
3. In the main pane, choose the training job name for which you want to see more details.
4. Browse the main pane and find the **Monitor** section to see the automated visualization.
5. To see training job logs, choose **View logs** in the **Monitor** section. You can access the distributed training job logs of the training job in CloudWatch. If you launched multi-node distributed training, you should see multiple log streams with tags in the format of `algo-n-1234567890`. The `algo-1` log stream tracks training logs from the main (0th) node.

For more information, see [Monitor and Analyze Training Jobs Using Amazon CloudWatch Metrics](#).

### Permissions

To run a SageMaker training job with model parallelism, make sure you have the right permissions in your IAM role, such as the following:

- To use [FSx for Lustre](#), add `AmazonFSxFullAccess`.
- To use Amazon S3 as a data channel, add `AmazonS3FullAccess`.
To use Docker, build your own container, and push it to Amazon ECR, add AmazonEC2ContainerRegistryFullAccess.

To have a full access to use the entire suite of SageMaker features, add AmazonSageMakerFullAccess.

The SageMaker model parallel library v2 reference

The following are references for the SageMaker model parallel library v2 (SMP v2).

Topics

- SMP v2 core feature configuration parameters
- Reference for the SMP v2 torch.sagemaker package
- Upgrade from SMP v1 to SMP v2

SMP v2 core feature configuration parameters

The following is a complete list of parameters to activate and configure the the section called “Core features of SMP v2”. These must be written in JSON format and passed to the PyTorch estimator in the SageMaker Python SDK or saved as a JSON file for SageMaker HyperPod.

```json
{
    "hybrid_shard_degree":  Integer,
    "sm_activation_offloading":  Boolean,
    "activation_loading_horizon":  Integer,
    "fsdp_cache_flush_warnings":  Boolean,
    "allow_empty_shards":  Boolean,
    "tensor_parallel_degree":  Integer,
    "tensor_parallel_seed":  Integer
}
```

- hybrid_shard_degree (Integer) – Specifies a sharded parallelism degree. The value must be an integer between 0 and world_size. The default value is 0.
- If set to 0, it falls back to the native PyTorch implementation and API in the script when tensor_parallel_degree is 1. Otherwise, it computes the largest possible hybrid_shard_degree based on tensor_parallel_degree and world_size. When falling back to the native PyTorch FSDP use cases, if FULL_SHARD is the strategy you use, it shards across the whole cluster of GPUs. If HYBRID_SHARD or _HYBRID_SHARD_ZERO2
was the strategy, it is equivalent to `hybrid_shard_degree` of 8. When tensor parallelism is enabled, it shards based on the revised `hybrid_shard_degree`.

- If set to 1, it falls back to the native PyTorch implementation and API for NO_SHARD in the script when `tensor_parallel_degree` is 1. Otherwise, it's equivalent to NO_SHARD within any given tensor parallel groups.

- If set to an integer between 2 and `world_size`, sharding happens across the specified number of GPUs. If you don't set up `sharding_strategy` in the FSDP script, it gets overridden to HYBRID_SHARD. If you set _HYBRID_SHARD_ZERO2, the `sharding_strategy` you specify is used.

- `sm_activation_offloading` (Boolean) – Specifies whether to enable the SMP activation offloading implementation. If False, offloading uses the native PyTorch implementation. If True, it uses the SMP activation offloading implementation. You also need to use the PyTorch activation offload wrapper (`torch.distributed.algorithms._checkpoint.checkpoint.checkpoint_wrapper.offload_wrapper`) in your script. To learn more, see the section called “Activation offloading”. The default value is True.

- `activation_loading_horizon` (Integer) – An integer specifying the activation offloading horizon type for FSDP. This is the maximum number of checkpointed or offloaded layers whose inputs can be in the GPU memory simultaneously. To learn more, see the section called “Activation offloading”. The input value must be a positive integer. The default value is 2.

- `fsdp_cache_flush_warnings` (Boolean) – Detects and warns if cache flushes happen in the PyTorch memory manager, because they can degrade computational performance. The default value is True.

- `allow_empty_shards` (Boolean) – Whether to allow empty shards when sharding tensors if tensor is not divisible. This is an experimental fix for crash during checkpointing in certain scenarios. Disabling this falls back to the original PyTorch behavior. The default value is False.

- `tensor_parallel_degree` (Integer) – Specifies a tensor parallelism degree. The value must be between 1 and `world_size`. The default value is 1. Passing a value greater than 1 does not enable tensor parallelism automatically. You also need to use the the section called “torch.sagemaker.transform” API to wrap the model in your training script. To learn more, see the section called “Tensor parallelism”.

- `tensor_parallel_seed` (Integer) – A seed number for the random operations in tensor-parallel distributed modules. This seed will be added to the tensor parallel rank to set the actual seed for each rank. It is unique for each tensor parallel rank. SMP v2 makes sure that random
number generation across tensor parallel ranks matches the non-tensor parallelism case. To learn more, see the section called “Tensor parallelism”.

Reference for the SMP v2 torch.sagemaker package

This section is a reference for the torch.sagemaker package provided by SMP v2.

Topics

- torch.sagemaker.delayed_param.DelayedParamIniter
- torch.sagemaker.nn.attn.FlashSelfAttention
- torch.sagemaker.nn.attn.FlashGroupedQueryAttention
- torch.sagemaker.nn.huggingface.llama_flashattn.LlamaFlashAttention
- torch.sagemaker.transform
- torch.sagemaker util functions and properties

torch.sagemaker.delayed_param.DelayedParamIniter

An API for applying the section called “Delayed parameter initialization” to a PyTorch model.

```python
class torch.sagemaker.delayed_param.DelayedParamIniter(
    model: nn.Module,
    init_method_using_config : Callable = None,
    verbose: bool = False,
)
```

Parameters

- model (nn.Module) – A PyTorch model to wrap and apply the delayed parameter initialization functionality of SMP v2.
- init_method_using_config (Callable) – If you use the tensor parallel implementation of SMP v2 or supported the section called “Hugging Face Transformer models compatible with the SMP tensor parallelism”, keep this parameter at the default value, which is None. By default, the DelayedParamIniter API finds out how to initialize the given model correctly. For any other models, you need to create a custom parameter initialization function and add it to your script. The following code snippet is the default init_method_using_config function that SMP v2 implemented for the the section called “Hugging Face Transformer models compatible with the SMP tensor parallelism”. Use the following code snippet as a reference for creating
your own initialization configuration function, adding it to your script, and passing it to the `init_method_using_config` parameter of the SMP DelayedParamIniter API.

```python
from torch.sagemaker.utils.module_utils import empty_module_params, move_buffers_to_device

# Define a custom init config function.
def custom_init_method_using_config(module):
    d = torch.cuda.current_device()
    empty_module_params(module, device=d)
    if isinstance(module, (nn.Linear, Conv1D)):
        module.weight.data.normal_(mean=0.0, std=config.initializer_range)
        if module.bias is not None:
            module.bias.data.zero_()
    elif isinstance(module, nn.Embedding):
        module.weight.data.normal_(mean=0.0, std=config.initializer_range)
        if module.padding_idx is not None:
            module.weight.data[module.padding_idx].zero_()
    elif isinstance(module, nn.LayerNorm):
        module.weight.data.fill_(1.0)
        module.bias.data.zero_()
    elif isinstance(module, LlamaRMSNorm):
        module.weight.data.fill_(1.0)
    move_buffers_to_device(module, device=d)

delayed_initer = DelayedParamIniter(model,
    init_method_using_config=custom_init_method_using_config)
```

For more information about the `torch.sagemaker.module_util` functions in the preceding code snippet, see the section called “tor\sagemaker util functions and properties”.

- **verbose** (Boolean) – Whether to enable more detailed logging during initialization and validation. The default value is False.

**Methods**

- **get_param_init_fn()** – Returns the parameter initialization function that you can pass to the `param_init_fn` argument of the PyTorch FSDP wrapper class.

- **get_post_param_init_fn()** – Returns the parameter initialization function that you can pass to the `post_param_init_fn` argument of the PyTorch FSDP wrapper class. This is needed when you have tied weights in the model. The model must implement the method
tie_weights. For more information, see the **Notes on tied weight** in the section called “Delayed parameter initialization”.

  - Tracks how many parameters are being initialized by the parameter initialization function. This helps implement the following validate_params_and_buffers_inited method. You usually don’t need to call this function explicitly, because the validate_params_and_buffers_inited method implicitly calls this method in the backend.

- **validate_params_and_buffers_inited** (enabled: bool=True) – This is a context manager that helps validate that the number of parameters initialized matches the total number of parameters in the model. It also validates that all parameters and buffers are now on GPU devices instead of meta devices. It raises AssertionError if these conditions are not met. This context manager is only optional and you’re not required to use this context manager to initialize parameters.

**torch.sagemaker.nn.attn.FlashSelfAttention**

An API for using the section called “FlashAttention” with SMP v2.

```python
class torch.sagemaker.nn.attn.FlashSelfAttention(
    attention_dropout_prob: float = 0.0,
    scale: Optional[float] = None,
    triton_flash_attention: bool = False,
    use_alibi: bool = False,
)
```

**Parameters**

- **attention_dropout_prob** (float) – The dropout probability to apply to attention. The default value is 0.0.

- **scale** (float) – If passed, this scale factor will be applied for softmax. If set to None (which is also the default value), the scale factor is 1 / sqrt(attention_head_size). The default value is None.

- **triton_flash_attention** (bool) – If passed, Triton implementation of flash attention will be used. This is necessary to support Attention with Linear Biases (ALiBi) (see the following use_alibi parameter). This version of the kernel doesn’t support dropout. The default value is False.
- `use_alibi` (bool) – If passed, it enables Attention with Linear Biases (ALiBi) using the mask provided. When using ALiBi, it needs an attention mask prepared as follows. The default value is False.

```python
def generate_alibi_attn_mask(attention_mask, batch_size, seq_length, num_attention_heads, alibi_bias_max=8):
    device, dtype = attention_mask.device, attention_mask.dtype
    alibi_attention_mask = torch.zeros(1, num_attention_heads, 1, seq_length, dtype=dtype, device=device)
    alibi_bias = torch.arange(1 - seq_length, 1, dtype=dtype, device=device).view(1, 1, 1, seq_length)
    m = torch.arange(1, num_attention_heads + 1, dtype=dtype, device=device)
    m.mul_(alibi_bias_max / num_attention_heads)
    alibi_bias = alibi_bias * (1.0 / (2 ** m.view(1, num_attention_heads, 1, 1)))
    alibi_attention_mask.add_(alibi_bias)
    alibi_attention_mask = alibi_attention_mask[..., :seq_length, :seq_length]
    if attention_mask is not None and attention_mask.bool().any():
        attention_mask.bool().view(batch_size, 1, 1, seq_length).masked_fill(float("-inf")
    return alibi_attention_mask
```

**Methods**

- `forward(self, qkv, attn_mask=None, causal=False, cast_dtype=None, layout="b h s d")` – A regular PyTorch module function. When a module(x) is called, SMP runs this function automatically.

  - `qkv` – torch.Tensor of the following form: `(batch_size x seqlen x (3 x num_heads) x head_size)` or `(batch_size, (3 x num_heads) x seqlen x head_size)`, a tuple of torch.Tensors each of which might be of shape `(batch_size x seqlen x num_heads x head_size)`, or `(batch_size x num_heads x seqlen x head_size)`. An appropriate layout arg must be passed based on the shape.

  - `attn_mask` – torch.Tensor of the following form `(batch_size x 1 x 1 x seqlen)`. To enable this attention mask parameter, it requires `triton_flash_attention=True` and
use_alibi=True. To learn how to generate an attention mask using this method, see the code examples at the section called “FlashAttention”. The default value is None.

- **causal** – When set to False, which is the default value of the argument, no mask is applied. When set to True, the forward method uses the standard lower triangular mask. The default value is False.

- **cast_dtype** – When set to a particular dtype, it casts the qkv tensors to that dtype before attn. This is useful for implementations such as the Hugging Face Transformer GPT-NeoX model, which has q and k with fp32 after rotary embeddings. If set to None, no cast is applied. The default value is None.

- **layout** (string) – Available values are b h s d or b s h d. This should be set to the layout of qkv tensors passed, so appropriate transformations can be applied for attn. The default value is b h s d.

**Returns**

A single torch.Tensor with shape (batch_size x num_heads x seq_len x head_size).

torch.sagemaker.nn.attn.FlashGroupedQueryAttention

An API for using FlashGroupedQueryAttention with SMP v2. To learn more about the usage of this API, see the section called “Use FlashAttention kernels for grouped-query attention”.

class torch.sagemaker.nn.attn.FlashGroupedQueryAttention(
    attention_dropout_prob: float = 0.0,
    scale: Optional[float] = None,
)

**Parameters**

- **attention_dropout_prob** (float) – The dropout probability to apply to attention. The default value is 0.0.

- **scale** (float) – If passed, this scale factor is applied for softmax. If set to None, 1 / sqrt(attention_head_size) is used as the scale factor. The default value is None.

**Methods**
• `forward(self, q, kv, causal=False, cast_dtype=None, layout="b s h d")`
  – A regular PyTorch module function. When a module(x) is called, SMP runs this function automatically.

  • `q` – `torch.Tensor` of the following form `(batch_size x seqlen x num_heads x head_size)` or `(batch_size x num_heads x seqlen x head_size)`. Appropriate layout arg must be passed based on the shape.

  • `kv` – `torch.Tensor` of the following form `(batch_size x seqlen x (2 x num_heads) x head_size)` or `(batch_size, (2 x num_heads) x seqlen x head_size)`, or a tuple of two `torch.Tensors`, each of which might be of shape `(batch_size x seqlen x num_heads x head_size)` or `(batch_size x num_heads x seqlen x head_size)`. Appropriate layout argument must also be passed based on the shape.

  • `causal` – When set to False, which is the default value of the argument, no mask is applied. When set to True, the forward method uses the standard lower triangular mask. The default value is False.

  • `cast_dtype` – When set to a particular dtype, it casts the qkv tensors to that dtype before attn. This is useful for implementations such as Hugging Face Transformers GPT-NeoX, which has q, k with fp32 after rotary embeddings. If set to None, no cast is applied. The default value is None.

  • `layout` (string) – Available values are "b h s d" or "b s h d". This should be set to the layout of qkv tensors passed, so appropriate transformations can be applied for attn. The default value is "b h s d".

Returns

Returns a single `torch.Tensor` `(batch_size x num_heads x seq_len x head_size)` that represents the output of attention computation.

torch.sagemaker.nn.huggingface.llama_flashattn.LlamaFlashAttention

An API that supports FlashAttention for the Llama model. This API uses the [section called "torch.sagemaker.nn.attn.FlashGroupedQueryAttention"](https://sagemaker.readthedocs.io/en/stable/) API at low level. To learn how to use this, see the [section called “Use FlashAttention kernels for grouped-query attention”](https://sagemaker.readthedocs.io/en/stable/).

class torch.sagemaker.nn.huggingface.llama_flashattn.LlamaFlashAttention(
    config: LlamaConfig
)
Parameters

- **config** – A FlashAttention configuration for the Llama model.

Methods

- **forward**(*self, hidden_states, attention_mask, position_ids, past_key_value, output_attentions, use_cache*)
  - **hidden_states** (*torch.Tensor*) – Hidden states of a tensor in form of (batch_size x seq_len x num_heads x head_size).
  - **attention_mask** (*torch.LongTensor*) – Mask to avoid performing attention on padding token indices in form of (batch_size x seqlen). The default value is None.
  - **position_ids** (*torch.LongTensor*) – When not being None, it is in form of (batch_size x seqlen), indicating the indices of positions of each input sequence token in the position embeddings. The default value is None.
  - **past_key_value** (*Cache*) – Pre-computed hidden-states (key and values in the self-attention blocks and in the cross-attention blocks). The default value is None.
  - **output_attentions** (*bool*) – Indicates whether to return the attentions tensors of all attention layers. The default value is False.
  - **use_cache** (*bool*) – Indicates whether to return past_key_values key value states. The default value is False.

Returns

Returns a single *torch.Tensor* (batch_size x num_heads x seq_len x head_size) that represents the output of attention computation.

**torch.sagemaker.transform**

SMP v2 provides this `torch.sagemaker.transform()` API for transforming Hugging Face Transformer models to SMP model implementations and enabling the SMP tensor parallelism.

```python
torch.sagemaker.transform(
    model: nn.Module,
    device: Optional[torch.device] = None,
    dtype: Optional[torch.dtype] = None,
    config: Optional[Dict] = None,
    load_state_dict_from_rank0: bool = False
)```
SMP v2 maintains transformation policies for the models compatible with the SMP tensor parallelism by converting the configuration of the Hugging Face Transformer models to the SMP transformer configuration.

**Parameters**

- **model (torch.nn.Module)** – A model from the section called “Hugging Face Transformer models compatible with the SMP tensor parallelism” to transform and apply the tensor parallelism feature of the SMP library.

- **device (torch.device)** – If passed, a new model is created on this device. If the original module has any parameter on meta device (see the section called “Delayed parameter initialization”), then the transformed module will also be created on meta device, ignoring the argument passed here. The default value is None.

- **dtype (torch.dtype)** – If passed, sets this as the dtype context manager for the creation of the model and creates a model with this dtype. This is typically unnecessary, as we want to create the model with fp32 when using MixedPrecision, and fp32 is the default dtype in PyTorch. The default value is None.

- **config (dict)** – This is a dictionary for configuring the SMP transformer. The following are the available keys. The default value is None.
  - **sequence_parallel (Boolean)** – A Boolean specifying whether to use sequence parallelism. Sequence parallelism partitions input sequences along the sequence dimensions to improve memory efficiency. The default value is True.
  - **load_state_dict_from_rank0 (Boolean)** – By default, this module creates a new instance of the model with new weights. When this argument is set to True, SMP tries to load the state dictionary of the original PyTorch model from the 0th rank into transformed model for the tensor parallel group that the 0th rank is part of. When this is set to True, rank 0 can't have any parameters on meta device. Only the first tensor parallel group populates the weights from the 0th rank after this transform call. You need to set sync_module_states to True in the FSDP wrapper to get these weights from the first tensor parallel group to all other processes. With this activated, the SMP library loads the state dictionary from the original model. The SMP library takes the state_dict of the model before transform, converts it to match the structure of the transformed model, shards it for each tensor parallel rank, communicates this state from the 0th rank to other ranks in the tensor parallel group that the 0th rank is part of, and loads it. The default value is False.
Returns

Returns a transformed model that you can wrap with PyTorch FSDP. When `load_state_dict_from_rank0` is set to `True`, the tensor parallel group that involves rank 0 has weights loaded from the original state dictionary on rank 0. When using the section called “Delayed parameter initialization” on the original model, only these ranks have the actual tensors on CPUs for the parameters and buffers of the transformed model. The rest of the ranks continue to have the parameters and buffers on the meta device to save memory.

`torch.sagemaker` util functions and properties

`torch.sagemaker` util functions

- `torch.sagemaker.init(config: Optional[Union[str, Dict[str, Any]]] = None) -> None` – Initializes the PyTorch training job with SMP.
- `torch.sagemaker.is_initialized() -> bool` – Checks whether the training job is initialized with SMP. When falling back to the native PyTorch while the job is initialized with SMP, some of the properties are not relevant and become `None`, as indicated in the following `Properties` list.
- `torch.sagemaker.utils.module_utils.empty_module_params(module: nn.Module, device: Optional[torch.device] = None, recurse: bool = False) -> nn.Module` – Creates empty parameters on the given device if any, and it can be recursive for all nested modules if specified.
- `torch.sagemaker.utils.module_utils.move_buffers_to_device(module: nn.Module, device: torch.device, recurse: bool = False) -> nn.Module` – Moves module buffers to the given device, and it can be recursive for all nested modules if specified.

Properties

`torch.sagemaker.state` holds multiple useful properties after the initialization of SMP with `torch.sagemaker.init`.

- `torch.sagemaker.state.hybrid_shard_degree (int)` – The sharded data parallelism degree, a copy from user input in the SMP configuration passed to `torch.sagemaker.init()`. To learn more, see the section called “Get started with SMP v2”.
- `torch.sagemaker.state.rank (int)` – The global rank for the device, in the range of `[0, world_size)`.
• torch.sagemaker.state.rep_rank_process_group
  (torch.distributed.ProcessGroup) – The process group including all devices
  with the same replication rank. Note the subtle but fundamental difference with
  torch.sagemaker.state.tp_process_group. When falling back to native PyTorch, it
  returns None.

• torch.sagemaker.state.tensor_parallel_degree (int) – The tensor parallelism degree,
  a copy from user input in the SMP configuration passed to torch.sagemaker.init(). To learn
  more, see the section called “Get started with SMP v2”.

• torch.sagemaker.state.tp_size (int) – An alias to
  torch.sagemaker.state.tensor_parallel_degree.

• torch.sagemaker.state.tp_rank (int) – The tensor parallelism rank for the device in
  the range of \([0, \text{tp_size})\), determined by the tensor parallelism degree and the ranking
  mechanism.

• torch.sagemaker.state.tp_process_group (torch.distributed.ProcessGroup) –
  The tensor parallel process group including all devices with the same rank in other dimensions
  (for example, sharded data parallelism and replication) but unique tensor parallel ranks. When
  falling back to native PyTorch, it returns None.

• torch.sagemaker.state.world_size (int) – The total number of devices used in training.

**Upgrade from SMP v1 to SMP v2**

To move from SMP v1 to SMP v2, you must make script changes to remove the SMP v1 APIs and
apply the SMP v2 APIs. Instead of starting from your SMP v1 script, we recommend you start from
a PyTorch FSDP script, and follow the instructions at the section called “Get started with SMP v2”.

To bring SMP v1 models to SMP v2, in SMP v1 you must collect the full model state dictionary
and apply the translation functions on the model state dictionary to convert it into the Hugging
Face Transformers model checkpoint format. Then in SMP v2, as discussed in the section called
“Save and load checkpoints while using SMP”, you can load the Hugging Face Transformers model
checkpoints, and then continue with using the PyTorch checkpoint APIs with SMP v2. To use SMP
with your PyTorch FSDP model, make sure that you move to SMP v2 and make changes to your
training script to use PyTorch FSDP and other latest features.

```python
import smdistributed.modelparallel.torch as smp

# Create model
```
model = ...
model = smp.DistributedModel(model)

# Run training
...

# Save v1 full checkpoint
if smp.rdp_rank() == 0:
    model_dict = model.state_dict(gather_to_rank0=True)  # save the full model
    # Get the corresponding translation function in smp v1 and translate
    if model_type == "gpt_neox":  # save the full model
        from smdistributed.modelparallel.torch.nn.huggingface.gptneox import translate_state_dict_to_hf_gptneox
        translated_state_dict = translate_state_dict_to_hf_gptneox(state_dict, max_seq_len=None)

    # Save the checkpoint
    checkpoint_path = "checkpoint.pt"
    if smp.rank() == 0:
        smp.save(
            {"model_state_dict": translated_state_dict},
            checkpoint_path,
            partial=False,
        )

To find available translation functions in SMP v1, see the section called “Support for Hugging Face Transformer Models”.

For instruction on model checkpoints saving and loading in SMP v2, see the section called “Save and load checkpoints while using SMP”.

Release notes for the SageMaker model parallelism library

See the following release notes to track the latest updates for the SageMaker model parallelism (SMP) library. If you have further questions about the SMP library, contact the SMP service team at sm-model-parallel-feedback@amazon.com.

The SageMaker model parallelism library v2.2.0

Date: March 7, 2024

New Features
• Added support for **FP8 training** of the following Hugging Face transformer models on P5 instances with Transformer Engine integration:
  • GPT-NeoX
  • Llama 2

**Bug Fixes**

• Fixed a bug where tensors were not guaranteed to be contiguous before the `AllGather` collective call during tensor parallelism training.

**Currency Updates**

• Added support for PyTorch v2.2.0.
• Upgraded the SMDDP library to v2.2.0.
• Upgraded the FlashAttention library to v2.3.3.
• Upgraded the NCCL library to v2.19.4.

**Deprecation**

• Discontinued support for Transformer Engine versions before v1.2.0.

**Known issues**

• The SMP [the section called “Activation offloading”](https://github.com/pytorch/pytorch/issues/117748) feature currently does not work. Use the native PyTorch activation offloading instead.

**Other changes**

• Included a patch to fix the performance regression discussed in the issue thread at [https://github.com/pytorch/pytorch/issues/117748](https://github.com/pytorch/pytorch/issues/117748) in the PyTorch GitHub repository.

**SMP Docker container**

The SMP library team distributes Docker containers in replacement of the SageMaker PyTorch framework containers. If you use the PyTorch estimator class in the SageMaker Python SDK and specify distribution configuration to use SMP v2, SageMaker automatically picks up the SMP
Docker containers. To use this release of SMP v2, upgrade your SageMaker Python SDK to v2.212.0 or later.

- SMP Docker container for PyTorch v2.2.0 with CUDA v12.1
  
  658645717510.dkr.ecr.us-west-2.amazonaws.com/smdistributed-modelparallel:2.2.0-gpu-py310-cu121

- Available for P4d, P4de, and P5 instances
- Pre-installed packages in this Docker container
  - The SMDDP library v2.2.0
  - CUDNN v8.9.5.29
  - FlashAttention v2.3.3
  - TransformerEngine v1.2.1
  - Hugging Face Transformers v4.37.1
  - Hugging Face Datasets library v2.16.1
  - EFA v1.30.0
  - NCCL v2.19.4

**The SageMaker model parallelism library v2.1.0**

*Date: February 6, 2024*

**Currency Updates**

- Added support for PyTorch v2.1.2.

**Deprecation**

- Discontinued support for Hugging Face Transformers v4.31.0.

**Known issues**

- An issue is discovered that fine-tuning of the Hugging Face Llama 2 model with `attn_implementation=flash_attention_2` and FSDP causes the model to diverge. For reference, see the issue ticket in the Hugging Face Transformers GitHub repository. To avoid the
divergence issue, use attn_implementation=sdpa. Alternatively, use the SMP transformer model implementation by setting up use_smp_implementation=True.

**SMP Docker container**

The SMP library team distributes Docker containers in replacement of the SageMaker PyTorch framework containers. If you use the PyTorch estimator class in the SageMaker Python SDK and specify distribution configuration to use SMP v2, SageMaker automatically picks up the SMP Docker containers. To use this release of SMP v2, upgrade your SageMaker Python SDK to v2.207.0 or later.

- SMP Docker container for PyTorch v2.1.2 with CUDA v12.1

  658645717510.dkr.ecr.us-west-2.amazonaws.com/smdistributed-modelparallel:2.1.2-gpu-py310-cu121

- Available for P4d, P4de, and P5 instances
- Pre-installed packages in this Docker container
  - The SMDDP library v2.1.0
  - CUDNN v8.9.5.29
  - FlashAttention v2.3.3
  - TransformerEngine v1.2.1
  - Hugging Face Transformers v4.37.1
  - Hugging Face Datasets library v2.16.1
  - EFA v1.30.0

**SMP Conda channel**

The following S3 bucket is a public Conda channel hosted by the SMP service team. If you want to install the SMP v2 library in an environment of highly customizable compute resources such as SageMaker HyperPod clusters, use this Conda channel to properly install the SMP library.


For more information about Conda channels in general, see [Channels](https://conda.readthedocs.io/en/latest/channel_architecture.html) in the *Conda documentation*. 
The SageMaker model parallelism library v2.0.0

Date: December 19, 2023

New features

Released the SageMaker model parallelism (SMP) library v2.0.0 with the following new offerings.

- A new `torch.sagemaker` package, entirely revamped from the previous `smdistributed.modelparallel.torch` package in SMP v1.x.
- Support for PyTorch 2.0.1.
- Support for PyTorch FSDP.
- Tensor parallelism implementation by integrating with the Transformer Engine library.
- Support for both SageMaker Training and SageMaker HyperPod.

Breaking changes

- SMP v2 revamped the APIs entirely and provides the `torch.sagemaker` package. Mostly, you only need to initialize with the `torch.sagemaker.init()` module and pass model parallel configuration parameters. With this new package, you can significantly simplify code modifications in your training script. To learn more about adapting your training script to use SMP v2, see the section called “Get started with SMP v2”.
- If you've used SMP v1 for training Hugging Face Transformer models and want to reuse the models in SMP v2, see the section called “Upgrade from SMP v1 to SMP v2”.
- For PyTorch FSDP training, you should use SMP v2.

Known issues

- Activation checkpointing currently only works with the following wrapping policies with FSDP.
  - `auto_wrap_policy = functools.partial(transformer_auto_wrap_policy, ...)`
- To use the section called “Activation offloading”, FSDP activation checkpointing type must be `REENTRANT`.
- When running with tensor parallel enabled with the sharded data parallel degree set to 1, you must use `backend = ncc1`. The `smddp` backend option is not supported in this scenario.
- Transformer Engine is required to use PyTorch with the SMP library even when not using tensor parallelism.
Other changes

- Starting from this release, the documentation for the SageMaker model parallelism library is fully available in this Amazon SageMaker Developer Guide. In favor of this complete developer guide for SMP v2 in the Amazon SageMaker Developer Guide, the additional reference for SMP v1.x in the SageMaker Python SDK documentation is deprecated. If you still need the documentation for SMP v1.x, the developer guide for SMP v1.x is available at the section called "(Archived) SageMaker model parallelism library v1.x", and the SMP Python library v1.x reference is available in the SageMaker Python SDK v2.199.0 documentation.

Deprecations

- Discontinued support for TensorFlow.
- There is no pipeline parallelism support in SMP v2.
- There is no support for the DeepSpeed library in favor of native PyTorch FSDP.

SMP Docker container

The SMP library team distributes Docker containers in replacement of the SageMaker PyTorch framework containers. If you use the PyTorch estimator class in the SageMaker Python SDK and specify distribution configuration to use SMP v2, SageMaker automatically picks up the SMP Docker containers. To use this release of SMP v2, upgrade your SageMaker Python SDK to v2.207.0 or later.

- SMP Docker container for PyTorch v2.0.1 with CUDA v12.1

(Archived) SageMaker model parallelism library v1.x

⚠️ Important

As of December 19, 2023, the SageMaker model parallelism (SMP) library v2 is released. In favor of the SMP library v2, the SMP v1 capabilities are no longer supported in future releases. The following section and topics are archived and specific to using the SMP library.
Use Amazon SageMaker's model parallel library to train large deep learning (DL) models that are difficult to train due to GPU memory limitations. The library automatically and efficiently splits a model across multiple GPUs and instances. Using the library, you can achieve a target prediction accuracy faster by efficiently training larger DL models with billions or trillions of parameters.

You can use the library to automatically partition your own TensorFlow and PyTorch models across multiple GPUs and multiple nodes with minimal code changes. You can access the library's API through the SageMaker Python SDK.

Use the following sections to learn more about model parallelism and the SageMaker model parallel library. This library's API documentation is located at Distributed Training APIs in the SageMaker Python SDK v2.199.0 documentation.

Topics

- Introduction to Model Parallelism
- Supported Frameworks and AWS Regions
- Core Features of the SageMaker Model Parallelism Library
- Run a SageMaker Distributed Training Job with Model Parallelism
- Checkpointing and Fine-Tuning a Model with Model Parallelism
- Amazon SageMaker model parallelism library v1 examples
- SageMaker Distributed Model Parallelism Best Practices
- The SageMaker Distributed Model Parallelism Library Configuration Tips and Pitfalls
- Model Parallel Troubleshooting

Introduction to Model Parallelism

Model parallelism is a distributed training method in which the deep learning model is partitioned across multiple devices, within or across instances. This introduction page provides a high-level overview about model parallelism, a description of how it can help overcome issues that arise when training DL models that are typically very large in size, and examples of what the SageMaker model parallel library offers to help manage model parallel strategies as well as memory consumption.
What is Model Parallelism?

Increasing the size of deep learning models (layers and parameters) yields better accuracy for complex tasks such as computer vision and natural language processing. However, there is a limit to the maximum model size you can fit in the memory of a single GPU. When training DL models, GPU memory limitations can be bottlenecks in the following ways:

- They limit the size of the model you can train, since the memory footprint of a model scales proportionally to the number of parameters.
- They limit the per-GPU batch size during training, driving down GPU utilization and training efficiency.

To overcome the limitations associated with training a model on a single GPU, SageMaker provides the model parallel library to help distribute and train DL models efficiently on multiple compute nodes. Furthermore, with the library, you can achieve most optimized distributed training using EFA-supported devices, which enhance the performance of inter-node communication with low latency, high throughput, and OS bypass.

Estimate Memory Requirements Before Using Model Parallelism

Before you use the SageMaker model parallel library, consider the following to get a sense of the memory requirements of training large DL models.

For a training job that uses AMP (FP16) and Adam optimizers, the required GPU memory per parameter is about 20 bytes, which we can break down as follows:

- An FP16 parameter ~ 2 bytes
- An FP16 gradient ~ 2 bytes
- An FP32 optimizer state ~ 8 bytes based on the Adam optimizers
- An FP32 copy of parameter ~ 4 bytes (needed for the `optimizer apply` (OA) operation)
- An FP32 copy of gradient ~ 4 bytes (needed for the OA operation)

Even for a relatively small DL model with 10 billion parameters, it can require at least 200GB of memory, which is much larger than the typical GPU memory (for example, NVIDIA A100 with 40GB/80GB memory and V100 with 16/32 GB) available on a single GPU. Note that on top of the memory requirements for model and optimizer states, there are other memory consumers such as activations generated in the forward pass. The memory required can be a lot greater than 200GB.
For distributed training, we recommend that you use Amazon EC2 P3 and P4 instances that have NVIDIA V100 and A100 Tensor Core GPUs respectively. For more details about specifications such as CPU cores, RAM, attached storage volume, and network bandwidth, see the *Accelerated Computing* section in the *Amazon EC2 Instance Types* page.

Even with the accelerated computing instances, it is obvious that models with about 10 billion parameters such as Megatron-LM and T5 and even larger models with hundreds of billions of parameters such as GPT-3 cannot fit model replicas in each GPU device.

**How the Library Employs Model Parallelism and Memory Saving Techniques**

The library consists of various types of model parallelism features and memory-saving features such as optimizer state sharding, activation checkpointing, and activation offloading. All these techniques can be combined to efficiently train large models that consist of hundreds of billions of parameters.

**Topics**

- Sharded data parallelism *(available for PyTorch)*
- Pipeline parallelism *(available for PyTorch and TensorFlow)*
- Tensor parallelism *(available for PyTorch)*
- Optimizer state sharding *(available for PyTorch)*
- Activation offloading and checkpointing *(available for PyTorch)*
- Choosing the right techniques for your model

**Sharded data parallelism (available for PyTorch)**

*Sharded data parallelism* is a memory-saving distributed training technique that splits the state of a model (model parameters, gradients, and optimizer states) across GPUs within a data-parallel group.

SageMaker implements sharded data parallelism through the implementation of MiCS, which is a library that minimizes communication scale and discussed in the blog post *Near-linear scaling of gigantic-model training on AWS*.

You can apply sharded data parallelism to your model as a stand-alone strategy. Furthermore, if you are using the most performant GPU instances equipped with NVIDIA A100 Tensor Core GPUs, `ml.p4d.24xlarge`, you can take the advantage of improved training speed from the `AllGather` operation offered by SMDDP Collectives.
To dive deep into sharded data parallelism and learn how to set it up or use a combination of sharded data parallelism with other techniques like tensor parallelism and FP16 training, see the section called “Sharded Data Parallelism”.

**Pipeline parallelism (available for PyTorch and TensorFlow)**

*Pipeline parallelism* partitions the set of layers or operations across the set of devices, leaving each operation intact. When you specify a value for the number of model partitions (pipeline_parallel_degree), the total number of GPUs (processes_per_host) must be divisible by the number of the model partitions. To set this up properly, you have to specify the correct values for the pipeline_parallel_degree and processes_per_host parameters. The simple math is as follows:

\[
(pipeline\_parallel\_degree) \times (data\_parallel\_degree) = processes\_per\_host
\]

The library takes care of calculating the number of model replicas (also called data_parallel_degree) given the two input parameters you provide.

For example, if you set "pipeline_parallel_degree": 2 and "processes_per_host": 8 to use an ML instance with eight GPU workers such as `ml.p3.16xlarge`, the library automatically sets up the distributed model across the GPUs and four-way data parallelism. The following image illustrates how a model is distributed across the eight GPUs achieving four-way data parallelism and two-way pipeline parallelism. Each model replica, where we define it as a *pipeline parallel group* and label it as PP_GROUP, is partitioned across two GPUs. Each partition of the model is assigned to four GPUs, where the four partition replicas are in a *data parallel group* and labeled as DP_GROUP. Without tensor parallelism, the pipeline parallel group is essentially the model parallel group.
Sample model with four layers

Model parallel configuration

Batch 1
- microbatch 1
- microbatch 2
- microbatch 3
- microbatch 4

Batch 2
- microbatch 1
- microbatch 2
- microbatch 3
- microbatch 4

Batch 3
- microbatch 1
- microbatch 2
- microbatch 3
- microbatch 4

Batch 4
- microbatch 1
- microbatch 2
- microbatch 3
- microbatch 4

DP_GROUP

GPU 0
- L1
- L2

GPU 1
- L3
- L4

GPU 2
- L1
- L2

GPU 3
- L3
- L4

GPU 4
- L1
- L2

GPU 5
- L3
- L4

GPU 6
- L1
- L2

GPU 7
- L3
- L4

"pipeline_parallel_degree": 2,
"microbatches": 4,
"ddp": True,
To dive deep into pipeline parallelism, see Core Features of the SageMaker Model Parallelism Library.

To get started with running your model using pipeline parallelism, see Run a SageMaker Distributed Training Job with the SageMaker Model Parallel Library.

Tensor parallelism (available for PyTorch)

Tensor parallelism splits individual layers, or `nn.Module`s, across devices, to be run in parallel. The following figure shows the simplest example of how the library splits a model with four layers to achieve two-way tensor parallelism ("tensor_parallel_degree": 2). The layers of each model replica are bisected and distributed into two GPUs. In this example case, the model parallel configuration also includes "pipeline_parallel_degree": 1 and "ddp": True (uses PyTorch DistributedDataParallel package in the background), so the degree of data parallelism becomes eight. The library manages communication across the tensor-distributed model replicas.
The usefulness of this feature is in the fact that you can select specific layers or a subset of layers to apply tensor parallelism. To dive deep into tensor parallelism and other memory-saving features for PyTorch, and to learn how to set a combination of pipeline and tensor parallelism, see Tensor Parallelism.

**Optimizer state sharding (available for PyTorch)**

To understand how the library performs optimizer state sharding, consider a simple example model with four layers. The key idea in optimizing state sharding is you don't need to replicate your optimizer state in all of your GPUs. Instead, a single replica of the optimizer state is sharded across data-parallel ranks, with no redundancy across devices. For example, GPU 0 holds the optimizer state for layer one, the next GPU 1 holds the optimizer state for L2, and so on. The following animated figure shows a backward propagation with the optimizer state sharding technique. At the end of the backward propagation, there's compute and network time for the optimizer apply (OA) operation to update optimizer states and the all-gather (AG) operation to update the model parameters for the next iteration. Most importantly, the reduce operation can overlap with the compute on GPU 0, resulting in a more memory-efficient and faster backward propagation. In the current implementation, AG and OA operations do not overlap with compute. It can result in an extended computation during the AG operation, so there might be a tradeoff.
For more information about how to use this feature, see Optimizer State Sharding.

**Activation offloading and checkpointing (available for PyTorch)**

To save GPU memory, the library supports activation checkpointing to avoid storing internal activations in the GPU memory for user-specified modules during the forward pass. The library recomputes these activations during the backward pass. In addition, the activation offloading feature offloads the stored activations to CPU memory and fetches back to GPU during the backward pass to further reduce activation memory footprint. For more information about how to use these features, see Activation Checkpointing and Activation Offloading.
Choosing the right techniques for your model

For more information about choosing the right techniques and configurations, see SageMaker Distributed Model Parallel Best Practices and Configuration Tips and Pitfalls.

Supported Frameworks and AWS Regions

Before using the SageMaker model parallelism library, check the supported frameworks and instance types, and determine if there are enough quotas in your AWS account and AWS Region.

Note

To check the latest updates and release notes of the library, see the SageMaker Model Parallel Release Notes in the SageMaker Python SDK documentation.

Supported Frameworks

The SageMaker model parallelism library supports the following deep learning frameworks and is available in AWS Deep Learning Containers (DLC) or downloadable as a binary file.

PyTorch versions supported by SageMaker and the SageMaker model parallelism library

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<tr>
<th>PyTorch version</th>
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<th>smdistributed-modelparallel-integrated DLC image URI</th>
<th>URL of the binary file**</th>
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** Note  
The SageMaker model parallelism library v1.6.0 and later provides extended features for PyTorch. For more information, see [Core Features of the SageMaker Model Parallelism Library](https://aws.amazon.com/sagemaker/).**  

** The URLs of the binary files are for installing the SageMaker model parallelism library in custom containers. For more information, see the section called “Create Your Own Docker Container with the Library”.**
TensorFlow versions supported by SageMaker and the SageMaker model parallelism library

<table>
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</tr>
</tbody>
</table>

Hugging Face Transformers versions supported by SageMaker and the SageMaker distributed data parallel library

The AWS Deep Learning Containers for Hugging Face use the SageMaker Training Containers for PyTorch and TensorFlow as their base images. To look up the Hugging Face Transformers library versions and paired PyTorch and TensorFlow versions, see the latest Hugging Face Containers and the Prior Hugging Face Container Versions.

AWS Regions

The SageMaker data parallel library is available in all of the AWS Regions where the AWS Deep Learning Containers for SageMaker are in service. For more information, see Available Deep Learning Containers Images.

Supported Instance Types

The SageMaker model parallelism library requires one of the following ML instance types.
### Instance type

- ml.g4dn.12xlarge
- ml.p3.16xlarge
- ml.p3dn.24xlarge
- ml.p4d.24xlarge
- ml.p4de.24xlarge

For specs of the instance types, see the **Accelerated Computing** section in the [Amazon EC2 Instance Types page](https://aws.amazon.com/ec2/instance-types/). For information about instance pricing, see [Amazon SageMaker Pricing](https://aws.amazon.com/sagemaker/pricing/).

If you encountered an error message similar to the following, follow the instructions at [Request a service quota increase for SageMaker resources](https).

```
ResourceLimitExceeded: An error occurred (ResourceLimitExceeded) when calling the CreateTrainingJob operation: The account-level service limit 'ml.p3dn.24xlarge for training job usage' is 0 Instances, with current utilization of 0 Instances and a request delta of 1 Instances. Please contact AWS support to request an increase for this limit.
```

### Core Features of the SageMaker Model Parallelism Library

Amazon SageMaker's model parallelism library offers distribution strategies and memory-saving techniques, such as sharded data parallelism, tensor parallelism, model partitioning by layers for pipeline scheduling, and checkpointing. The model parallelism strategies and techniques help distribute large models across multiple devices while optimizing training speed and memory consumption. The library also provides Python helper functions, context managers, and wrapper functions to adapt your training script for automated or manual partitioning of your model.

When you implement model parallelism to your training job, you keep the same two-step workflow shown in the [Run a SageMaker Distributed Training Job with Model Parallelism](https) section. For adapting your training script, you'll add zero or few additional code lines to your training script. For launching a training job of the adapted training script, you'll need to set the distribution configuration parameters to activate the memory-saving features or to pass values for the degree of parallelism.
To get started with examples, see the following Jupyter notebooks that demonstrate how to use the SageMaker model parallelism library.

- **PyTorch example notebooks**
- **TensorFlow example notebooks**

To dive deep into the core features of the library, see the following topics.

**Note**

The SageMaker distributed training libraries are available through the AWS deep learning containers for PyTorch, Hugging Face, and TensorFlow within the SageMaker Training platform. To utilize the features of the distributed training libraries, we recommend that you use the SageMaker Python SDK. You can also manually configure in JSON request syntax if you use SageMaker APIs through SDK for Python (Boto3) or AWS Command Line Interface. Throughout the documentation, instructions and examples focus on how to use the distributed training libraries with the SageMaker Python SDK.

**Important**

The SageMaker model parallelism library supports all the core features for PyTorch, and supports pipeline parallelism for TensorFlow.

**Topics**

- **Sharded Data Parallelism**
- **Pipelining a Model**
- **Tensor Parallelism**
- **Optimizer State Sharding**
- **Activation Checkpointing**
- **Activation Offloading**
- **FP16 Training with Model Parallelism**
- **Support for FlashAttention**
Sharded Data Parallelism

*Sharded data parallelism* is a memory-saving distributed training technique that splits the state of a model (model parameters, gradients, and optimizer states) across GPUs in a data parallel group.

**Note**

Sharded data parallelism is available for PyTorch in the SageMaker model parallelism library v1.11.0 and later.

When scaling up your training job to a large GPU cluster, you can reduce the per-GPU memory footprint of the model by sharding the training state of the model over multiple GPUs. This returns two benefits: you can fit larger models, which would otherwise run out of memory with standard data parallelism, or you can increase the batch size using the freed-up GPU memory.

The standard data parallelism technique replicates the training states across the GPUs in the data parallel group, and performs gradient aggregation based on the *AllReduce* operation. Sharded data parallelism modifies the standard data-parallel distributed training procedure to account for the sharded nature of the optimizer states. A group of ranks over which the model and optimizer states are sharded is called a *sharding group*. The sharded data parallelism technique shards the trainable parameters of a model and corresponding gradients and optimizer states across the GPUs in the *sharding group*.

SageMaker achieves sharded data parallelism through the implementation of MiCS, which is discussed in the AWS blog post [Near-linear scaling of gigantic-model training on AWS](https://aws.amazon.com/blogs/machine-learning/near-linear-scaling-of-gigantic-model-training-on-aws/). In this implementation, you can set the sharding degree as a configurable parameter, which must be less than the data parallelism degree. During each forward and backward pass, MiCS temporarily recombines the model parameters in all GPUs through the *AllGather* operation. After the forward or backward pass of each layer, MiCS shards the parameters again to save GPU memory. During the backward pass, MiCS reduces gradients and simultaneously shards them across GPUs through the *ReduceScatter* operation. Finally, MiCS applies the local reduced and sharded gradients to their corresponding local parameter shards, using the local shards of optimizer states. To bring down communication overhead, the SageMaker model parallelism library prefetches the upcoming layers in the forward or backward pass, and overlaps the network communication with the computation.

The training state of the model is replicated across the sharding groups. This means that before gradients are applied to the parameters, the *AllReduce* operation must take place across the
sharding groups, in addition to the ReduceScatter operation that takes place within the sharding group.

In effect, sharded data parallelism introduces a tradeoff between the communication overhead and GPU memory efficiency. Using sharded data parallelism increases the communication cost, but the memory footprint per GPU (excluding the memory usage due to activations) is divided by the sharded data parallelism degree, thus larger models can be fit in the GPU cluster.

**Selecting the degree of sharded data parallelism**

When you select a value for the degree of sharded data parallelism, the value must evenly divide the degree of data parallelism. For example, for an 8-way data parallelism job, choose 2, 4, or 8 for the sharded data parallelism degree. While choosing the sharded data parallelism degree, we recommend that you start with a small number, and gradually increase it until the model fits in the memory together with the desired batch size.

**Selecting the batch size**

After setting up sharded data parallelism, make sure you find the most optimal training configuration that can successfully run on the GPU cluster. For training large language models (LLM), start from the batch size 1, and gradually increase it until you reach the point to receive the out-of-memory (OOM) error. If you encounter the OOM error even with the smallest batch size, apply a higher degree of sharded data parallelism or a combination of sharded data parallelism and tensor parallelism.

**Topics**

- [How to apply sharded data parallelism to your training job](#)
- [Reference configurations](#)
- [Sharded data parallelism with SMDDP Collectives](#)
- [Mixed precision training with sharded data parallelism](#)
- [Sharded data parallelism with tensor parallelism](#)
- [Tips and considerations for using sharded data parallelism](#)

**How to apply sharded data parallelism to your training job**

To get started with sharded data parallelism, apply required modifications to your training script, and set up the SageMaker PyTorch estimator with the sharded-data-parallelism-specific parameters. Also consider to take reference values and example notebooks as a starting point.
Adapt your PyTorch training script

Follow the instructions at [Step 1: Modify a PyTorch Training Script](#) to wrap the model and optimizer objects with the `smdistributed.modelparallel.torch` wrappers of the `torch.nn.parallel` and `torch.distributed` modules.

(Optional) Additional modification to register external model parameters

If your model is built with `torch.nn.Module` and uses parameters that is not defined within the module class, you should register them to the module manually for SMP to gather the full parameters while . To register parameters to a module, use `smp.register_parameter(module, parameter)`.

```python
class Module(torch.nn.Module):
    def __init__(self, *args):
        super().__init__(self, *args)
        self.layer1 = Layer1()
        self.layer2 = Layer2()
        smp.register_parameter(self, self.layer1.weight)

    def forward(self, input):
        x = self.layer1(input)
        # self.layer1.weight is required by self.layer2.forward
        y = self.layer2(x, self.layer1.weight)
        return y
```

Set up the SageMaker PyTorch estimator

When configuring a SageMaker PyTorch estimator in the section called “Step 2: Launch a Training Job”, add the parameters for sharded data parallelism.

To turn on sharded data parallelism, add the `sharded_data_parallel_degree` parameter to the SageMaker PyTorch Estimator. This parameter specifies the number of GPUs over which the training state is sharded. The value for `sharded_data_parallel_degree` must be an integer between one and the data parallelism degree and must evenly divide the data parallelism degree. Note that the library automatically detects the number of GPUs so thus the data parallel degree. The following additional parameters are available for configuring sharded data parallelism.

- "sdp_reduce_bucket_size" *(int, default: 5e8)* – Specifies the size of PyTorch DDP gradient buckets in number of elements of the default dtype.
• "sdp_param_persistence_threshold" (int, default: 1e6) – Specifies the size of a parameter tensor in number of elements that can persist at each GPU. Sharded data parallelism splits each parameter tensor across GPUs of a data parallel group. If the number of elements in the parameter tensor is smaller than this threshold, the parameter tensor is not split; this helps reduce communication overhead because the parameter tensor is replicated across data-parallel GPUs.

• "sdp_max_live_parameters" (int, default: 1e9) – Specifies the maximum number of parameters that can simultaneously be in a recombined training state during the forward and backward pass. Parameter fetching with the AllGather operation pauses when the number of active parameters reaches the given threshold. Note that increasing this parameter increases the memory footprint.

• "sdp_hierarchical_allgather" (bool, default: True) – If set to True, the AllGather operation runs hierarchically: it runs within each node first, and then runs across nodes. For multi-node distributed training jobs, the hierarchical AllGather operation is automatically activated.

• "sdp_gradient_clipping" (float, default: 1.0) – Specifies a threshold for gradient clipping the L2 norm of the gradients before propagating them backward through the model parameters. When sharded data parallelism is activated, gradient clipping is also activated. The default threshold is 1.0. Adjust this parameter if you have the exploding gradients problem.

The following code shows an example of how to configure sharded data parallelism.

```python
import sagemaker
from sagemaker.pytorch import PyTorch

smp_options = {
    "enabled": True,
    "parameters": {
        # "pipeline_parallel_degree": 1,  # Optional, default is 1
        # "tensor_parallel_degree": 1,   # Optional, default is 1
        "ddp": True,
        # parameters for sharded data parallelism
        "sharded_data_parallel_degree": 2,  # Add this to activate sharded data parallelism
        "sdp_reduce_bucket_size": int(5e8),  # Optional
        "sdp_param_persistence_threshold": int(1e6),  # Optional
        "sdp_max_live_parameters": int(1e9),  # Optional
        "sdp_hierarchical_allgather": True,  # Optional
    }
}
```
"sdp_gradient_clipping": 1.0  # Optional
}
}

mpi_options = {
    "enabled": True,  # Required
    "processes_per_host": 8  # Required
}

smp_estimator = PyTorch(
    entry_point="your_training_script.py",  # Specify your train script
    role=sagemaker.get_execution_role(),
    instance_count=1,
    instance_type='ml.p3.16xlarge',
    framework_version='1.13.1',
    py_version='py3',
    distribution={
        "smdistributed": {"modelparallel": smp_options},
        "mpi": mpi_options
    },
    base_job_name="sharded-data-parallel-job"
)

smp_estimator.fit('s3://my_bucket/my_training_data/')

Reference configurations

The SageMaker distributed training team provides the following reference configurations that you can use as a starting point. You can extrapolate from the following configurations to experiment and estimate the GPU memory usage for your model configuration.

Sharded data parallelism with SMDDP Collectives

<table>
<thead>
<tr>
<th>Model/the number of parameters</th>
<th>Num instances</th>
<th>Instance type</th>
<th>Sequence length</th>
<th>Global batch size</th>
<th>Mini batch size</th>
<th>Sharded data parallel degree</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPT-NEOX-20B</td>
<td>2</td>
<td>ml.p4d.24 xlarge</td>
<td>2048</td>
<td>64</td>
<td>4</td>
<td>16</td>
</tr>
</tbody>
</table>
For example, if you increase the sequence length for a 20-billion-parameter model or increase the size of the model to 65 billion parameters, you need to try reducing the batch size first. If the model still doesn’t fit with the smallest batch size (the batch size of 1), try increasing the degree of model parallelism.

### Sharded data parallelism with tensor parallelism and NCCL Collectives

<table>
<thead>
<tr>
<th>Model/the number of parameters</th>
<th>Num instances</th>
<th>Instance type</th>
<th>Sequence length</th>
<th>Global batch size</th>
<th>Mini batch size</th>
<th>Sharded data parallel degree</th>
<th>Tensor parallel degree</th>
<th>Activation offloading</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPT-NEOX-20B</td>
<td>8</td>
<td>ml.p4d.24 xlarge</td>
<td>2048</td>
<td>768</td>
<td>12</td>
<td>32</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GPT-NEOX-65B</td>
<td>64</td>
<td>ml.p4d.24 xlarge</td>
<td>2048</td>
<td>512</td>
<td>8</td>
<td>16</td>
<td>8</td>
<td>Y</td>
</tr>
<tr>
<td>GPT-NEOX-65B</td>
<td>64</td>
<td>ml.p4d.24 xlarge</td>
<td>4096</td>
<td>512</td>
<td>2</td>
<td>64</td>
<td>2</td>
<td>Y</td>
</tr>
</tbody>
</table>

The combined usage of sharded data parallelism and tensor parallelism is useful when you want to fit a large language model (LLM) into a large-scale cluster while using text data with a longer sequence length, which leads to use a smaller batch size, and consequently handling the GPU memory usage to train LLMs against longer text sequences. To learn more, see the section called “Sharded data parallelism with tensor parallelism”.
For case studies, benchmarks, and more configuration examples, see the blog post *New performance improvements in Amazon SageMaker model parallel library.*

**Sharded data parallelism with SMDDP Collectives**

The SageMaker data parallelism library offers collective communication primitives (SMDDP collectives) optimized for the AWS infrastructure. It achieves optimization by adopting an all-to-all-type communication pattern by making use of *Elastic Fabric Adapter (EFA)*, resulting in high-throughput and less latency-sensitive collectives, offloading the communication-related processing to the CPU, and freeing up GPU cycles for computation. On large clusters, SMDDP Collectives can offer improvements in distributed training performance by up to 40% compared to NCCL. For case studies and benchmark results, see the blog *New performance improvements in the Amazon SageMaker model parallelism library.*

**Note**

Sharded data parallelism with SMDDP Collectives is available in the SageMaker model parallelism library v1.13.0 and later, and the SageMaker data parallelism library v1.6.0 and later. See also *Supported configurations* to use sharded data parallelism with SMDDP Collectives.

In sharded data parallelism, which is a commonly used technique in large-scale distributed training, the *AllGather* collective is used to reconstitute the sharded layer parameters for forward and backward pass computations, in parallel with GPU computation. For large models, performing the *AllGather* operation efficiently is critical to avoid GPU bottleneck problems and slowing down training speed. When sharded data parallelism is activated, SMDDP Collectives drops into these performance-critical *AllGather* collectives, improving training throughput.

**Train with SMDDP Collectives**

When your training job has sharded data parallelism activated and meets the *Supported configurations*, SMDDP Collectives are automatically activated. Internally, SMDDP Collectives optimize the *AllGather* collective to be performant on the AWS infrastructure and falls back to NCCL for all other collectives. Furthermore, under unsupported configurations, all collectives, including *AllGather*, automatically use the NCCL backend.

Since the SageMaker model parallelism library version 1.13.0, the "ddp_dist_backend" parameter is added to the *modelparallel options*. The default value for this configuration
parameter is "auto", which uses SMDDP Collectives whenever possible, and falls back to NCCL otherwise. To force the library to always use NCCL, specify "nccl" to the "ddp_dist_backend" configuration parameter.

The following code example shows how to set up a PyTorch estimator using the sharded data parallelism with the "ddp_dist_backend" parameter, which is set to "auto" by default and, therefore, optional to add.

```python
import sagemaker
from sagemaker.pytorch import PyTorch

smp_options = {
    "enabled": True,
    "parameters": {
        "partitions": 1,
        "ddp": True,
        "sharded_data_parallel_degree": 64
        "bf16": True,
        "ddp_dist_backend": "auto"  # Specify "nccl" to force to use NCCL.
    }
}

mpi_options = {
    "enabled": True,       # Required
    "processes_per_host": 8  # Required
}

smd_mp_estimator = PyTorch(
    entry_point="your_training_script.py",  # Specify your train script
    source_dir="location_to_your_script",
    role=sagemaker.get_execution_role(),
    instance_count=8,
    instance_type='ml.p4d.24xlarge',
    framework_version='1.13.1',
    py_version='py3',
    distribution={
        "smdistributed": {"modelparallel": smp_options},
        "mpi": mpi_options
    },
    base_job_name="sharded-data-parallel-demo",
)
smd_mp_estimator.fit('s3://my_bucket/my_training_data/')

**Supported configurations**

The `AllGather` operation with SMDDP Collectives are activated in training jobs when all the following configuration requirements are met.

- The sharded data parallelism degree greater than 1
- `Instance_count` greater than 1
- `Instance_type` equal to `ml.p4d.24xlarge`
- SageMaker training container for PyTorch v1.12.1 or later
- The SageMaker data parallelism library v1.6.0 or later
- The SageMaker model parallelism library v1.13.0 or later

**Performance and memory tuning**

SMDDP Collectives utilize additional GPU memory. There are two environment variables to configure the GPU memory usage depending on different model training use cases.

- `SMDDP_AG_SCRATCH_BUFFER_SIZE_BYTES` – During the SMDDP `AllGather` operation, the `AllGather` input buffer is copied into a temporary buffer for inter-node communication. The `SMDDP_AG_SCRATCH_BUFFER_SIZE_BYTES` variable controls the size (in bytes) of this temporary buffer. If the size of the temporary buffer is smaller than the `AllGather` input buffer size, the `AllGather` collective falls back to use NCCL.
  - Default value: 16 * 1024 * 1024 (16 MB)
  - Acceptable values: any multiple of 8192

- `SMDDP_AG_SORT_BUFFER_SIZE_BYTES` – The `SMDDP_AG_SORT_BUFFER_SIZE_BYTES` variable is to size the temporary buffer (in bytes) to hold data gathered from inter-node communication. If the size of this temporary buffer is smaller than $\frac{1}{8} \times \text{sharded_data_parallel_degree} \times \text{AllGather input size}$, the `AllGather` collective falls back to use NCCL.
  - Default value: 128 * 1024 * 1024 (128 MB)
  - Acceptable values: any multiple of 8192
Tuning guidance on the buffer size variables

The default values for the environment variables should work well for most use cases. We recommend tuning these variables only if training runs into the out-of-memory (OOM) error.

The following list discusses some tuning tips to reduce the GPU memory footprint of SMDDP Collectives while retaining the performance gain from them.

- Tuning SMDDP_AG_SCRATCH_BUFFER_SIZE_BYTES
  - The AllGather input buffer size is smaller for smaller models. Hence, the required size for SMDDP_AG_SCRATCH_BUFFER_SIZEBYTES can be smaller for models with fewer parameters.
  - The AllGather input buffer size decreases as sharded_data_parallel_degree increases, because the model gets sharded across more GPUs. Hence, the required size for SMDDP_AG_SCRATCH_BUFFER_SIZEBYTES can be smaller for training jobs with large values for sharded_data_parallel_degree.

- Tuning SMDDP_AG_SORT_BUFFER_SIZE_BYTES
  - The amount of data gathered from inter-node communication is less for models with fewer parameters. Hence, the required size for SMDDP_AG_SORT_BUFFER_SIZEBYTES can be smaller for such models with fewer number of parameters.

Some collectives might fall back to use NCCL; hence, you might not get the performance gain from the optimized SMDDP collectives. If additional GPU memory is available for use, you can consider increasing the values of SMDDP_AG_SCRATCH_BUFFER_SIZEBYTES and SMDDP_AG_SORT_BUFFER_SIZEBYTES to benefit from the performance gain.

The following code shows how you can configure the environment variables by appending them to mpi_options in the distribution parameter for the PyTorch estimator.

```python
import sagemaker
from sagemaker.pytorch import PyTorch

smp_options = {
    .... # All modelparallel configuration options go here
}

mpi_options = {
    "enabled" : True, # Required
    "processes_per_host" : 8 # Required
}
```
Mixed precision training with sharded data parallelism

To further save GPU memory with half-precision floating point numbers and sharded data parallelism, you can activate 16-bit floating point format (FP16) or Brain floating point format (BF16) by adding one additional parameter to the distributed training configuration.

Note
Mixed precision training with sharded data parallelism is available in the SageMaker model parallelism library v1.11.0 and later.

For FP16 Training with Sharded Data Parallelism

To run FP16 training with sharded data parallelism, add "fp16": True to the smp_options configuration dictionary. In your training script, you can choose between the static and dynamic loss scaling options through the smp.DistributedOptimizer module. For more information, see the section called “FP16 Training with Model Parallelism”.

SageMaker model parallelism library v2
smp_options = {
    "enabled": True,
    "parameters": {
        "ddp": True,
        "sharded_data_parallel_degree": 2,
        "fp16": True
    }
}

---

**For BF16 Training with Sharded Data Parallelism**

The sharded data parallelism feature of SageMaker supports training in BF16 data type. The BF16 data type uses 8 bits to represent the exponent of a floating point number, while the FP16 data type uses 5 bits. Preserving the 8 bits for the exponent allows to keep the same representation of the exponent of a 32-bit single precision floating point (FP32) number. This makes the conversion between FP32 and BF16 simpler and significantly less prone to cause overflow and underflow issues that arise often in FP16 training, especially when training larger models. While both data types use 16 bits in total, this increased representation range for the exponent in the BF16 format comes at the expense of reduced precision. For training large models, this reduced precision is often considered an acceptable trade-off for the range and training stability.

---

**Note**

Currently, BF16 training works only when sharded data parallelism is activated.

---

To run BF16 training with sharded data parallelism, add "bf16": True to the smp_options configuration dictionary.

smp_options = {
    "enabled": True,
    "parameters": {
        "ddp": True,
        "sharded_data_parallel_degree": 2,
        "bf16": True
    }
}
**Sharded data parallelism with tensor parallelism**

If you use sharded data parallelism and also need to reduce the global batch size, consider using **tensor parallelism** with sharded data parallelism. When training a large model with sharded data parallelism on a very large compute cluster (typically 128 nodes or beyond), even a small batch size per GPU results in a very large global batch size. It might lead to convergence issues or low computational performance issues. Reducing the batch size per GPU sometimes is not possible with sharded data parallelism alone when a single batch is already large and cannot be reduced further. In such cases, using sharded data parallelism in combination with tensor parallelism helps reduce the global batch size.

Choosing the optimal sharded data parallel and tensor parallel degrees depends on the scale of the model, the instance type, and the global batch size that is reasonable for the model to converge. We recommend that you start from a low tensor parallel degree to fit the global batch size into the compute cluster to resolve CUDA out-of-memory errors and achieve the best performance. See the following two example cases to learn how the combination of tensor parallelism and sharded data parallelism helps you adjust the global batch size by grouping GPUs for model parallelism, resulting in a lower number of model replicas and a smaller global batch size.

### Note

This feature is available from the SageMaker model parallelism library v1.15, and supports PyTorch v1.13.1.

### Note

This feature is available for the supported models by the tensor parallelism functionality of the library. To find the list of the supported models, see [Support for Hugging Face Transformer Models](https://github.com/aws/sagemaker-model-parallelism/blob/master/docs/models.md). Also note that you need to pass `tensor_parallelism=True` to the `smp.model_creation` argument while modifying your training script. To learn more, see the training script `train_gpt_simple.py` in the *SageMaker Examples GitHub repository*.

#### Example 1

Assume that we want to train a model over a cluster of 1536 GPUs (192 nodes with 8 GPUs in each), setting the degree of sharded data parallelism to 32...
(sharded_data_parallel_degree=32) and the batch size per GPU to 1, where each batch has a sequence length of 4096 tokens. In this case, there are 1536 model replicas, the global batch size becomes 1536, and each global batch contains about 6 million tokens.

Adding tensor parallelism to it can lower the global batch size. One configuration example can be setting the tensor parallel degree to 8 and the batch size per GPU to 4. This forms 192 tensor parallel groups or 192 model replicas, where each model replica is distributed across 8 GPUs. The batch size of 4 is the amount of training data per iteration and per tensor parallel group; that is, each model replica consumes 4 batches per iteration. In this case, the global batch size becomes 768, and each global batch contains about 3 million tokens. Hence, the global batch size is reduced by half compared to the previous case with sharded data parallelism only.

Example 2

When both sharded data parallelism and tensor parallelism are activated, the library first applies tensor parallelism and shards the model across this dimension. For each tensor parallel rank, the data parallelism is applied as per sharded_data_parallel_degree.

For example, assume that we want to set 32 GPUs with a tensor parallel degree of 4 (forming groups of 4 GPUs), a sharded data parallel degree of 4, ending up with a replication degree of 2. The assignment creates eight GPU groups based on the tensor parallel degree as follows: (0,1,2,3), (4,5,6,7), (8,9,10,11), (12,13,14,15), (16,17,18,19), (20,21,22,23), (24,25,26,27), (28,29,30,31). That is, four GPUs form one tensor parallel group. In this case, the reduced data parallel group for the 0th rank GPUs of the tensor parallel groups would be (0,4,8,12,16,20,24,28). The reduced data parallel group is sharded based on the sharded data parallel degree of 4, resulting in two replication groups for data parallelism. GPUs (0,4,8,12) form one sharding group, which collectively hold a complete copy of all parameters for the 0th tensor parallel rank, and GPUs (16,20,24,28) form another such group. Other tensor parallel ranks also have similar sharding and replication groups.
Figure 1: Tensor parallelism groups for (nodes, sharded data parallel degree, tensor parallel degree) = (4, 4, 4), where each rectangle represents a GPU with indices from 0 to 31. The GPUs form tensor parallelism groups from $TPG_0$ to $TPG_7$. Replication groups are ($\{TPG_0, TPG_4\}$, $\{TPG_1, TPG_5\}$, $\{TPG_2, TPG_6\}$ and $\{TPG_3, TPG_7\}$); each replication group pair shares the same color but filled differently.

Figure 2: Sharded data parallelism groups for (nodes, sharded data parallel degree, tensor parallel degree) = (4, 4, 4), where each rectangle represents a GPU with indices from 0 to 31. The GPUs form sharded data parallelism groups from $SDPG_0$ to $SDPG_7$. Replication groups are ($\{SDPG_0, SDPG_4\}$, $\{SDPG_1, SDPG_5\}$, $\{SDPG_2, SDPG_6\}$ and $\{SDPG_3, SDPG_7\}$); each replication group pair shares the same color but filled differently.

How to activate sharded data parallelism with tensor parallelism

To use sharded data parallelism with tensor parallelism, you need to set both `sharded_data_parallel_degree` and `tensor_parallel_degree` in the configuration for distribution while creating an object of the SageMaker PyTorch estimator class.
You also need to activate `prescaled_batch`. This means that, instead of each GPU reading its own batch of data, each tensor parallel group collectively reads a combined batch of the chosen batch size. Effectively, instead of dividing the dataset into parts equal to the number of GPUs (or data parallel size, `smp.dp_size()`), it divides into parts equal to the number of GPUs divided by `tensor_parallel_degree` (also called reduced data parallel size, `smp.rdp_size()`). For more details on prescaled batch, see Prescaled Batch in the SageMaker Python SDK documentation. See also the example training script `train_gpt_simple.py` for GPT-2 in the SageMaker Examples GitHub repository.

The following code snippet shows an example of creating a PyTorch estimator object based on the aforementioned scenario in the section called “Example 2”.

```python
mpi_options = "-verbose --mca orte_base_help_aggregate 0 "
smp_parameters = {
    "ddp": True,
    "fp16": True,
    "prescaled_batch": True,
    "sharded_data_parallel_degree": 4,
    "tensor_parallel_degree": 4
}

pytorch_estimator = PyTorch(
    entry_point="your_training_script.py",
    role=role,
    instance_type="ml.p4d.24xlarge",
    volume_size=200,
    instance_count=4,
    sagemaker_session=sagemaker_session,
    py_version="py3",
    framework_version="1.13.1",
    distribution={
        "smdistributed": {
            "modelparallel": {
                "enabled": True,
                "parameters": smp_parameters,
            }
        },
        "mpi": {
            "enabled": True,
            "processes_per_host": 8,
            "custom_mpi_options": mpi_options,
        },
    },
Tips and considerations for using sharded data parallelism

Consider the following when using the SageMaker model parallelism library's sharded data parallelism.

- Sharded data parallelism is compatible with FP16 training. To run FP16 training, see the **section called “FP16 Training with Model Parallelism”** section.
- Sharded data parallelism is compatible with tensor parallelism. The following items are what you might need to consider for using sharded data parallelism with tensor parallelism.
  - When using sharded data parallelism with tensor parallelism, the embedding layers are also automatically distributed across the tensor parallel group. In other words, the `distribute_embedding` parameter is automatically set to `True`. For more information about tensor parallelism, see the **section called “Tensor Parallelism”**.
  - Note that sharded data parallelism with tensor parallelism currently uses the NCCL collectives as the backend of the distributed training strategy.

To learn more, see the **section called “Sharded data parallelism with tensor parallelism”** section.

- Sharded data parallelism currently is not compatible with pipeline parallelism or optimizer state sharding. To activate sharded data parallelism, turn off optimizer state sharding and set the pipeline parallel degree to 1.
- The activation checkpointing and activation offloading features are compatible with sharded data parallelism.
- To use sharded data parallelism with gradient accumulation, set the `backward_passes_per_step` argument to the number of accumulation steps while wrapping your model with the `smdistributed.modelparallel.torch.DistributedModel` module. This ensures that the gradient AllReduce operation across the model replication groups (sharding groups) takes place at the boundary of gradient accumulation.
- You can checkpoint your models trained with sharded data parallelism using the library's checkpointing APIs, `smp.save_checkpoint` and `smp.resume_from_checkpoint`. For more information, see the **section called “Checkpointing a distributed PyTorch model (for the SageMaker model parallelism library v1.10.0 and later)”**.
• The behavior of the `delayed_parameter_initialization` configuration parameter changes under sharded data parallelism. When these two features are simultaneously turned on, parameters are immediately initialized upon model creation in a sharded manner instead of delaying the parameter initialization, so that each rank initializes and stores its own shard of parameters.

• When sharded data parallelism is activated, the library performs gradient clipping internally when the `optimizer.step()` call runs. You don't need to use utility APIs for gradient clipping, such as `torch.nn.utils.clip_grad_norm_()`. To adjust the threshold value for gradient clipping, you can set it through the `sdp_gradient_clipping` parameter for the distribution parameter configuration when you construct the SageMaker PyTorch estimator, as shown in the the section called “How to apply sharded data parallelism to your training job” section.

**Pipelining a Model**

One of the core features of SageMaker's model parallelism library is *pipeline parallelism*, which determines the order in which computations are made and data is processed across devices during model training. Pipelining is a technique to achieve true parallelization in model parallelism, by having the GPUs compute simultaneously on different data samples, and to overcome the performance loss due to sequential computation. When you use pipeline parallelism, training job is executed in a pipelined fashion over microbatches to maximize GPU usage.

**Note**

Pipeline parallelism, also called model partitioning, is available for both PyTorch and TensorFlow. For supported versions of the frameworks, see the section called “Supported Frameworks and AWS Regions”.

**Pipeline Execution Schedule**

Pipelining is based on splitting a mini-batch into microbatches, which are fed into the training pipeline one-by-one and follow an execution schedule defined by the library runtime. A *microbatch* is a smaller subset of a given training mini-batch. The pipeline schedule determines which microbatch is executed by which device for every time slot.

For example, depending on the pipeline schedule and the model partition, GPU $i$ might perform (forward or backward) computation on microbatch $b$ while GPU $i+1$ performs computation on
microbatch b+1, thereby keeping both GPUs active at the same time. During a single forward or backward pass, execution flow for a single microbatch might visit the same device multiple times, depending on the partitioning decision. For instance, an operation that is at the beginning of the model might be placed on the same device as an operation at the end of the model, while the operations in between are on different devices, which means this device is visited twice.

The library offers two different pipeline schedules, *simple* and *interleaved*, which can be configured using the `pipeline` parameter in the SageMaker Python SDK. In most cases, interleaved pipeline can achieve better performance by utilizing the GPUs more efficiently.

**Interleaved Pipeline**

In an interleaved pipeline, backward execution of the microbatches is prioritized whenever possible. This allows quicker release of the memory used for activations, using memory more efficiently. It also allows for scaling the number of microbatches higher, reducing the idle time of the GPUs. At steady-state, each device alternates between running forward and backward passes. This means that the backward pass of one microbatch may run before the forward pass of another microbatch finishes.

The preceding figure illustrates an example execution schedule for the interleaved pipeline over 2 GPUs. In the figure, F0 represents the forward pass for microbatch 0, and B1 represents the backward pass for microbatch 1. **Update** represents the optimizer update of the parameters. GPU0 always prioritizes backward passes whenever possible (for instance, executes B0 before F2), which allows for clearing of the memory used for activations earlier.

**Simple Pipeline**

A simple pipeline, by contrast, finishes running the forward pass for each microbatch before starting the backward pass. This means that it only pipelines the forward pass and backward pass stages within themselves. The following figure illustrates an example of how this works, over 2 GPUs.
Pipelining Execution in Specific Frameworks

Use the following sections to learn about the framework-specific pipeline scheduling decisions SageMaker's model parallelism library makes for TensorFlow and PyTorch.

Pipeline Execution with TensorFlow

The following image is an example of a TensorFlow graph partitioned by the model parallelism library, using automated model splitting. When a graph is split, each resulting subgraph is replicated B times (except for the variables), where B is the number of microbatches. In this figure, each subgraph is replicated 2 times (B=2). An SMPInput operation is inserted at each input of a subgraph, and an SMPOutput operation is inserted at each output. These operations communicate with the library backend to transfer tensors to and from each other.

The following image is an example of 2 subgraphs split with B=2 with gradient operations added. The gradient of a SMPInput op is a SMPOutput op, and vice versa. This enables the gradients to flow backwards during back-propagation.
This GIF demonstrates an example interleaved pipeline execution schedule with B=2 microbatches and 2 subgraphs. Each device sequentially executes one of the subgraph replicas to improve GPU utilization. As B grows larger, the fraction of idle time slots goes to zero. Whenever it is time to do (forward or backward) computation on a specific subgraph replica, the pipeline layer signals to the corresponding blue SMPInput operations to start executing.

Once the gradients from all microbatches in a single mini-batch are computed, the library combines the gradients across microbatches, which can then be applied to the parameters.

**Pipeline Execution with PyTorch**

Conceptually, pipelining follows a similar idea in PyTorch. However, since PyTorch does not involve static graphs and so the model parallelism library's PyTorch feature uses a more dynamic pipelining paradigm.

As in TensorFlow, each batch is split into a number of microbatches, which are executed one at a time on each device. However, the execution schedule is handled via execution servers launched on each device. Whenever the output of a submodule that is placed on another device is needed on the current device, an execution request is sent to the execution server of the remote device along with the input tensors to the submodule. The server then executes this module with the given inputs and returns the response to the current device.

Since the current device is idle during the remote submodule execution, the local execution for the current microbatch pauses, and the library runtime switches execution to another microbatch.
which the current device can actively work on. The prioritization of microbatches is determined by the chosen pipeline schedule. For an interleaved pipeline schedule, microbatches that are in the backward stage of the computation are prioritized whenever possible.

Tensor Parallelism

Tensor parallelism is a type of model parallelism in which specific model weights, gradients, and optimizer states are split across devices. In contrast to pipeline parallelism, which keeps individual weights intact but partitions the set of weights, tensor parallelism splits individual weights. This typically involves distributed computation of specific operations, modules, or layers of the model.

Tensor parallelism is required in cases in which a single parameter consumes most of the GPU memory (such as large embedding tables with a large vocabulary size or a large softmax layer with a large number of classes). In this case, treating this large tensor or operation as an atomic unit is inefficient and impedes balance of the memory load.

Tensor parallelism is also useful for extremely large models in which a pure pipelining is simply not enough. For example, with GPT-3-scale models that require partitioning over tens of instances, a pure microbatch pipelining is inefficient because the pipeline depth becomes too high and the overhead becomes prohibitively large.

Note

Tensor parallelism is available for PyTorch in the SageMaker model parallelism library v1.6.0 and later.

Topics

- How Tensor Parallelism Works
- Run a SageMaker Distributed Model Parallel Training Job with Tensor Parallelism
- Support for Hugging Face Transformer Models
- Ranking Mechanism when Using a Combination of Pipeline Parallelism and Tensor Parallelism

How Tensor Parallelism Works

Tensor parallelism takes place at the level of nn.Modules; it partitions specific modules in the model across tensor parallel ranks. This is in addition to the existing partition of the set of modules used in pipeline parallelism.
When a module is partitioned through tensor parallelism, its forward and backward propagation are distributed. The library handles the necessary communication across devices to implement the distributed execution of these modules. The modules are partitioned across multiple data parallel ranks. Contrary to the traditional distribution of workloads, each data parallel rank does **not** have the complete model replica when the library’s tensor parallelism is used. Instead, each data parallel rank may have only a partition of the distributed modules, in addition to the entirety of the modules that are not distributed.

**Example:** Consider tensor parallelism across data parallel ranks, where the degree of data parallelism is 4 and the degree of tensor parallelism is 2. Assume that you have a data parallel group that holds the following module tree, after partitioning the set of modules.

```
A
### B
|   ### E
|   ### F
### C
### D
    ### G
    ### H
```

Assume that tensor parallelism is supported for the modules B, G, and H. One possible outcome of tensor parallel partition of this model could be:

```
dp_rank 0 (tensor parallel rank 0): A, B:0, C, D, G:0, H
dp_rank 1 (tensor parallel rank 1): A, B:1, C, D, G:1, H
dp_rank 2 (tensor parallel rank 0): A, B:0, C, D, G:0, H
dp_rank 3 (tensor parallel rank 1): A, B:1, C, D, G:1, H
```

Each line represents the set of modules stored in that dp_rank, and the notation X:y represents the yth fraction of the module X. Note the following:

1. Partitioning takes place across subsets of data parallel ranks, which we call TP_GROUP, not the entire DP_GROUP, so that the exact model partition is replicated across dp_rank 0 and dp_rank 2, and similarly across dp_rank 1 and dp_rank 3.
2. The modules E and F are no longer part of the model, since their parent module B is partitioned, and any execution that is normally a part of E and F takes place within the (partitioned) B module.
3. Even though H is supported for tensor parallelism, in this example it is not partitioned, which highlights that whether to partition a module depends on user input. The fact that a module is supported for tensor parallelism does not necessarily mean it is partitioned.

**How the library adapts tensor parallelism to PyTorch `nn.Linear` module**

When tensor parallelism is performed over data parallel ranks, a subset of the parameters, gradients, and optimizer states are partitioned across the tensor parallel devices *for the modules that are partitioned*. For the rest of the modules, the tensor parallel devices operate in a regular data parallel manner. To execute the partitioned module, a device first collects the necessary parts of *all data samples* across peer devices in the same tensor parallelism group. The device then runs the local fraction of the module on all these data samples, followed by another round of synchronization which both combines the parts of the output for each data sample and returns the combined data samples to the GPUs from which the data sample first originated. The following figure shows an example of this process over a partitioned `nn.Linear` module.
The first figure shows a small model with a large `nn.Linear` module with data parallelism over the two tensor parallelism ranks. The `nn.Linear` module is replicated into the two parallel ranks.

The second figure shows tensor parallelism applied on a larger model while splitting the `nn.Linear` module. Each `tp_rank` holds half the linear module, and the entirety of the rest of the operations. While the linear module runs, each `tp_rank` collects the relevant half of all data samples and passes it through their half of the `nn.Linear` module. The result needs to be reduce-scattered (with summation as the reduction operation) so that each rank has the final linear output for their own data samples. The rest of the model runs in the typical data parallel manner.

### Run a SageMaker Distributed Model Parallel Training Job with Tensor Parallelism

In this section, you learn:

- How to configure a SageMaker PyTorch estimator and the SageMaker model parallelism option to use tensor parallelism.
- How to adapt your training script using the extended `smdistributed.modelparallel` modules for tensor parallelism.

To learn more about the `smdistributed.modelparallel` modules, see the [SageMaker model parallel APIs](https://docs.aws.amazon.com/sagemaker/latest/dg/model-parallel-api-ref.html) in the *SageMaker Python SDK documentation*.

### Topics

- **Tensor parallelism alone**
- **Tensor parallelism combined with pipeline parallelism**

### Tensor parallelism alone

The following is an example of a distributed training option to activate tensor parallelism alone, without pipeline parallelism. Configure the `mpi_options` and `smp_options` dictionaries to specify distributed training options to the SageMaker PyTorch estimator.

---

**Note**

Extended memory-saving features are available through Deep Learning Containers for PyTorch, which implements the SageMaker model parallelism library v1.6.0 or later.
Configure a SageMaker PyTorch estimator

```python
mpi_options = {
    "enabled" : True,
    "processes_per_host" : 8, # 8 processes
    "custom_mpi_options" : "--mca btl_vader_single_copy_mechanism none"
}

smp_options = {
    "enabled":True,
    "parameters": {
        "pipeline_parallel_degree": 1, # alias for "partitions"
        "placement_strategy": "cluster",
        "tensor_parallel_degree": 4, # tp over 4 devices
        "ddp": True
    }
}

smp_estimator = PyTorch(
    entry_point='your_training_script.py', # Specify
    role=role,
    instance_type='ml.p3.16xlarge',
    sagemaker_session=sagemaker_session,
    framework_version='1.13.1',
    py_version='py36',
    instance_count=1,
    distribution={
        "smdistributed": {
            "modelparallel": smp_options,
        },
        "mpi": mpi_options
    },
    base_job_name="SMD-MP-demo",
)

smp_estimator.fit('s3://my_bucket/my_training_data/')
```

**Tip**

To find a complete list of parameters for distribution, see [Configuration Parameters for Model Parallelism](#) in the SageMaker Python SDK documentation.

Adapt your PyTorch training script

---

SageMaker model parallelism library v2 3726
The following example training script shows how to adapt the SageMaker model parallelism library to a training script. In this example, it is assumed that the script is named `your_training_script.py`.

```python
import torch
import torch.nn as nn
import torch.nn.functional as F
import torch.optim as optim
from torchvision import datasets
import smdistributed.modelparallel.torch as smp

class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
        self.conv1 = nn.Conv2d(1, 32, 3, 1)
        self.conv2 = nn.Conv2d(32, 64, 3, 1)
        self.fc1 = nn.Linear(9216, 128)
        self.fc2 = nn.Linear(128, 10)

    def forward(self, x):
        x = self.conv1(x)
        x = F.relu(x)
        x = self.conv2(x)
        x = F.relu(x)
        x = F.max_pool2d(x, 2)
        x = torch.flatten(x, 1)
        x = self.fc1(x)
        x = F.relu(x)
        x = self.fc2(x)
        return F.log_softmax(x, 1)

def train(model, device, train_loader, optimizer):
    model.train()
    for batch_idx, (data, target) in enumerate(train_loader):
        # smdistributed: Move input tensors to the GPU ID used by
        # the current process, based on the set_device call.
        data, target = data.to(device), target.to(device)
        optimizer.zero_grad()
        output = model(data)
        loss = F.nll_loss(output, target, reduction="mean")
        loss.backward()
```
optimizer.step()

# smdistributed: Initialize the backend
smp.init()

# smdistributed: Set the device to the GPU ID used by the current process.
# Input tensors should be transferred to this device.
torch.cuda.set_device(smp.local_rank())
device = torch.device("cuda")

# smdistributed: Download only on a single process per instance.
# When this is not present, the file is corrupted by multiple processes trying
# to download and extract at the same time
if smp.local_rank() == 0:
    dataset = datasets.MNIST("../data", train=True, download=False)
smp.barrier()

# smdistributed: Shard the dataset based on data parallel ranks
if smp.dp_size() > 1:
    partitions_dict = {f"{i}": 1 / smp.dp_size() for i in range(smp.dp_size())}
    dataset = SplitDataset(dataset, partitions=partitions_dict)
    dataset.select(f"{smp.dp_rank()}")
train_loader = torch.utils.data.DataLoader(dataset, batch_size=64)

# smdistributed: Enable tensor parallelism for all supported modules in the model
# i.e., nn.Linear in this case. Alternatively, we can use
# smp.set_tensor_parallelism(model.fc1, True)
# to enable it only for model.fc1
with smp.tensor_parallelism():
    model = Net()

# smdistributed: Use the DistributedModel wrapper to distribute the
# modules for which tensor parallelism is enabled
model = smp.DistributedModel(model)

optimizer = optim.AdaDelta(model.parameters(), lr=4.0)
optimizer = smp.DistributedOptimizer(optimizer)

train(model, device, train_loader, optimizer)
Tensor parallelism combined with pipeline parallelism

The following is an example of a distributed training option that enables tensor parallelism combined with pipeline parallelism. Set up the `mpi_options` and `smp_options` parameters to specify model parallel options with tensor parallelism when you configure a SageMaker PyTorch estimator.

```python
mpi_options = {
    "enabled": True,
    "processes_per_host": 8, # 8 processes
    "custom_mpi_options": "--mca btl_vader_single_copy_mechanism none "
}

smp_options = {
    "enabled": True,
    "parameters": {
        "microbatches": 4,
        "pipeline_parallel_degree": 2, # alias for "partitions"
        "placement_strategy": "cluster",
        "tensor_parallel_degree": 2, # tp over 2 devices
        "ddp": True
    }
}

smp_estimator = PyTorch(  
    entry_point='your_training_script.py', # Specify  
    role=role,  
    instance_type='ml.p3.16xlarge',  
    sagemaker_session=sagemaker_session,  
    framework_version='1.13.1',  
    py_version='py36',  
    instance_count=1,  
    distribution={  
        "smdistributed": {"modelparallel": smp_options},
    }
)
```

Note

Extended memory-saving features are available through Deep Learning Containers for PyTorch, which implements the SageMaker model parallelism library v1.6.0 or later.
Adapt your PyTorch training script

The following example training script shows how to adapt the SageMaker model parallelism library to a training script. Note that the training script now includes the `smp.step` decorator:

```python
import torch
import torch.nn as nn
import torch.nn.functional as F
import torch.optim as optim
from torchnet.dataset import SplitDataset
from torchvision import datasets

import smdistributed.modelparallel.torch as smp

class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
        self.conv1 = nn.Conv2d(1, 32, 3, 1)
        self.conv2 = nn.Conv2d(32, 64, 3, 1)
        self.fc1 = nn.Linear(9216, 128)
        self.fc2 = nn.Linear(128, 10)

    def forward(self, x):
        x = self.conv1(x)
        x = F.relu(x)
        x = self.conv2(x)
        x = F.relu(x)
        x = F.max_pool2d(x, 2)
        x = torch.flatten(x, 1)
        x = self.fc1(x)
        x = F.relu(x)
        x = self.fc2(x)
        return F.log_softmax(x, 1)

    # smdistributed: Define smp.step. Return any tensors needed outside.
```

smp Estimator.fit('s3://my_bucket/my_training_data/')
```python
@smp.step
def train_step(model, data, target):
    output = model(data)
    loss = F.nll_loss(output, target, reduction="mean")
    model.backward(loss)
    return output, loss

def train(model, device, train_loader, optimizer):
    model.train()
    for batch_idx, (data, target) in enumerate(train_loader):
        # smdistributed: Move input tensors to the GPU ID used by
        # the current process, based on the set_device call.
        data, target = data.to(device), target.to(device)
        optimizer.zero_grad()
        # Return value, loss_mb is a StepOutput object
        _, loss_mb = train_step(model, data, target)

        # smdistributed: Average the loss across microbatches.
        loss = loss_mb.reduce_mean()

        optimizer.step()

    # smdistributed: Initialize the backend
    smp.init()

    # smdistributed: Set the device to the GPU ID used by the current process.
    # Input tensors should be transferred to this device.
    torch.cuda.set_device(smp.local_rank())
    device = torch.device("cuda")

    # smdistributed: Download only on a single process per instance.
    # When this is not present, the file is corrupted by multiple processes trying
    # to download and extract at the same time
    if smp.local_rank() == 0:
        dataset = datasets.MNIST("../data", train=True, download=False)
        smp.barrier()

    # smdistributed: Shard the dataset based on data parallel ranks
    if smp.dp_size() > 1:
        partitions_dict = {f"{i}" : 1 / smp.dp_size() for i in range(smp.dp_size())}
        dataset = SplitDataset(dataset, partitions=partitions_dict)
        dataset.select(f"{smp.dp_rank()}")

    # smdistributed: Set drop_last=True to ensure that batch size is always divisible
```
# by the number of microbatches
train_loader = torch.utils.data.DataLoader(dataset, batch_size=64, drop_last=True)

model = Net()

# smdistributed: enable tensor parallelism only for model.fc1
smp.set_tensor_parallelism(model.fc1, True)

# smdistributed: Use the DistributedModel container to provide the model
# to be partitioned across different ranks. For the rest of the script,
# the returned DistributedModel object should be used in place of
# the model provided for DistributedModel class instantiation.
model = smp.DistributedModel(model)

optimizer = optim.AdaDelta(model.parameters(), lr=4.0)
optimizer = smp.DistributedOptimizer(optimizer)

train(model, device, train_loader, optimizer)

---

**Support for Hugging Face Transformer Models**

The SageMaker model parallelism library's tensor parallelism offers out-of-the-box support for the following Hugging Face Transformer models:

- GPT-2, BERT, and RoBERTa (Available in the SageMaker model parallelism library v1.7.0 and later)
- GPT-J (Available in the SageMaker model parallelism library v1.8.0 and later)
- GPT-Neo (Available in the SageMaker model parallelism library v1.10.0 and later)

---

**Note**

For any other Transformers models, you need to use the `smdistributed.modelparallel.torch.tp_register_with_module()` API to apply tensor parallelism.

**Note**

To use tensor parallelism for training Hugging Face Transformer models, make sure you use Hugging Face Deep Learning Containers for PyTorch that has the SageMaker model parallelism library v2.
Supported Models Out of the Box

For the Hugging Face transformer models supported by the library out of the box, you don't need to manually implement hooks to translate Transformer APIs to smdistributed transformer layers. You can activate tensor parallelism by using the context manager `smdistributed.modelparallel.torch.tensor_parallelism()` and wrapping the model by `smdistributed.modelparallel.torch.DistributedModel()`. You don't need to manually register hooks for tensor parallelism using the smp.tp_register API.

The state_dict translation functions between Hugging Face Transformers and smdistributed.modelparallel can be accessed as follows.

- `smdistributed.modelparallel.torch.nn.huggingface.gpt2.translate_state_dict_to_hf_gpt2(state_dict, max_seq_len=None)`
- `smdistributed.modelparallel.torch.nn.huggingface.gpt2.translate_hf_state_dict_to_smdistributed_gpt2(state_dict)`
- `smdistributed.modelparallel.torch.nn.huggingface.bert.translate_state_dict_to_hf_bert(state_dict, max_seq_len=None)`
- `smdistributed.modelparallel.torch.nn.huggingface.bert.translate_hf_state_dict_to_smdistributed_bert(state_dict)`
- `smdistributed.modelparallel.torch.nn.huggingface.roberta.translate_state_dict_to_hf_roberta(state_dict, max_seq_len=None)`
- `smdistributed.modelparallel.torch.nn.huggingface.roberta.translate_hf_state_dict_to_smdistributed_roberta(state_dict)`
- `smdistributed.modelparallel.torch.nn.huggingface.gptj.translate_state_dict_to_hf_gptj(state_dict, max_seq_len=None)` (Available in the SageMaker model parallelism library v1.8.0 and later)
- `smdistributed.modelparallel.torch.nn.huggingface.gptj.translate_hf_gptj_state_dict_to_smdistributed_gptj` (Available in the SageMaker model parallelism library v1.8.0 and later)
- `smdistributed.modelparallel.torch.nn.huggingface.gptneo.translate_state_dict_to_hf_gptneo(state_dict, max_seq_len=None)` (Available in the SageMaker model parallelism library v1.10.0 and later)
- `smdistributed.modelparallel.torch.nn.huggingface.gptneo.translate_hf_state_dict_to_smdistributed_gptneo(state_dict)` (Available in the SageMaker model parallelism library v1.10.0 and later)

Example usage of the GPT-2 translation function
Start with wrapping the model as shown in the following code.

```python
from transformers import AutoModelForCausalLM

with smp.tensor_parallelism():
    model = AutoModelForCausalLM.from_config(hf_gpt2_config)

model = smp.DistributedModel(model)
```

Given a `state_dict` from the `DistributedModel` object, you can load the weights into the original Hugging Face GPT-2 model using the `translate_state_dict_to_hf_gpt2` function as shown in the following code.

```python
from smdistributed.modelparallel.torch.nn.huggingface.gpt2 import translate_state_dict_to_hf_gpt2
max_seq_len = 1024

# [... code block for training ...]

if smp.rdp_rank() == 0:
    state_dict = dist_model.state_dict()
    hf_state_dict = translate_state_dict_to_hf_gpt2(state_dict, max_seq_len)

    # can now call model.load_state_dict(hf_state_dict) to the original HF model
```

**Example usage of the RoBERTa translation function**

Similarly, given a supported HuggingFace model `state_dict`, you can use the `translate_hf_state_dict_to_smdistributed` function to convert it to a format readable by `smp.DistributedModel`. This can be useful in transfer learning use cases, where a pre-trained model is loaded into a `smp.DistributedModel` for model-parallel fine-tuning:

```python
from smdistributed.modelparallel.torch.nn.huggingface.roberta import translate_state_dict_to_smdistributed

model = AutoModelForMaskedLM.from_config(roberta_config)
model = smp.DistributedModel(model)
pretrained_model = AutoModelForMaskedLM.from_pretrained("roberta-large")
translated_state_dict = translate_state_dict_to_smdistributed(pretrained_model.state_dict())
```
# load the translated pretrained weights into the smp.DistributedModel
model.load_state_dict(translated_state_dict)

# start fine-tuning...

## Ranking Mechanism when Using a Combination of Pipeline Parallelism and Tensor Parallelism

This section explains how the ranking mechanism of model parallelism works with tensor parallelism. This is extended from the [Ranking Basics](#) for [Core Features of the SageMaker Model Parallelism Library](#). With tensor parallelism, the library introduces three types of ranking and process group APIs: `smp.tp_rank()` for tensor parallel rank, `smp.pp_rank()` for pipeline parallel rank, and `smp.rdp_rank()` for reduced-data parallel rank. The corresponding communication process groups are tensor parallel group (TP_GROUP), pipeline parallel group (PP_GROUP), and reduced-data parallel group (RDP_GROUP). These groups are defined as follows:

- A **tensor parallel group** (TP_GROUP) is an evenly divisible subset of the data parallel group, over which tensor parallel distribution of modules takes place. When the degree of pipeline parallelism is 1, TP_GROUP is the same as **model parallel group** (MP_GROUP).

- A **pipeline parallel group** (PP_GROUP) is the group of processes over which pipeline parallelism takes place. When the tensor parallelism degree is 1, PP_GROUP is the same as MP_GROUP.

- A **reduced-data parallel group** (RDP_GROUP) is a set of processes that hold both the same pipeline parallelism partitions and the same tensor parallel partitions, and perform data parallelism among themselves. This is called the reduced data parallel group because it is a subset of the entire data parallelism group, DP_GROUP. For the model parameters that are distributed within the TP_GROUP, the gradient allreduce operation is performed only for reduced-data parallel group, while for the parameters that are not distributed, the gradient allreduce takes place over the entire DP_GROUP.

- A model parallel group (MP_GROUP) refers to a group of processes that collectively store the entire model. It consists of the union of the PP_GROUPs of all the ranks that are in the TP_GROUP of the current process. When the degree of tensor parallelism is 1, MP_GROUP is equivalent to PP_GROUP. It is also consistent with the existing definition of MP_GROUP from previous smdistributed releases. Note that the current TP_GROUP is a subset of both the current DP_GROUP and the current MP_GROUP.

To learn more about the communication process APIs in the SageMaker model parallelism library, see the [Common API](#) and the [PyTorch-specific APIs](#) in the [SageMaker Python SDK documentation](#).
This figure shows ranking mechanism, parameter distribution, and associated AllReduce operations of tensor parallelism.

For example, consider process groups for a single node with 8 GPUs, where the degree of tensor parallelism is 2, the degree of pipeline parallelism is 2, and the degree of data parallelism is 4. The upper center part of the preceding figure shows an example of a model with 4 layers. The lower left and lower right parts of figure illustrate the 4-layer model distributed across 4 GPUs using both pipeline parallelism and tensor parallelism, where tensor parallelism is used for the middle two layers. These two lower figures are simple copies to illustrate different group boundary lines. The partitioned model is replicated for data parallelism across GPUs 0-3 and 4-7. The lower left figure shows the definitions of MP_GROUP, PP_GROUP, and TP_GROUP. The lower right figure shows RDP_GROUP, DP_GROUP, and WORLD over the same set of GPUs. The gradients for the layers and layer slices that have the same color are allreduced together for data parallelism. For example, the first layer (light blue) gets the allreduce operations across DP_GROUP, whereas the dark...
orange slice in the second layer only gets the `allreduce` operations within the RDP_GROUP of its process. The bold dark red arrows represent tensors with the batch of its entire TP_GROUP.

In this example, pipeline parallelism occurs across the GPU pairs (0,1); (2,3); (4,5) and (6,7). In addition, data parallelism (`allreduce`) takes place across GPUs 0, 2, 4, 6, and independently over GPUs 1, 3, 5, 7. Tensor parallelism happens over subsets of DP_GROUPS, across the GPU pairs (0,2); (1,3); (4,6) and (5,7).

**Optimizer State Sharding**

*Optimizer state sharding* is a useful memory-saving technique that shards the optimizer state (the set of weights that describes the state of optimizer) across data parallel device groups. You can use optimizer state sharding whenever you use a stateful optimizer (such as Adam) or an FP16 optimizer (which stores both FP16 and FP32 copies of the parameters).

**Note**

Optimizer state sharding is available for PyTorch in the SageMaker model parallelism library v1.6.0 and later.

**How to Use Optimizer State Sharding**

You can turn on optimizer state sharding by setting "shard_optimizer_state": True in the modelparallel configuration.

When this feature is turned on, the library partitions the set of model parameters based on the data parallelism degree. The gradients corresponding to the i\text{th} partition get reduced only at the i\text{th} data parallel rank. At the end of the first call to an smp.step decorator function, the optimizer wrapped by smp.DistributedOptimizer redefines its parameters to be only
limited to those parameters corresponding to the partition of the current data parallel rank. The redefined parameters are called *virtual parameters* and share underlying storage with the original parameters. During the first call to `optimizer.step`, the optimizer states are created based on these redefined parameters, which are sharded because of the original partition. After the optimizer update, the AllGather operation (as part of the `optimizer.step` call) runs across the data parallel ranks to achieve consistent parameter states.

**Tip**

Optimizer state sharding can be useful when the degree of data parallelism is greater than 1 and the model has more than a billion parameters. The degree of data parallelism is calculated by `(processes_per_host * instance_count / pipeline_parallel_degree)`, and the `smp.dp_size()` function handles the sizing in the background.

Configure a SageMaker PyTorch estimator

```python
mpi_options = {
    "enabled": True,
    "processes_per_host": 8,  # 8 processes
    "custom_mpi_options": "--mca btl_vader_single_copy_mechanism none"
}

smp_options = {
    "enabled": True,
    "parameters": {
        "microbatches": 4,
        "pipeline_parallel_degree": 2,  # alias for "partitions"
        "placement_strategy": "cluster",
        "tensor_parallel_degree": 2,  # tp over 2 devices
        "ddp": True,
        "shard_optimizer_state": True
    }
}
```

Adapt your PyTorch training script

See Adapt your PyTorch training script in the Tensor parallelism combined with pipeline parallelism section. There's no additional modification required for the script.
Activation Checkpointing

*Activation checkpointing (or gradient checkpointing)* is a technique to reduce memory usage by clearing activations of certain layers and recomputing them during a backward pass. Effectively, this trades extra computation time for reduced memory usage. If a module is checkpointed, at the end of a forward pass, the inputs to and outputs from the module stay in memory. Any intermediate tensors that would have been part of the computation inside that module are freed up during the forward pass. During the backward pass of checkpointed modules, these tensors are recomputed. At this point, the layers beyond this checkpointed module have finished their backward pass, so the peak memory usage with checkpointing can be lower.

**Note**

This feature is available for PyTorch in the SageMaker model parallelism library v1.6.0 and later.

How to Use Activation Checkpointing

With `smdistributed.modelparallel`, you can use activation checkpointing at the granularity of a module. For all `torch.nn` modules except `torch.nn.Sequential`, you can only checkpoint a module tree if it lies within one partition from the perspective of pipeline parallelism. In case of the `torch.nn.Sequential` module, each module tree inside the sequential module must lie completely within one partition for activation checkpointing to work. When you use manual partitioning, be aware of these restrictions.

When you use [automated model partitioning](https://docs.aws.amazon.com/sagemaker/latest/dg/model-parallelism.html), you can find the partitioning assignment logs starting with `Partition assignments:` in the training job logs. If a module is partitioned across multiple ranks (for example, with one descendant on one rank and another descendant on a different rank), the library ignores the attempt to checkpoint the module and raises a warning message that the module won't be checkpointed.

**Note**

The SageMaker model parallelism library supports both overlapping and non-overlapping allreduce operation in combination with checkpointing.
**Note**

PyTorch’s native checkpointing API is not compatible with `smdistributed.modelparallel`.

**Example 1:** The following sample code shows how to use activation checkpointing when you have a model definition in your script.

```python
import torch.nn as nn
import torch.nn.functional as F
from smdistributed.modelparallel.torch.patches.checkpoint import checkpoint

class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
        self.conv1 = nn.Conv2d(1, 32, 3, 1)
        self.conv2 = nn.Conv2d(32, 64, 3, 1)
        self.fc1 = nn.Linear(9216, 128)
        self.fc2 = nn.Linear(128, 10)

    def forward(self, x):
        x = self.conv1(x)
        x = self.conv2(x)
        x = F.max_pool2d(x, 2)
        x = torch.flatten(x, 1)
        # This call of fc1 will be checkpointed
        x = checkpoint(self.fc1, x)
        x = self.fc2(x)
        return F.log_softmax(x, 1)
```

**Example 2:** The following sample code shows how to use activation checkpointing when you have a sequential model in your script.

```python
import torch.nn as nn
from smdistributed.modelparallel.torch.patches.checkpoint import checkpoint_sequential

class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
```
self.seq = nn.Sequential(
    nn.Conv2d(1,20,5),
    nn.ReLU(),
    nn.Conv2d(20,64,5),
    nn.ReLU()
)

def forward(self, x):
    # This call of self.seq will be checkpointed
    x = checkpoint_sequential(self.seq, x)
    return F.log_softmax(x, 1)

Example 3: The following sample code shows how to use activation checkpointing when you import a prebuilt model from a library, such as PyTorch and Hugging Face Transformers. Whether you checkpoint sequential modules or not, do the following:

1. Wrap the model by `smp.DistributedModel()`.
2. Define an object for sequential layers.
3. Wrap the sequential layer object by `smp.set_activation_checkpointing()`.

```python
import smdistributed.modelparallel.torch as smp
from transformers import AutoModelForCausalLM

smp.init()
model = AutoModelForCausalLM(*args, **kwargs)
model = smp.DistributedModel(model)

# Call set_activation_checkpointing API
transformer_layers = model.module.module.module.transformer.seq_layers
smp.set_activation_checkpointing(
    transformer_layers, pack_args_as_tuple=True, strategy='each')
```

Activation Offloading

When activation checkpointing and pipeline parallelism are turned on and the number of microbatches is greater than one, *activation offloading* is an additional feature that can further reduce memory usage. Activation offloading asynchronously moves the checkpointed activations corresponding to their microbatches that are not currently running in the CPU. Right before the GPU needs the activations for the microbatch’s backward pass, this functionality prefetches the offloaded activations back from the CPU.
How to Use Activation Offloading

Use activation offloading to reduce memory usage when the number of microbatches is greater than 1, and activation checkpointing is turned on (see Activation Checkpointing). When the activation checkpointing is not used, activation offloading has no effect. When it is used with only one microbatch, it does not save memory.

To use activation offloading, set "offload_activations": True in the modelparallel configuration.

Activation offloading moves the checkpointed activations in nn.Sequential modules to CPU asynchronously. The data transfer over the PCIe link overlaps with GPU computation. The offloading happens immediately, as soon as the forward pass for a particular checkpointed layer is computed. The activations are loaded back to the GPU shortly before they are needed for the backward pass of a particular microbatch. The CPU-GPU transfer similarly overlaps with computation.

To adjust how early the activations are loaded back into the GPU, you can use the configuration parameter "activation_loading_horizon" (default is set to 4, must be int larger than 0). A larger activation loading horizon would cause the activations to be loaded back to the GPU earlier. If the horizon is too large, the memory-saving impact of activation offloading might be diminished. If the horizon is too small, the activations may not be loaded back in time, reducing the amount of overlap and degrading performance.

Tip

Activation offloading can be useful for large models with over a hundred billion parameters.

Configure a SageMaker PyTorch estimator

```python
mpi_options = {
```
FP16 Training with Model Parallelism

For FP16 training, apply the following modifications to your training script and estimator.

Note

This feature is available for PyTorch in the SageMaker model parallelism library v1.10.0 and later.

Adapt your PyTorch training script

1. Wrap your model using the `smdistributed.modelparallel.torch.model_creation()` context manager.

```python
# fp16_training_script.py

import torch
import smdistributed.modelparallel.torch as smp

with smp.model_creation(
    dtype=torch.float16 if args.fp16 else torch.get_default_dtype()
):
    model = ...
```
Tip

If you are using tensor parallelism, add `tensor_parallelism=smp.tp_size() > 1` to the `smp.model_creation` context manager. Adding this line also helps automatically detect whether tensor parallelism is activated or not.

```python
with smp.model_creation(
    ...
    tensor_parallelism=smp.tp_size() > 1
):
    model = ...
```

2. When you wrap the optimizer with `smdistributed.modelparallel.torch.DistributedOptimizer`, set either the `static_loss_scaling` or `dynamic_loss_scaling` argument. By default, `static_loss_scaling` is set to 1.0, and `dynamic_loss_scaling` is set to False. If you set `dynamic_loss_scale=True`, you can feed dynamic loss scaling options as a dictionary through the `dynamic_loss_args` argument. In most cases, we recommend you use dynamic loss scaling with the default options. For more information, options, and examples of the optimizer wrapper function, see the `smdistributed.modelparallel.torch.DistributedOptimizer API`.

The following code is an example of wrapping an Adadelta optimizer object with dynamic loss scaling for FP16 training.

```python
optimizer = torch.optim.Adadelta(...)  
optimizer = smp.DistributedOptimizer(
    optimizer,
    static_loss_scale=None,
    dynamic_loss_scale=True,
    dynamic_loss_args={
        "scale_window": 1000,
        "min_scale": 1,
        "delayed_shift": 2
    }
)
```

Configure a SageMaker PyTorch estimator
Add the FP16 parameter ("fp16") to the distribution configuration for model parallelism when creating a SageMaker PyTorch estimator object. For a complete list of the configuration parameters for model parallelism, see Parameters for smdistributed.

```python
from sagemaker.pytorch import PyTorch

smp_options = {
    "enabled": True,
    "parameters": {
        "microbatches": 4,
        "pipeline_parallel_degree": 2,
        "tensor_parallel_degree": 2,
        ...
        "fp16": True
    }
}

fp16_estimator = PyTorch(
    entry_point="fp16_training_script.py", # Specify your train script
    ...
    distribution={
        "smdistributed": {"modelparallel": smp_options},
        "mpi": {...}
    }
)

fp16_estimator.fit(...)
```

When FP16 training starts, the model and the optimizer are wrapped by FP16_Module and FP16_Optimizer respectively, which are modified smdistributed versions of the Apex utils. FP16_Module converts the model to FP16 dtype and deals with the forward pass in FP16.

**Tip**

You can apply gradient clipping by calling `clip_master_grads` before `optimizer.step`.

```
optimizer.clip_master_grads(max_norm)  # max_norm(float or int): max norm of the gradients
```
Tip

When using `torch.optim.lr_scheduler` and FP16 training, you need to pass `optimizer.optimizer` to the LR scheduler rather than the optimizer. See the following example code.

```python
from torch.optim.lr_scheduler import StepLR

scheduler = StepLR(
    optimizer.optimizer if smp.state.cfg.fp16 else optimizer,
    step_size=1,
    gamma=args.gamma
)
```

Support for FlashAttention

Support for FlashAttention is a feature of the library only applicable for the `distributed transformer` model, which is a Transformer model wrapped by `smp.DistributedModel()` for model-parallel training. This feature is also compatible with the section called “Tensor Parallelism”.

The FlashAttention library only supports models when `attention_head_size` is set to a value that’s a multiple of 8 and less than 128. Therefore, when you train a distributed transformer and make sure that FlashAttention works properly, you should adjust parameters to make the attention head size comply the requirements. For more information, see also Installation and features in the FlashAttention GitHub repository.

For example, assume that you configure a Transformer model with `hidden_width=864` and `num_heads=48`. The head size of FlashAttention is calculated as `attention_head_size = hidden_width / num_heads = 864 / 48 = 18`. To enable FlashAttention, you need to adjust the `num_heads` parameter to 54, so that `attention_head_size = hidden_width / num_heads = 864 / 54 = 16`, which is a multiple of 8.

Run a SageMaker Distributed Training Job with Model Parallelism

Learn how to run a model-parallel training job of your own training script using the SageMaker Python SDK with the SageMaker model parallelism library.

There are three use-case scenarios for running a SageMaker training job.
1. You can use one of the pre-built AWS Deep Learning Container for TensorFlow and PyTorch. This option is recommended if it is the first time for you to use the model parallel library. To find a tutorial for how to run a SageMaker model parallel training job, see the example notebooks at [PyTorch training with Amazon SageMaker's model parallelism library](#).

2. You can extend the pre-built containers to handle any additional functional requirements for your algorithm or model that the pre-built SageMaker Docker image doesn't support. To find an example of how you can extend a pre-built container, see [Extend a Pre-built Container](#).

3. You can adapt your own Docker container to work with SageMaker using the [SageMaker Training toolkit](#). For an example, see [Adapting Your Own Training Container](#).

For options 2 and 3 in the preceding list, refer to [Extend a Pre-built Docker Container that Contains SageMaker's Distributed Model Parallel Library](#) to learn how to install the model parallel library in an extended or customized Docker container.

In all cases, you launch your training job configuring a SageMaker TensorFlow or PyTorch estimator to activate the library. To learn more, see the following topics.

### Topics
- [Step 1: Modify Your Own Training Script Using SageMaker's Distributed Model Parallel Library](#)
- [Step 2: Launch a Training Job Using the SageMaker Python SDK](#)

### Step 1: Modify Your Own Training Script Using SageMaker's Distributed Model Parallel Library

Use this section to learn how to customize your training script to use the core features of the Amazon SageMaker model parallelism library. To use the library-specific API functions and parameters, we recommend you use this documentation alongside the [SageMaker model parallel library APIs](#) in the [SageMaker Python SDK documentation](#).

The training script examples provided in these sections are simplified and designed to highlight the required changes you must make to use the library. For end-to-end, runnable notebook examples that demonstrate how to use a TensorFlow or PyTorch training script with the SageMaker model parallelism library, see [Amazon SageMaker model parallelism library v2 examples](#).

### Topics
- [Split the model of your training script using the SageMaker model parallelism library](#)
- [Modify a TensorFlow training script](#)
Modify a PyTorch Training Script

Split the model of your training script using the SageMaker model parallelism library

There are two ways to modify your training script to set up model splitting: automated splitting or manual splitting.

Automated model splitting

When you use SageMaker's model parallelism library, you can take advantage of automated model splitting, also referred to as automated model partitioning. The library uses a partitioning algorithm that balances memory, minimizes communication between devices, and optimizes performance. You can configure the automated partitioning algorithm to optimize for speed or memory.

Alternatively, you can use manual model splitting. We recommend automated model splitting, unless you are very familiar with the model architecture and have a good idea of how to efficiently partition your model.

How it works

Auto-partitioning occurs during the first training step, when the `smp.step`-decorated function is first called. During this call, the library first constructs a version of the model on the CPU RAM (to avoid GPU memory limitations), and then analyzes the model graph and makes a partitioning decision. Based on this decision, each model partition is loaded on a GPU, and only then the first step is executed. Because of these analysis and partitioning steps, the first training step might take longer.

In either framework, the library manages the communication between devices through its own backend, which is optimized for AWS infrastructure.

The auto-partition design adapts to the characteristics of the framework, and the library does the partitioning at the granularity level that is more natural in each framework. For instance, in TensorFlow, each specific operation can be assigned to a different device, whereas in PyTorch, the assignment is done at the module level, where each module consists of multiple operations. The following section reviews the specifics of the design in each framework.

Automated model splitting with PyTorch

During the first training step, the model parallelism library internally runs a tracing step that is meant to construct the model graph and determine the tensor and parameter shapes. After
this tracing step, the library constructs a tree, which consists of the nested nn.Module objects in the model, as well as additional data gathered from tracing, such as the amount of stored nn.Parameter, and execution time for each nn.Module.

Next, the library traverses this tree from the root and runs a partitioning algorithm that assigns each nn.Module to a device, which balances computational load (measured by module execution time) and memory use (measured by the total stored nn.Parameter size and activations). If multiple nn.Modules share the same nn.Parameter, then these modules are placed on the same device to avoid maintaining multiple versions of the same parameter. Once the partitioning decision is made, the assigned modules and weights are loaded to their devices.

For instructions on how to register the smp.step decorator to your PyTorch training script, see the section called “Automated splitting with PyTorch”.

Automated model splitting with TensorFlow

The model parallelism library analyzes the sizes of the trainable variables and the graph structure, and internally uses a graph partitioning algorithm. This algorithm comes up with a device assignment for each operation, with the objective of minimizing the amount of communication needed across devices, subject to two constraints:

- Balancing the number of variables stored in each device
- Balancing the number of operations executed in each device

If you specify speed for optimize (in the model parallelism parameters in the Python SDK), the library tries to balance the number of operations and tf.Variable objects in each device. Otherwise, it tries to balance the total size of tf.Variables.

Once the partitioning decision is made, the library creates a serialized representation of the subgraph that each device needs to execute and imports them onto each device. While partitioning, the library places operations that consume the same tf.Variable and operations that are part of the same Keras layer onto the same device. It also respects the colocation constraints imposed by TensorFlow. This means that, for example, if there are two Keras layers that share a tf.Variable, then all operations that are part of these layers are placed on a single device.

For instructions on how to register the smp.step decorator to your PyTorch training script, see the section called “Automated splitting with TensorFlow”.
Comparison of automated model splitting between frameworks

In TensorFlow, the fundamental unit of computation is a `tf.Operation`, and TensorFlow represents the model as a directed acyclic graph (DAG) of `tf.Operations`, and therefore the model parallelism library partitions this DAG so that each node goes to one device. Crucially, `tf.Operation` objects are sufficiently rich with customizable attributes, and they are universal in the sense that every model is guaranteed to consist of a graph of such objects.

PyTorch on the other hand, does not have an equivalent notion of operation that is sufficiently rich and universal. The closest unit of computation in PyTorch that has these characteristics is an `nn.Module`, which is at a much higher granularity level, and this is why the library does partitioning at this level in PyTorch.

Manual Model Splitting

If you want to manually specify how to partition your model across devices, use the `smp.partition` context manager. For instructions on how to set the context manager for manual partitioning, see the following pages.

- the section called “Manual splitting with TensorFlow”
- the section called “Manual splitting with PyTorch”

To use this option after making modifications, in Step 2, you'll need to set `auto_partition` to `False`, and define a `default_partition` in the framework estimator class of the SageMaker Python SDK. Any operation that is not explicitly placed on a partition through the `smp.partition` context manager is executed on the `default_partition`. In this case, the automated splitting logic is bypassed, and each operation is placed based on your specification. Based on the resulting graph structure, the model parallelism library creates a pipelined execution schedule automatically.

Modify a TensorFlow training script

In this section, you learn how to modify TensorFlow training scripts to configure the SageMaker model parallelism library for auto-partitioning and manual partitioning. This selection of examples also includes an example integrated with Horovod for hybrid model and data parallelism.

Note

To find which TensorFlow versions are supported by the library, see the section called “Supported Frameworks and AWS Regions”.

SageMaker model parallelism library v2
The required modifications you must make to your training script to use the library are listed in [Automated splitting with TensorFlow](#).

To learn how to modify your training script to use hybrid model and data parallelism with Horovod, see [Automated splitting with TensorFlow and Horovod for hybrid model and data parallelism](#).

If you want to use manual partitioning, also review [Manual splitting with TensorFlow](#).

The following topics show examples of training scripts that you can use to configure SageMaker's model parallelism library for auto-partitioning and manual partitioning TensorFlow models.

### Note

Auto-partitioning is enabled by default. Unless otherwise specified, the example scripts use auto-partitioning.

### Topics

- [Automated splitting with TensorFlow](#)
- [Automated splitting with TensorFlow and Horovod for hybrid model and data parallelism](#)
- [Manual splitting with TensorFlow](#)
- [Unsupported framework features](#)

### Automated splitting with TensorFlow

The following training script changes are required to run a TensorFlow model with SageMaker's model parallelism library:

1. Import and initialize the library with `smp.init()`.

2. Define a Keras model by inheriting from `smp.DistributedModel` instead of the Keras Model class. Return the model outputs from the call method of the `smp.DistributedModel` object. Be mindful that any tensors returned from the call method will be broadcast across model-parallel devices, incurring communication overhead, so any tensors that are not needed outside the call method (such as intermediate activations) should not be returned.

3. Set `drop_remainder=True` in `tf.Dataset.batch()` method. This is to ensure that the batch size is always divisible by the number of microbatches.
4. Seed the random operations in the data pipeline using `smp.dp_rank()`, e.g., `shuffle(ds, seed=smp.dp_rank())` to ensure consistency of data samples across GPUs that hold different model partitions.

5. Put the forward and backward logic in a step function and decorate it with `smp.step`.

6. Perform post-processing on the outputs across microbatches using `StepOutput` methods such as `reduce_mean`. The `smp.step` function must have a return value that depends on the output of `smp.DistributedModel`.

7. If there is an evaluation step, similarly place the forward logic inside an `smp.step`-decorated function and post-process the outputs using `StepOutput API`.

To learn more about the SageMaker's model parallelism library API, refer to the API documentation.

The following Python script is an example of a training script after the changes are made.

```python
import tensorflow as tf

# smdistributed: Import TF2.x API
import smdistributed.modelparallel.tensorflow as smp

# smdistributed: Initialize
smp.init()

# Download and load MNIST dataset.
(x_train, y_train), (x_test, y_test) = tf.keras.datasets.mnist.load_data(
    "MNIST-data-%d" % smp.rank()
)
x_train, x_test = x_train / 255.0, x_test / 255.0

# Add a channels dimension
x_train = x_train[..., tf.newaxis]
x_test = x_test[..., tf.newaxis]

# smdistributed: If needed, seed the shuffle with smp.dp_rank(), and drop_remainder
# in batching to make sure batch size is always divisible by number of microbatches
train_ds = (tf.data.Dataset.from_tensor_slices((x_train, y_train))
            .shuffle(10000, seed=smp.dp_rank())
            .batch(256, drop_remainder=True)
          )
```

SageMaker model parallelism library v2
# smdistributed: Define smp.DistributedModel the same way as Keras sub-classing API

class MyModel(smp.DistributedModel):
    def __init__(self):
        super(MyModel, self).__init__()
        # define layers

    def call(self, x, training=None):
        # define forward pass and return the model output

model = MyModel()

loss_object = tf.keras.losses.SparseCategoricalCrossentropy(from_logits=True)
optimizer = tf.keras.optimizers.Adam()
train_accuracy = tf.keras.metrics.SparseCategoricalAccuracy(name="train_accuracy")

# smdistributed: Define smp.step. Return any tensors needed outside
@smp.step
def get_grads(images, labels):
    predictions = model(images, training=True)
    loss = loss_object(labels, predictions)

    gradients = optimizer.get_gradients(loss, model.trainable_variables)
    return gradients, loss, predictions

@tf.function
def train_step(images, labels):
    gradients, loss, predictions = get_grads(images, labels)

    # smdistributed: Accumulate the gradients across microbatches
    gradients = [g.accumulate() for g in gradients]
    optimizer.apply_gradients(zip(gradients, model.trainable_variables))

    # smdistributed: Merge predictions and average losses across microbatches
    train_accuracy(labels, predictions.merge())
    return loss.reduce_mean()

for epoch in range(5):
    # Reset the metrics at the start of the next epoch
    train_accuracy.reset_states()
    for images, labels in train_ds:
        loss = train_step(images, labels)
accuracy = train_accuracy.result()

If you are done preparing your training script, proceed to Step 2: Launch a Training Job Using the SageMaker Python SDK. If you want to run a hybrid model and data parallel training job, continue to the next section.

**Automated splitting with TensorFlow and Horovod for hybrid model and data parallelism**

You can use the SageMaker model parallelism library with Horovod for hybrid model and data parallelism. To read more about how the library splits a model for hybrid parallelism, see [Pipeline parallelism (available for PyTorch and TensorFlow)](https://aws.amazon.com/sagemaker/)

In this step, we focus on how to modify your training script to adapt the SageMaker model parallelism library.

To properly set up your training script to pick up the hybrid parallelism configuration that you'll set in Step 2: Launch a Training Job Using the SageMaker Python SDK, use the library's helper functions, `smp.dp_rank()` and `smp.mp_rank()`, which automatically detect the data parallel rank and model parallel rank respectively.

To find all MPI primitives the library supports, see [MPI Basics](https://aws.amazon.com/sagemaker/) in the SageMaker Python SDK documentation.

The required changes needed in the script are:

- Adding `hvd.allreduce`
- Broadcasting variables after the first batch, as required by Horovod
- Seeding shuffling and/or sharding operations in the data pipeline with `smp.dp_rank()`.

⚠️ **Note**

When you use Horovod, you must not directly call `hvd.init` in your training script. Instead, you'll have to set "horovod" to `True` in the SageMaker Python SDK modelparallel parameters in Step 2: Launch a Training Job Using the SageMaker Python SDK. This allows the library to internally initialize Horovod based on the device assignments of model partitions. Calling `hvd.init()` directly in your training script can cause problems.
**Note**

Using the `hvd.DistributedOptimizer` API directly in your training script might result in a poor training performance and speed, because the API implicitly places the `AllReduce` operation inside `smp.step`. We recommend you to use the model parallelism library with Horovod by directly calling `hvd.allreduce` after calling `accumulate()` or `reduce_mean()` on the gradients returned from `smp.step`, as will be shown in the following example.

To learn more about the SageMaker's model parallelism library API, refer to the [API documentation](#).

```python
import tensorflow as tf
import horovod.tensorflow as hvd

# smdistributed: Import TF2.x API
import smdistributed.modelparallel.tensorflow as smp

# smdistributed: Initialize
smp.init()

# Download and load MNIST dataset.
(x_train, y_train), (x_test, y_test) = tf.keras.datasets.mnist.load_data('MNIST-data-%d' % smp.rank())
x_train, x_test = x_train / 255.0, x_test / 255.0

# Add a channels dimension
x_train = x_train[..., tf.newaxis]
x_test = x_test[..., tf.newaxis]

# smdistributed: Seed the shuffle with smp.dp_rank(), and drop_remainder
# in batching to make sure batch size is always divisible by number of microbatches
train_ds = (tf.data.Dataset.from_tensor_slices((x_train, y_train))
    .shuffle(10000, seed=smp.dp_rank())
    .batch(256, drop_remainder=True)
)

# smdistributed: Define smp.DistributedModel the same way as Keras sub-classing API
```
class MyModel(smp.DistributedModel):
    def __init__(self):
        super(MyModel, self).__init__()
        # define layers

        def call(self, x, training=None):
            # define forward pass and return model outputs

    model = MyModel()

    loss_object = tf.keras.losses.SparseCategoricalCrossentropy(from_logits=True)
    optimizer = tf.keras.optimizers.Adam()
    train_accuracy = tf.keras.metrics.SparseCategoricalAccuracy(name="train_accuracy")

    # smdistributed: Define smp.step. Return any tensors needed outside
    @smp.step
    def get_grads(images, labels):
        predictions = model(images, training=True)
        loss = loss_object(labels, predictions)

        grads = optimizer.get_gradients(loss, model.trainable_variables)
        return grads, loss, predictions

    @tf.function
    def train_step(images, labels, first_batch):
        gradients, loss, predictions = get_grads(images, labels)

        # smdistributed: Accumulate the gradients across microbatches
        # Horovod: AllReduce the accumulated gradients
        gradients = [hvd.allreduce(g.accumulate()) for g in gradients]
        optimizer.apply_gradients(zip(gradients, model.trainable_variables))

        # Horovod: Broadcast the variables after first batch
        if first_batch:
            hvd.broadcast_variables(model.variables, root_rank=0)
            hvd.broadcast_variables(optimizer.variables(), root_rank=0)

        # smdistributed: Merge predictions across microbatches
        train_accuracy(labels, predictions.merge())

        return loss.reduce_mean()
for epoch in range(5):
    # Reset the metrics at the start of the next epoch
    train_accuracy.reset_states()

    for batch, (images, labels) in enumerate(train_ds):
        loss = train_step(images, labels, tf.constant(batch == 0))

Manual splitting with TensorFlow

Use `smp.partition` context managers to place operations in specific partition. Any operation not placed in any `smp.partition` contexts is placed in the `default_partition`. To learn more about the SageMaker's model parallelism library API, refer to the [API documentation](#).

```python
import tensorflow as tf

# smdistributed: Import TF2.x API.
import smdistributed.modelparallel.tensorflow as smp

# smdistributed: Initialize
smp.init()

# Download and load MNIST dataset.
(x_train, y_train), (x_test, y_test) = tf.keras.datasets.mnist.load_data(
    "MNIST-data-%d" % smp.rank()
)

x_train, x_test = x_train / 255.0, x_test / 255.0

# Add a channels dimension
x_train = x_train[..., tf.newaxis]
x_test = x_test[..., tf.newaxis]

# smdistributed: If needed, seed the shuffle with smp.dp_rank(), and drop_remainder
# in batching to make sure batch size is always divisible by number of microbatches.
train_ds = (tf.data.Dataset.from_tensor_slices((x_train, y_train))
                   .shuffle(10000, seed=smp.dp_rank())
                   .batch(256, drop_remainder=True)
)

# smdistributed: Define `smp.DistributedModel` the same way as Keras sub-classing API.
class MyModel(smp.DistributedModel):
    def __init__(self):
        # define layers
```
```python
def call(self, x):
    with smp.partition(0):
        x = self.layer0(x)
    with smp.partition(1):
        return self.layer1(x)

model = MyModel()

loss_object = tf.keras.losses.SparseCategoricalCrossentropy(from_logits=True)
optimizer = tf.keras.optimizers.Adam()
train_accuracy = tf.keras.metrics.SparseCategoricalAccuracy(name="train_accuracy")

# smdistributed: Define smp.step. Return any tensors needed outside
@smp.step
def get_grads(images, labels):
    predictions = model(images, training=True)
    loss = loss_object(labels, predictions)
    
    grads = optimizer.get_gradients(loss, model.trainable_variables)
    return grads, loss, predictions

@tf.function
def train_step(images, labels):
    gradients, loss, predictions = get_grads(images, labels)
    
    # smdistributed: Accumulate the gradients across microbatches
    gradients = [g.accumulate() for g in gradients]
    optimizer.apply_gradients(zip(gradients, model.trainable_variables))
    
    # smdistributed: Merge predictions and average losses across microbatches
    train_accuracy(labels, predictions.merge())
    return loss.reduce_mean()

for epoch in range(5):
    # Reset the metrics at the start of the next epoch
    train_accuracy.reset_states()
    for images, labels in train_ds:
        loss = train_step(images, labels)
        accuracy = train_accuracy.result()
```

Unsupported framework features

The following TensorFlow features are not supported by the library:

- `tf.GradientTape()` is currently not supported. You can use `Optimizer.get_gradients()` or `Optimizer.compute_gradients()` instead to compute gradients.
- The `tf.train.Checkpoint.restore()` API is currently not supported. For checkpointing, use `smp.CheckpointManager` instead, which provides the same API and functionality. Note that checkpoint restores with `smp.CheckpointManager` should take place after the first step.

Modify a PyTorch Training Script

In this section, you learn how to modify PyTorch training scripts to configure the SageMaker model parallelism library for auto-partitioning and manual partitioning.

**Note**

To find which PyTorch versions are supported by the library, see the section called “Supported Frameworks and AWS Regions”.

**Tip**

For end-to-end notebook examples that demonstrate how to use a PyTorch training script with the SageMaker model parallelism library, see Amazon SageMaker model parallelism library v1 examples.

Note that auto-partitioning is enabled by default. Unless otherwise specified, the following scripts use auto-partitioning.

Topics

- Automated splitting with PyTorch
- Manual splitting with PyTorch
- Considerations
- Unsupported framework features
Automated splitting with PyTorch

The following training script changes are required to run a PyTorch training script with SageMaker's model parallelism library:

1. Import and initialize the library with `smdistributed.modelparallel.torch.init()`.  
2. Wrap the model with `smdistributed.modelparallel.torch.DistributedModel`. Be mindful that any tensors returned from the `forward` method of the underlying `nn.Module` object will be broadcast across model-parallel devices, incurring communication overhead, so any tensors that are not needed outside the call method (such as intermediate activations) should not be returned.

   **Note**  
   For FP16 training, you need to use the `smdistributed.modelparallel.torch.model_creation()` context manager to wrap the model. For more information, see [FP16 Training with Model Parallelism](#).  

3. Wrap the optimizer with `smdistributed.modelparallel.torch.DistributedOptimizer`.

   **Note**  
   For FP16 training, you need to set up static or dynamic loss scaling. For more information, see [FP16 Training with Model Parallelism](#).  

4. Use the returned `DistributedModel` object instead of a user model.

5. Put the forward and backward logic in a step function and decorate it with `smdistributed.modelparallel.torch.step`.

6. Restrict each process to its own device through `torch.cuda.set_device(smp.local_rank())`.

7. Move the input tensors to the GPU using the `.to()` API before the `smp.step` call (see example below).


9. Perform post-processing on the outputs across microbatches using `StepOutput` methods such as `reduce_mean`.
10. If there is an evaluation step, similarly place the forward logic inside an `smp.step`-decorated function and post-process the outputs using `StepOutput API`.

11. Set `drop_last=True` in `DataLoader`. Alternatively, manually skip a batch in the training loop if the batch size is not divisible by the number of microbatches.

To learn more about the SageMaker's model parallelism library API, refer to the [API documentation](#).

```python
import torch
import torch.nn as nn
import torch.nn.functional as F
import torch.optim as optim
from torchtext.dataset import SplitDataset
from torchvision import datasets
import smdistributed.modelparallel.torch as smp

class GroupedNet(nn.Module):
    def __init__(self):
        super(GroupedNet, self).__init__()
        # define layers

        def forward(self, x):
            # define forward pass and return model outputs

    @smp.step
    def train_step(model, data, target):
        output = model(data)
        loss = F.nll_loss(output, target, reduction="mean")
        model.backward(loss)
        return output, loss

def train(model, device, train_loader, optimizer):
    model.train()
    for batch_idx, (data, target) in enumerate(train_loader):
        # smdistributed: Move input tensors to the GPU ID used by the current process, based on the set_device call.
        data, target = data.to(device), target.to(device)
        optimizer.zero_grad()
# Return value, loss_mb is a StepOutput object
_, loss_mb = train_step(model, data, target)

# smdistributed: Average the loss across microbatches.
loss = loss_mb.reduce_mean()

optimizer.step()

# smdistributed: initialize the backend
smp.init()

# smdistributed: Set the device to the GPU ID used by the current process.
# Input tensors should be transferred to this device.
torch.cuda.set_device(smp.local_rank())
device = torch.device("cuda")

# smdistributed: Download only on a single process per instance.
# When this is not present, the file is corrupted by multiple processes trying
# to download and extract at the same time
dataset = datasets.MNIST("../data", train=True, download=False)

# smdistributed: Shard the dataset based on data-parallel ranks
if smp.dp_size() > 1:
    partitions_dict = {f"{i}": 1 / smp.dp_size() for i in range(smp.dp_size())}
dataset = SplitDataset(dataset, partitions=partitions_dict)
dataset.select(f"{smp.dp_rank()}")

# smdistributed: Set drop_last=True to ensure that batch size is always divisible
# by the number of microbatches
train_loader = torch.utils.data.DataLoader(dataset, batch_size=64, drop_last=True)

model = GroupedNet()
optimizer = optim.Adadelta(model.parameters(), lr=4.0)

# smdistributed: Use the DistributedModel container to provide the model
# to be partitioned across different ranks. For the rest of the script,
# the returned DistributedModel object should be used in place of
# the model provided for DistributedModel class instantiation.
model = smp.DistributedModel(model)
optimizer = smp.DistributedOptimizer(optimizer)

train(model, device, train_loader, optimizer)
Manual splitting with PyTorch

Use `smp.partition` context managers to place modules in specific devices. Any module not placed in any `smp.partition` contexts is placed in the `default_partition`. The `default_partition` needs to be provided if `auto_partition` is set to `False`. The modules that are created within a specific `smp.partition` context are placed on the corresponding partition.

To learn more about the SageMaker's model parallelism library API, refer to the [API documentation](#).

```python
import torch
import torch.nn as nn
import torch.nn.functional as F
import torch.optim as optim
from torchtext.dataset import SplitDataset
from torchvision import datasets
import smdistributed.modelparallel.torch as smp

class GroupedNet(nn.Module):
    def __init__(self):
        super(GroupedNet, self).__init__()
        with smp.partition(0):
            # define child modules on device 0
            with smp.partition(1):
                # define child modules on device 1

        def forward(self, x):
            # define forward pass and return model outputs

    # smdistributed: Define smp.step. Return any tensors needed outside.
    @smp.step
    def train_step(model, data, target):
        output = model(data)
        loss = F.nll_loss(output, target, reduction="mean")
        model.backward(loss)
        return output, loss

    def train(model, device, train_loader, optimizer):
        model.train()```
for batch_idx, (data, target) in enumerate(train_loader):
    # smdistributed: Move input tensors to the GPU ID used by the current process,
    # based on the set_device call.
    data, target = data.to(device), target.to(device)
    optimizer.zero_grad()
    # Return value, loss_mb is a StepOutput object
    _, loss_mb = train_step(model, data, target)

    # smdistributed: Average the loss across microbatches.
    loss = loss_mb.reduce_mean()

    optimizer.step()

    # smdistributed: Initialize the backend
    smp.init()

    # smdistributed: Set the device to the GPU ID used by the current process.
    # Input tensors should be transferred to this device.
    torch.cuda.set_device(smp.local_rank())
    device = torch.device("cuda")

    # smdistributed: Download only on a single process per instance.
    # When this is not present, the file is corrupted by multiple processes trying
    # to download and extract at the same time
    dataset = datasets.MNIST("../data", train=True, download=False)

    # smdistributed: Shard the dataset based on data-parallel ranks
    if smp.dp_size() > 1:
        partitions_dict = {f"{i}" : 1 / smp.dp_size() for i in range(smp.dp_size())}
        dataset = SplitDataset(dataset, partitions=partitions_dict)
        dataset.select(f"{smp.dp_rank()}"")

    # smdistributed: Set drop_last=True to ensure that batch size is always divisible
    # by the number of microbatches
    train_loader = torch.utils.data.DataLoader(dataset, batch_size=64, drop_last=True)

model = GroupedNet()
optimizer = optim.Adadelta(model.parameters(), lr=4.0)

# smdistributed: Use the DistributedModel container to provide the model
# to be partitioned across different ranks. For the rest of the script,
# the returned DistributedModel object should be used in place of
# the model provided for DistributedModel class instantiation.
model = smp.DistributedModel(model)
optimizer = smp.DistributedOptimizer(optimizer)

train(model, device, train_loader, optimizer)

Considerations

When you configure a PyTorch training script using SageMaker's model parallelism library, you should be aware of the following:

• If you are using an optimization technique that relies on global gradient norms, for example gradient norm from the entire model, such as some variants of LAMB optimizer or global gradient clipping, you need to gather all the norms across the model partitions for correctness. You can use the library's communication basic data types to do this.

• All torch.Tensor arguments to the forward methods of the nn.Modules in your model must be used in the computation of the module output. In other words, the library does not support the case where there is a torch.Tensor argument to a module on which the module output does not depend.

• The argument to the smp.DistributedModel.backward() call must depend on all model outputs. In other words, there cannot be an output from the smp.DistributedModel.forward call that is not used in the computation of the tensor that is fed into the smp.DistributedModel.backward call.

• If there are torch.cuda.synchronize() calls in your code, you might need to call torch.cuda.set_device(smp.local_rank()) immediately before the synchronize call. Otherwise unnecessary CUDA contexts might be created in device 0, which will needlessly consume memory.

• Since the library places nn.Modules on different devices, the modules in the model must not depend on any global state that is modified inside smp.step. Any state that remains fixed throughout training, or that is modified outside smp.step in a way that is visible to all processes, is allowed.

• You don't need to move the model to GPU (for example, using model.to(device)) when using the library. If you try to move the model to GPU before the model is partitioned (before the first smp.step call), the move call is ignored. The library automatically moves the part of the model assigned to a rank to its GPU. Once training with the library starts, don’t move the model to CPU and use it, as it won't have correct parameters for modules not assigned to the partition held by the process. If you want to retrain a model or use it for inference without the library after it was trained using the model parallelism library, the recommended way is to save the full model using our checkpointing API and load it back to a regular PyTorch Module.
• If you have a list of modules such that output of one feeds into another, replacing that list with `nn.Sequential` can significantly improve performance.

• The weight update (`optimizer.step()`) needs to happen outside of `smp.step` because that is when the entire backward pass is done and gradients are ready. When using a hybrid model with model and data parallelism, at this point, AllReduce of gradients is also guaranteed to finish.

• When using the library in combination with data parallelism, make sure that the number of batches on all data parallel ranks is the same so that AllReduce does not hang waiting for a rank which is not participating in the step.

• If you launch a training job using an ml.p4d instance type (such as ml.p4d.24xlarge), you must set the data loader variable `num_workers=0`. For example, you may define your `DataLoader` as follows:

```python
dataloader = torch.utils.data.DataLoader(
    data,
    batch_size=batch_size,
    num_workers=0,
    pin_memory=True,
    drop_last=True,
    shuffle=shuffle,
)
```

• The inputs to `smp.step` must be the model inputs generated by `DataLoader`. This is because `smp.step` internally splits the input tensors along the batch dimension and pipelines them. This means that passing `DataLoader` itself to the `smp.step` function to generate the model inputs inside does not work.

For example, if you define a `DataLoader` as follows:

```python
train_loader = torch.utils.data.DataLoader(dataset, batch_size=64, drop_last=True)
```

You should access the model inputs generated by `train_loader` and pass those to an `smp.step` decorated function. Do not pass `train_loader` directly to the `smp.step` function.

```python
def train(model, device, train_loader, optimizer):
    model.train()
    for batch_idx, (data, target) in enumerate(train_loader):
        ...
        _, loss_mb = train_step(model, data, target)
        ...
@smp.step
def train_step(model, data, target):
    ...
    return output, loss

- The input tensors to smp.step must be moved to the current device using .to() API, which must take place after the torch.cuda.set_device(local_rank()) call.

For example, you may define the train function as follows. This function adds data and target to the current device using .to() API before using those input tensors to call train_step.

```python
def train(model, device, train_loader, optimizer):
    model.train()
    for batch_idx, (data, target) in enumerate(train_loader):
        # smdistributed: Move input tensors to the GPU ID used by the current process,
        # based on the set_device call.
        data, target = data.to(device), target.to(device)
        optimizer.zero_grad()
        # Return value, loss_mb is a StepOutput object
        _, loss_mb = train_step(model, data, target)

        # smdistributed: Average the loss across microbatches.
        loss = loss_mb.reduce_mean()

        optimizer.step()
```

The input tensors to this smp.set decorated function have been moved to the current device in the train function above. The model does not need to be moved to the current device. The library automatically moves the part of the model assigned to a rank to its GPU.

```python
@smp.step
def train_step(model, data, target):
    output = model(data)
    loss = F.nll_loss(output, target, reduction="mean")
    model.backward(loss)
    return output, loss
```
Unsupported framework features

The following PyTorch features are unsupported by SageMaker's model parallelism library:

- If you use data parallelism with the native PyTorch DDP, the torch.nn.parallel.DistributedDataParallel wrapper module is not supported by the library. The library internally manages integrating with PyTorch DDP, including parameter broadcast and gradient AllReduce. When using the library, module buffers are only broadcast once at the start of training. If your model has module buffers that need to be synchronized across data parallel groups at each step, you can do so through the torch.distributed API, using the process group that can be obtained via smp.get_dp_process_group().
- For mixed precision training, the apex.amp module is not supported. The recommended way to use the library with automatic mixed-precision is to use torch.cuda.amp, with the exception of using smp.amp.GradScaler instead of the implementation in torch.
- torch.jit.ScriptModules or ScriptFunctions are not supported by smp.DistributedModel.
- apex:FusedLayerNorm, FusedAdam, FusedLAMB, and FusedNovoGrad from apex are not supported. You can use the library implementations of these through smp.optimizers and smp.nn APIs instead.

Step 2: Launch a Training Job Using the SageMaker Python SDK

The SageMaker Python SDK supports managed training of models with ML frameworks such as TensorFlow and PyTorch. To launch a training job using one of these frameworks, you define a SageMaker TensorFlow estimator, a SageMaker PyTorch estimator, or a SageMaker generic Estimator to use the modified training script and model parallelism configuration.

Topics

- Using the SageMaker TensorFlow and PyTorch Estimators
- Extend a Pre-built Docker Container that Contains SageMaker's Distributed Model Parallel Library
- Create Your Own Docker Container with the SageMaker Distributed Model Parallel Library

Using the SageMaker TensorFlow and PyTorch Estimators

The TensorFlow and PyTorch estimator classes contain the distribution parameter, which you can use to specify configuration parameters for using distributed training frameworks. The
SageMaker model parallel library internally uses MPI for hybrid data and model parallelism, so you must use the MPI option with the library.

The following template of a TensorFlow or PyTorch estimator shows how to configure the distribution parameter for using the SageMaker model parallel library with MPI.

Using the SageMaker TensorFlow estimator

```python
import sagemaker
from sagemaker.tensorflow import TensorFlow

smp_options = {
    "enabled": True,       # Required
    "parameters": {
        "partitions": 2,    # Required
        "microbatches": 4,
        "placement_strategy": "spread",
        "pipeline": "interleaved",
        "optimize": "speed",
        "horovod": True,     # Use this for hybrid model and data parallelism
    }
}

mpi_options = {
    "enabled": True,       # Required
    "processes_per_host": 8,  # Required
    # "custom_mpi_options": "--mca btl_vader_single_copy_mechanism none"
}

smd_mp_estimator = TensorFlow(
    entry_point="your_training_script.py",  # Specify your train script
    source_dir="location_to_your_script",
    role=sagemaker.get_execution_role(),
    instance_count=1,
    instance_type='ml.p3.16xlarge',
    framework_version='2.6.3',
    py_version='py38',
    distribution={
        "smdistributed": {"modelparallel": smp_options},
        "mpi": mpi_options
    },
    base_job_name="SMD-MP-demo",
)
import sagemaker
from sagemaker.pytorch import PyTorch

smp_options = {
    "enabled": True,
    "parameters": {
        "pipeline_parallel_degree": 2,  # Required
        "microbatches": 4,              # Required
        "placement_strategy": "spread",
        "pipeline": "interleaved",
        "optimize": "speed",
        "ddp": True,
    }
}

mpi_options = {
    "enabled": True,  # Required
    "processes_per_host": 8,  # Required
    # "custom_mpi_options": "--mca btl_vader_single_copy_mechanism none"
}

smd_mp_estimator = PyTorch(
    entry_point="your_training_script.py",  # Specify your train script
    source_dir="location_to_your_script",
    role=sagemaker.get_execution_role(),
    instance_count=1,
    instance_type='ml.p3.16xlarge',
    framework_version='1.13.1',
    py_version='py38',
    distribution={
        "smdistributed": {"modelparallel": smp_options},
        "mpi": mpi_options
    },
    base_job_name="SMD-MP-demo",
)

smd_mp_estimator.fit('s3://my_bucket/my_training_data/')}
To enable the library, you need to pass configuration dictionaries to the "smdistributed" and "mpi" keys through the distribution argument of the SageMaker estimator constructors.

**Configuration parameters for SageMaker model parallelism**

- For the "smdistributed" key, pass a dictionary with the "modelparallel" key and the following inner dictionaries.

  
  ![Note]
  Using "modelparallel" and "dataparallel" in one training job is not supported.

- "enabled" – Required. To enable model parallelism, set "enabled": True.
- "parameters" – Required. Specify a set of parameters for SageMaker model parallelism.
  - For a complete list of common parameters, see [Parameters for smdistributed](#) in the [SageMaker Python SDK documentation](#).

  For TensorFlow, see [TensorFlow-specific Parameters](#).

  For PyTorch, see [PyTorch-specific Parameters](#).

- "pipeline_parallel_degree" (or "partitions" in smdistributed-modelparallel<v1.6.0) – Required. Among the [parameters for smdistributed](#), this parameter is required to specify how many model partitions you want to split into.

  
  ![Important]
  There is a breaking change in the parameter name. The "pipeline_parallel_degree" parameter replaces the "partitions" since smdistributed-modelparallel v1.6.0. For more information, see [Common Parameters](#) for SageMaker model parallelism configuration and [SageMaker Distributed Model Parallel Release Notes](#) in the [SageMaker Python SDK documentation](#).

- For the "mpi" key, pass a dictionary that contains the following:
  - "enabled" – Required. Set True to launch the distributed training job with MPI.
  - "processes_per_host" – Required. Specify the number of processes MPI should launch on each host. In SageMaker, a host is a single Amazon EC2 ML instance. The SageMaker Python
SDK maintains a one-to-one mapping between processes and GPUs across model and data parallelism. This means that SageMaker schedules each process on a single, separate GPU and no GPU contains more than one process. If you are using PyTorch, you must restrict each process to its own device through `torch.cuda.set_device(smp.local_rank())`. To learn more, see [Automated splitting with PyTorch](#).

**Important**

`process_per_host` *must* not be greater than the number of GPUs per instance and typically will be equal to the number of GPUs per instance.

- "custom_mpi_options" (optional) – Use this key to pass any custom MPI options you might need. If you do not pass any MPI custom options to the key, the MPI option is set by default to the following flag.

```
--mca btl_vader_single_copy_mechanism none
```

**Note**

You do not need to explicitly specify this default flag to the key. If you explicitly specify it, your distributed model parallel training job might fail with the following error:

```
The following MCA parameter has been listed multiple times on the command line:
MCA param: btl_vader_single_copy_mechanism MCA parameters can only be listed once
on a command line to ensure there is no ambiguity as to its value. Please correct the situation and try again.
```

**Tip**

If you launch a training job using an EFA-enabled instance type, such as `ml.p4d.24xlarge` and `ml.p3dn.24xlarge`, use the following flag for best performance:
To launch the training job using the estimator and your SageMaker model parallel configured training script, run the `estimator.fit()` function.

Use the following resources to learn more about using the model parallelism features in the SageMaker Python SDK:

- Use TensorFlow with the SageMaker Python SDK
- Use PyTorch with the SageMaker Python SDK

We recommend you use a SageMaker notebook instance if you are new users. To see an example of how you can launch a training job using a SageMaker notebook instance, see Amazon SageMaker model parallelism library v2 examples.

You can also submit a distributed training job from your machine using AWS CLI. To set up AWS CLI on your machine, see set up your AWS credentials and Region for development.

### Extend a Pre-built Docker Container that Contains SageMaker's Distributed Model Parallel Library

To extend a pre-built container and use SageMaker's model parallelism library, you must use one of the available AWS Deep Learning Containers (DLC) images for PyTorch or TensorFlow. The SageMaker model parallelism library is included in the TensorFlow (2.3.0 and later) and PyTorch (1.6.0 and later) DLC images with CUDA (cuxyz). For a complete list of DLC images, see Available Deep Learning Containers Images in the AWS Deep Learning Containers GitHub repository.

⚠️ Tip

We recommend that you use the image that contains the latest version of TensorFlow or PyTorch to access the most up-to-date version of the SageMaker model parallelism library.

For example, your Dockerfile should contain a FROM statement similar to the following:

```bash
# Use the SageMaker DLC image URI for TensorFlow or PyTorch
```
FROM aws-dlc-account-id.dkr.ecr.aws-region.amazonaws.com/framework-training:{framework-version-tag}

# Add your dependencies here
RUN ...

ENV PATH="/opt/ml/code:${PATH}"

# this environment variable is used by the SageMaker container to determine our user code directory.
ENV SAGEMAKER_SUBMIT_DIRECTORY /opt/ml/code

Additionally, when you define a PyTorch or TensorFlow estimator, you must specify that the entry_point for your training script. This should be the same path identified with ENV SAGEMAKER_SUBMIT_DIRECTORY in your Dockerfile.

⚠️ Tip

You must push this Docker container to Amazon Elastic Container Registry (Amazon ECR) and use the image URI (image_uri) to define a SageMaker estimator for training. For more information, see Extend a Pre-built Container.

After you finish hosting the Docker container and retrieving the image URI of the container, create a SageMaker PyTorch estimator object as follows. This example assumes that you have already defined smp_options and mpi_options.

```python
smd_mp_estimator = Estimator(
    entry_point="your_training_script.py",
    role=sagemaker.get_execution_role(),
    instance_type='ml.p3.16xlarge',
    sagemaker_session=sagemaker_session,
    image_uri='your_aws_account_id.dkr.ecr.region.amazonaws.com/name:tag'
    instance_count=1,
    distribution={
        "smdistributed": smp_options,
        "mpi": mpi_options
    },
    base_job_name="SMD-MP-demo",
)```

SageMaker model parallelism library v2
Create Your Own Docker Container with the SageMaker Distributed Model Parallel Library

To build your own Docker container for training and use the SageMaker model parallel library, you must include the correct dependencies and the binary files of the SageMaker distributed parallel libraries in your Dockerfile. This section provides the minimum set of code blocks you must include to properly prepare a SageMaker training environment and the model parallel library in your own Docker container.

Note
This custom Docker option with the SageMaker model parallel library as a binary is available only for PyTorch.

To create a Dockerfile with the SageMaker training toolkit and the model parallel library

1. Start with one of the NVIDIA CUDA base images.

   FROM <cuda-cudnn-base-image>

Tip
The official AWS Deep Learning Container (DLC) images are built from the NVIDIA CUDA base images. We recommend you look into the official Dockerfiles of AWS Deep Learning Container for PyTorch to find which versions of the libraries you need to install and how to configure them. The official Dockerfiles are complete, benchmark tested, and managed by the SageMaker and Deep Learning Container service teams. In the provided link, choose the PyTorch version you use, choose the CUDA (cuxyz) folder, and choose the Dockerfile ending with .gpu or .sagemaker.gpu.

2. To set up a distributed training environment, you need to install software for communication and network devices, such as Elastic Fabric Adapter (EFA), NVIDIA Collective Communications Library (NCCL), and Open MPI. Depending on the PyTorch and CUDA versions you choose, you must install compatible versions of the libraries.
**Important**

Because the SageMaker model parallel library requires the SageMaker data parallel library in the subsequent steps, we highly recommend that you follow the instructions at [Create your own Docker container with the SageMaker distributed data parallel library](#) to properly set up a SageMaker training environment for distributed training.

For more information about setting up EFA with NCCL and Open MPI, see [Get started with EFA and MPI](#) and [Get started with EFA and NCCL](#).

3. Add the following arguments to specify the URLs of the SageMaker distributed training packages for PyTorch. The SageMaker model parallel library requires the SageMaker data parallel library to use the cross-node Remote Direct Memory Access (RDMA).

ARG SMDATAPARALLEL_BINARY=https://smdataparallel.s3.amazonaws.com/binary/pytorch/1.10.2/cu113/2022-02-18/smdistributed_dataparallel-1.4.0-cp38-cp38-linux_x86_64.whl

4. Install dependencies that the SageMaker model parallel library requires.

   a. Install the [METIS](#) library.

ARG METIS=metis-5.1.0

RUN rm /etc/apt/sources.list.d/* \
    && wget -nv http://glaros.dtc.umn.edu/gkhome/fetch/sw/metis/${METIS}.tar.gz \ 
    && gunzip -f ${METIS}.tar.gz \ 
    && tar -xvf ${METIS}.tar \ 
    && cd ${METIS} \ 
    && apt-get update \ 
    && make config shared=1 \ 
    && make install \ 
    && cd .. \ 
    && rm -rf ${METIS}.tar* \ 
    && rm -rf ${METIS} \ 
    && rm -rf /var/lib/apt/lists/* \ 
    && apt-get clean
b. Install the RAPIDS Memory Manager library. This requires CMake 3.14 or later.

```
ARG RMM_VERSION=0.15.0

RUN wget -nv https://github.com/rapidsai/rmm/archive/v${RMM_VERSION}.tar.gz 
    && tar -xvf v${RMM_VERSION}.tar.gz 
    && cd rmm-${RMM_VERSION} 
    && INSTALL_PREFIX=/usr/local ./build.sh librmm 
    && cd .. 
    && rm -rf v${RMM_VERSION}.tar* 
    && rm -rf rmm-${RMM_VERSION}
```

5. Install the SageMaker model parallel library.

```
RUN pip install --no-cache-dir -U ${SMD_MODEL_PARALLEL_URL}
```

6. Install the SageMaker data parallel library.

```
RUN SMDATAPARALLEL_PT=1 pip install --no-cache-dir ${SMDATAPARALLEL_BINARY}
```

7. Install the sagemaker-training toolkit. The toolkit contains the common functionality that's necessary to create a container compatible with the SageMaker training platform and the SageMaker Python SDK.

```
RUN pip install sagemaker-training
```

8. After you finish creating the Dockerfile, see Adapting Your Own Training Container to learn how to build the Docker container and host it in Amazon ECR.

Tip

For more general information about creating a custom Dockerfile for training in SageMaker, see Use Your Own Training Algorithms.

Checkpointing and Fine-Tuning a Model with Model Parallelism

The SageMaker model parallelism library provides checkpointing APIs to save the model state and the optimizer state split by the various model parallelism strategies, and to load checkpoints for
continuous training from where you want to restart training and fine-tune. The APIs also support options to save the model and optimizer states partially or fully.

Topics

- Checkpointing a distributed model
- Fine-tuning a distributed model

Checkpointing a distributed model

Choose one of the following topics depending on the framework between PyTorch and TensorFlow and the version of the SageMaker model parallelism library you use.

Topics

- Checkpointing a distributed PyTorch model (for the SageMaker model parallelism library v1.10.0 and later)
- Checkpointing a distributed PyTorch model (for the SageMaker model parallelism library between v1.6.0 and v1.9.0)
- Checkpointing a distributed TensorFlow model

Checkpointing a distributed PyTorch model (for the SageMaker model parallelism library v1.10.0 and later)

The SageMaker model parallelism library provides checkpoint APIs to save and load full or partial checkpoints of the distributed model state and its optimizer state.

Note

This checkpointing method is recommended if you use PyTorch and the SageMaker model parallelism library v1.10.0 or later.

Partial checkpointing

To save checkpoints of a model trained with model parallelism, use the `smdistributed.modelparallel.torch.save_checkpoint` API with the partial checkpointing option set to true (`partial=True`). This saves each model partition individually. In addition to the model and the optimizer state, you can also save any additional custom data.
through the `user_content` argument. The checkpointed model, optimizer, and user content are saved as separate files. The `save_checkpoint` API call creates checkpoint folders in the following structure.

- path
- `${tag}_partial` (folder for partial checkpoints)
- model_rankinfo.pt
- optimizer_rankinfo.pt
- fp16_states_rankinfo.pt
- user_content.pt
- `$tag` (checkpoint file for full checkpoints)
- user_content_${tag} (user_content file for full checkpoints)
- newest (a file that indicates the newest checkpoint)

To resume training from partial checkpoints, use the `smdistributed.modelparallel.torch.resume_from_checkpoint` API with `partial=True`, and specify the checkpoint directory and the tag used while saving the partial checkpoints. Note that the actual loading of model weights happens after model partitioning, during the first run of the `smdistributed.modelparallel.torch.step`-decorated training step function.

When saving a partial checkpoint, the library also saves the model partition decision as files with `.pt` file extension. Conversely, when resuming from the partial checkpoint, the library loads the partition decision files together. Once the partition decision is loaded, you can't change the partition.

The following code snippet shows how to set the checkpoint APIs in a PyTorch training script.

```python
import smdistributed.modelparallel.torch as smp

model = ...
model = smp.DistributedModel(model)
optimizer = ...
optimizer = smp.DistributedOptimizer(optimizer)
user_content = ...

checkpoint_path = "[/opt/ml/checkpoint/model_parallel"

# Save a checkpoint.
smp.save_checkpoint(
    path=checkpoint_path,
    tag=f"total_steps{total_steps}"),
```
partial=True,
model=model,
optimizer=optimizer,
user_content=user_content
num_kept_partial_checkpoints=5
)

# Load a checkpoint.
# This automatically loads the most recently saved checkpoint.
smp_checkpoint = smp.resume_from_checkpoint(
    path=checkpoint_path,
    partial=True
)

Full checkpointing

To save the final model artifact for inference purposes, use the
smdistributed.modelparallel.torch.save_checkpoint API with partial=False, which
combines the model partitions to create a single model artifact. Note that this does not combine
the optimizer states.

To initialize training with particular weights, given a full model checkpoint, you can use
the smdistributed.modelparallel.torch.resume_from_checkpoint API with
partial=False. Note that this does not load optimizer states.

Note

With tensor parallelism, in general, the state_dict must be translated between
the original model implementation and the DistributedModel implementation.
Optionally, you can provide the state_dict translation function as an argument to the
smdistributed.modelparallel.torch.resume_from_checkpoint. However,
for the section called “Supported Models Out of the Box”, the library takes care of this
translation automatically.

The following code shows an example of how to use the checkpoint APIs for fully checkpointing a
PyTorch model trained with model parallelism.

import smdistributed.modelparallel.torch as smp
model = ...
model = smp.DistributedModel(model)
optimizer = ...
optimizer = smp.DistributedOptimizer(optimizer)
user_content = ...  # additional custom data
checkpoint_path = "/opt/ml/checkpoint/model_parallel"

# Save a checkpoint.
smp.save_checkpoint(
    path=checkpoint_path,
    tag=f"total_steps{total_steps}"
    partial=False,
    model=model,
    optimizer=optimizer,
    user_content=user_content
    num_kept_partial_checkpoints=5
)

# Load a checkpoint.
# This automatically loads the most recently saved checkpoint.
smp_checkpoint = smp.resume_from_checkpoint(
    path=checkpoint_path,
    partial=False
)

Checkpointing a distributed PyTorch model (for the SageMaker model parallelism library between v1.6.0 and v1.9.0)

The SageMaker model parallelism library provides Python functions for saving partial or full checkpoints for training jobs with tensor parallelism. The following procedure shows how to use `smp.save()` and `smp.load()` to save and load a checkpoint when you use tensor parallelism.

Note

This checkpointing method is recommended if you use PyTorch, the section called “Tensor Parallelism”, and the SageMaker model parallelism library between v1.6.0 and v1.9.0.

1. Prepare a model object and wrap it with the library's wrapper function `smp.DistributedModel()`.  

   ```python
   model = MyModel(...)  
   ```
2. Prepare an optimizer for the model. A set of model parameters is an iterable argument required by optimizer functions. To prepare a set of model parameters, you must process `model.parameters()` to assign unique IDs to individual model parameters.

If there are parameters with duplicated IDs in the model parameter iterable, loading the checkpointed optimizer state fails. To create an iterable of model parameters with unique IDs for your optimizer, see the following:

```python
unique_params = []
unique_params_set = set()
for p in model.parameters():
    if p not in unique_params_set:
        unique_params.append(p)
        unique_params_set.add(p)
del unique_params_set
optimizer = MyOpt(unique_params, ...)
```

3. Wrap the optimizer using the library's wrapper function `smp.DistributedOptimizer()`.

```python
optimizer = smp.DistributedOptimizer(optimizer)
```

4. Save the model and the optimizer state using `smp.save()`. Depending on how you want to save checkpoints, choose one of the following two options:

- **Option 1**: Save a partial model on each `mp_rank` for a single `MP_GROUP`.

```python
model_dict = model.local_state_dict() # save a partial model
opt_dict = optimizer.local_state_dict() # save a partial optimizer state
# Save the dictionaries at rdp_rank 0 as a checkpoint
if smp.rdp_rank() == 0:
    smp.save(
        {"model_state_dict": model_dict, "optimizer_state_dict": opt_dict},
        f"/checkpoint.pt",
        partial=True,
    )
```

With tensor parallelism, the library saves checkpointed files named in the following format: `checkpoint.pt_{pp_rank}_{tp_rank}`.
Note

With tensor parallelism, make sure you set the if statement as

```python
if smp.rdp_rank() == 0
```

instead of

```python
if smp.dp_rank() == 0
```

When the optimizer state is sharded with tensor parallelism, all reduced-data parallel ranks must save their own partition of the optimizer state. Using a wrong if statement for checkpointing might result in a stalling training job. For more information about using

```python
if smp.dp_rank() == 0
```

without tensor parallelism, see General Instruction for Saving and Loading in the SageMaker Python SDK documentation.

• Option 2: Save the full model.

```python
if smp.rdp_rank() == 0:
    model_dict = model.state_dict(gather_to_rank0=True) # save the full model
if smp.rank() == 0:
    smp.save(
        {"model_state_dict": model_dict},
        "/checkpoint.pt",
        partial=False,
    )
```

Note

Consider the following for full checkpointing:

• If you set `gather_to_rank0=True`, all ranks other than 0 return empty dictionaries.

• For full checkpointing, you can only checkpoint the model. Full checkpointing of optimizer states is currently not supported.

• The full model only needs to be saved at `smp.rank() == 0`.

5. Load the checkpoints using `smp.load()`. Depending on how you checkpointed in the previous step, choose one of the following two options:

• Option 1: Load the partial checkpoints.

```python
checkpoint = smp.load("/checkpoint.pt", partial=True)
model.load_state_dict(checkpoint["model_state_dict"], same_partition_load=False)
```
You can set `same_partition_load=True` in `model.load_state_dict()` for a faster load, if you know that the partition will not change.

- **Option 2**: Load the full checkpoints.

```python
if smp.rdp_rank() == 0:
    checkpoint = smp.load("/checkpoint.pt", partial=False)
    model.load_state_dict(checkpoint["model_state_dict"])  
```

The `if smp.rdp_rank() == 0` condition is not required, but it can help avoid redundant loading among different MP_GROUPS. Full checkpointing optimizer state dict is currently not supported with tensor parallelism.

### Checkpointing a distributed TensorFlow model

To save a TensorFlow model while training with model parallelism, use the following functions provided by the SageMaker model parallelism library.

- `smdistributed.modelparallel.tensorflow.DistributedModel.save_model`
- `smdistributed.modelparallel.tensorflow.CheckpointManager`

### Fine-tuning a distributed model

The fine-tuning needs to be configured in your training script. The following code snippet shows an example structure of a training script using the `AutoModelForCausalLM` class of Hugging Face Transformers with modifications for registering the `smdistributed.model.parallel.torch` modules and settings for fine-tuning.

#### Note

Fine-tuning a distributed transformer (a Transformer model wrapped by `smp.DistributedModel()`) with the `smp.delayed_param_initialization` function activated requires the fine-tuning job to be configured with an FSx for Lustre file system. In cases where you want to fine-tune a large-scale model with the delayed parameter initialization option, you should set up an FSx for Lustre file system.
import argparse
from transformers import AutoModelForCausalLM
import smdistributed.modelparallel
import smdistributed.modelparallel.torch as smp

def parse_args():
    parser = argparse.ArgumentParser()

    # set an arg group for model
    model_grp = parser.add_argument_group(
        title="model", description="arguments to describe model configuration"
    )

    ... # set up numerous args to parse from the configuration dictionary to the script
    for training

    # add arg for activating fine-tuning
    model_grp.add_argument(
        "--fine_tune",
        type=int,
        default=0,
        help="Fine-tune model from checkpoint or pretrained model",
    )

def main():
    """Main function to train GPT.""
    args = parse_args()

    ... # parse numerous args

    if args.fine_tune > 0 and args.delayed_param > 0 and smp.rank() == 0:
        pretrained_model = AutoModelForCausalLM.from_pretrained(
            args.model_name or args.model_dir
        )
        model_state_dict = pretrained_model.state_dict()
        path = os.path.join(args.model_dir, "fullmodel.pt")
        torch.save(model_state_dict, path)

        # create a Transformer model and wrap by smp.model_creation()
        # with options to configure model parallelism parameters offered by SageMaker
        with smp.model_creation(
            tensor_parallelism=smp.tp_size() > 1 or args.use_distributed_transformer > 0,
import torch.nn as nn

zero_init=args.use_distributed_transformer == 0,
dtype=dtype,
distribute_embedding=0 and smp.tp_size() > 1,
use_alibi=0,
attention_in_fp32=0,
fp32_residual_addition=0,
query_key_layer_scaling=0 and args.bf16 < 1,
fused_softmax=0,
fused_dropout=0,
fused_bias_gelu=0,
flash_attention=0,
)

if args.fine_tune > 0 and args.delayed_param == 0:
    model = AutoModelForCausalLM.from_pretrained(
        args.model_name or args.model_dir
    )
else:
    model = AutoModelForCausalLM.from_config(model_config)

# wrap the model by smp.DistributedModel() to apply SageMaker model parallelism
model = smp.DistributedModel(
    model, trace_device="gpu", backward_passes_per_step=args.gradient_accumulation
)

# wrap the optimizer by smp.DistributedOptimizer() to apply SageMaker model parallelism
optimizer= ... # define an optimizer
optimizer = smp.DistributedOptimizer(
    optimizer,
    static_loss_scale=0,
    dynamic_loss_scale=True,
    dynamic_loss_args={"scale_window": 1000, "min_scale": 1, "delayed_shift": 2},
)

# for fine-tuning, use smp.resume_from_checkpoint() to load a pre-trained model
if args.fine_tune > 0 and args.delayed_param > 0:
    smp.resume_from_checkpoint(args.model_dir, tag="fullmodel.pt", partial=False)

For a complete example of training scripts and Jupyter notebooks, see the GPT-2 examples for PyTorch in the SageMaker Examples GitHub repository.
Amazon SageMaker model parallelism library v1 examples

This page provides a list of blogs and Jupyter notebooks that present practical examples of implementing the SageMaker model parallelism (SMP) library v1 to run distributed training jobs on SageMaker.

Blogs and Case Studies

The following blogs discuss case studies about using SMP v1.

- New performance improvements in the Amazon SageMaker model parallelism library, AWS Machine Learning Blog (December 16, 2022)
- Train gigantic models with near-linear scaling using sharded data parallelism on Amazon SageMaker, AWS Machine Learning Blog (October 31, 2022)

Example notebooks

Example notebooks are provided in the SageMaker examples GitHub repository. To download the examples, run the following command to clone the repository and go to `training/distributed_training/pytorch/model_parallel`.

```bash
git clone https://github.com/aws/amazon-sagemaker-examples.git
cd amazon-sagemaker-examples/training/distributed_training/pytorch/model_parallel
```

Note

Clone and run the example notebooks in the following SageMaker ML IDEs.

- SageMaker JupyterLab (available in Studio created after December 2023)
- SageMaker Code Editor (available in Studio created after December 2023)
- Studio Classic (available as an application in Studio created after December 2023)
- SageMaker Notebook Instances

SMP v1 example notebooks for PyTorch

- Train GPT-2 with near-linear scaling using the sharded data parallelism technique in the SageMaker model parallelism library
• Fine-tune GPT-2 with near-linear scaling using sharded data parallelism technique in the SageMaker model parallelism library

• Train GPT-NeoX-20B with near-linear scaling using the sharded data parallelism technique in the SageMaker model parallelism library

• Train GPT-J 6B using the sharded data parallelism and tensor parallelism techniques in the SageMaker model parallelism library

• Train FLAN-T5 with near-linear scaling using sharded data parallelism technique in the SageMaker model parallelism library

• Train Falcon with near-linear scaling using sharded data parallelism technique in the SageMaker model parallelism library

SMP v1 example notebooks for TensorFlow

• CNN with TensorFlow 2.3.1 and the SageMaker model parallelism library

• HuggingFace with TensorFlow Distributed model parallelism library Training on SageMaker

SageMaker Distributed Model Parallelism Best Practices

Use the following guidelines when you run a distributed training job with the SageMaker model parallel library.

Setting Up the Right Configuration for a Given Model

When scaling up a model, we recommend you to go over the following list in order. Each list item discusses the advantage of using the library's techniques along with the tradeoffs that might arise.

Tip

If a model can fit well using a subset of the library's features, adding more model parallelism or memory saving features does not usually improve performance.

Using large GPU instance types

• In the realm of model parallelism, it is best to use powerful instances with large GPU memories to handle overhead from model parallelism operations such as partitioning models across multiple GPUs. We recommend using ml.p4d or ml.p3dn instances for training large DL
models. These instances are also equipped with Elastic Fabric Adapter (EFA), which provides higher network bandwidth and enables large-scale training with model parallelism.

**Sharding optimizer state**

- The impact of sharding optimizer state depends on the number of data parallel ranks. Typically, a higher degree of data parallelism (proportional to the size of compute node) can improve the efficiency of memory usage.

When you want to downsize a cluster, make sure you check the optimizer state sharding configuration. For example, a large DL model with optimizer state sharding that fits on a compute cluster with 16 GPUs (for example, two P4d or P4de instances) might not always fit on a node with 8 GPUs (for example, a single P4d or P4de instance). This is because the combined memory of 8 GPUs is lower than the combined memory of 16 GPUs, and the required memory per GPU for sharding over 8 GPUs is also higher than the memory per GPU for sharding over the 16-GPU scenario. As a result, the increased memory requirement might not fit into the smaller cluster.

For more information, see [Optimizer State Sharding](#).

**Activation checkpointing**

- Memory efficiency can be improved by using activation checkpointing for a group of modules. The more you group the modules, the more efficient the memory usage. When checkpointing sequential modules for layers, the `strategy` argument of the `smp.set_activation_checkpointing` function groups the layers together for checkpointing. For example, grouping two or more layers together for checkpointing is more memory efficient than checkpointing one layer at a time, and this trades extra computation time for reduced memory usage.

For more information, see [Activation Checkpointing](#).

**Tensor parallelism**

- The degree of tensor parallelism should be a power of two (2, 4, 8, ..., $2^n$), where the maximum degree must be equal to the number of GPUs per node. For example, if you use a node with 8 GPUs, possible numbers for the degree of tensor parallelism are 2, 4, and 8. We don’t
recommend arbitrary numbers (such as 3, 5, 6, and 7) for the degree of tensor parallelism. When you use multiple nodes, misconfiguring the degree of tensor parallelism might result in running tensor parallelism across the nodes; this adds significant overhead from communication of activations across the nodes and can become computationally expensive.

For more information, see Tensor Parallelism.

**Pipeline parallelism across nodes**

- You can run pipeline parallelism both within a single node and across multiple nodes. When you use pipeline parallelism in combination with tensor parallelism, we recommend running pipeline parallelism across multiple nodes and keeping tensor parallelism within individual nodes.

- Pipeline parallelism comes with the following three knobs: microbatches, active_micobatches, and prescaled_batch.

- When you use tensor parallelism with pipeline parallelism, we recommend activating prescaled_batch so that the batch size per model parallel group can be increased for efficient pipelining. With prescaled_batch activated, the batch size set in the training script becomes tp_size times the batch size set for each rank without prescaled_batch.

- Increasing the number of microbatches helps achieve efficient pipelining and better performance. Note that the effective microbatch size is the batch size divided by number of microbatches. If you increase the number of microbatches while keeping batch size constant, each microbatch processes fewer samples.

- The number of active_micobatches is the maximum number of microbatches that are simultaneously in process during pipelining. For each active microbatch in process, its activations and gradients take up GPU memory. Therefore, increasing active_micobatches takes up more GPU memory.

- If both GPU and GPU memory are underutilized, increase active_micobatches for better parallelization during pipelining.

- For more information about how to use tensor parallelism with pipeline parallelism, see Tensor parallelism combined with pipeline parallelism.

- To find descriptions of the aforementioned parameters, see Parameters for smdistributed in the SageMaker Python SDK documentation.
Offloading activations to CPU

- Make sure that this is used in combination with activation checkpointing and pipeline parallelism. To ensure that the offloading and preloading happen in the background, specify a value greater than 1 to the microbatches parameter.

- When offloading activations, you might be able to increase active_microbatches and sometimes match with the total number of microbatches. This depends on which modules are checkpointed and how the model is partitioned.

For more information, see Activation Offloading.

Reference configurations

The SageMaker model parallelism training team provides the following reference points based on experiments with the GPT-2 model, the sequence length of 512, and the vocabulary size of 50,000.

<table>
<thead>
<tr>
<th>The number of model parameters</th>
<th>Instance type</th>
<th>Pipeline parallelism</th>
<th>Tensor parallelism</th>
<th>Optimizer state sharding</th>
<th>Activation checkpointing</th>
<th>Prescaled batch</th>
<th>Batch size</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 billion</td>
<td>ml.p4d.2xlarge</td>
<td>1</td>
<td>4</td>
<td>True</td>
<td>Each transformer layer</td>
<td>True</td>
<td>batch_size=40</td>
</tr>
<tr>
<td>30 billion</td>
<td>ml.p4d.2xlarge</td>
<td>1</td>
<td>8</td>
<td>True</td>
<td>Each transformer layer</td>
<td>True</td>
<td>batch_size=32</td>
</tr>
<tr>
<td>60 billion</td>
<td>ml.p4d.2xlarge</td>
<td>2</td>
<td>8</td>
<td>True</td>
<td>Each transformer layer</td>
<td>True</td>
<td>batch_size=56, microbatches=4, active_microbatches=2</td>
</tr>
</tbody>
</table>
You can extrapolate from the preceding configurations to estimate GPU memory usage for your model configuration. For example, if you increase the sequence length for a 10-billion-parameter model or increase the size of the model to 20 billion, you might want to lower batch size first. If the model still doesn’t fit, try increasing the degree of tensor parallelism.

**Modifying Your Training Script**

- Before you use the SageMaker model parallel library's features in your training script, review The SageMaker Distributed Model Parallelism Library Configuration Tips and Pitfalls.

- To launch a training job faster, use the SageMaker local mode. This helps you quickly run a training job locally on a SageMaker notebook instance. Depending on the scale of the ML instance on which your SageMaker notebook instance is running, you might need to adjust the size of your model by changing the model configurations, such as the hidden width, number of transformer layers, and attention heads. Validate if the reduced model runs well on the notebook instance before using a large cluster for training the full model.

**Monitoring and Logging a Training Job Using the SageMaker Console and Amazon CloudWatch**

To monitor system-level metrics such as CPU memory utilization, GPU memory utilization, and GPU utilization, use visualization provided through the SageMaker console.

1. In the left navigation pane, choose Training.

2. Choose Training jobs.

3. In the main pane, choose the training job name for which you want to see more details.

4. Browse the main pane and find the Monitor section to see the automated visualization.

5. To see training job logs, choose View logs in the Monitor section. You can access the distributed training job logs of the training job in CloudWatch. If you launched multi-node distributed training, you should see multiple log streams with tags in the format of algo-n-1234567890. The algo-1 log stream tracks training logs from the main (0th) node.

For more information, see Monitor and Analyze Training Jobs Using Amazon CloudWatch Metrics.

**Permissions**

To run a SageMaker training job with model parallelism or the SageMaker distributed training example notebooks, make sure you have the right permissions in your IAM role, such as the following:
To use FSx for Lustre, add AmazonFSxFullAccess.

To use Amazon S3 as a data channel, add AmazonS3FullAccess.

To use Docker, build your own container, and push it to Amazon ECR, add AmazonEC2ContainerRegistryFullAccess.

To have a full access to use the entire suite of SageMaker features, add AmazonSageMakerFullAccess.

The SageMaker Distributed Model Parallelism Library Configuration Tips and Pitfalls

Review the following tips and pitfalls before using Amazon SageMaker's model parallelism library. This list includes tips that are applicable across frameworks. For TensorFlow and PyTorch specific tips, see Modify a TensorFlow training script and Modify a PyTorch Training Script, respectively.

**Batch Size and Number of Microbatches**

- The library is most efficient when the batch size is increased. For use cases where the model fits within a single device, but can only be trained with a small batch size, batch size can and should be increased after the library is integrated. Model parallelism saves memory for large models, enabling you to train using batch sizes that previously did not fit in memory.

- Choosing a number of microbatches that is too small or too large can lower performance. The library executes each microbatch sequentially in each device, so microbatch size (batch size divided by number of microbatches) must be large enough to fully utilize each GPU. At the same time, pipeline efficiency increases with the number of microbatches, so striking the right balance is important. Typically, a good starting point is to try 2 or 4 microbatches, increasing the batch size to the memory limit, and then experiment with larger batch sizes and numbers of microbatches. As the number of microbatches is increased, larger batch sizes might become feasible if an interleaved pipeline is used.

- Your batch size must be always divisible by the number of microbatches. Note that depending on the size of the dataset, sometimes the last batch of every epoch can be of a smaller size than the rest, and this smaller batch needs to be divisible by the number of microbatches as well. If it is not, you can set drop_remainder=True in the tf.Dataset.batch() call (in TensorFlow), or set drop_last=True in DataLoader (in PyTorch), so that this last, small batch is not used. If you are using a different API for the data pipeline, you might need to manually skip the last batch whenever it is not divisible by the number of microbatches.
Manual Partitioning

• If you use manual partitioning, be mindful of the parameters that are consumed by multiple operations and modules in your model, such as the embedding table in transformer architectures. Modules that share the same parameter must be placed in the same device for correctness. When auto-partitioning is used, the library automatically enforces this constraint.

Data Preparation

• If the model takes multiple inputs, make sure you seed the random operations in your data pipeline (e.g., shuffling) with `smp.dp_rank()`. If the dataset is being deterministically sharded across data parallel devices, make sure that the shard is indexed by `smp.dp_rank()`. This is to make sure that the order of the data seen on all ranks that form a model partition is consistent.

Returning Tensors from `smp.DistributedModel`

• Any tensor that is returned from the `smp.DistributedModel.call` (for TensorFlow) or `smp.DistributedModel.forward` (for PyTorch) function is broadcast to all other ranks, from the rank that computed that particular tensor. As a result, any tensor that is not needed outside the call and forward methods (intermediate activations, for example) should not be returned, as this causes needless communication and memory overhead and hurts performance.

The `@smp.step` Decorator

• If an `smp.step`-decorated function has a tensor argument that does not have a batch dimension, the argument name must be provided in the `non_split_inputs` list when calling `smp.step`. This prevents the library from attempting to split the tensor into microbatches. For more information see `smp.step` in the API documentation.

Delaying Parameter Initialization

For very large models over 100 billion parameters, weight initialization through the CPU memory might result in an out-of-memory error. To get around this, the library offers `smp.delay_param_initialization` context manager. This delays the physical allocation of parameters until they move to GPU during the first execution of a `smp.step`-decorated function. This avoids unnecessary memory usage of the CPU during the initialization of training. Use the context manager when you create a model object as shown in the following code.
with smp.delay_param_initialization(enabled=True):
    model = MyModel()

Tensor Parallelism for PyTorch

- If you are using a seed for deterministic results, set the seed based on smp.dp_rank() (for example, torch.manual_seed(42 + smp.dp_rank())). If you do not do this, different partitions of an nn.Parameter are initialized in the same way, impacting convergence.

- SageMaker's model parallelism library uses NCCL to implement collectives needed for the distribution of the modules. Especially for smaller models, if too many NCCL calls are scheduled on the GPU at the same time, memory usage might increase because of additional space used by NCCL. To counteract this, smp throttles the NCCL calls so that the number of ongoing NCCL operations at any given time is less than or equal to a given limit. The default limit is 8, but this can be adjusted using the environment variable SMP_NCCL_THROTTLE_LIMIT. If you observe more memory usage than you expect while using tensor parallelism, you can try reducing this limit. However, choosing a limit that is too small might cause throughput loss. To disable throttling altogether, you can set SMP_NCCL_THROTTLE_LIMIT=-1.

- The following identity, which holds when the degree of tensor parallelism is 1, does not hold when the degree of tensor parallelism is greater than 1: smp.mp_size() * smp.dp_size() == smp.size(). This is because the tensor parallel group is part of both the model parallelism group and the data parallelism group. If your code has existing references to mp_rank, mp_size, MP_GROUP, and so on, and if you want to work with only the pipeline parallel group, you might need to replace the references with smp.pp_size(). The following identities are always true:
  - smp.mp_size() * smp.rdp_size() == smp.size()
  - smp.pp_size() * smp.dp_size() == smp.size()
  - smp.pp_size() * smp.tp_size() * smp.rdp_size() == smp.size()

- Since the smp.DistributedModel wrapper modifies the model parameters when tensor parallelism is enabled, the optimizer should be created after calling smp.DistributedModel, with the distributed parameters. For example, the following does not work:

  ```python
  ## WRONG
  model = MyModel()
  optimizer = SomeOptimizer(model.parameters())
  model = smp.DistributedModel(model)  # optimizer now has outdated parameters!
  ```
Instead, the optimizer should be created with the parameters of the `smp.DistributedModel` as follows:

```python
## CORRECT
model = smp.DistributedModel(MyModel())
optimizer = SomeOptimizer(model.optimizers())
```

- When a module is replaced with its distributed counterpart through tensor parallelism, the distributed module does not inherit its weights from the original module, and initializes new weights. This means that, for instance, if the weights need to be initialized in a particular call (for example, through a `load_state_dict` call), this needs to happen after the `smp.DistributedModel` call, once the module distribution takes place.

- When accessing the parameters of distributed modules directly, note that the weight does not have the same shape as the original module. For instance,

```python
with smp.tensor_parallelism():
    linear = nn.Linear(60, 60)

    # will pass
    assert tuple(linear.weight.shape) == (60, 60)

distributed_linear = smp.DistributedModel(linear)

    # will fail. the number of input channels will have been divided by smp.tp_size()
    assert tuple(distributed_linear.module.weight.shape) == (60, 60)
```

- Using `torch.utils.data.distributed.DistributedSampler` is strongly recommended for tensor parallelism. This ensures that every data parallel rank receives the same number of data samples, which prevents hangs that might result from different `dp_ranks` taking a different number of steps.

- If you use the `join` API of PyTorch's `DistributedDataParallel` class to handle cases in which different data parallel ranks have different numbers of batches, you still need to make sure that ranks that are in the same `TP_GROUP` have the same number of batches; otherwise the communication collectives used in distributed execution of modules may hang. Ranks that are in different `TP_GROUPS` can have different numbers of batches, as long as `join` API is used.

- If you want to checkpoint your model and use tensor parallelism, consider the following:
To avoid stalling and race conditions while saving and loading models when you use tensor parallelism, make sure you call appropriate functions from the following model and optimizer states inside a reduced-data parallelism rank.

If you are transitioning an existing pipeline parallel script and enabling tensor parallel for the script, ensure that you modify any `if smp.dp_rank() == 0` block used for saving and loading with `if smp.rdp_rank() == 0` blocks. Otherwise, it might cause your training job to stall.

For more information about checkpointing a model with tensor parallelism, see the section called “Checkpointing a distributed model”.

Model Parallel Troubleshooting

If you run into an error, you can use the following list to try to troubleshoot your training job. If the problem persists, contact AWS Support.

Topics

- Considerations for Using SageMaker Debugger with the SageMaker Model Parallelism Library
- Saving Checkpoints
- Convergence Using Model Parallel and TensorFlow
- Stalling or Crashing Distributed Training Jobs
- Receiving NCCL Error for a PyTorch Training Job
- Receiving RecursionError for a PyTorch Training Job

Considerations for Using SageMaker Debugger with the SageMaker Model Parallelism Library

SageMaker Debugger is not available for the SageMaker model parallelism library. Debugger is enabled by default for all SageMaker TensorFlow and PyTorch training jobs, and you might see an error that looks like the following:

```
FileNotFoundError: [Errno 2] No such file or directory: '/opt/ml/checkpoints/metadata.json.sagemaker-uploading'
```

To fix this issue, disable Debugger by passing `debugger_hook_config=False` when creating a framework estimator as shown in the following example.
bucket=sagemaker.Session().default_bucket()
base_job_name="sagemaker-checkpoint-test"
checkpoint_in_bucket="checkpoints"

# The S3 URI to store the checkpoints
checkpoint_s3_bucket="s3://{}/{}/{}".format(bucket, base_job_name, checkpoint_in_bucket)
estimator = TensorFlow(
    ...
    distribution={"smdistributed": {"modelparallel": { "enabled": True }}},
    checkpoint_s3_uri=checkpoint_s3_bucket,
    checkpoint_local_path="/opt/ml/checkpoints",
    debugger_hook_config=False
)

**Saving Checkpoints**

You might run into the following error when saving checkpoints of a large model on SageMaker:

```
InternalServerError: We encountered an internal error. Please try again
```

This could be caused by a SageMaker limitation while uploading the local checkpoint to Amazon S3 during training. To disable checkpointing in SageMaker, use the following example to explicitly upload the checkpoints.

If you run into the preceding error, do not use `checkpoint_s3_uri` with the SageMaker `estimator` call. While saving checkpoints for larger models, we recommend saving checkpoints to a custom directory and passing the same to the helper function (as a `local_path` argument).

```python
import os
def aws_s3_sync(source, destination):
    """aws s3 sync in quiet mode and time profile""
    import time, subprocess
    cmd = ["aws", "s3", "sync", "--quiet", source, destination]
    print(f"Syncing files from {source} to {destination}""
    start_time = time.time()
    p = subprocess.Popen(cmd, stdout=subprocess.PIPE, stderr=subprocess.PIPE)
    p.wait()
    end_time = time.time()
```
print("Time Taken to Sync: ", (end_time-start_time))
return

def sync_local_checkpoints_to_s3(local_path="/opt/ml/checkpoints",
s3_uri=os.path.dirname(os.path.dirname(os.getenv('SM_MODULE_DIR', '')))+'/checkpoints'):
    """ sample function to sync checkpoints from local path to s3 """

    import boto3
    #check if local path exists
    if not os.path.exists(local_path):
        raise RuntimeError("Provided local path {local_path} does not exist. Please check")

    #check if s3 bucket exists
    s3 = boto3.resource('s3')
    if not s3_uri.startswith("s3://"):  
        raise ValueError(f"Provided s3 uri {s3_uri} is not valid.")

    s3_bucket = s3_uri.replace('s3://', '').split('/')[0]
    print(f"S3 Bucket: {s3_bucket}")
    try:
        s3.meta.client.head_bucket(Bucket=s3_bucket)
    except Exception as e:
        raise e
    aws_s3_sync(local_path, s3_uri)
    return

def sync_s3_checkpoints_to_local(local_path="/opt/ml/checkpoints",
s3_uri=os.path.dirname(os.path.dirname(os.getenv('SM_MODULE_DIR', '')))+'/checkpoints'):
    """ sample function to sync checkpoints from s3 to local path """

    import boto3
    #try to create local path if it does not exist
    if not os.path.exists(local_path):
        print(f"Provided local path {local_path} does not exist. Creating...")
        try:
            os.makedirs(local_path)
        except Exception as e:
            raise RuntimeError(f"Failed to create {local_path}")

    #check if s3 bucket exists
    s3 = boto3.resource('s3')
if not s3_uri.startswith("s3://"):  
    raise ValueError(f"Provided s3 uri {s3_uri} is not valid.")

s3_bucket = s3_uri.replace('s3://','').split('/')[0]
print(f"S3 Bucket: {s3_bucket}"
try:
    s3.meta.client.head_bucket(Bucket=s3_bucket)
except Exception as e:
    raise e
aws_s3_sync(s3_uri, local_path)
return

Usage of helper functions:

#base_s3_uri - user input s3 uri or save to model directory (default)
#curr_host - to save checkpoints of current host
#iteration - current step/epoch during which checkpoint is saved

# save checkpoints on every node using local_rank
if smp.local_rank() == 0:
    base_s3_uri = os.path.dirname(os.path.dirname(os.getenv('SM_MODULE_DIR', '')))
    curr_host = os.environ['SM_CURRENT_HOST']
    full_s3_uri = f'{base_s3_uri}/checkpoints/{curr_host}/{iteration}'
    sync_local_checkpoints_to_s3(local_path=checkpoint_dir, s3_uri=full_s3_uri)

Convergence Using Model Parallel and TensorFlow

When you use SageMaker multi-node training with TensorFlow and the model parallelism library, the loss may not converge as expected because the order of training input files may be different on each node. This may cause different ranks in the same model parallel group to work on different input files, causing inconsistencies. To prevent this, ensure the input files are ordered the same way in all the ranks before they get converted to TensorFlow datasets. One way to achieve this is to sort the input file names in the training script.

Stalling or Crashing Distributed Training Jobs

If your training job has stalling, crashing, or not responding issues, read the following troubleshooting items to identify what's the cause of the issue. If you need any further support, reach out to the SageMaker distributed training team through AWS Support.

• If you see a distributed training job stalling at the NCCL initialization step, consider the following:
• If you are using one of the EFA-enabled instances (ml.p4d or ml.p3dn instances) with a custom VPC and its subnet, ensure that the security group used has inbound and outbound connections for all ports to and from the same SG. You also generally need outbound connections to any IP as a separate rule (for internet access). To find instructions on how to add inbound and outbound rules for EFA communication, refer to SageMaker distributed training job stalling during initialization.

• If you see a distributed training job stalling when checkpointing the full model, this might be because the state_dict() call on the model or optimizer was not made on all ranks with rdp_rank()==0 (when using tensor parallelism) or dp_rank()==0 (when using only pipeline parallelism). These ranks need to communicate to construct the checkpoint to be saved. Similar stalling issues can also happen when checkpointing partial optimizer if shard_optimizer_state is enabled.

For more information about checkpointing a model with model parallelism, see General Instruction for Saving and Loading and Checkpointing a distributed PyTorch model (for the SageMaker model parallelism library between v1.6.0 and v1.9.0).

• If the training job crashes with a CUDA Out of Memory error, this means that the distributed training configuration needs to be adjusted to fit the model on the GPU cluster. For more information and best practices, see Setting Up the Right Configuration for a Given Model.

• If the training job crashes with an uncorrectable ECC error, this means that one of the GPUs in the cluster has gone bad. If you need technical support, share the job ARN with the AWS team and restart your training job from a checkpoint if possible.

• In rare cases, a job configuration that worked previously but is close to the limits of GPU memory might fail later with a different cluster due to a CUDA Out of Memory error. This could be because some GPU has lower available memory than usual due to ECC errors.

• Network timeout crash might happen when running a multinode job which doesn’t use all GPUs in the node. To get around this, use all GPUs on the node by ensuring that the processes_per_host parameter is set to the number of GPUs in each instance. For example, this is processes_per_host=8 for ml.p3.16xlarge, ml.p3dn.24xlarge, and ml.p4d.24xlarge instances.

• If you find that your training job takes a long time during the data downloading stage, make sure the Amazon S3 path you provided to checkpoint_s3_uri for the SageMaker Estimator class is unique for the current training job. If this path is reused across multiple training jobs running simultaneously, all those checkpoints are uploaded and downloaded to the same Amazon S3 path and might significantly increase checkpoint loading time.
• Use FSx for Lustre when you deal with large data and models.
  • If your dataset is large and fetching it takes a long time, we recommend keeping your dataset in FSx for Lustre.
  • When training models are beyond 10 billion parameters, we recommend using FSx for Lustre for checkpointing.
  • After you create a file system, make sure to wait for the status to become available before starting a training job using it.

**Receiving NCCL Error for a PyTorch Training Job**

If you encountered the following error, it might be due to a process running out of GPU memory.

```plaintext
NCCL error in: ../torch/lib/tensor/core/ProcessGroupNCCL.cpp:825, unhandled system error, NCCL version 2.7.8
ncclSystemError: System call (socket, malloc, munmap, etc) failed.
```

You can resolve this by reducing the batch size or active_microbatches. If auto partitioning is not resulting in a well-balanced partitioning, you might have to consider manual partitioning. For more information, see [Pipeline parallelism across nodes](#).

**Receiving RecursionError for a PyTorch Training Job**

The library does not support calling `super.forward()` inside a module's forward call. If you use `super.forward()`, you might receive the following error message.

```plaintext
RecursionError: maximum recursion depth exceeded
```

To fix the error, instead of calling `super.forward()`, you should call `super._orig_forward()`.

**Distributed computing with SageMaker best practices**

This best practices page presents various flavors of distributed computing for machine learning (ML) jobs in general. The term *distributed computing* in this page encompasses distributed training for machine learning tasks and parallel computing for data processing, data generation, feature engineering, and reinforcement learning. In this page, we discuss about common challenges in distributed computing, and available options in SageMaker Training and SageMaker Processing. For additional reading materials about distributed computing, see [What Is Distributed Computing?](#).
You can configure ML tasks to run in a distributed manner across multiple nodes (instances), accelerators (NVIDIA GPUs, AWS Trainium chips), and vCPU cores. By running distributed computation, you can achieve a variety of goals such as computing operations faster, handling large datasets, or training large ML models.

The following list covers common challenges that you might face when you run an ML training job at scale.

- You need to make decisions on how to distribute computation depending on ML tasks, software libraries you want to use, and compute resources.
- Not all ML tasks are straightforward to distribute. Also, not all ML libraries support distributed computation.
- Distributed computation might not always result in a linear increase in compute efficiency. In particular, you need to identify if data I/O and inter-GPU communication have bottlenecks or cause overhead.
- Distributed computation might disturb numerical processes and change model accuracy. Specifically to data-parallel neural network training, when you change the global batch size while scaling up to a larger compute cluster, you also need to adjust the learning rate accordingly.

SageMaker provides distributed training solutions to ease such challenges for various use cases. Choose one of the following options that best fits your use case.

**Topics**
- **Option 1: Use a SageMaker built-in algorithm that supports distributed training**
- **Option 2: Run a custom ML code in the SageMaker managed training or processing environment**
- **Option 3: Write your own custom distributed training code**
- **Option 4: Launch multiple jobs in parallel or sequentially**

**Option 1: Use a SageMaker built-in algorithm that supports distributed training**

SageMaker provides built-in algorithms that you can use out of the box through the SageMaker console or the SageMaker Python SDK. Using the built-in algorithms, you don’t need to spend time for code customization, understanding science behind the models, or running Docker on provisioned Amazon EC2 instances.
A subset of the SageMaker built-in algorithms support distributed training. To check if the algorithm of your choice supports distributed training, see the **Parallelizable** column in the [Common Information About Built-in Algorithms](#) table. Some of the algorithms support multi-instance distributed training, while the rest of the parallelizable algorithms support parallelization across multiple GPUs in a single instance, as indicated in the **Parallelizable** column.

**Option 2: Run a custom ML code in the SageMaker managed training or processing environment**

SageMaker jobs can instantiate distributed training environment for specific use cases and frameworks. This environment acts as a ready-to-use whiteboard, where you can bring and run your own ML code.

**If your ML code uses a deep learning framework**

You can launch distributed training jobs using the [Deep Learning Containers (DLC)](#) for SageMaker Training, which you can orchestrate either through the dedicated Python modules in the [SageMaker Python SDK](#), or through the SageMaker APIs with [AWS CLI](#), [AWS SDK for Python (Boto3)](#). SageMaker provides training containers for machine learning frameworks, including [PyTorch](#), [TensorFlow](#), [Hugging Face Transformers](#), and [Apache MXNet](#). You have two options to write deep learning code for distributed training.

- **The SageMaker distributed training libraries**

  The SageMaker distributed training libraries propose AWS-managed code for neural network data parallelism and model parallelism. SageMaker distributed training also comes with launcher clients built into the SageMaker Python SDK, and you don’t need to author parallel launch code. To learn more, see [SageMaker’s data parallelism library](#) and [SageMaker’s model parallelism library](#).

- **Open-source distributed training libraries**

  Open source frameworks have their own distribution mechanisms such as [DistributedDataParallelism (DDP) in PyTorch](#) or `tf.distribute` modules in TensorFlow. You can choose to run these distributed training frameworks in the SageMaker-managed framework containers. For example, the sample code for [training MaskRCNN in SageMaker](#) shows how to use both PyTorch DDP in the SageMaker PyTorch framework container and [Horovod](#) in the SageMaker TensorFlow framework container.
SageMaker ML containers also come with MPI preinstalled, so you can parallelize your entry point script using mpi4py. Using the MPI integrated training containers is a great option when you launch a third-party distributed training launcher or write ad-hoc parallel code in the SageMaker managed training environment.

**Notes for data-parallel neural network training on GPUs**

- **Scale to multi-GPU and multi-machine parallelism when appropriate**

  We often run neural network training jobs on multiple-CPU or multiple-GPU instances. Each GPU-based instance usually contains multiple GPU devices. Consequently, distributed GPU computing can happen either within a single GPU instance with multiple GPUs (single-node multi-GPU training), or across multiple GPU instances with multiple GPU cores in each (multi-node multi-GPU training). Single-instance training is easier to write code and debug, and the intra-node GPU-to-GPU throughput is usually faster than the inter-node GPU-to-GPU throughput. Therefore, it is a good idea to scale data parallelism vertically first (use one GPU instance with multiple GPUs) and expand to multiple GPU instances if needed. This might not apply to cases where the CPU budget is high (for example, a massive workload for data pre-processing) and when the CPU-to-GPU ratio of a multi-GPU instance is too low. In all cases, you need to experiment with different combinations of instance types based on your own ML training needs and workload.

- **Monitor the quality of convergence**

  When training a neural network with data parallelism, increasing the number of GPUs while keeping the mini-batch size per GPU constant leads to increasing the size of global mini-batch for the mini-batch stochastic gradient descent (MSGD) process. The size of mini-batches for MSGD is known to impact the descent noise and convergence. For properly scaling while preserving accuracy, you need to adjust other hyperparameters such as the learning rate [Goyal et al. (2017)].

- **Monitor I/O bottlenecks**

  As you increase the number of GPUs, the throughput for reading and writing storage should also increase. Make sure that your data source and pipeline don’t become bottlenecks.

- **Modify your training script as needed**

  Training scripts written for single-GPU training must be modified for multi-node multi-GPU training. In most data parallelism libraries, script modification is required to do the following.
  - Assign batches of training data to each GPU.
• Use an optimizer that can deal with gradient computation and parameter updates across multiple GPUs.

• Assign responsibility of checkpointing to a specific host and GPU.

If your ML code involves tabular data processing

PySpark is a Python frontend of Apache Spark, which is an open-source distributed computing framework. PySpark has been widely adopted for distributed tabular data processing for large-scale production workloads. If you want to run tabular data processing code, consider using the SageMaker Processing PySpark containers and running parallel jobs. You can also run data processing jobs in parallel using SageMaker Training and SageMaker Processing APIs in Amazon SageMaker Studio Classic, which is integrated with Amazon EMR and AWS Glue.

Option 3: Write your own custom distributed training code

When you submit a training or processing job to SageMaker, SageMaker Training and SageMaker Processing APIs launch Amazon EC2 compute instances. You can customize training and processing environment in the instances by running your own Docker container or installing additional libraries in the AWS managed containers. For more information about Docker with SageMaker Training, see Adapting your own Docker container to work with SageMaker and Create a container with your own algorithms and models. For more information about Docker with SageMaker Processing, see Use Your Own Processing Code.

Every SageMaker training job environment contains a configuration file at /opt/ml/input/config/resourceconfig.json, and every SageMaker processing job environment contains a similar configuration file at /opt/ml/config/resourceconfig.json. Your code can read this file to find hostnames and establish inter-node communications. To learn more, including the schema of the JSON file, see Distributed Training Configuration and How Amazon SageMaker Processing Configures Your Processing Container. You can also install and use third-party distributed computing libraries such as Ray or DeepSpeed in SageMaker.

You can also use SageMaker Training and SageMaker Processing to run custom distributed computations that do not require inter-worker communication. In the computing literature, those tasks are often described as embarrassingly parallel or share-nothing. Examples include parallel processing of data files, training models in parallel on different configurations, or running batch inference on a collection of records. You can trivially parallelize such share-nothing use cases with Amazon SageMaker. When you launch a SageMaker Training or SageMaker Processing job on a
cluster with multiple nodes, SageMaker by default replicates and launches your training code (in Python or Docker) on all the nodes. Tasks requiring random spread of input data across such multiple nodes can be facilitated by setting `S3DataDistributionType=ShardedByS3Key` in the data input configuration of the SageMaker TrainingInput API.

**Option 4: Launch multiple jobs in parallel or sequentially**

You can also distribute an ML compute workflow into smaller parallel or sequential compute tasks, each represented by its own SageMaker Training or SageMaker Processing job. Splitting a task into multiple jobs can be beneficial for the following situations or tasks:

- When you have specific data channels and metadata entries (such as hyperparameters, model configuration, or instance types) for each sub-tasks.
- When you implement retry steps at a sub-task level.
- When you vary the configuration of the sub-tasks over the course of the workload, such as when training on increasing batch sizes.
- When you need to run an ML task that takes longer than the maximum training time allowed for a single training job (28 days maximum).
- When different steps of a compute workflow require different instance types.

For the specific case of hyperparameter search, use [SageMaker Automated Model Tuning](https://docs.aws.amazon.com/sagemaker/latest/dg/model-optimization-baselines.html). SageMaker Automated Model Tuning is a serverless parameter search orchestrator that launches multiple training jobs on your behalf, according to a search logic that can be random, Bayesian, or HyperBand.

Additionally, to orchestrate multiple training jobs, you can also consider workflow orchestration tools, such as [SageMaker Pipelines](https://docs.aws.amazon.com/sagemaker/latest/dg/pipelines.html), [AWS Step Functions](https://docs.aws.amazon.com/stepfunctions/latest/dg/), and Apache Airflow supported by [Amazon Managed Workflows for Apache Airflow (MWAA)](https://aws.amazon.com/mwaa/) and [SageMaker Workflows](https://docs.aws.amazon.com/sagemaker/latest/dg/workflows.html).

**Amazon SageMaker Training Compiler**

Use Amazon SageMaker Training Compiler to train deep learning (DL) models faster on scalable GPU instances managed by SageMaker.

**What Is SageMaker Training Compiler?**

State-of-the-art deep learning (DL) models consist of complex multi-layered neural networks with billions of parameters that can take thousands of GPU hours to train. Optimizing such
models on training infrastructure requires extensive knowledge of DL and systems engineering; this is challenging even for narrow use cases. Although there are open-source implementations of compilers that optimize the DL training process, they can lack the flexibility to integrate DL frameworks with some hardware such as GPU instances.

SageMaker Training Compiler is a capability of SageMaker that makes these hard-to-implement optimizations to reduce training time on GPU instances. The compiler optimizes DL models to accelerate training by more efficiently using SageMaker machine learning (ML) GPU instances. SageMaker Training Compiler is available at no additional charge within SageMaker and can help reduce total billable time as it accelerates training.

SageMaker Training Compiler is integrated into the AWS Deep Learning Containers (DLCs). Using the SageMaker Training Compiler–enabled AWS DLCs, you can compile and optimize training jobs on GPU instances with minimal changes to your code. Bring your deep learning models to SageMaker and enable SageMaker Training Compiler to accelerate the speed of your training job on SageMaker ML instances for accelerated computing.

**How It Works**

SageMaker Training Compiler converts DL models from their high-level language representation to hardware-optimized instructions. Specifically, SageMaker Training Compiler applies graph-level optimizations, dataflow-level optimizations, and backend optimizations to produce an optimized model that efficiently uses hardware resources. As a result, you can train your models faster than when you train them without compilation.

It is a two-step process to activate SageMaker Training Compiler for your training job:
1. Bring your own DL script and, if needed, adapt to compile and train with SageMaker Training Compiler. To learn more, see Bring Your Own Deep Learning Model.

2. Create a SageMaker estimator object with the compiler configuration parameter using the SageMaker Python SDK.
   
a. Turn on SageMaker Training Compiler by adding 
   
   ```python
   compiler_config=TrainingCompilerConfig()
   ```
   
to the SageMaker estimator class.

b. Adjust hyperparameters (batch_size and learning_rate) to maximize the benefit that SageMaker Training Compiler provides.

Compilation through SageMaker Training Compiler changes the memory footprint of the model. Most commonly, this manifests as a reduction in memory utilization and a consequent increase in the largest batch size that can fit on the GPU. In some cases, the compiler intelligently promotes caching which leads to a decrease in the largest batch size that can fit on the GPU. Note that if you want to change the batch size, you must adjust the learning rate appropriately.

For a reference for batch_size tested for popular models, see Tested Models.

When you adjust the batch size, you also have to adjust the learning_rate appropriately. For best practices for adjusting the learning rate along with the change in batch size, see the section called “Best Practices and Considerations”.

c. By running the estimator.fit() class method, SageMaker compiles your model and starts the training job.

For instructions on how to launch a training job, see Enable SageMaker Training Compiler.

SageMaker Training Compiler does not alter the final trained model, while allowing you to accelerate the training job by more efficiently using the GPU memory and fitting a larger batch size per iteration. The final trained model from the compiler-accelerated training job is identical to the one from the ordinary training job.

### Tip

SageMaker Training Compiler only compiles DL models for training on supported GPU instances managed by SageMaker. To compile your model for inference and deploy it to run anywhere in the cloud and at the edge, use SageMaker Neo compiler.
Topics

- Supported Frameworks, AWS Regions, Instance Types, and Tested Models
- Bring Your Own Deep Learning Model
- Enable SageMaker Training Compiler
- SageMaker Training Compiler Example Notebooks and Blogs
- SageMaker Training Compiler Best Practices and Considerations
- SageMaker Training Compiler FAQ
- SageMaker Training Compiler Troubleshooting
- Amazon SageMaker Training Compiler Release Notes

**Supported Frameworks, AWS Regions, Instance Types, and Tested Models**

Before using SageMaker Training Compiler, check if your framework of choice is supported, the instance types are available in your AWS account, and your AWS account is in one of the supported AWS Regions.

ℹ️ **Note**

SageMaker Training Compiler is available in the SageMaker Python SDK v2.70.0 or later.

**Supported Frameworks**

SageMaker Training Compiler supports the following deep learning frameworks and is available through AWS Deep Learning Containers.

**Topics**

- PyTorch
- TensorFlow
## PyTorch

<table>
<thead>
<tr>
<th>Framework</th>
<th>Framework version</th>
<th>Deep Learning Container URI</th>
<th>Extendable for Docker customization</th>
</tr>
</thead>
<tbody>
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**TensorFlow**

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### Supported Frameworks, AWS Regions, Instance Types, and Tested Models

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For more information, see [Available Images](#) in the [AWS Deep Learning Containers GitHub repository](#).

**AWS Regions**

The [SageMaker Training Compiler Containers](#) are available in the AWS Regions where [AWS Deep Learning Containers](#) are in service except the China regions.
Supported Instance Types

SageMaker Training Compiler is tested on and supports the following ML instance types.

- P4 instances
- P3 instances
- G4dn instances
- G5 instances

For specs of the instance types, see the **Accelerated Computing** section in the [Amazon EC2 Instance Types page](#). For information about instance pricing, see [Amazon SageMaker Pricing](#).

If you encountered an error message similar to the following, follow the instructions at [Request a service quota increase for SageMaker resources](#).

```
ResourceLimitExceeded: An error occurred (ResourceLimitExceeded) when calling the CreateTrainingJob operation: The account-level service limit 'ml.p3dn.24xlarge for training job usage' is 0 Instances, with current utilization of 0 Instances and a request delta of 1 Instances. Please contact AWS support to request an increase for this limit.
```

Tested Models

The following table includes a list of the models that have been tested with SageMaker Training Compiler. For reference, the largest batch size that is able to fit into memory is also included alongside other training parameters. SageMaker Training Compiler can change the memory footprint of the model training process; as a result, a larger batch size can often be used during the training process, further decreasing total training time. In some cases, SageMaker Training Compiler intelligently promotes caching which leads to a decrease in the largest batch size that can fit on the GPU. You must retune your model hyperparameters and find an optimal batch size for your case. To save time, use the following reference tables to look up a batch size that can be a good starting point for your use case.

---

**Note**

The batch sizes are local batch size that fit into each individual GPU in the respective instance type. You should also adjust the learning rate when changing the batch size.
## Natural language processing (NLP) models

The following models are tested for training jobs for all combinations of single-node and multi-node with single or multi GPU cores and Automatic Mixed Precision (AMP) as indicated.

<table>
<thead>
<tr>
<th>Model</th>
<th>Dataset</th>
<th>Instance type</th>
<th>Precision</th>
<th>Sequence Length</th>
<th>Batch size for native frameworks</th>
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Computer Vision (CV) models

Tested using TensorFlow Model Garden with Automatic Mixed Precision (AMP) as indicated.

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PyTorch 1.12.0

Natural language processing (NLP) models

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# Single-node/multi-node single-GPU/multi-GPU

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<th>Precision</th>
<th>Sequence Length</th>
<th>Batch size for native frameworks</th>
<th>Batch size for SageMaker Training Compiler</th>
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## TensorFlow 2.11.0

### Computer Vision (CV) models

Tested using [TensorFlow Model Garden](https://github.com/tensorflow/models) with Automatic Mixed Precision (AMP) as indicated.
### Single/multi-node single/multi-GPU

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<th>Dataset</th>
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<th>Batch size for native frameworks</th>
<th>Batch size for SageMaker Training Compiler</th>
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### Natural Language Processing (NLP) models

Tested using [Transformer models](#) with Sequence_len=128 and Automatic Mixed Precision (AMP) as indicated.
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### TensorFlow 2.10.0

#### Computer Vision (CV) models

Tested using [TensorFlow Model Garden](https://github.com/tensorflow/models) with Automatic Mixed Precision (AMP) as indicated.

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### Natural Language Processing (NLP) models

Tested using [Transformer models](#) with `Sequence_Len=128` and Automatic Mixed Precision (AMP) as indicated.

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**TensorFlow 2.9.1**

Tested using [TensorFlow Model Garden](https://github.com/tensorflow/models) with Automatic Mixed Precision (AMP).
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* The batch sizes marked with the asterisk symbol (*) indicate the largest batch size tested by the SageMaker Training Compiler developer team. For the marked cells, the instance may be able to fit a larger batch size than what is indicated.

#### Transformers 4.21.1 with PyTorch 1.11.0

Tested with Sequence Len=512 and Automatic Mixed Precision (AMP).

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**Transformers 4.17.0 with PyTorch 1.10.2**

Tested with `Sequence_Len=512` and Automatic Mixed Precision (AMP).

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### Transformers 4.11.0 with PyTorch 1.9.0

Tested with Sequence Len=512 and Automatic Mixed Precision (AMP).
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### Single-node single-GPU

<table>
<thead>
<tr>
<th>Model</th>
<th>Instance type</th>
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<th>Batch size for Training Compiler</th>
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### Single-node multi-GPU

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<th>Model</th>
<th>Instance type</th>
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## Transformers 4.17.0 with TensorFlow 2.6.3

Tested with Sequence.Len=128 and Automatic Mixed Precision (AMP).

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<td>Model</td>
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### Model Instance type | Batch size for native frameworks | Batch size for Training Compiler
--- | --- | ---
microsoft_mpnet-base | ml.p3.8xlarge | 130 | 92
roberta-base | ml.g4dn.16xlarge | 108 | 64
roberta-base | ml.g5.4xlarge | 176 | 142
roberta-base | ml.p3.2xlarge | 118 | 100
roberta-base | ml.p3.8xlarge | 112 | 88

**Transformers 4.11.0 with TensorFlow 2.5.1**

Tested with Sequence_Len=128 and Automatic Mixed Precision (AMP).

### Single-node single-GPU

<table>
<thead>
<tr>
<th>Model</th>
<th>Instance type</th>
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## Single-node single-GPU

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## Bring Your Own Deep Learning Model

This guide walks you through how to adapt your training script for a compiler-accelerated training job. The preparation of your training script depends on the following:
• Training settings such as single-core or distributed training.
• Frameworks and libraries that you use to create the training script.

Choose one of the following topics depending on the framework you use.

Topics
• PyTorch
• TensorFlow

Note
After you finish preparing your training script, you can run a SageMaker training job using the SageMaker framework estimator classes. For more information, see the previous topic at Enable SageMaker Training Compiler.

PyTorch
Bring your own PyTorch model to SageMaker, and run the training job with SageMaker Training Compiler.

Topics
• PyTorch Models with Hugging Face Transformers

PyTorch Models with Hugging Face Transformers
PyTorch models with Hugging Face Transformers are based on PyTorch's torch.nn.Module API. Hugging Face Transformers also provides Trainer and pretrained model classes for PyTorch to help reduce the effort for configuring natural language processing (NLP) models. After preparing your training script, you can launch a training job using the SageMaker PyTorch or HuggingFace estimator with the SageMaker Training Compiler configuration when you'll proceed to the next topic at Enable SageMaker Training Compiler.

Tip
When you create a tokenizer for an NLP model using Transformers in your training script, make sure that you use a static input tensor shape by specifying padding='max_length'.
Do not use `padding='longest'` because padding to the longest sequence in the batch can change the tensor shape for each training batch. The dynamic input shape can trigger recompilation of the model and might increase total training time. For more information about padding options of the Transformers tokenizers, see [Padding and truncation](#) in the [Hugging Face Transformers documentation](#).

**Topics**

- [Large Language Models Using the Hugging Face Transformers Trainer Class](#)
- [Large Language Models Using PyTorch Directly (without the Hugging Face Transformers Trainer API)](#)

**Large Language Models Using the Hugging Face Transformers Trainer Class**

If you use the transformers library's Trainer class, you don’t need to make any additional changes to your training script. SageMaker Training Compiler automatically compiles your Trainer model if you enable it through the estimator class. The following code shows the basic form of a PyTorch training script with Hugging Face Trainer API.

```python
from transformers import Trainer, TrainingArguments

training_args=TrainingArguments(**kwargs)
trainer=Trainer(args=training_args, **kwargs)
```

**Topics**

- [For single GPU training](#)
- [For distributed training](#)
- [Best Practices to Use SageMaker Training Compiler with Trainer](#)

**For single GPU training**

You don't need to change your code when you use the `transformers.Trainer` class.

**For distributed training**

*PyTorch v1.11.0 and later*
To run distributed training with SageMaker Training Compiler, you must add the following `_mp_fn()` function in your training script and wrap the `main()` function. It redirects the `_mp_fn(index)` function calls from the SageMaker distributed runtime for PyTorch (`pytorchxla`) to the `main()` function of your training script.

```python
def _mp_fn(index):
    main()
```

This function accepts the `index` argument to indicate the rank of the current GPU in the cluster for distributed training. To find more example scripts, see the [Hugging Face Transformers language modeling example scripts](#).

For Transformers v4.17 and before with PyTorch v1.10.2 and before

SageMaker Training Compiler uses an alternate mechanism for launching a distributed training job, and you don't need to make any modification in your training script. Instead, SageMaker Training Compiler requires you to pass a SageMaker distributed training launcher script to the `entry_point` argument and pass your training script to the `hyperparameters` argument in the SageMaker Hugging Face estimator.

Best Practices to Use SageMaker Training Compiler with Trainer

- Make sure that you use SyncFree optimizers by setting the `optim` argument to `adamw_torch_xla` while setting up `transformers.TrainingArguments`. See also [Optimizer](#) in the [Hugging Face Transformers documentation](#).

- Ensure that the throughput of the data processing pipeline is higher than the training throughput. You can tweak the `dataloader_num_workers` and `preprocessing_num_workers` arguments of the `transformers.TrainingArguments` class to achieve this. Typically, these need to be greater than or equal to the number of GPUs but less than the number of CPUs.

After you have completed adapting your training script, proceed to [the section called “Run PyTorch Training Jobs with Training Compiler”](#).
Large Language Models Using PyTorch Directly (without the Hugging Face Transformers Trainer API)

If you have a training script that uses PyTorch directly, you need to make additional changes to your PyTorch training script to implement PyTorch/XLA. Follow the instructions to modify your script to properly set up the PyTorch/XLA primatives.

Topics

- For single GPU training
- For distributed training
- Best Practices to Use SageMaker Training Compiler with PyTorch/XLA

For single GPU training

1. Import the optimization libraries.

```python
import torch_xla
import torch_xla.core.xla_model as xm
```

2. Change the target device to be XLA instead of `torch.device("cuda")`

```python
device=xm.xla_device()
```

3. If you're using PyTorch's **Automatic Mixed Precision** (AMP), do the following:
   a. Replace `torch.cuda.amp` with the following:

   ```python
   import torch_xla.amp
   ```

   b. Replace `torch.optim.SGD` and `torch.optim.Adam` with the following:

   ```python
   import torch_xla.amp.syncfree.Adam as adam
   import torch_xla.amp.syncfree.SGD as SGD
   ```

   c. Replace `torch.cuda.amp.GradScaler` with the following:

   ```python
   import torch_xla.amp.GradScaler as grad_scaler
   ```

4. If you're not using AMP, replace `optimizer.step()` with the following:
xm.optimizer_step(optimizer)

5. If you're using a distributed dataloader, wrap your dataloader in the PyTorch/XLA's ParallelLoader class:

```python
import torch_xla.distributed.parallel_loader as pl
parallel_loader=pl.ParallelLoader(dataloader, [device]).per_device_loader(device)
```

6. Add `mark_step` at the end of the training loop when you're not using `parallel_loader`:

```python
xm.mark_step()
```

7. To checkpoint your training, use the PyTorch/XLA's model checkpoint method:

```python
xm.save(model.state_dict(), path_to_save)
```

After you have completed adapting your training script, proceed to the section called “Run PyTorch Training Jobs with Training Compiler”.

**For distributed training**

In addition to the changes listed in the previous For single GPU training section, add the following changes to properly distribute workload across GPUs.

1. If you're using AMP, add `all_reduce` after `scaler.scale(loss).backward()`:

```python
gradients=xm._fetch_gradients(optimizer)
xm.all_reduce('sum', gradients, scale=1.0/xm.xrt_world_size())
```

2. If you need to set variables for `local_ranks` and `world_size`, use similar code to the following:

```python
local_rank=xm.get_local_ordinal()
world_size=xm.xrt_world_size()
```

3. For any `world_size` (num_gpus_per_node*num_nodes) greater than 1, you must define a train sampler which should look similar to the following:

```python
import torch_xla.core.xla_model as xm
```
if xm.xrt_world_size() > 1:
    train_sampler=torch.utils.data.distributed.DistributedSampler(
        train_dataset,
        num_replicas=xm.xrt_world_size(),
        rank=xm.get_ordinal(),
        shuffle=True
    )

train_loader=torch.utils.data.DataLoader(
    train_dataset,
    batch_size=args.batch_size,
    sampler=train_sampler,
    drop_last=args.drop_last,
    shuffle=False if train_sampler else True,
    num_workers=args.num_workers
)

4. Make the following changes to make sure you use the parallel_loader provided by the torch_xla distributed module.

   import torch_xla.distributed.parallel_loader as pl
   train_device_loader=pl.MpDeviceLoader(train_loader, device)

   The train_device_loader functions like a regular PyTorch loader as follows:

   for step, (data, target) in enumerate(train_device_loader):
       optimizer.zero_grad()
       output=model(data)
       loss=torch.nn.NLLLoss(output, target)
       loss.backward()

With all of these changes, you should be able to launch distributed training with any PyTorch model without the Transformer Trainer API. Note that these instructions can be used for both single-node multi-GPU and multi-node multi-GPU.

5. For PyTorch v1.11.0 and later

   To run distributed training with SageMaker Training Compiler, you must add the following _mp_fn() function in your training script and wrap the main() function. It redirects the _mp_fn(index) function calls from the SageMaker distributed runtime for PyTorch (pytorchxla) to the main() function of your training script.
This function accepts the `index` argument to indicate the rank of the current GPU in the cluster for distributed training. To find more example scripts, see the Hugging Face Transformers language modeling example scripts.

**For Transformers v4.17 and before with PyTorch v1.10.2 and before**

SageMaker Training Compiler uses an alternate mechanism for launching a distributed training job and requires you to pass a SageMaker distributed training launcher script to the `entry_point` argument and pass your training script to the `hyperparameters` argument in the SageMaker Hugging Face estimator.

After you have completed adapting your training script, proceed to the section called “Run PyTorch Training Jobs with Training Compiler”.

**Best Practices to Use SageMaker Training Compiler with PyTorch/XLA**

If you want to leverage the SageMaker Training Compiler on your native PyTorch training script, you may want to first get familiar with PyTorch on XLA devices. The following sections list some best practices to enable XLA for PyTorch.

**Note**

This section for best practices assumes that you use the following PyTorch/XLA modules:

```python
import torch_xla.core.xla_model as xm
import torch_xla.distributed.parallel_loader as pl
```

**Understand the lazy mode in PyTorch/XLA**

One significant difference between PyTorch/XLA and native PyTorch is that the PyTorch/XLA system runs in lazy mode while the native PyTorch runs in eager mode. Tensors in lazy mode are placeholders for building the computational graph until they are materialized after the compilation and evaluation are complete. The PyTorch/XLA system builds the computational graph on the fly when you call PyTorch APIs to build the computation using tensors and operators. The
computational graph gets compiled and executed when `xm.mark_step()` is called explicitly or implicitly by `plMpDeviceLoader/plParallelLoader`, or when you explicitly request the value of a tensor such as by calling `loss.item()` or `print(loss)`.

**Minimize the number of compilation-and-executions using `plMpDeviceLoader/plParallelLoader` and `xm.step_closure`**

For best performance, you should keep in mind the possible ways to initiate compilation-and-executions as described in [Understand the lazy mode in PyTorch/XLA](#) and should try to minimize the number of compilation-and-executions. Ideally, only one compilation-and-execution is necessary per training iteration and is initiated automatically by `plMpDeviceLoader/plParallelLoader`. The `MpDeviceLoader` is optimized for XLA and should always be used if possible for best performance. During training, you might want to examine some intermediate results such as loss values. In such case, the printing of lazy tensors should be wrapped using `xm.add_step_closure()` to avoid unnecessary compilation-and-executions.

**Use AMP and syncfree optimizers**

Training in Automatic Mixed Precision (AMP) mode significantly accelerates your training speed by leveraging the Tensor cores of NVIDIA GPUs. SageMaker Training Compiler provides syncfree optimizers that are optimized for XLA to improve AMP performance. Currently, the following three syncfree optimizers are available and should be used if possible for best performance.

```python
torch_xla.amp.syncfree.SGD
torch_xla.amp.syncfree.Adam
torch_xla.amp.syncfree.AdamW
```

These syncfree optimizers should be paired with `torch_xla.amp.GradScaler` for gradient scaling/unscaling.

**Tip**

Starting PyTorch 1.13.1, SageMaker Training Compiler improves performance by letting PyTorch/XLA to automatically override the optimizers (such as SGD, Adam, AdamW) in `torch.optim` or `transformers.optimization` with the syncfree versions of them in `torch_xla.amp.syncfree` (such as `torch_xla.amp.syncfree.SGD`, `torch_xla.amp.syncfree.Adam`, `torch_xla.amp.syncfree.AdamW`). You don't need to change those code lines where you define optimizers in your training script.
**TensorFlow**

Bring your own TensorFlow model to SageMaker, and run the training job with SageMaker Training Compiler.

**TensorFlow Models**

SageMaker Training Compiler automatically optimizes model training workloads that are built on top of the native TensorFlow API or the high-level Keras API.

### Tip

For preprocessing your input dataset, ensure that you use a static input shape. Dynamic input shape can initiate recompilation of the model and might increase total training time.

### Using Keras (Recommended)

For the best compiler acceleration, we recommend using models that are subclasses of TensorFlow Keras (`tf.keras.Model`).

#### For single GPU training

There's no additional change you need to make in the training script.

### Without Keras

SageMaker Training Compiler does not support eager execution in TensorFlow. Accordingly, you should wrap your model and training loops with the TensorFlow function decorator (`@tf.function`) to leverage compiler acceleration.

SageMaker Training Compiler performs a graph-level optimization, and uses the decorator to make sure your TensorFlow functions are set to run in **graph mode**.

#### For single GPU training

TensorFlow 2.0 or later has the eager execution on by default, so you should add the `@tf.function` decorator in front of every function that you use for constructing a TensorFlow model.
TensorFlow Models with Hugging Face Transformers

TensorFlow models with Hugging Face Transformers are based on TensorFlow's `tf.keras.Model` API. Hugging Face Transformers also provides pretrained model classes for TensorFlow to help reduce the effort for configuring natural language processing (NLP) models. After creating your own training script using the Transformers library, you can run the training script using the SageMaker HuggingFace estimator with the SageMaker Training Compiler configuration class as shown in the previous topic at `Run TensorFlow Training Jobs with SageMaker Training Compiler`.

SageMaker Training Compiler automatically optimizes model training workloads that are built on top of the native TensorFlow API or the high-level Keras API, such as the TensorFlow transformer models.

⚠️ Tip

When you create a tokenizer for an NLP model using Transformers in your training script, make sure that you use a static input tensor shape by specifying `padding='max_length'`. Do not use `padding='longest'` because padding to the longest sequence in the batch can change the tensor shape for each training batch. The dynamic input shape can initiate recompilation of the model and might increase total training time. For more information about padding options of the Transformers tokenizers, see Padding and truncation in the Hugging Face Transformers documentation.

Topics

- Using Keras
- Without Keras

Using Keras

For the best compiler acceleration, we recommend using models that are subclasses of TensorFlow Keras (`tf.keras.Model`). As noted in the Quick tour page in the Hugging Face Transformers documentation, you can use the models as regular TensorFlow Keras models.

For single GPU training

There's no additional change you need to make in the training script.
For distributed training

SageMaker Training Compiler acceleration works transparently for multi-GPU workloads when the model is constructed and trained using Keras APIs within the scope of `tf.distribute.Strategy.scope()` call.

1. Choose the right distributed training strategy.
   a. For single-node multi-GPU, use `tf.distribute.MirroredStrategy` to set the strategy.

   ```python
   strategy = tf.distribute.MirroredStrategy()
   ```

   b. For multi-node multi-GPU, add the following code to properly set the TensorFlow distributed training configuration before creating the strategy.

   ```python
def set_sm_dist_config():
    DEFAULT_PORT = '8890'
    DEFAULT_CONFIG_FILE = '/opt/ml/input/config/resourceconfig.json'
    with open(DEFAULT_CONFIG_FILE) as f:
        config = json.loads(f.read())
        current_host = config['current_host']
    tf_config = {
        'cluster': {
            'worker': []
        },
        'task': {'type': 'worker', 'index': -1}
    }
    for i, host in enumerate(config['hosts']):
        tf_config['cluster']['worker'].append("%s:%s" % (host, DEFAULT_PORT))
        if current_host == host:
            tf_config['task']['index'] = i
    os.environ['TF_CONFIG'] = json.dumps(tf_config)

set_sm_dist_config()
```

Use `tf.distribute.MultiWorkerMirroredStrategy` to set the strategy.

```python
strategy = tf.distribute.MultiWorkerMirroredStrategy()
```

2. Using the strategy of your choice, wrap the model.

```python
with strategy.scope():
```
# create a model and do fit

## Without Keras

If you want to bring custom models with custom training loops using TensorFlow without Keras, you should wrap the model and the training loop with the TensorFlow function decorator (`@tf.function`) to leverage compiler acceleration.

SageMaker Training Compiler performs a graph-level optimization, and uses the decorator to make sure your TensorFlow functions are set to run in graph mode.

### For single GPU training

TensorFlow 2.0 or later has the eager execution on by default, so you should add the `@tf.function` decorator in front of every function that you use for constructing a TensorFlow model.

### For distributed training

In addition to the changes needed for [Using Keras for distributed training](#), you need to ensure that functions to be run on each GPU are annotated with `@tf.function`, while cross-GPU communication functions are not annotated. An example training code should look like the following:

```python
@tf.function()
def compiled_step(inputs, outputs):
    with tf.GradientTape() as tape:
        pred=model(inputs, training=True)
        total_loss=loss_object(outputs, pred)/args.batch_size
        gradients=tape.gradient(total_loss, model.trainable_variables)
    return total_loss, pred, gradients

def train_step(inputs, outputs):
    total_loss, pred, gradients=compiled_step(inputs, outputs)
    if args.weight_decay > 0.:
        gradients=[g+v*args.weight_decay for g,v in zip(gradients, model.trainable_variables)]
    optimizer.apply_gradients(zip(gradients, model.trainable_variables))
    train_loss.update_state(total_loss)
```
train_accuracy.update_state(outputs, pred)

@tf.function()
def train_step_dist(inputs, outputs):
    strategy.run(train_step, args= (inputs, outputs))

Note that this instruction can be used for both single-node multi-GPU and multi-node multi-GPU.

Enable SageMaker Training Compiler

SageMaker Training Compiler is built into the SageMaker Python SDK and AWS Deep Learning Containers so that you don’t need to change your workflows to enable Training Compiler. Choose one of the following topics that matches with your use case.

Topics

- Run PyTorch Training Jobs with SageMaker Training Compiler
- Run TensorFlow Training Jobs with SageMaker Training Compiler

Run PyTorch Training Jobs with SageMaker Training Compiler

You can use any of the SageMaker interfaces to run a training job with SageMaker Training Compiler: Amazon SageMaker Studio Classic, Amazon SageMaker notebook instances, AWS SDK for Python (Boto3), and AWS Command Line Interface.

Topics

- Using the SageMaker Python SDK
- Using the SageMaker CreateTrainingJob API Operation

Using the SageMaker Python SDK

SageMaker Training Compiler for PyTorch is available through the SageMaker PyTorch and HuggingFace framework estimator classes. To turn on SageMaker Training Compiler, add the compiler_config parameter to the SageMaker estimators. Import the TrainingCompilerConfig class and pass an instance of it to the compiler_config parameter. The following code examples show the structure of SageMaker estimator classes with SageMaker Training Compiler turned on.
Tip
To get started with prebuilt models provided by PyTorch or Transformers, try using the batch sizes provided in the reference table at Tested Models.

Note
The native PyTorch support is available in the SageMaker Python SDK v2.121.0 and later. Make sure that you update the SageMaker Python SDK accordingly.

Note
Starting PyTorch v1.12.0, SageMaker Training Compiler containers for PyTorch are available. Note that the SageMaker Training Compiler containers for PyTorch are not prepackaged with Hugging Face Transformers. If you need to install the library in the container, make sure that you add the requirements.txt file under the source directory when submitting a training job.
For PyTorch v1.11.0 and before, use the previous versions of the SageMaker Training Compiler containers for Hugging Face and PyTorch.
For a complete list of framework versions and corresponding container information, see the section called “Supported Frameworks”.

For information that fits your use case, see one of the following options.

For single GPU training

PyTorch v1.12.0 and later

To compile and train a PyTorch model, configure a SageMaker PyTorch estimator with SageMaker Training Compiler as shown in the following code example.

Note
This native PyTorch support is available in the SageMaker Python SDK v2.120.0 and later. Make sure that you update the SageMaker Python SDK.
```python
from sagemaker.pytorch import PyTorch, TrainingCompilerConfig

# the original max batch size that can fit into GPU memory without compiler
batch_size_native=12
learning_rate_native=float('5e-5')

# an updated max batch size that can fit into GPU memory with compiler
batch_size=64

# update learning rate
learning_rate=learning_rate_native/batch_size_native*batch_size

hyperparameters={
    "n_gpus": 1,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

pytorch_estimator=PyTorch(
    entry_point='train.py',
    source_dir='path-to-requirements-file', # Optional. Add this if need to install additional packages.
    instance_count=1,
    instance_type='ml.p3.2xlarge',
    framework_version='1.13.1',
    py_version='py3',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)

pytorch_estimator.fit()
```

### Hugging Face Transformers with PyTorch v1.11.0 and before

To compile and train a transformer model with PyTorch, configure a SageMaker Hugging Face estimator with SageMaker Training Compiler as shown in the following code example.

```python
from sagemaker.huggingface import HuggingFace, TrainingCompilerConfig

# the original max batch size that can fit into GPU memory without compiler
batch_size_native=12
```

Enable Training Compiler
learning_rate_native=float('5e-5')

# an updated max batch size that can fit into GPU memory with compiler
batch_size=64

# update learning rate
learning_rate=learning_rate_native/batch_size_native*batch_size

hyperparameters={
    "n_gpus": 1,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

pytorch_huggingface_estimator=HuggingFace(
    entry_point='train.py',
    instance_count=1,
    instance_type='ml.p3.2xlarge',
    transformers_version='4.21.1',
    pytorch_version='1.11.0',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)

pytorch_huggingface_estimator.fit()

To prepare your training script, see the following pages.

- [For single GPU training](#) of a PyTorch model using Hugging Face Transformers' [Trainer API](#)
- [For single GPU training](#) of a PyTorch model without Hugging Face Transformers' [Trainer API](#)

To find end-to-end examples, see the following notebooks:

- [Compile and Train a Hugging Face Transformers Trainer Model for Question and Answering with the SQuAD dataset](#)
- [Compile and Train a Hugging Face Transformer BERT Model with the SST Dataset using SageMaker Training Compiler](#)
- [Compile and Train a Binary Classification Trainer Model with the SST2 Dataset for Single-Node Single-GPU Training](#)
For distributed training

PyTorch v1.12

For PyTorch v1.12, you can run distributed training with SageMaker Training Compiler by adding the `pytorch_xla` option specified to the `distribution` parameter of the SageMaker PyTorch estimator class.

⚠️ Note

This native PyTorch support is available in the SageMaker Python SDK v2.121.0 and later. Make sure that you update the SageMaker Python SDK.

```python
from sagemaker.pytorch import PyTorch, TrainingCompilerConfig

# choose an instance type, specify the number of instances you want to use, # and set the num_gpus variable the number of GPUs per instance.
instance_count=1
instance_type='ml.p3.8xlarge'
num_gpus=4

# the original max batch size that can fit to GPU memory without compiler
batch_size_native=16
learning_rate_native=float('5e-5')

# an updated max batch size that can fit to GPU memory with compiler
batch_size=26

# update learning rate
learning_rate=learning_rate_native/
batch_size_native*batch_size*num_gpus*instance_count

hyperparameters={
    "n_gpus": num_gpus,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

pytorch_estimator=PyTorch(
    entry_point='your_training_script.py',
)```
source_dir='path-to-requirements-file', # Optional. Add this if need to install additional packages.
    instance_count=instance_count,
    instance_type=instance_type,
    framework_version='1.13.1',
    py_version='py3',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    distribution ={'pytorchxla' : { 'enabled': True }},
    disable_profiler=True,
    debugger_hook_config=False
)

pytorch_estimator.fit()

---

Tip

To prepare your training script, see PyTorch

Transformers v4.21 with PyTorch v1.11

For PyTorch v1.11 and later, SageMaker Training Compiler is available for distributed training with the pytorch_xla option specified to the distribution parameter.

from sagemaker.huggingface import HuggingFace, TrainingCompilerConfig

# choose an instance type, specify the number of instances you want to use,
# and set the num_gpus variable the number of GPUs per instance.
instance_count=1
instance_type='ml.p3.8xlarge'
num_gpus=4

# the original max batch size that can fit to GPU memory without compiler
batch_size_native=16
learning_rate_native=float('5e-5')

# an updated max batch size that can fit to GPU memory with compiler
batch_size=26

# update learning rate
learning_rate=learning_rate_native/
batch_size_native*batch_size*num_gpus*instance_count

hyperparameters={
    "n_gpus": num_gpus,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

pysotorch_huggingface_estimatoren=HuggingFace(
    entry_point='your_training_script.py',
    instance_count=instance_count,
    instance_type=instance_type,
    transformers_version='4.21.1',
    pytorch_version='1.11.0',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    distribution ={'pytorchxla' : { 'enabled': True }},
    disable_profiler=True,
    debugger_hook_config=False
)

pytorch_huggingface_estimatoren.fit()

**Tip**

To prepare your training script, see the following pages.

- For distributed training of a PyTorch model using Hugging Face Transformers' Trainer API
- For distributed training of a PyTorch model without Hugging Face Transformers' Trainer API

Transformers v4.17 with PyTorch v1.10.2 and before

For the supported version of PyTorch v1.10.2 and before, SageMaker Training Compiler requires an alternate mechanism for launching a distributed training job. To run distributed training, SageMaker Training Compiler requires you to pass a SageMaker distributed training launcher script to the entry_point argument, and pass your training script to the hyperparameters
argument. The following code example shows how to configure a SageMaker Hugging Face estimator applying the required changes.

```python
from sagemaker.huggingface import HuggingFace, TrainingCompilerConfig

# choose an instance type, specify the number of instances you want to use, # and set the num_gpus variable the number of GPUs per instance.
instance_count=1
instance_type='ml.p3.8xlarge'
num_gpus=4

# the original max batch size that can fit to GPU memory without compiler
batch_size_native=16
learning_rate_native=float('5e-5')

# an updated max batch size that can fit to GPU memory with compiler
batch_size=26

# update learning rate
learning_rate=learning_rate_native/
batch_size_native*batch_size*num_gpus*instance_count

training_script="your_training_script.py"

hyperparameters={
    "n_gpus": num_gpus,
    "batch_size": batch_size,
    "learning_rate": learning_rate,
    "training_script": training_script  # Specify the file name of your training
    script.
}

cpytorch_huggingface_estimator=HuggingFace(
    entry_point='distributed_training_launcher.py',  # Specify the distributed
    training launcher script.
    instance_count=instance_count,
    instance_type=instance_type,
    transformers_version='4.17.0',
    pytorch_version='1.10.2',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)
The launcher script should look like the following. It wraps your training script and configures the distributed training environment depending on the size of the training instance of your choice.

```python
#!/bin/python
import subprocess
import sys
if __name__ == '__main__':
    arguments_command = ' '.join(sys.argv[1:])
    subprocess.check_call("python -m torch_xla.distributed.sm_dist " + arguments_command, shell=True)

Tip
To prepare your training script, see the following pages.

- For distributed training of a PyTorch model using Hugging Face Transformers' Trainer API
- For distributed training of a PyTorch model without Hugging Face Transformers' Trainer API

Tip
To find end-to-end examples, see the following notebooks:

- Compile and Train the GPT2 Model using the Transformers Trainer API with the SST2 Dataset for Single-Node Multi-GPU Training
The following list is the minimal set of parameters required to run a SageMaker training job with the compiler.

### Note

When using the SageMaker Hugging Face estimator, you must specify the `transformers_version`, `pytorch_version`, `hyperparameters`, and `compiler_config` parameters to enable SageMaker Training Compiler. You cannot use `image_uri` to manually specify the Training Compiler integrated Deep Learning Containers that are listed at [Supported Frameworks](https://docs.aws.amazon.com/sagemaker/latest/dg/supportedframeworks.html).

- `entry_point` (str) – Required. Specify the file name of your training script.

### Note

To run a distributed training with SageMaker Training Compiler and PyTorch v1.10.2 and before, specify the file name of a launcher script to this parameter. The launcher script should be prepared to wrap your training script and configure the distributed training environment. For more information, see the following example notebooks:

- [Compile and Train the GPT2 Model using the Transformers Trainer API with the SST2 Dataset for Single-Node Multi-GPU Training](https://aws.amazon.com/sagemaker)
- [Compile and Train the GPT2 Model using the Transformers Trainer API with the SST2 Dataset for Multi-Node Multi-GPU Training](https://aws.amazon.com/sagemaker)

- `source_dir` (str) – Optional. Add this if need to install additional packages. To install packages, you need to prepare a `requirements.txt` file under this directory.

- `instance_count` (int) – Required. Specify the number of instances.

- `instance_type` (str) – Required. Specify the instance type.

- `transformers_version` (str) – Required only when using the SageMaker Hugging Face estimator. Specify the Hugging Face Transformers library version supported by SageMaker Training Compiler. To find available versions, see [Supported Frameworks](https://docs.aws.amazon.com/sagemaker/latest/dg/supportedframeworks.html).
• framework_version or pytorch_version (str) – Required. Specify the PyTorch version supported by SageMaker Training Compiler. To find available versions, see Supported Frameworks.

Note
When using the SageMaker Hugging Face estimator, you must specify both transformers_version and pytorch_version.

• hyperparameters (dict) – Optional. Specify hyperparameters for the training job, such as n_gpus, batch_size, and learning_rate. When you enable SageMaker Training Compiler, try larger batch sizes and adjust the learning rate accordingly. To find case studies of using the compiler and adjusted batch sizes to improve training speed, see the section called “Tested Models” and SageMaker Training Compiler Example Notebooks and Blogs.

Note
To run a distributed training with SageMaker Training Compiler and PyTorch v1.10.2 and before, you need to add an additional parameter, "training_script", to specify your training script, as shown in the preceding code example.

• compiler_config (TrainingCompilerConfig object) – Required to activate SageMaker Training Compiler. Include this parameter to turn on SageMaker Training Compiler. The following are parameters for the TrainingCompilerConfig class.
  • enabled (bool) – Optional. Specify True or False to turn on or turn off SageMaker Training Compiler. The default value is True.
  • debug (bool) – Optional. To receive more detailed training logs from your compiler-accelerated training jobs, change it to True. However, the additional logging might add overhead and slow down the compiled training job. The default value is False.
  • distribution (dict) – Optional. To run a distributed training job with SageMaker Training Compiler, add distribution = { 'pytorchxla' : { 'enabled': True }}.

Warning
If you turn on SageMaker Debugger, it might impact the performance of SageMaker Training Compiler. We recommend that you turn off Debugger when running SageMaker
Training Compiler to make sure there's no impact on performance. For more information, see the section called “Considerations”. To turn the Debugger functionalities off, add the following two arguments to the estimator:

```python
disable_profiler=True,
ddebugger_hook_config=False
```

If the training job with the compiler is launched successfully, you receive the following logs during the job initialization phase:

- **With TrainingCompilerConfig(debug=False)**

  ```shell
  Found configuration for Training Compiler
  Configuring SM Training Compiler...
  ```

- **With TrainingCompilerConfig(debug=True)**

  ```shell
  Found configuration for Training Compiler
  Configuring SM Training Compiler...
  Training Compiler set to debug mode
  ```

### Using the SageMaker CreateTrainingJob API Operation

SageMaker Training Compiler configuration options must be specified through the `AlgorithmSpecification` and `HyperParameters` field in the request syntax for the `CreateTrainingJob API operation`.

```json
"AlgorithmSpecification": {
    "TrainingImage": "<sagemaker-training-compiler-enabled-dlc-image>"
},

"HyperParameters": {
    "sagemaker_training_compiler_enabled": "true",
    "sagemaker_training_compiler_debug_mode": "false",
    "sagemaker_pytorch_xla_multi_worker_enabled": "false" // set to "true" for distributed training
}
```
To find a complete list of deep learning container image URIs that have SageMaker Training Compiler implemented, see Supported Frameworks.

Run TensorFlow Training Jobs with SageMaker Training Compiler

You can use any of the SageMaker interfaces to run a training job with SageMaker Training Compiler: Amazon SageMaker Studio Classic, Amazon SageMaker notebook instances, AWS SDK for Python (Boto3), and AWS Command Line Interface.

Topics

- Using the SageMaker Python SDK
- Using the SageMaker Python SDK and Extending SageMaker Framework Deep Learning Containers
- Enable SageMaker Training Compiler Using the SageMaker CreateTrainingJob API Operation

Using the SageMaker Python SDK

To turn on SageMaker Training Compiler, add the `compiler_config` parameter to the SageMaker TensorFlow or Hugging Face estimator. Import the `TrainingCompilerConfig` class and pass an instance of it to the `compiler_config` parameter. The following code examples show the structure of the SageMaker estimator classes with SageMaker Training Compiler turned on.

Tip

To get started with prebuilt models provided by the TensorFlow and Transformers libraries, try using the batch sizes provided in the reference table at Tested Models.

Note

SageMaker Training Compiler for TensorFlow is available through the SageMaker TensorFlow and Hugging Face framework estimators.

For information that fits your use case, see one of the following options.
For single GPU training

TensorFlow

```python
from sagemaker.tensorflow import TensorFlow, TrainingCompilerConfig

# the original max batch size that can fit into GPU memory without compiler
batch_size_native=12
learning_rate_native=float('5e-5')

# an updated max batch size that can fit into GPU memory with compiler
batch_size=64

# update the global learning rate
learning_rate=learning_rate_native/batch_size_native*batch_size

hyperparameters={
    "n_gpus": 1,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

tensorflow_estimator=TensorFlow(
    entry_point='train.py',
    instance_count=1,
    instance_type='ml.p3.2xlarge',
    framework_version='2.9.1',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)

tensorflow_estimator.fit()
```

To prepare your training script, see the following pages.

- **For single GPU training** of a model constructed using TensorFlow Keras (tf.keras.*).
- **For single GPU training** of a model constructed using TensorFlow modules (tf.* excluding the TensorFlow Keras modules).
Hugging Face Estimator with TensorFlow

```python
from sagemaker.huggingface import HuggingFace, TrainingCompilerConfig

# the original max batch size that can fit into GPU memory without compiler
batch_size_native=12
learning_rate_native=float('5e-5')

# an updated max batch size that can fit into GPU memory with compiler
batch_size=64

# update the global learning rate
learning_rate=learning_rate_native/batch_size_native*batch_size

hyperparameters={
    "n_gpus": 1,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

tensorflow_huggingface_estimator=HuggingFace(
    entry_point='train.py',
    instance_count=1,
    instance_type='ml.p3.2xlarge',
    transformers_version='4.21.1',
    tensorflow_version='2.6.3',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)

tensorflow_huggingface_estimator.fit()
```

To prepare your training script, see the following pages.

- **For single GPU training** of a TensorFlow Keras model with Hugging Face Transformers
- **For single GPU training** of a TensorFlow model with Hugging Face Transformers
For distributed training

Hugging Face Estimator with TensorFlow

```
from sagemaker.huggingface import HuggingFace, TrainingCompilerConfig

# choose an instance type, specify the number of instances you want to use, # and set the num_gpus variable the number of GPUs per instance.
instance_count=1
instance_type='ml.p3.8xlarge'
num_gpus=4

# the original max batch size that can fit to GPU memory without compiler
batch_size_native=16
learning_rate_native=float('5e-5')

# an updated max batch size that can fit to GPU memory with compiler
batch_size=26

# update learning rate
learning_rate=learning_rate_native/batch_size_native*batch_size*num_gpus*instance_count

hyperparameters={
    "n_gpus": num_gpus,
    "batch_size": batch_size,
    "learning_rate": learning_rate
}

tensorflow_huggingface_estimator=HuggingFace(
    entry_point='train.py',
    instance_count=instance_count,
    instance_type=instance_type,
    transformers_version='4.21.1',
    tensorflow_version='2.6.3',
    hyperparameters=hyperparameters,
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)

tensorflow_huggingface_estimator.fit()
```
Tip
To prepare your training script, see the following pages.

- For distributed training of a TensorFlow Keras model with Hugging Face Transformers
- For distributed training of a TensorFlow model with Hugging Face Transformers

The following list is the minimal set of parameters required to run a SageMaker training job with the compiler.

Note
When using the SageMaker Hugging Face estimator, you must specify the transformers_version, tensorflow_version, hyperparameters, and compiler_config parameters to enable SageMaker Training Compiler. You cannot use image_uri to manually specify the Training Compiler integrated Deep Learning Containers that are listed at Supported Frameworks.

- entry_point (str) – Required. Specify the file name of your training script.
- instance_count (int) – Required. Specify the number of instances.
- instance_type (str) – Required. Specify the instance type.
- transformers_version (str) – Required only when using the SageMaker Hugging Face estimator. Specify the Hugging Face Transformers library version supported by SageMaker Training Compiler. To find available versions, see Supported Frameworks.
- framework_version or tensorflow_version (str) – Required. Specify the TensorFlow version supported by SageMaker Training Compiler. To find available versions, see Supported Frameworks.

Note
When using the SageMaker TensorFlow estimator, you must specify framework_version.
When using the SageMaker Hugging Face estimator, you must specify both transformers_version and tensorflow_version.
hyperparameters (dict) – Optional. Specify hyperparameters for the training job, such as `n_gpus`, `batch_size`, and `learning_rate`. When you enable SageMaker Training Compiler, try larger batch sizes and adjust the learning rate accordingly. To find case studies of using the compiler and adjusted batch sizes to improve training speed, see the section called “Tested Models” and SageMaker Training Compiler Example Notebooks and Blogs.

compiler_config (TrainingCompilerConfig object) – Required. Include this parameter to turn on SageMaker Training Compiler. The following are parameters for the TrainingCompilerConfig class.

- enabled (bool) – Optional. Specify `True` or `False` to turn on or turn off SageMaker Training Compiler. The default value is `True`.
- debug (bool) – Optional. To receive more detailed training logs from your compiler-accelerated training jobs, change it to `True`. However, the additional logging might add overhead and slow down the compiled training job. The default value is `False`.

⚠️ Warning
If you turn on SageMaker Debugger, it might impact the performance of SageMaker Training Compiler. We recommend that you turn off Debugger when running SageMaker Training Compiler to make sure there's no impact on performance. For more information, see the section called “Considerations”. To turn the Debugger functionalities off, add the following two arguments to the estimator:

```python
disable_profiler=True,
debugger_hook_config=False
```

If the training job with the compiler is launched successfully, you receive the following logs during the job initialization phase:

- With `TrainingCompilerConfig(debug=False)`

```
Found configuration for Training Compiler
Configuring SM Training Compiler...
```

- With `TrainingCompilerConfig(debug=True)`

```
Found configuration for Training Compiler
```
Using the SageMaker Python SDK and Extending SageMaker Framework Deep Learning Containers

AWS Deep Learning Containers (DLC) for TensorFlow use adapted versions of TensorFlow that include changes on top of the open-source TensorFlow framework. The SageMaker Framework Deep Learning Containers are optimized for the underlying AWS infrastructure and Amazon SageMaker. With the advantage of using the DLCs, SageMaker Training Compiler integration adds more performance improvements over the native TensorFlow. Furthermore, you can create a custom training container by extending the DLC image.

⚠️ Note

This Docker customization feature is currently available only for TensorFlow.

To extend and customize the SageMaker TensorFlow DLCs for your use-case, use the following instructions.

Create a Dockerfile

Use the following Dockerfile template to extend the SageMaker TensorFlow DLC. You must use the SageMaker TensorFlow DLC image as the base image of your Docker container. To find the SageMaker TensorFlow DLC image URIs, see Supported Frameworks.

```
# SageMaker TensorFlow Deep Learning Container image
FROM 763104351884.dkr.ecr.<aws-region>.amazonaws.com/tensorflow-training:<image-tag>

ENV PATH="/opt/ml/code:${PATH}"

# This environment variable is used by the SageMaker container
# to determine user code directory.
ENV SAGEMAKER_SUBMIT_DIRECTORY /opt/ml/code

# Add more code lines to customize for your use-case
...
```

For more information, see Step 2: Create and upload the Dockerfile and Python training scripts.
Consider the following pitfalls when extending SageMaker Framework DLCs:

- Do not explicitly uninstall or change the version of TensorFlow packages in SageMaker containers. Doing so causes the AWS optimized TensorFlow packages to be overwritten by open-source TensorFlow packages, which might result in performance degradation.

- Watch out for packages that have a particular TensorFlow version or flavor as a dependency. These packages might implicitly uninstall the AWS optimized TensorFlow and install open-source TensorFlow packages.

For example, there's a known issue that the `tensorflow/models` and `tensorflow/text` libraries always attempt to reinstall open source TensorFlow. If you need to install these libraries to choose a specific version for your use case, we recommend that you look into the SageMaker TensorFlow DLC Dockerfiles for v2.9 or later. The paths to the Dockerfiles are typically in the following format: `tensorflow/training/docker/<tensorflow-version>/py3/<cuda-version>/Dockerfile.gpu`. In the Dockerfiles, you should find the code lines to reinstall AWS managed TensorFlow binary (specified to the TF_URL environment variable) and other dependencies in order. The reinstallation section should look like the following example:

```bash
# tf-models does not respect existing installations of TensorFlow
# and always installs open source TensorFlow

RUN pip3 install --no-cache-dir -U \
    tf-models-official==x.y.z

RUN pip3 uninstall -y tensorflow tensorflow-gpu \
    ; pip3 install --no-cache-dir -U \
    ${TF_URL} \
    tensorflow-io==x.y.z \
    tensorflow-datasets==x.y.z
```

**Build and push to ECR**

To build and push your Docker container to Amazon ECR, follow the instructions in the following links:

- **Step 3: Build the container**
- **Step 4: Test the container**
- **Step 5: Push the container to Amazon ECR**
Run using the SageMaker Python SDK Estimator

Use the SageMaker TensorFlow framework estimator as usual. You must specify `image_uri` to use the new container you hosted in Amazon ECR.

```python
import sagemaker, boto3
from sagemaker import get_execution_role
from sagemaker.tensorflow import TensorFlow, TrainingCompilerConfig

account_id = boto3.client('sts').get_caller_identity().get('Account')
ecr_repository = 'tf-custom-container-test'
tag = ':latest'

region = boto3.session.Session().region_name
uri_suffix = 'amazonaws.com'

byoc_image_uri = '{}.dkr.ecr.{}.amazonaws.com/{}'.format(
    account_id, region, uri_suffix, ecr_repository + tag
)

estimator = TensorFlow(
    image_uri=byoc_image_uri,
    role=get_execution_role(),
    base_job_name='tf-custom-container-test-job',
    instance_count=1,
    instance_type='ml.p3.8xlarge',
    compiler_config=TrainingCompilerConfig(),
    disable_profiler=True,
    debugger_hook_config=False
)

# Start training
estimator.fit()
```
Enable SageMaker Training Compiler Using the SageMaker CreateTrainingJob API Operation

SageMaker Training Compiler configuration options must be specified through the AlgorithmSpecification and HyperParameters field in the request syntax for the CreateTrainingJob API operation.

```json
"AlgorithmSpecification": {
    "TrainingImage": "<sagemaker-training-compiler-enabled-dlc-image>"
},

"HyperParameters": {
    "sagemaker_training_compiler_enabled": "true",
    "sagemaker_training_compiler_debug_mode": "false"
}
```

To find a complete list of deep learning container image URIs that have SageMaker Training Compiler implemented, see Supported Frameworks.

**SageMaker Training Compiler Example Notebooks and Blogs**

The following blogs, case studies, and notebooks provide examples of how to implement SageMaker Training Compiler.

Example notebooks are provided in the SageMaker examples GitHub repository, and you can also browse them on the SageMaker examples website.

**Blogs and Case Studies**

The following blogs discuss case studies about using SageMaker Training Compiler.

- [New – Introducing SageMaker Training Compiler](#)
- [Hugging Face Transformers BERT fine-tuning using Amazon SageMaker Training Compiler](#)
- [Speed up Hugging Face Training Jobs on AWS by Up to 50% with SageMaker Training Compiler](#)

**Examples Notebooks**

To find examples of using SageMaker Training Compiler, see the Training Compiler page in the Amazon SageMaker Example Read the Docs website.
SageMaker Training Compiler Best Practices and Considerations

Review the following best practices and considerations when using SageMaker Training Compiler.

Best Practices

Use the following guidelines to achieve the best results when you run training jobs with SageMaker Training Compiler.

General Best Practices

- Make sure that you use one of the [Supported Instance Types](#) and [Tested Models](#).

- When you create a tokenizer for an NLP model using the Hugging Face Transformers library in your training script, make sure that you use a static input tensor shape by specifying `padding='max_length'`. Do not use `padding='longest'` because padding to the longest sequence in the batch can change the tensor shape for each training batch. The dynamic input shape can initiate recompilation of the model and might increase total training time. For more information about padding options of the Transformers tokenizers, see [Padding and truncation](#) in the *Hugging Face Transformers documentation*.

- Measure GPU memory utilization to make sure that you use the maximum batch size that can fit into the GPU memory. Amazon SageMaker Training Compiler reduces the memory footprint of your model during training, which typically allows you to fit a larger `batch_size` in the GPU memory. Using a larger `batch_size` results in a better GPU utilization and reduces the total training time.

  When you adjust the batch size, you also have to adjust the `learning_rate` appropriately. For example, if you increased the batch size by a factor of $k$, you need to adjust `learning_rate` linearly (simple multiplication by $k$) or multiply by the square root of $k$. This is to achieve the same or similar convergence behavior in the reduced training time. For reference of `batch_size` tested for popular models, see [Tested Models](#).

- To debug the compiler-accelerated training job, enable the debug flag in the `compiler_config` parameter. This enables SageMaker to put the debugging logs into SageMaker training job logs.

```python
huggingface_estimator=HuggingFace(
    ...,
    compiler_config=TrainingCompilerConfig(debug=True)
)
```
Note that if you enable full debugging of the training job with the compiler, this might add some overhead.

**Best Practices for PyTorch**

- If you bring a PyTorch model and want to checkpoint it, make sure you use PyTorch/XLA's model save function to properly checkpoint your model. For more information about the function, see `torch_xla.core.xla_model.save` in the *PyTorch on XLA Devices documentation*.

  To learn how to add the modifications to your PyTorch script, see *Large Language Models Using PyTorch Directly (without the Hugging Face Transformers Trainer API)*.

  For more information about the actual application of using the model save function, see *Checkpoint Writing and Loading* in the *Hugging Face on PyTorch/XLA TPUs: Faster and cheaper training blog*.

- To achieve the most optimal training time for distributed training, consider the following.
  - Use instances with multiple GPUs instead of using single-gpu instances. For example, a single `ml.p3dn.24xlarge` instance has faster training time compared to 8 x `ml.p3.2xlarge` instances.
  - Use instances with EFA support such as `ml.p3dn.24xlarge` and `ml.p4d.24xlarge`. These instance types have accelerated networking speed and reduce training time.
  - Tune the `preprocessing_num_workers` parameter for datasets, so that model training is not delayed by slow preprocessing.

**Considerations**

Consider the following when using SageMaker Training Compiler.

**Performance degradation due to logging, checkpointing, and profiling**

- Avoid logging, checkpointing, and profiling model tensors that lead to explicit evaluations. To understand what an explicit evaluation is, consider the following code compiling example.

  ```python
  a = b+c
  e = a+d
  ```

  A compiler interprets the code as follows and reduces the memory footprint for the variable `a`: 
Now consider the following case in which the code is changed to add a print function for the variable \(a\).

\[
\begin{align*}
\text{e} &= \text{b} + \text{c} + \text{d} \\
\text{a} &= \text{b} + \text{c} \\
\text{e} &= \text{a} + \text{d} \\
\text{print(a)}
\end{align*}
\]

The compiler makes an explicit evaluation of the variable \(a\) as follows.

\[
\begin{align*}
\text{e} &= \text{b} + \text{c} + \text{d} \\
\text{a} &= \text{b} + \text{c} \quad \# \text{Explicit evaluation} \\
\text{print(a)}
\end{align*}
\]

In PyTorch, for example, avoid using `torch.tensor.items()`, which might introduce explicit evaluations. In deep learning, such explicit evaluations can cause overhead because they break fused operations in a compilation graph of a model and lead to recomputation of the tensors.

If you still want to periodically evaluate the model during training while using SageMaker Training Compiler, we recommend logging and checkpointing at a lower frequency to reduce overhead due to explicit evaluations. For example, log every 10 epochs instead of every epoch.

- Graph compilation runs during the first few steps of training. As a result, the first few steps are expected to be exceptionally slow. However, this is a one-time compilation cost and can be amortized by training for a longer duration because compilation makes future steps much faster. The initial compilation overhead depends on the size of the model, the size of the input tensors, and the distribution of input tensor shapes.

**Incorrect use of the PyTorch/XLA APIs when using PyTorch directly**

PyTorch/XLA defines a set of APIs to replace some of the existing PyTorch training APIs. Failing to use them properly leads PyTorch training to fail.

- One of the most typical errors when compiling a PyTorch model is due to a wrong device type for operators and tensors. To properly compile a PyTorch model, make sure you use XLA devices (`xm.xla_device()`) instead of using CUDA or mixing CUDA devices and XLA devices.
- `mark_step()` is a barrier just for XLA. Failing to set it correctly causes a training job to stall.
• PyTorch/XLA provides additional distributed training APIs. Failing to program the APIs properly causes gradients to be collected incorrectly, which causes a training convergence failure.

To properly set up your PyTorch script and avoid the aforementioned incorrect API uses, see Large Language Models Using PyTorch Directly (without the Hugging Face Transformers Trainer API).

SageMaker Training Compiler FAQ

Use the following FAQ items to find answers to commonly asked questions about SageMaker Training Compiler.

Q. How do I know SageMaker Training Compiler is working?

If you successfully launched your training job with SageMaker Training Compiler, you receive the following log messages:

• With `TrainingCompilerConfig(debug=False)`
  
  Found configuration for Training Compiler
  Configuring SM Training Compiler...

• With `TrainingCompilerConfig(debug=True)`
  
  Found configuration for Training Compiler
  Configuring SM Training Compiler...
  Training Compiler set to debug mode

Q. Which models does SageMaker Training Compiler accelerate?

SageMaker Training Compiler supports the most popular deep learning models from the Hugging Face transformers library. With most of the operators that the compiler supports, these models can be trained faster with SageMaker Training Compiler. Compilable models include but are not limited to the following: bert-base-cased, bert-base-chinese, bert-base-uncased, distilbert-base-uncased, distilbert-base-uncased-finetuned-sst-2-english, gpt2, roberta-base, roberta-large, t5-base, and xlm-roberta-base. The compiler works with most DL operators and data structures and can accelerate many other DL models beyond those that have been tested.

Q. What happens if I enable SageMaker Training Compiler with a model that isn't tested?
For an untested model, you might need to first modify the training script to be compatible with SageMaker Training Compiler. For more information, see Bring Your Own Deep Learning Model and follow the instructions on how to prepare your training script.

Once you have updated your training script, you can start the training job. The compiler proceeds to compile the model. However, training speed may not increase and might even decrease relative to the baseline with an untested model. You might need to retune training parameters such as batch_size and learning_rate to achieve any speedup benefits.

If compilation of the untested model fails, the compiler returns an error. See SageMaker Training Compiler Troubleshooting for detailed information about the failure types and error messages.

**Q. Will I always get a faster training job with SageMaker Training Compiler?**

No, not necessarily. First, SageMaker Training Compiler adds some compilation overhead before the ongoing training process can be accelerated. The optimized training job must run sufficiently long to amortize and make up for this incremental compilation overhead at the beginning of the training job.

Additionally, as with any model training process, training with suboptimal parameters can increase training time. SageMaker Training Compiler can change the characteristics of the training job by, for example, changing the memory footprint of the job. Because of these differences, you might need to retune your training job parameters to speed up training. A reference table specifying the best performing parameters for training jobs with different instance types and models can be found at Tested Models.

Finally, some code in a training script might add additional overhead or disrupt the compiled computation graph and slow training. If working with a customized or untested model, see the instructions at Best Practices to Use SageMaker Training Compiler with PyTorch/XLA.

**Q. Can I always use a larger batch size with SageMaker Training Compiler?**

Batch size increases in most, but not all, cases. The optimizations made by SageMaker Training Compiler can change the characteristics of your training job, such as the memory footprint. Typically, a Training Compiler job occupies less memory than an uncompiled training job with the native framework, which allows for a larger batch size during training. A larger batch size, and a corresponding adjustment to the learning rate, increases training throughput and can decrease total training time.

However, there could be cases where SageMaker Training Compiler might actually increase memory footprint based on its optimization scheme. The compiler uses an analytical cost model to predict
the execution schedule with the lowest cost of execution for any compute-intensive operator. This model could find an optimal schedule that increases memory use. In this case, you won’t be able to increase batch sizes, but your sample throughput is still higher.

**Q. Does SageMaker Training Compiler work with other SageMaker training features, such as the SageMaker distributed training libraries and SageMaker Debugger?**

SageMaker Training Compiler is currently not compatible with SageMaker’s distributed training libraries.

SageMaker Training Compiler is compatible with SageMaker Debugger, but Debugger might degrade computational performance by adding overhead.

**Q. Does SageMaker Training Compiler support custom containers (bring your own container)?**

SageMaker Training Compiler is provided through AWS Deep Learning Containers, and you can extend a subset of the containers to customize for your use-case. Containers that are extended from AWS DLCs are supported by SageMaker Training Compiler. For more information, see [Supported Frameworks](#) and [Using the SageMaker Python SDK and Extending SageMaker Framework Deep Learning Containers](#). If you need further support, reach out to the SageMaker team through [AWS Support](#) or [AWS Developer Forums for Amazon SageMaker](#).

**SageMaker Training Compiler Troubleshooting**

If you run into an error, you can use the following list to try to troubleshoot your training job. If you need further support, reach out to the SageMaker team through [AWS Support](#) or [AWS Developer Forums for Amazon SageMaker](#).

### Training job is not converging as expected when compared to the native framework training job

Convergence issues range from “the model is not learning when SageMaker Training Compiler is turned on” to “the model is learning but slower than the native framework”. In this troubleshooting guide, we assume your convergence is fine without SageMaker Training Compiler (in the native framework) and consider this the baseline.

When faced with such convergence issues, the first step is to identify if the issue is limited to distributed training or stems from single-GPU training. Distributed training with SageMaker Training Compiler is an extension of single-GPU training with additional steps.
1. Set up a cluster with multiple instances or GPUs.
2. Distribute input data to all workers.
3. Synchronize the model updates from all workers.

Therefore, any convergence issue in single-GPU training propagates to distributed training with multiple workers.
A flow chart to troubleshoot convergence issues in training jobs when using SageMaker Training Compiler. Descriptions are in the following sections.

**Convergence issues occurring in single-GPU training**

If your convergence issue stems from single-GPU training, this is likely due to improper settings for hyperparameters or the `torch_xla` APIs.

**Check the hyperparameters**

Training with SageMaker Training Compiler leads to change in the memory footprint of a model. The compiler intelligently arbitrates between re-use and re-compute leading to a corresponding increase or decrease in memory consumption. To leverage this, it is essential to re-tune the batch size and associated hyperparameters when migrating a training job to SageMaker Training Compiler. However, incorrect hyperparameter settings often cause oscillation in training loss and possibly a slower convergence as a result. In rare cases, aggressive hyperparameters might result in the model not learning (the training loss metric doesn’t decrease or returns NaN). To identify if the convergence issue is due to the hyperparameters, do a side-by-side test of two training jobs with and without SageMaker Training Compiler while keeping all the hyperparameters the same.

**Check if the `torch_xla` APIs are properly set up for single-GPU training**

If the convergence issue persists with the baseline hyperparameters, you need to check if there's any improper usage of the `torch_xla` APIs, specifically the ones for updating the model. Fundamentally, `torch_xla` continues to accumulate instructions (deferring execution) in the form of graph until it is explicitly instructed to run the accumulated graph. The `torch_xla.core.xla_model.mark_step()` function facilitates the execution of the accumulated graph. The graph execution should be synchronized using this function after each model update and before printing and logging any variables. If it lacks the synchronization step, the model might use stale values from memory during prints, logs, and the subsequent forward passes, instead of using the most recent values that have to be synchronized after every iteration and model update.

It can be more complicated when using SageMaker Training Compiler with gradient scaling (possibly from the use of AMP) or gradient clipping techniques. The appropriate order of gradient computation with AMP is as follows.

1. Gradient computation with scaling
2. Gradient un-scaling, gradient clipping, and then scaling
3. Model update
4. Synchronizing the graph execution with `mark_step()`

To find the right APIs for the operations mentioned in the list, see the guide for [migrating your training script to SageMaker Training Compiler](https://docs.aws.amazon.com/sagemaker/latest/dg/).

**Consider using Automatic Model Tuning**

If the convergence issue arises when re-tuning the batch size and associated hyperparameters such as the learning rate while using SageMaker Training Compiler, consider using [Automatic Model Tuning](https://docs.aws.amazon.com/sagemaker/latest/dg/) to tune your hyperparameters. You can refer to the [example notebook on tuning hyperparameters with SageMaker Training Compiler](https://docs.aws.amazon.com/sagemaker/latest/dg/).

**Convergence issues occurring in distributed training**

If your convergence issue persists in distributed training, this is likely due to improper settings for weight initialization or the `torch_xla` APIs.

**Check weight initialization across the workers**

If the convergence issue arises when running a distributed training job with multiple workers, ensure there is a uniform deterministic behavior across all workers by setting a constant seed where applicable. Beware of techniques such as weight initialization, which involves randomization. Each worker might end up training a different model in the absence of a constant seed.

**Check if the `torch_xla` APIs are properly set up for distributed training**

If the issue still persists, this is likely due to improper use of the `torch_xla` APIs for distributed training. Make sure that you add the following in your estimator to set up a cluster for distributed training with SageMaker Training Compiler.

```python
distribution={'torchxla': {'enabled': True}}
```

This should be accompanied by a function `_mp_fn(index)` in your training script, which is invoked once per worker. Without the `_mp_fn(index)` function, you might end up letting each of the workers train the model independently without sharing model updates.

Next, make sure that you use the `torch_xla.distributed.parallel_loader.MpDeviceLoader` API along with the distributed data sampler, as guided in the documentation about [migrating your training script to SageMaker Training Compiler](https://docs.aws.amazon.com/sagemaker/latest/dg/), as in the following example.
This ensures that the input data is properly distributed across all workers.

Finally, to synchronize model updates from all workers, use `torch_xla.core.xla_model._fetch_gradients` to gather gradients from all workers and `torch_xla.core.xla_model.all_reduce` to combine all the gathered gradients into a single update.

It can be more complicated when using SageMaker Training Compiler with gradient scaling (possibly from use of AMP) or gradient clipping techniques. The appropriate order of gradient computation with AMP is as follows.

1. Gradient computation with scaling
2. Gradient synchronization across all workers
3. Gradient un-scaling, gradient clipping, and then gradient scaling
4. Model update
5. Synchronizing the graph execution with `mark_step()`

Note that this checklist has an additional item for synchronizing all workers, compared to the checklist for single-GPU training.

**Training job fails due to missing PyTorch/XLA configuration**

If a training job fails with the `Missing XLA configuration` error message, it might be due to a misconfiguration in the number of GPUs per instance that you use.

XLA requires additional environment variables to compile the training job. The most common missing environment variable is `GPU_NUM_DEVICES`. For the compiler to work properly, you must set this environment variable equal to the number of GPUs per instance.

There are three approaches to set the `GPU_NUM_DEVICES` environment variable:

- **Approach 1** – Use the `environment` argument of the SageMaker estimator class. For example, if you use an `ml.p3.8xlarge` instance that has four GPUs, do the following:

```python
# Using the SageMaker Python SDK's HuggingFace estimator
hf_estimator=HuggingFace()
```
instance_type="ml.p3.8xlarge",
hyperparameters={...},
environment={
    ...
    "GPU_NUM_DEVICES": "4" # corresponds to number of GPUs on the specified instance
},
)

• **Approach 2** – Use the hyperparameters argument of the SageMaker estimator class and parse it in your training script.

1. To specify the number of GPUs, add a key-value pair to the hyperparameters argument.

   For example, if you use an `ml.p3.8xlarge` instance that has four GPUs, do the following:

   ```python
   # Using the SageMaker Python SDK's HuggingFace estimator
   hf_estimator=HuggingFace(
       ...
       entry_point = "train.py"
       instance_type= "ml.p3.8xlarge",
       hyperparameters = {
           ...
           "n_gpus": 4 # corresponds to number of GPUs on specified instance
       }
   )
   hf_estimator.fit()
   ```

2. In your training script, parse the `n_gpus` hyperparameter and specify it as an input for the `GPU_NUM_DEVICES` environment variable.

   ```python
   # train.py
   import os, argparse

   if __name__ == "__main__":
       parser = argparse.ArgumentParser()

   # Data, model, and output directories
   parser.add_argument("--output_data_dir", type=str,
       default=os.environ["SM_OUTPUT_DATA_DIR"])
   parser.add_argument("--model_dir", type=str,
       default=os.environ["SM_MODEL_DIR"])
   ```
Approach 3 – Hard-code the GPU_NUM_DEVICES environment variable in your training script. For example, add the following to your script if you use an instance that has four GPUs.

```python
# train.py
import os
os.environ['GPU_NUM_DEVICES'] = 4
```

Tip

To find the number of GPU devices on machine learning instances that you want to use, see Accelerated Computing in the Amazon EC2 Instance Types page.

SageMaker Training Compiler doesn't reduce the total training time

If the total training time does not decrease with SageMaker Training Compiler, we highly recommend you to go over the SageMaker Training Compiler Best Practices and Considerations page to check your training configuration, padding strategy for the input tensor shape, and hyperparameters.

Amazon SageMaker Training Compiler Release Notes

See the following release notes to track the latest updates for Amazon SageMaker Training Compiler.
SageMaker Training Compiler Release Notes: February 13, 2023

Currency Updates

- Added support for PyTorch v1.13.1

Bug Fixes

- Fixed a race condition issue on GPU which was causing NAN loss in some models like vision transformer (ViT) models.

Other Changes

- SageMaker Training Compiler improves performance by letting PyTorch/ XLA to automatically override the optimizers (such as SGD, Adam, AdamW) in torch.optim or transformers.optimization with the syncfree versions of them in torch_xla.amp.syncfree (such as torch_xla.amp.syncfree.SGD, torch_xla.amp.syncfree.Adam, torch_xla.amp.syncfree.AdamW). You don't need to change those code lines where you define optimizers in your training script.

Migration to AWS Deep Learning Containers

This release passed benchmark testing and is migrated to the following AWS Deep Learning Container:

- PyTorch v1.13.1


To find a complete list of the prebuilt containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.

SageMaker Training Compiler Release Notes: January 9, 2023

Breaking Changes
tf.keras.optimizers.Optimizer points to a new optimizer in TensorFlow 2.11.0 and later. The old optimizers are moved to tf.keras.optimizers.legacy. You might encounter job failure due to the breaking change when you do the following.

- Load checkpoints from an old optimizer. We recommend you to switch to use the legacy optimizers.
- Use TensorFlow v1. We recommend you to migrate to TensorFlow v2, or switch to the legacy optimizers if you need to continue using TensorFlow v1.

For more detailed list of breaking changes from the optimizer changes, see the official TensorFlow v2.11.0 release notes in the TensorFlow GitHub repository.

Migration to AWS Deep Learning Containers

This release passed benchmark testing and is migrated to the following AWS Deep Learning Container:

- TensorFlow v2.11.0

763104351884.dkr.ecr.<region>.amazonaws.com/tensorflow-training:2.11.0-gpu-py39-cu112-ubuntu20.04-sagemaker

To find a complete list of the prebuilt containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.

SageMaker Training Compiler Release Notes: December 8, 2022

Bug Fixes

- Fixed the seed for PyTorch training jobs starting PyTorch v1.12 to ensure that there is no discrepancy in model initialization across different processes. See also PyTorch Reproducibility.
- Fixed the issue causing PyTorch distributed training jobs on G4dn and G5 instances to not default to communication through PCIe.

Known Issues

- Improper use of PyTorch/XLA APIs in Hugging Face’s vision transformers might lead to convergence issues.
Other Changes

- When using the Hugging Face Transformers Trainer class, make sure that you use SyncFree optimizers by setting the optim argument to adamw_torch_xla. For more information, see Large Language Models Using the Hugging Face Transformers Trainer Class. See also Optimizer in the Hugging Face Transformers documentation.

Migration to AWS Deep Learning Containers

This release passed benchmark testing and is migrated to the following AWS Deep Learning Container:

- PyTorch v1.12.0


To find a complete list of the prebuilt containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.

SageMaker Training Compiler Release Notes: October 4, 2022

Currency Updates

- Added support for TensorFlow v2.10.0.

Other Changes

- Added Hugging Face NLP models using the Transformers library to TensorFlow framework tests. To find the tested Transformer models, see the section called “Tested Models”.

Migration to AWS Deep Learning Containers

This release passed benchmark testing and is migrated to the following AWS Deep Learning Container:

- TensorFlow v2.10.0
To find a complete list of the prebuilt containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.

SageMaker Training Compiler Release Notes: September 1, 2022

Currency Updates

- Added support for Hugging Face Transformers v4.21.1 with PyTorch v1.11.0.

Improvements

- Implemented a new distributed training launcher mechanism to activate SageMaker Training Compiler for Hugging Face Transformer models with PyTorch. To learn more, see Run PyTorch Training Jobs with SageMaker Training Compiler for Distributed Training.
- Integrated with EFA to improve the collective communication in distributed training.
- Added support for G5 instances for PyTorch training jobs. For more information, see the section called “Supported Frameworks, AWS Regions, Instance Types, and Tested Models”.

Migration to AWS Deep Learning Containers

This release passed benchmark testing and is migrated to the following AWS Deep Learning Container:

- HuggingFace v4.21.1 with PyTorch v1.11.0

To find a complete list of the prebuilt containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.
SageMaker Training Compiler Release Notes: June 14, 2022

New Features

• Added support for TensorFlow v2.9.1. SageMaker Training Compiler fully supports compiling TensorFlow modules (tf.*) and TensorFlow Keras modules (tf.keras.*).

• Added support for custom containers created by extending AWS Deep Learning Containers for TensorFlow. For more information, see Enable SageMaker Training Compiler Using the SageMaker Python SDK and Extending SageMaker Framework Deep Learning Containers.

• Added support for G5 instances for TensorFlow training jobs.

Migration to AWS Deep Learning Containers

This release passed benchmark testing and is migrated to the following AWS Deep Learning Container:

• TensorFlow 2.9.1

763104351884.dkr.ecr.<region>.amazonaws.com/tensorflow-training:2.9.1-gpu-py39-cu112-ubuntu20.04-sagemaker

To find a complete list of the pre-built containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.

SageMaker Training Compiler Release Notes: April 26, 2022

Improvements

• Added support for all of the AWS Regions where AWS Deep Learning Containers are in service except the China regions.

SageMaker Training Compiler Release Notes: April 12, 2022

Currency Updates

• Added support for Hugging Face Transformers v4.17.0 with TensorFlow v2.6.3 and PyTorch v1.10.2.
SageMaker Training Compiler Release Notes: February 21, 2022

Improvements

• Completed benchmark test and confirmed training speed-ups on the ml.g4dn instance types. To find a complete list of tested ml instances, see Supported Instance Types.

SageMaker Training Compiler Release Notes: December 01, 2021

New Features

• Launched Amazon SageMaker Training Compiler at AWS re:Invent 2021.

Migration to AWS Deep Learning Containers

• Amazon SageMaker Training Compiler passed benchmark testing and is migrated to AWS Deep Learning Containers. To find a complete list of the prebuilt containers with Amazon SageMaker Training Compiler, see Supported Frameworks, AWS Regions, Instance Types, and Tested Models.

Access Training Data

When you create a training job, you specify the location of a training dataset and an input mode for accessing the dataset. For data location, Amazon SageMaker supports Amazon Simple Storage Service (Amazon S3), Amazon Elastic File System (Amazon EFS), and Amazon FSx for Lustre. The input modes determine whether to stream data files of the dataset in real time or download the whole dataset at the start of the training job.

ℹ️ Note

Your input dataset must be in the same AWS Region as your training job.

SageMaker Input Modes and AWS Cloud Storage

This section summarizes SageMaker input modes for Amazon S3 and file systems in Amazon EFS and Amazon FSx for Lustre.
- **File mode** presents a file system view of the dataset to the training container. This is the default input mode if you don't explicitly specify one of the other two options. If you use file mode, SageMaker downloads the training data from the storage location to a local directory in the Docker container. Training starts after the full dataset has been downloaded. In file mode, the training instance must have enough storage space to fit the entire dataset. File mode download speed depends on the size of dataset, the average size of files, and the number of files. You can configure the dataset for file mode by providing either an Amazon S3 prefix, manifest file, or augmented manifest file. You should use an S3 prefix when all your dataset files are located within a common S3 prefix. File mode is compatible with **SageMaker local mode** (starting a SageMaker training container interactively in seconds). For distributed training, you can shard the dataset across multiple instances with the `ShardedByS3Key` option.

- **Fast file mode** provides file system access to an Amazon S3 data source while leveraging the performance advantage of pipe mode. At the start of training, fast file mode identifies the data files but does not download them. Training can start without waiting for the entire dataset to download. This means that the training startup takes less time when there are fewer files in the Amazon S3 prefix provided.
In contrast to pipe mode, fast file mode works with random access to the data. However, it works best when data is read sequentially. Fast file mode doesn't support augmented manifest files.

Fast file mode exposes S3 objects using a POSIX-compliant file system interface, as if the files are available on the local disk of your training instance. It streams S3 content on demand as your training script consumes data. This means that your dataset no longer needs to fit into the training instance storage space as a whole, and you don't need to wait for the dataset to be downloaded to the training instance before training starts. Fast file currently supports S3 prefixes only (it does not support manifest and augmented manifest). Fast file mode is compatible with SageMaker local mode.

- **Pipe mode** streams data directly from an Amazon S3 data source. Streaming can provide faster start times and better throughput than file mode.

When you stream the data directly, you can reduce the size of the Amazon EBS volumes used by the training instance. Pipe mode needs only enough disk space to store the final model artifacts.

It is another streaming mode that is largely replaced by the newer and simpler-to-use fast file mode. In pipe mode, data is pre-fetched from Amazon S3 at high concurrency and throughput, and streamed into a named pipe, which also known as a First-In-First-Out (FIFO) pipe for its behavior. Each pipe may only be read by a single process. A SageMaker specific extension to TensorFlow conveniently integrates Pipe mode into the native TensorFlow data loader for streaming text, TFRecords, or RecordIO file formats. Pipe mode also supports managed sharding and shuffling of data.

- **Amazon S3 Express One Zone** is a high-performance, single Availability Zone storage class that can deliver consistent, single-digit millisecond data access for the most latency-sensitive applications including SageMaker model training. Amazon S3 Express One Zone allows customers to collocate their object storage and compute resources in a single AWS Availability Zone, optimizing both compute performance and costs with increased data processing speed. To further increase access speed and support hundreds of thousands of requests per second, data is stored in a new bucket type, an Amazon S3 directory bucket.

SageMaker model training supports high-performance Amazon S3 Express One Zone directory buckets as a data input location for file mode, fast file mode, and pipe mode. To use Amazon S3 Express One Zone, input the location of the Amazon S3 Express One Zone directory bucket instead of an Amazon S3 bucket. Provide the ARN for the IAM role with the required access control and permissions policy. Refer to [AmazonSageMakerFullAccesspolicy](https://aws.amazon.com/service-quotas-docs/) for details. For more information, see [Amazon S3 Express One Zone](https://aws.amazon.com/aws-solutions/1357/).
Amazon FSx for Lustre – FSx for Lustre can scale to hundreds of gigabytes of throughput and millions of IOPS with low-latency file retrieval. When starting a training job, SageMaker mounts the FSx for Lustre file system to the training instance file system, then starts your training script. Mounting itself is a relatively fast operation that doesn't depend on the size of the dataset stored in FSx for Lustre.

To access FSx for Lustre, your training job must connect to an Amazon Virtual Private Cloud (VPC), which requires DevOps setup and involvement. To avoid data transfer costs, the file system uses a single Availability Zone, and you need to specify a VPC subnet which maps to this Availability Zone ID when running the training job.

Amazon EFS – To use Amazon EFS as a data source, the data must already reside in Amazon EFS prior to training. SageMaker mounts the specified Amazon EFS file system to the training instance, then starts your training script. Your training job must connect to a VPC to access Amazon EFS.

Tip

To learn more about how to specify your VPC configuration to SageMaker estimators, see Use File Systems as Training Inputs in the SageMaker Python SDK documentation.

Choosing Data Input Mode Using the SageMaker Python SDK

SageMaker Python SDK provides the generic Estimator class and its variations for ML frameworks for launching training jobs. You can specify one of the data input modes while configuring the SageMaker Estimator class or the Estimator.fit method. The following code templates show the two ways to specify input modes.

To specify the input mode using the Estimator class

```python
from sagemaker.estimator import Estimator
from sagemaker.inputs import TrainingInput

estimator = Estimator(
    checkpoint_s3_uri='s3://my-bucket/checkpoint-destination/",
    output_path='s3://my-bucket/output-path/",
    base_job_name='job-name",
    input_mode='File'  # Available options: File | Pipe | FastFile
    ...
```
# Run the training job
estimator.fit(
    inputs=TrainingInput(s3_data="s3://my-bucket/my-data/train")
)

For more information, see the `sagemaker.estimator.Estimator` class in the *SageMaker Python SDK documentation*.

**To specify the input mode through the Estimator fit method**

```python
from sagemaker.estimator import Estimator
from sagemaker.inputs import TrainingInput

estimator = Estimator(
    checkpoint_s3_uri='s3://my-bucket/checkpoint-destination/',
    output_path='s3://my-bucket/output-path/',
    base_job_name='job-name',
    ...
)

# Run the training job
estimator.fit(
    inputs=TrainingInput(
        s3_data="s3://my-bucket/my-data/train",
        input_mode='File'  # Available options: File | Pipe | FastFile
    )
)
```

For more information, see the `sagemaker.estimator.Estimator.fit` class method and the `sagemaker.inputs.TrainingInput` class in the *SageMaker Python SDK documentation*.

**Tip**

To learn more about how to configure Amazon FSx for Lustre or Amazon EFS with your VPC configuration using the SageMaker Python SDK estimators, see [Use File Systems as Training Inputs](#) in the *SageMaker Python SDK documentation*. 
Tip
The data input mode integrations with Amazon S3, Amazon EFS, and FSx for Lustre are recommended ways to optimally configure data source for the best practices. You can strategically improve data loading performance using the SageMaker managed storage options and input modes, but it's not strictly constrained. You can write your own data reading logic directly in your training container. For example, you can set to read from a different data source, write your own S3 data loader class, or use third-party frameworks' data loading functions within your training script. However, you must make sure that you specify the right paths that SageMaker can recognize.

Tip
If you use a custom training container, make sure you install the SageMaker training toolkit that helps set up the environment for SageMaker training jobs. Otherwise, you must specify the environment variables explicitly in your Dockerfile. For more information, see Create a container with your own algorithms and models.

For more information about how to set the data input modes using the low-level SageMaker APIs, see How Amazon SageMaker Provides Training Information, the CreateTrainingJob API, and the TrainingInputMode in AlgorithmSpecification.

Configure Data Input Channel to Use Amazon FSx for Lustre

Learn how to use Amazon FSx for Lustre as your data source for higher throughput and faster training by reducing the time for data loading.

Sync Amazon S3 and Amazon FSx for Lustre

To link your Amazon S3 to Amazon FSx for Lustre and upload your training datasets, do the following.

1. Prepare your dataset and upload to an Amazon S3 bucket. For example, assume that the Amazon S3 paths for a train dataset and a test dataset are in the following format.

```
s3://my-bucket/data/train
```
2. To create an FSx for Lustre file system linked with the Amazon S3 bucket with the training data, follow the steps at Linking your file system to an Amazon S3 bucket in the Amazon FSx for Lustre User Guide. Make sure that you add an endpoint to your VPC allowing Amazon S3 access. For more information, see the section called “Create an Amazon S3 VPC Endpoint”. When you specify Data repository path, provide the Amazon S3 bucket URI of the folder that contains your datasets. For example, based on the example S3 paths in step 1, the data repository path should be the following.

s3://my-bucket/data

3. After the FSx for Lustre file system is created, check the configuration information by running the following commands.

```bash
aws fsx describe-file-systems &&
aws fsx describe-data-repository-association
```

These commands return FileSystemId, MountName, FileSystemPath, and DataRepositoryPath. For example, the outputs should look like the following.

```
# Output of aws fsx describe-file-systems
"FileSystemId": "fs-0123456789abcdef0"
"MountName": "1234abcd"

# Output of aws fsx describe-data-repository-association
"FileSystemPath": "/ns1",
"DataRepositoryPath": "s3://my-bucket/data/"
```

After the sync between Amazon S3 and Amazon FSx has completed, your datasets are saved in Amazon FSx in the following directories.

```
/ns1/train  # synced with s3://my-bucket/data/train
/ns1/test   # synced with s3://my-bucket/data/test
```
Set the Amazon FSx file system path as the data input channel for SageMaker training

The following procedures walk you through the process of setting the Amazon FSx file system as the data source for SageMaker training jobs.

Using the SageMaker Python SDK

To properly set the Amazon FSx file system as the data source, configure the SageMaker estimator classes and FileSystemInput using the following instruction.

1. Configure a FileSystemInput class object.

```python
from sagemaker.inputs import FileSystemInput

train_fs = FileSystemInput(
    file_system_id="fs-0123456789abcdef0",
    file_system_type="FSxLustre",
    directory_path="/1234abcd/ns1/",
    file_system_access_mode="ro",
)
```

**Tip**

When you specify directory_path, make sure that you provide the Amazon FSx file system path starting with MountName.

2. Configure a SageMaker estimator with the VPC configuration used for the Amazon FSx file system.

```python
from sagemaker.estimator import Estimator

estimator = Estimator(
    ...
    role="your-iam-role-with-access-to-your-fsx",
    subnets=["subnet-id"],  # Should be the same as the subnet used for Amazon FSx
    security_group_ids="security-group-id"
)
```

3. Launch the training job by running the estimator.fit method with the Amazon FSx file system.
estimator.fit(train_fs)

To find more code examples, see Use File Systems as Training Inputs in the SageMaker Python SDK documentation.

Using the SageMaker CreateTrainingJob API

As part of the CreateTrainingJob request JSON, configure InputDataConfig as follows.

```json
"InputDataConfig": [
  {
    "ChannelName": "string",
    "DataSource": {
      "FileSystemDataSource": {
        "DirectoryPath": "/1234abcd/ns1/",
        "FileSystemAccessMode": "ro",
        "FileSystemId": "fs-0123456789abcdef0",
        "FileSystemType": "FSxLustre"
      }
    }
  }
],
```

Tip
---

When you specify DirectoryPath, make sure that you provide the Amazon FSx file system path starting with MountName.

Tips and Considerations When Configuring FSx for Lustre

1. When you use EFA-enabled instances such as P4d and P3dn, make sure that you set appropriate inbound and output rules in the security group. Specially, opening up these ports is necessary for SageMaker to access the Amazon FSx file system in the training job. To learn more, see File System Access Control with Amazon VPC.

2. Make sure the IAM Role used to launch the SageMaker training job has access to Amazon FSx.
Best Practices for Choosing Data Source and Input Mode

The best data source for your training job depends on workload characteristics such as the size of the dataset, the file format, the average size of files, the training duration, a sequential or random data loader read pattern, and how fast your model can consume the training data. The following best practices provide guidelines to get started with the most suitable input mode and data storage for your use case.

This flowchart summarizes and visualizes best practices of choosing the best storage as the data source and input file mode. All of the cases in the flowchart are described in the following sections.
When to use Amazon EFS

If your dataset is stored in Amazon Elastic File System, you might have a preprocessing or annotations application that uses Amazon EFS for storage. You can run a training job configured with a data channel that points to the Amazon EFS file system. For more information, see Speed up training on Amazon SageMaker using Amazon FSx for Lustre and Amazon EFS file systems. If you cannot achieve better performance, check your optimization options following the Amazon Elastic File System performance guide or consider using different input modes or data storage.

Use file mode for small datasets

If the dataset is stored in Amazon Simple Storage Service and its overall volume is relatively small (for example, less than 50-100 GB), try using file mode. The overhead of downloading a 50 GB dataset can vary based on the total number of files. For example, it takes about 5 minutes if a dataset is chunked into 100 MB shards. Whether this startup overhead is acceptable primarily depends on the overall duration of your training job, because a longer training phase means a proportionally smaller download phase.

Serializing many small files

If your dataset size is small (less than 50-100 GB), but is made up of many small files (less than 50 MB per file), the file mode download overhead grows, because each file needs to be downloaded individually from Amazon Simple Storage Service to the training instance volume. To reduce this overhead and data traversal time in general, consider serializing groups of such small files into fewer larger file containers (such as 150 MB per file) by using file formats, such as TFRecord for TensorFlow, WebDataset for PyTorch, and RecordIO for MXNet.

When to use fast file mode

For larger datasets with larger files (more than 50 MB per file), the first option is to try fast file mode, which is more straightforward to use than FSx for Lustre because it doesn't require creating a file system, or connecting to a VPC. Fast file mode is ideal for large file containers (more than 150 MB), and might also do well with files more than 50 MB. Because fast file mode provides a POSIX interface, it supports random reads (reading non-sequential byte-ranges). However, this is not the ideal use case, and your throughput might be lower than with the sequential reads. However, if you have a relatively large and computationally intensive ML model, fast file mode might still be able to saturate the effective bandwidth of the training pipeline and not result in an IO bottleneck. You'll need to experiment and see. To switch from file mode to fast file mode (and back), just add
(or remove) the input_mode='FastFile' parameter while defining your input channel using the SageMaker Python SDK:

```python
sagemaker.inputs.TrainingInput(S3_INPUT_FOLDER, input_mode = 'FastFile')
```

**When to use Amazon FSx for Lustre**

If your dataset is too large for file mode, has many small files that you can't serialize easily, or uses a random read access pattern, FSx for Lustre is a good option to consider. Its file system scales to hundreds of gigabytes per second (GB/s) of throughput and millions of IOPS, which is ideal when you have many small files. However, note that there might be the cold start issue due to lazy loading and the overhead of setting up and initializing the FSx for Lustre file system.

⚠️ **Tip**

To learn more, see Choose the best data source for your Amazon SageMaker training job. This AWS machine learning blog further discusses case studies and performance benchmark of data sources and input modes.

**Train Using a Heterogeneous Cluster**

Using the heterogeneous cluster feature of SageMaker Training, you can run a training job with multiple types of ML instances for a better resource scaling and utilization for different ML training tasks and purposes. For example, if your training job on a cluster with GPU instances suffers low GPU utilization and CPU bottleneck problems due to CPU-intensive tasks, using a heterogeneous cluster can help offload CPU-intensive tasks by adding more cost-efficient CPU instance groups, resolve such bottleneck problems, and achieve a better GPU utilization.

⚠️ **Note**

This feature is available in the SageMaker Python SDK v2.98.0 and later.
How to Configure a Heterogeneous Cluster

This section provides instructions on how to run a training job using a heterogeneous cluster that consists of multiple instance types.

Topics

- Using the SageMaker Python SDK
- Using the Low-Level SageMaker APIs

Using the SageMaker Python SDK

Follow instructions on how to configure instance groups for a heterogeneous cluster using the SageMaker Python SDK.

1. To configure instance groups of a heterogeneous cluster for a training job, use the sagemaker.instance_group.InstanceGroup class. You can specify a custom name for each instance group, the instance type, and the number of instances for each instance group. For more information, see sagemaker.instance_group.InstanceGroup in the SageMaker Python SDK documentation.
The following code example shows how to set up two instance groups that consists of two **ml.c5.18xlarge** CPU-only instances named `instance_group_1` and one **ml.p3dn.24xlarge** GPU instance named `instance_group_2`, as shown in the following diagram.

```python
from sagemaker.instance_group import InstanceGroup

instance_group_1 = InstanceGroup(
    "instance_group_1", "ml.c5.18xlarge", 2
)
instance_group_2 = InstanceGroup(
    "instance_group_2", "ml.p3dn.24xlarge", 1
)
```

2. Using the instance group objects, set up training input channels and assign instance groups to the channels through the `instance_group_names` argument of the `sagemaker.inputs.TrainingInput` class. The `instance_group_names` argument accepts a list of strings of instance group names.
The following example shows how to set two training input channels and assign the instance groups created in the example of the previous step. You can also specify Amazon S3 bucket paths to the `s3_data` argument for the instance groups to process data for your usage purposes.

```python
code_snippet
from sagemaker.inputs import TrainingInput

training_input_channel_1 = TrainingInput(  
    s3_data_type='S3Prefix', # Available Options: S3Prefix | ManifestFile | AugmentedManifestFile  
    s3_data='s3://your-training-data-storage/folder1',  
    distribution='FullyReplicated', # Available Options: FullyReplicated | ShardedByS3Key  
    input_mode='File', # Available Options: File | Pipe | FastFile  
    instance_groups=['instance_group_1']
)

training_input_channel_2 = TrainingInput(  
    s3_data_type='S3Prefix',  
    s3_data='s3://your-training-data-storage/folder2',  
    distribution='FullyReplicated',  
    input_mode='File',  
    instance_groups=['instance_group_2']
)
```

For more information about the arguments of `TrainingInput`, see the following links.

- The `sagemaker.inputs.TrainingInput` class in the *SageMaker Python SDK documentation*
- The `S3DataSource` API in the *SageMaker API Reference*

3. Configure a SageMaker estimator with the `instance_groups` argument as shown in the following code example. The `instance_groups` argument accepts a list of InstanceGroup objects.

**PyTorch**

```python
code_snippet
from sagemaker.pytorch import PyTorch

estimator = PyTorch(  
    ...  
    entry_point='my-training-script.py',  
    framework_version='x.y.z', # 1.10.0 or later
)
```

How to Configure a Heterogeneous Cluster
TensorFlow

```python
from sagemaker.tensorflow import TensorFlow

estimator = TensorFlow(
    ...
    entry_point='my-training-script.py',
    framework_version='x.y.z', # 2.6.0 or later
    py_version='pyxy',
    job_name='my-training-job-with-heterogeneous-cluster',
    instance_groups=[instance_group_1, instance_group_2]
)
```

**Note**

The instance_type and instance_count argument pair and the instance_groups argument of the SageMaker estimator class are mutually exclusive. For homogeneous cluster training, use the instance_type and instance_count argument pair. For heterogeneous cluster training, use instance_groups.

**Note**

To find a complete list of available framework containers, framework versions, and Python versions, see SageMaker Framework Containers in the AWS Deep Learning Container GitHub repository.

4. Configure the `estimator.fit` method with the training input channels configured with the instance groups and start the training job.

```python
estimator.fit(
    inputs={
        'training': training_input_channel_1,
        'dummy-input-channel': training_input_channel_2
    }
```
Using the Low-Level SageMaker APIs

If you use the AWS Command Line Interface or AWS SDK for Python (Boto3) and want to use low-level SageMaker APIs for submitting a training job request with a heterogeneous cluster, see the following API references.

- CreateTrainingJob
- ResourceConfig
- InstanceGroup
- S3DataSource

Distributed Training with a Heterogeneous Cluster

Through the distribution argument of the SageMaker estimator class, you can assign a specific instance group to run distributed training. For example, assume that you have the following two instance groups and want to run multi-GPU training on one of them.

```python
from sagemaker.instance_group import InstanceGroup

instance_group_1 = InstanceGroup("instance_group_1", "ml.c5.18xlarge", 1)
instance_group_2 = InstanceGroup("instance_group_2", "ml.p3dn.24xlarge", 2)
```

You can set the distributed training configuration for one of the instance groups. For example, the following code examples show how to assign `training_group_2` with two `ml.p3dn.24xlarge` instances to the distributed training configuration.

**Note**

Currently, only one instance group of a heterogeneous cluster can be specified to the distribution configuration.

With MPI
## PyTorch

```python
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    ...
    instance_groups=[instance_group_1, instance_group_2],
    distribution={
        "mpi": {
            "enabled": True, "processes_per_host": 8
        },
        "instance_groups": [instance_group_2]
    }
)
```

## TensorFlow

```python
from sagemaker.tensorflow import TensorFlow

estimator = TensorFlow(
    ...
    instance_groups=[instance_group_1, instance_group_2],
    distribution={
        "mpi": {
            "enabled": True, "processes_per_host": 8
        },
        "instance_groups": [instance_group_2]
    }
)
```

## With the SageMaker data parallel library

### PyTorch

```python
from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    ...
    instance_groups=[instance_group_1, instance_group_2],
    distribution={
        "smdistributed": {
            "dataparallel": {
                ""dataparallel": {
```
TensorFlow

from sagemaker.tensorflow import TensorFlow

estimator = TensorFlow(
    ...
    instance_groups=[instance_group_1, instance_group_2],
    distribution={
        "smdistributed": {
            "dataparallel": {
                "enabled": True
            }
        },
        "instance_groups": [instance_group_2]
    }
)

Note

When using the SageMaker data parallel library, make sure the instance group consists of the supported instance types by the library.

For more information about the SageMaker data parallel library, see SageMaker Data Parallel Training.

With the SageMaker model parallel library

PyTorch

from sagemaker.pytorch import PyTorch

estimator = PyTorch(
    ...
)
instance_groups=[instance_group_1, instance_group_2], distribution=
  "smdistributed": {
    "modelparallel": {
      "enabled": True,
      "parameters": {
        ...  # SageMaker model parallel parameters
      }
    }
  },
  "instance_groups": [instance_group_2]
)}

TensorFlow

from sagemaker.tensorflow import TensorFlow

estimator = TensorFlow(
  ...  
  instance_groups=[instance_group_1, instance_group_2],
  distribution=
    "smdistributed": {
      "modelparallel": {
        "enabled": True,
        "parameters": {
          ...  # SageMaker model parallel parameters
        }
      }
    },
    "instance_groups": [instance_group_2]
  )
)

For more information about the SageMaker model parallel library, see [SageMaker Model Parallel Training](#).

**Modify Your Training Script to Assign Instance Groups**

With the heterogeneous cluster configuration in the previous sections, you have prepared the SageMaker training environment and instances for your training job. To further assign the instance
groups to certain training and data processing tasks, the next step is to modify your training script. By default, the training job simply makes training script replicas for all nodes regardless the size of the instance, and this might lead to performance loss.

For example, if you mix CPU instances and GPU instances in a heterogeneous cluster while passing a deep neural network training script to the entry_point argument of the SageMaker estimator, the entry_point script is replicated to each instance. This means that, without proper task assignments, CPU instances also run the entire script and start the training job that’s designed for distributed training on GPU instances. Therefore, you must make changes in specific processing functions that you want to offload and run on the CPU instances. You can use the SageMaker environment variables to retrieve the information of the heterogeneous cluster and let specific processes to run accordingly.

**Query instance group information during the initialization phase of a SageMaker training job**

When your training job starts, your training script reads SageMaker training environment information that includes heterogeneous cluster configuration. The configuration contains information such as the current instance groups, the current hosts in each group, and in which group the current host resides.

You can retrieve instance group information in the following ways.

**(Recommended) Reading instance group information with the SageMaker training toolkit**

Use the environment Python module that the [SageMaker training toolkit library](https://sagemaker.readthedocs.io/en/stable/) provides. The toolkit library is preinstalled in the [SageMaker framework containers](https://github.com/aws/sagemaker) for TensorFlow and PyTorch, so you don’t need an additional installation step when using the prebuilt containers. This is the recommended way to retrieve the SageMaker environment variables with fewer code changes in your training script.

```python
from sagemaker_training import environment

env = environment.Environment()
```

Environment variables related to general SageMaker training and heterogeneous clusters:

- `env.is_hetero` – Returns a Boolean result whether a heterogeneous cluster is configured or not.
- `env.current_host` – Returns the current host.
• `env.current_instance_type` – Returns the type of instance of the current host.
• `env.current_instance_group` – Returns the name of the current instance group.
• `env.current_instance_group_hosts` – Returns a list of hosts in current instance group.
• `env.instance_groups` – Returns a list of instance group names used for training.
• `env.instance_groups_dict` – Returns the entire heterogeneous cluster configuration of the training job.
• `env.distribution_instance_groups` – Returns a list of instance groups assigned to the distribution parameter of the SageMaker estimator class.
• `env.distribution_hosts` – Returns a list of hosts belonging to the instance groups assigned to the distribution parameter of the SageMaker estimator class.

For example, consider the following example of a heterogeneous cluster that consists of two instance groups.

```python
from sagemaker.instance_group import InstanceGroup

instance_group_1 = InstanceGroup(
    "instance_group_1", "ml.c5.18xlarge", 1)
instance_group_2 = InstanceGroup(
    "instance_group_2", "ml.p3dn.24xlarge", 2)
```

The output of `env.instance_groups_dict` of the example heterogeneous cluster should be similar to the following.

```json
{
    "instance_group_1": {
        "hosts": [
            "algo-2"
        ],
        "instance_group_name": "instance_group_1",
        "instance_type": "ml.c5.18xlarge"
    },
    "instance_group_2": {
        "hosts": [
            "algo-3",
            "algo-1"
        ],
        "instance_group_name": "instance_group_2",
```
"instance_type": "ml.p3dn.24xlarge"
}
}

(Optional) Reading instance group information from the resource configuration JSON file

If you prefer to retrieve the environment variables in JSON format, you can directly use the resource configuration JSON file. The JSON file in a SageMaker training instance is located at /opt/ml/input/config/resourceconfig.json by default.

```python
file_path = '/opt/ml/input/config/resourceconfig.json'
config = read_file_as_json(file_path)
print(json.dumps(config, indent=4, sort_keys=True))
```

Considerations

Consider the following items when using the heterogeneous cluster feature.

- All instance groups share the same Docker image and training script. Therefore, your training script should be modified to detect which instance group it belongs to and fork execution accordingly.
- The heterogeneous cluster feature is not supported in SageMaker local mode.
- The Amazon CloudWatch log streams of a heterogeneous cluster training job are not grouped by instance groups. You need to figure out from the logs which nodes are in which group.
- The heterogeneous cluster feature is available through the SageMaker PyTorch and TensorFlow framework estimator classes. Supported frameworks are PyTorch v1.10 or later and TensorFlow v2.6 or later. To find a complete list of available framework containers, framework versions, and Python versions, see SageMaker Framework Containers in the AWS Deep Learning Container GitHub repository.
- A distributed training strategy can be applied only to one instance group.

Examples, Blogs, and Case Studies

The following blog discusses case studies about using the SageMaker heterogeneous cluster training.

- Improve price performance of your model training using Amazon SageMaker heterogeneous clusters (Oct 27, 2022)
Use Incremental Training in Amazon SageMaker

Over time, you might find that a model generates inference that are not as good as they were in the past. With incremental training, you can use the artifacts from an existing model and use an expanded dataset to train a new model. Incremental training saves both time and resources.

Use incremental training to:

- Train a new model using an expanded dataset that contains an underlying pattern that was not accounted for in the previous training and which resulted in poor model performance.
- Use the model artifacts or a portion of the model artifacts from a popular publicly available model in a training job. You don't need to train a new model from scratch.
- Resume a training job that was stopped.
- Train several variants of a model, either with different hyperparameter settings or using different datasets.

For more information about training jobs, see [Train a Model with Amazon SageMaker](#).

You can train incrementally using the SageMaker console or the [Amazon SageMaker Python SDK](#).

⚠️ Important

Only three built-in algorithms currently support incremental training: [Object Detection - MXNet](#), [Image Classification - MXNet](#), and [Semantic Segmentation Algorithm](#).

Topics

- [Perform Incremental Training (Console)](#)
- [Perform Incremental Training (API)](#)

Perform Incremental Training (Console)

To complete this procedure, you need:

- The Amazon Simple Storage Service (Amazon S3) bucket URI where you've stored the training data.
To perform incremental training (console)

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the navigation pane, choose Training, then choose Training jobs.
3. Choose Create training job.
4. Provide a name for the training job. The name must be unique within an AWS Region in an AWS account. The training job name must have 1 to 63 characters. Valid characters: a-z, A-Z, 0-9, and . : + = @ _ % - (hyphen).
5. Choose the algorithm that you want to use. For information about algorithms, see Use Amazon SageMaker Built-in Algorithms or Pre-trained Models.
6. (Optional) For Resource configuration, either leave the default values or increase the resource consumption to reduce computation time.
   a. (Optional) For Instance type, choose the ML compute instance type that you want to use. In most cases, ml.m4.xlarge is sufficient.
   b. For Instance count, use the default, 1.
   c. (Optional) For Additional volume per instance (GB), choose the size of the ML storage volume that you want to provision. In most cases, you can use the default, 1. If you are using a large dataset, use a larger size.
7. Provide information about the input data for the training dataset.
   a. For Channel name, either leave the default (train) or enter a more meaningful name for the training dataset, such as expanded-training-dataset.
   b. For InputMode, choose File. For incremental training, you need to use file input mode.
c. For **S3 data distribution type**, choose **FullyReplicated**. This causes each ML compute instance to use a full replicate of the expanded dataset when training incrementally.

d. If the expanded dataset is uncompressed, set the **Compression type** to **None**. If the expanded dataset is compressed using Gzip, set it to **Gzip**.

e. (Optional) If you are using File input mode, leave **Content type** empty. For Pipe input mode, specify the appropriate MIME type. **Content type** is the multipurpose internet mail extension (MIME) type of the data.

f. For **Record wrapper**, if the dataset is saved in RecordIO format, choose **RecordIO**. If your dataset is not saved as a RecordIO formatted file, choose **None**.

g. For **S3 data type**, if the dataset is stored as a single file, choose **S3Prefix**. If the dataset is stored as several files in a folder, choose **Manifest**.

h. For **S3 location**, provide the URL to the path where you stored the expanded dataset.

i. Choose **Done**.

8. To use model artifacts in a training job, you need to add a new channel and provide the needed information about the model artifacts.

a. For **Input data configuration**, choose **Add channel**.

b. For **Channel name**, enter `model` to identify this channel as the source of the model artifacts.

c. For **InputMode**, choose **File**. Model artifacts are stored as files.

d. For **S3 data distribution type**, choose **FullyReplicated**. This indicates that each ML compute instance should use all of the model artifacts for training.

e. For **Compression type**, choose **None** because we are using a model for the channel.

f. Leave **Content type** empty. Content type is the multipurpose internet mail extension (MIME) type of the data. For model artifacts, we leave it empty.

g. Set **Record wrapper** to **None** because model artifacts are not stored in RecordIO format.

h. For **S3 data type**, if you are using a built-in algorithm or an algorithm that stores the model as a single file, choose **S3Prefix**. If you are using an algorithm that stores the model as several files, choose **Manifest**.

i. For **S3 location**, provide the URL to the path where you stored the model artifacts. Typically, the model is stored with the name `model.tar.gz`. To find the URL for the model artifacts, in the navigation pane, choose **Inference**, then choose **Models**. From the list of models, choose a model to display its details page. The URL for the model artifacts is listed under **Primary container**.
j. Choose Done.

9. For Output data configuration, provide the following information:

   a. For S3 location, type the path to the S3 bucket where you want to store the output data.
   b. (Optional) For Encryption key, you can add your AWS Key Management Service (AWS KMS) encryption key to encrypt the output data at rest. Provide the key ID or its Amazon Resource Number (ARN). For more information, see KMS-Managed Encryption Keys.

10. (Optional) For Tags, add one or more tags to the training job. A tag is metadata that you can define and assign to AWS resources. In this case, you can use tags to help you manage your training jobs. A tag consists of a key and a value, which you define. For example, you might want to create a tag with Project as a key and a value referring to a project that is related to the training job, such as Home value forecasts.

11. Choose Create training job. SageMaker creates and runs training job.

   After the training job has completed, the newly trained model artifacts are stored under the S3 output path that you provided in the Output data configuration field. To deploy the model to get predictions, see Step 5: Deploy the Model to Amazon EC2.

Perform Incremental Training (API)

This example shows how to use SageMaker APIs to train a model using the SageMaker image classification algorithm and the Caltech 256 Image Dataset, then train a new model using the first one. It uses Amazon S3 for input and output sources. Please see the incremental training sample notebook for more details on using incremental training.

Note

In this example we used the original datasets in the incremental training, however you can use different datasets, such as ones that contain newly added samples. Upload the new datasets to S3 and make adjustments to the data_channels variable used to train the new model.

Get an AWS Identity and Access Management (IAM) role that grants required permissions and initialize environment variables:

```python
import sagemaker
```
from sagemaker import get_execution_role

role = get_execution_role()
print(role)

sess = sagemaker.Session()

bucket = sess.default_bucket()
print(bucket)
prefix = 'ic-incr-training'

Get the training image for the image classification algorithm:

from sagemaker.amazon.amazon_estimator import get_image_uri

training_image = get_image_uri(sess.boto_region_name, 'image-classification', repo_version="latest")
# Display the training image
print (training_image)

Download the training and validation datasets, then upload them to Amazon Simple Storage Service (Amazon S3):

import os
import urllib.request
import boto3

# Define a download function
def download(url):
    filename = url.split('/')[-1]
    if not os.path.exists(filename):
        urllib.request.urlretrieve(url, filename)

# Download the caltech-256 training and validation datasets
download('http://data.mxnet.io/data/caltech-256/caltech-256-60-train.rec')
download('http://data.mxnet.io/data/caltech-256/caltech-256-60-val.rec')

# Create four channels: train, validation, train_lst, and validation_lst
s3train = 's3://{}/{}'.format(bucket, prefix)
s3validation = 's3://{}/{}'.format(bucket, prefix)

# Upload the first files to the train and validation channels
!aws s3 cp caltech-256-60-train.rec $s3train --quiet
Define the training hyperparameters:

```python
# Define hyperparameters for the estimator
hyperparams = {
    "num_layers": "18",
    "resize": "32",
    "num_training_samples": "50000",
    "num_classes": "10",
    "image_shape": "3,28,28",
    "mini_batch_size": "128",
    "epochs": "3",
    "learning_rate": "0.1",
    "lr_scheduler_step": "2,3",
    "lr_scheduler_factor": "0.1",
    "augmentation_type": "crop_color",
    "optimizer": "sgd",
    "momentum": "0.9",
    "weight_decay": "0.0001",
    "beta_1": "0.9",
    "beta_2": "0.999",
    "gamma": "0.9",
    "eps": "1e-8",
    "top_k": "5",
    "checkpoint_frequency": "1",
    "use_pretrained_model": "0",
    "model_prefix": "" }
```

Create an estimator object and train the first model using the training and validation datasets:

```python
# Fit the base estimator
s3_output_location = 's3://{}/{}//{}/output'.format(bucket, prefix)
ic = sagemaker.estimator.Estimator(training_image,
    role,
    instance_count=1,
    instance_type='ml.p2.xlarge',
    volume_size=50,
    max_run=360000,
    input_mode='File',
    output_path=s3_output_location,
    sagemaker_session=sess,
    hyperparameters=hyperparams)
```
To use the model to incrementally train another model, create a new estimator object and use the model artifacts (ic.model_data, in this example) for the model_uri input argument:

```python
# Given the base estimator, create a new one for incremental training
incr_ic = sagemaker.estimator.Estimator(training_image,
    role,
    instance_count=1,
    instance_type='ml.p2.xlarge',
    volume_size=50,
    max_run=360000,
    input_mode='File',
    output_path=s3_output_location,
    sagemaker_session=sess,
    hyperparameters=hyperparams,
    model_uri=ic.model_data) # This parameter will ingest the previous job's model as a new channel
incr_ic.fit(inputs=data_channels, logs=True)
```

After the training job has completed, the newly trained model artifacts are stored under the S3 output path that you provided in Output_path. To deploy the model to get predictions, see Step 5: Deploy the Model to Amazon EC2.

### Use Managed Spot Training in Amazon SageMaker

Amazon SageMaker makes it easy to train machine learning models using managed Amazon EC2 Spot instances. Managed spot training can optimize the cost of training models up to 90% over on-demand instances. SageMaker manages the Spot interruptions on your behalf.
Managed Spot Training uses Amazon EC2 Spot instance to run training jobs instead of on-demand instances. You can specify which training jobs use spot instances and a stopping condition that specifies how long SageMaker waits for a job to run using Amazon EC2 Spot instances. Metrics and logs generated during training runs are available in CloudWatch.

Amazon SageMaker automatic model tuning, also known as hyperparameter tuning, can use managed spot training. For more information on automatic model tuning, see Perform Automatic Model Tuning with SageMaker.

Spot instances can be interrupted, causing jobs to take longer to start or finish. You can configure your managed spot training job to use checkpoints. SageMaker copies checkpoint data from a local path to Amazon S3. When the job is restarted, SageMaker copies the data from Amazon S3 back into the local path. The training job can then resume from the last checkpoint instead of restarting. For more information about checkpointing, see Use checkpoints in Amazon SageMaker.

Note

Unless your training job will complete quickly, we recommend you use checkpointing with managed spot training. SageMaker built-in algorithms and marketplace algorithms that do not checkpoint are currently limited to a MaxWaitTimeInSeconds of 3600 seconds (60 minutes).

Topics

• Using Managed Spot Training
• Managed Spot Training Lifecycle

Using Managed Spot Training

To use managed spot training, create a training job. Set EnableManagedSpotTraining to True and specify the MaxWaitTimeInSeconds. MaxWaitTimeInSeconds must be larger than MaxRuntimeInSeconds. For more information about creating a training job, see DescribeTrainingJob.

You can calculate the savings from using managed spot training using the formula \(1 - (\text{BillableTimeInSeconds} / \text{TrainingTimeInSeconds}) \times 100\). For example, if BillableTimeInSeconds is 100 and TrainingTimeInSeconds is 500, this means that your
A training job ran for 500 seconds, but you were billed for only 100 seconds. Your savings is \((1 - (100 / 500)) \times 100 = 80\%\).

To learn how to run training jobs on Amazon SageMaker spot instances and how managed spot training works and reduces the billable time, see the following example notebooks:

- Managed Spot Training with TensorFlow
- Managed Spot Training with PyTorch
- Managed Spot Training with XGBoost
- Managed Spot Training with MXNet
- Amazon SageMaker Managed Spot Training Examples GitHub repository

**Managed Spot Training Lifecycle**

You can monitor a training job using `TrainingJobStatus` and `SecondaryStatus` returned by `DescribeTrainingJob`. The list below shows how `TrainingJobStatus` and `SecondaryStatus` values change depending on the training scenario:

- **Spot instances acquired with no interruption during training**
  1. InProgress: Starting ↞ Downloading ↞ Training ↞ Uploading

- **Spot instances interrupted once. Later, enough spot instances were acquired to finish the training job.**
  1. InProgress: Starting ↞ Downloading ↞ Training ↞ Interrupted ↞ Starting ↞ Downloading ↞ Training ↞ Uploading

- **Spot instances interrupted twice and MaxWaitTimeInSeconds exceeded.**
  1. InProgress: Starting ↞ Downloading ↞ Training ↞ Interrupted ↞ Starting ↞ Downloading ↞ Training ↞ Interrupted ↞ Downloading ↞ Training
  2. Stopping: Stopping
  3. Stopped: MaxWaitTimeExceeded

- **Spot instances were never launched.**
  1. InProgress: Starting
  2. Stopping: Stopping
  3. Stopped: MaxWaitTimeExceeded
Train Using SageMaker Managed Warm Pools

SageMaker managed warm pools let you retain and reuse provisioned infrastructure after the completion of a training job to reduce latency for repetitive workloads, such as iterative experimentation or running many jobs consecutively. Subsequent training jobs that match specified parameters run on the retained warm pool infrastructure, which speeds up start times by reducing the time spent provisioning resources.

⚠️ Important
SageMaker managed warm pools are a billable resource. For more information, see Billing.

Topics
- How it works
- Warm pool resource limits
- How to use SageMaker managed warm pools
- Considerations

How it works

To use SageMaker managed warm pools and reduce latency between similar consecutive training jobs, create a training job that specifies a KeepAlivePeriodInSeconds value in its ResourceConfig. This value represents the duration of time in seconds to retain configured resources in a warm pool for subsequent training jobs. If you need to run several training jobs using similar configurations, you can further reduce latency and billable time by using a dedicated persistent cache directory to store and re-use your information in a different job.

Topics
- Warm pool lifecycle
- Warm pool creation
- Matching training jobs
- Maximum warm pool duration
- Using persistent cache
Warm pool lifecycle

1. Create an initial training job with a `KeepAlivePeriodInSeconds` value greater than 0. When you run this first training job, this “cold-starts” a cluster with typical startup times.

2. When the first training job completes, the provisioned resources are kept alive in a warm pool for the period specified in the `KeepAlivePeriodInSeconds` value. As long as the cluster is healthy and the warm pool is within the specified `KeepAlivePeriodInSeconds`, then the warm pool status is Available.

3. The warm pool stays Available until it either identifies a matching training job for reuse or it exceeds the specified `KeepAlivePeriodInSeconds` and is terminated. The maximum length of time allowed for the `KeepAlivePeriodInSeconds` is 3600 seconds (60 minutes). If the warm pool status is Terminated, then this is the end of the warm pool lifecycle.

4. If the warm pool identifies a second training job with matching specifications such as instance count or instance type, then the warm pool moves from the first training job to the second training job for reuse. The status of the first training job warm pool becomes Reused. This is the end of the warm pool lifecycle for the first training job.

5. The status of the second training job that reused the warm pool becomes InUse. After the second training job completes, the warm pool is Available for the `KeepAlivePeriodInSeconds` duration specified in the second training job. A warm pool can continue moving to subsequent matching training jobs for a maximum of 28 days.

6. If the warm pool is no longer available to reuse, the warm pool status is Terminated. Warm pools are no longer available if they are terminated by a user, for a patch update, or for exceeding the specified `KeepAlivePeriodInSeconds`.

For more information on warm pool status options, see `WarmPoolStatus` in the Amazon SageMaker API Reference.

Warm pool creation

If an initial training job successfully completes and has a `KeepAlivePeriodInSeconds` value greater than 0, this creates a warm pool. If you stop a training job after a cluster is already launched, a warm pool is still retained. If the training job fails due to an algorithm or client error, a warm pool is still retained. If the training job fails for any other reason that might compromise the health of the cluster, then the warm pool is not created.
To verify successful warm pool creation, check the warm pool status of your training job. If a warm pool successfully provisions, the warm pool status is Available. If a warm pool fails to provision, the warm pool status is Terminated.

**Matching training jobs**

For a warm pool to persist, it must find a matching training job within the time specified in the `KeepAlivePeriodIn Seconds` value. The next training job is a match if the following values are identical:

- `RoleArn`
- `ResourceConfig values`:
  - `InstanceCount`
  - `InstanceType`
  - `VolumeKmsKeyId`
  - `VolumeSizeInGB`
- `VpcConfig values`:
  - `SecurityGroupIds`
  - `Subnets`
  - `EnableInterContainerTrafficEncryption`
  - `EnableNetworkIsolation`

All of these values must be the same for a warm pool to move to a subsequent training job for reuse.

**Maximum warm pool duration**

The maximum `KeepAlivePeriodInSeconds` for a single training job is 3600 seconds (60 minutes) and the maximum length of time that a warm pool cluster can continue running consecutive training jobs is 28 days.

Each subsequent training job must also specify a `KeepAlivePeriodInSeconds` value. When the warm pool moves to the next training job, it inherits the new `KeepAlivePeriodInSeconds` value specified in that training job's `ResourceConfig`. In this way, you can keep a warm pool moving from training job to training job for a maximum of 28 days.
If no `KeepAlivePeriodInSeconds` is specified, then the warm pool spins down after the training job completes.

**Using persistent cache**

When you create a warm pool, SageMaker mounts a special directory on the volume that will persist throughout the lifecycle of the warm pool. This directory can also be used to store information that you want to re-use in another job.

Using persistent cache can reduce latency and billable time over using warm pools alone for jobs that require the following:

- multiple interactions with similar configurations
- incremental training jobs
- hyperparameter optimization

For example, you can avoid downloading the same Python dependencies on repeated runs by setting up a pip cache directory inside the persistent cache directory. You are fully responsible for managing the contents of this directory. The following are examples of types of information that you can put in your persistent cache to help reduce your latency and billable time.

- Dependencies managed by pip.
- Dependencies managed by conda.
- [Checkpoint information](#).
- Any additional information generated during training.

The location of the persistent cache is `/opt/ml/sagemaker/warmpoolcache`. The environment variable `SAGEMAKER_MANAGED_WARMPOOL_CACHE_DIRECTORY` points to the location of the persistent cache directory.

The following code example shows you how to set up a warm pool and use persistent cache to store your pip dependencies for use in a subsequent job. The subsequent job must run within the time frame given by the parameter `keep_alive_period_in_seconds`.

```python
import sagemaker
from sagemaker import get_execution_role
from sagemaker.tensorflow

# Creates a SageMaker session and gets execution role
session = sagemaker.Session()
```

---

**How it works**

3931
In the previous code example, using the `environment` parameter exports the environment variable `PIP_CACHE_DIRECTORY` to point to the directory `/opt/ml/sagemaker/warmpoolcache/pip`. Exporting this environment variable will change where pip stores its cache to the new location. Any directory, including nested directories, that you create inside the persistent cache directory will be available for re-use during a subsequent training run. In the previous code example, a directory called pip is changed to be the default location to cache any dependencies installed using pip.

The persistent cache location may also be accessed from within your Python training script using the environment variable as shown in the following code example.

```python
import os
import shutil
if __name__ == '__main__':
    PERSISTED_DIR = os.environ["SAGEMAKER_MANAGED_WARMPOOL_CACHE_DIRECTORY"]
    
    # create a file to be persisted
    open(os.path.join(PERSISTED_DIR, "test.txt"), 'a').close()
```
# create a directory to be persisted
os.mkdir(os.path.join(PERSISTED_DIR, "test_dir"))

# Move a file to be persisted
shutil.move("path/of/your/file.txt", PERSISTED_DIR)

## Billing

SageMaker managed warm pools are a billable resource. Retrieve the warm pool status for your training job to check the billable time for your warm pools. You can check the warm pool status either through the [Using the Amazon SageMaker console](https://console.aws.amazon.com/sagemaker/home?region=ap-south-1#/warm-pools) or directly through the `DescribeTrainingJob` API command. For more information, see [WarmPoolStatus](https://docs.aws.amazon.com/sagemaker/latest/dg/how-warm-pools-work.html) in the Amazon SageMaker API Reference.

**Note**

After the time specified by the parameter `KeepAlivePeriodInSeconds` has ended, both the warm pool and persistent cache will shut down, and the contents will be deleted.

---

### Warm pool resource limits

To get started, you must first request a service limit increase for SageMaker managed warm pools. The default resource limit for warm pools is 0.

If a training job is created with `KeepAlivePeriodInSeconds` specified, but you did not request a warm pool limit increase, then a warm pool is not retained after the completion of the training job. A warm pool is only created if your warm pool limit has sufficient resources. After a warm pool is created, the resources are released when they move to a matching training job or if the `KeepAlivePeriodInSeconds` expires (if the warm pool status is Reused or Terminated).

**Request a warm pool quota increase**

Request a warm pool quota increase using the AWS Service Quotas console.
1. Open the AWS Service Quotas console.
2. On the left-hand navigation panel, choose AWS services.
3. Search for and choose Amazon SageMaker.
4. Search for the keyword warm pool to see all available warm pool service quotas.
5. Find the instance type for which you want to increase your warm pool quota, select the warm pool service quota for that instance type, and choose Request quota increase.
6. Enter your requested instance limit number under Change quota value. The new value must be greater than the current Applied quota value.
7. Choose Request.

There is a limit on the number of instances that you can retain for each account, which is determined by instance type. You can check your resource limits in the AWS Service Quotas console or directly using the list-service-quotas AWS CLI command. For more information on AWS Service Quotas, see Requesting a quota increase in the Service Quotas User Guide.

You can also use AWS Support Center to request a warm pool quota increase. For a list of available instance types according to Region, see Amazon SageMaker Pricing and choose Training in the On-Demand Pricing table.

How to use SageMaker managed warm pools

You can use SageMaker managed warm pools through the SageMaker Python SDK, the Amazon SageMaker console, or through the low-level APIs. Administrators can optionally use the sagemaker:KeepAlivePeriod condition key to further restrict the KeepAlivePeriodInSeconds limits for certain users or groups.

Topics

- Using the SageMaker Python SDK
- Using the Amazon SageMaker console
Using the low-level SageMaker APIs

IAM condition key

Using the SageMaker Python SDK

Create, update, or terminate warm pools using the SageMaker Python SDK.

Note
This feature is available in the SageMaker Python SDK v2.110.0 and later.

Topics

• Create a warm pool
• Update a warm pool
• Terminate a warm pool

Create a warm pool

To create a warm pool, use the SageMaker Python SDK to create an estimator with a keep_alive_period_in_seconds value greater than 0 and call fit(). When the training job completes, a warm pool is retained. For more information on training scripts and estimators, see Train a Model with the SageMaker Python SDK. If your script does not create a warm pool, see Warm pool creation for possible explanations.

```python
import sagemaker
from sagemaker import get_execution_role
from sagemaker.tensorflow import TensorFlow

# Creates a SageMaker session and gets execution role
session = sagemaker.Session()
role = get_execution_role()

# Creates an example estimator
estimator = TensorFlow(
  ...
  entry_point='my-training-script.py',
  source_dir='code',
)```
Next, create a second matching training job. In this example, we create my-training-job-2, which has all of the necessary attributes to match with my-training-job-1, but has a different hyperparameter for experimentation. The second training job reuses the warm pool and starts up faster than the first training job. The following code example uses a Tensorflow estimator. The warm pool feature can be used with any training algorithm that runs on Amazon SageMaker. For more information on which attributes need to match, see Matching training jobs.

```python
# Creates an example estimator
estimator = TensorFlow(
...
    entry_point='my-training-script.py',
    source_dir='code',
    role=role,
    model_dir='model_dir',
    framework_version='2.2',
    py_version='py37',
    job_name='my-training-job-2',
    instance_type='ml.g4dn.xlarge',
    instance_count=1,
    volume_size=250,
    hyperparameters={
        "batch-size": 512,
        "epochs": 1,
        "learning-rate": 1e-3,
        "beta_1": 0.9,
        "beta_2": 0.999,
    },
    keep_alive_period_in_seconds=1800,
)

# Starts a SageMaker training job and waits until completion
estimator.fit('s3://my_bucket/my_training_data/')
```
"batch-size": 512,
"epochs": 2,
"learning-rate": 1e-3,
"beta_1": 0.9,
"beta_2": 0.999,
},
keep_alive_period_in_seconds=1800,
)

# Starts a SageMaker training job and waits until completion
estimator.fit('s3://my_bucket/my_training_data/

Check the warm pool status of both training jobs to confirm that the warm pool is Reused for my-training-job-1 and InUse for my-training-job-2.

**Note**

Training job names have date/time suffixes. The example training job names my-training-job-1 and my-training-job-2 should be replaced with actual training job names. You can use the estimator.latest_training_job.job_name command to fetch the actual training job name.

```
session.describe_training_job('my-training-job-1')
session.describe_training_job('my-training-job-2')
```

The result of describe_training_job provides all details about a given training job. Find the WarmPoolStatus attribute to check information about a training job's warm pool. Your output should look similar to the following example:

```
# Warm pool status for training-job-1
...
'WarmPoolStatus': {'Status': 'Reused',
  'ResourceRetainedBillableTimeInSeconds': 1000,
  'ReusedByName': my-training-job-2}
...

# Warm pool status for training-job-2
...
'WarmPoolStatus': {'Status': 'InUse'}
```
Update a warm pool

When the training job is complete and the warm pool status is Available, then you can update the KeepAlivePeriodInSeconds value.

```
session.update_training_job(job_name, 
resource_config={"KeepAlivePeriodInSeconds":3600})
```

Terminate a warm pool

To manually terminate a warm pool, set the KeepAlivePeriodInSeconds value to 0.

```
session.update_training_job(job_name, resource_config={"KeepAlivePeriodInSeconds":0})
```

The warm pool automatically terminates when it exceeds the designated KeepAlivePeriodInSeconds value or if there is a patch update for the cluster.

Using the Amazon SageMaker console

Through the console, you can create a warm pool, release a warm pool, or check the warm pool status and billable time of specific training jobs. You can also see which matching training job reused a warm pool.

1. Open the Amazon SageMaker console and choose Training jobs from the navigation pane. If applicable, the warm pool status of each training job is visible in the Warm pool status column and the time left for an active warm pool is visible in the Time left column.
2. To create a training job that uses a warm pool from the console, choose Create training job. Then, be sure to specify a value for the Keep alive period field when configuring your training job resources. This value must be an integer between 1 and 3600, which represents duration of time in seconds.
3. To release a warm pool from the console, select a specific training job and choose Release cluster from the Actions dropdown menu.
4. To see more information about a warm pool, choose a training job name. In the job details page, scroll down to the Warm pool status section to find the warm pool status, the time left if the warm pool status is Available, the warm pool billable seconds, and the name of the training job that reused the warm pool if the warm pool status is Reused.
Using the low-level SageMaker APIs

Use SageMaker managed warm pools with either the SageMaker API or the AWS CLI.

**SageMaker API**

Set up SageMaker managed warm pools using the SageMaker API with the following commands:

- `CreateTrainingJob`
- `UpdateTrainingJob`
- `ListTrainingJobs`
- `DescribeTrainingJob`

**AWS CLI**

Set up SageMaker managed warm pools using the AWS CLI with the following commands:

- `create-training-job`
- `update-training-job`
- `list-training-jobs`
- `describe-training-job`

**IAM condition key**

Administrators can optionally use the `sagemaker:KeepAlivePeriod` condition key to further restrict the `KeepAlivePeriodIn Seconds` limits for certain users or groups. SageMaker managed warm pools are limited to a `KeepAlivePeriodInSeconds` value of 3600 seconds (60 minutes), but administrators can lower this limit if needed.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "EnforceKeepAlivePeriodLimit",
         "Effect": "Allow",
         "Action": [
            "sagemaker:CreateTrainingJob"
         ]
      }
   ]
}
```
Considerations

Consider the following items when using SageMaker managed warm pools.

- SageMaker managed warm pools cannot be used with heterogeneous cluster training.
- SageMaker managed warm pools cannot be used with spot instances.
- SageMaker managed warm pools are limited to a `KeepAlivePeriodInSeconds` value of 3600 seconds (60 minutes).
- If a warm pool continues to successfully match training jobs within the specified `KeepAlivePeriodInSeconds` value, the cluster can only continue running for a maximum of 28 days.

Monitor and Analyze Training Jobs Using Amazon CloudWatch Metrics

An Amazon SageMaker training job is an iterative process that teaches a model to make predictions by presenting examples from a training dataset. Typically, a training algorithm computes several metrics, such as training error and prediction accuracy. These metrics help diagnose whether the model is learning well and will generalize well for making predictions on unseen data. The training algorithm writes the values of these metrics to logs, which SageMaker monitors and sends to Amazon CloudWatch in real time. To analyze the performance of your training job, you can view graphs of these metrics in CloudWatch. When a training job has completed, you can also get a list of the metric values that it computes in its final iteration by calling the `DescribeTrainingJob` operation.

For more information, see [Condition keys for Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/service-authorization-concepts.html) in the *Service Authorization Reference*. 
Note

Amazon CloudWatch supports high-resolution custom metrics, and its finest resolution is 1 second. However, the finer the resolution, the shorter the lifespan of the CloudWatch metrics. For the 1-second frequency resolution, the CloudWatch metrics are available for 3 hours. For more information about the resolution and the lifespan of the CloudWatch metrics, see GetMetricStatistics in the Amazon CloudWatch API Reference.

Tip

If you want to profile your training job with a finer resolution down to 100-millisecond (0.1 second) granularity and store the training metrics indefinitely in Amazon S3 for custom analysis at any time, consider using Amazon SageMaker Debugger. SageMaker Debugger provides built-in rules to automatically detect common training issues; it detects hardware resource utilization issues (such as CPU, GPU, and I/O bottlenecks) and non-converging model issues (such as overfit, vanishing gradients, and exploding tensors). SageMaker Debugger also provides visualizations through Studio Classic and its profiling report. To explore the Debugger visualizations, see SageMaker Debugger Insights Dashboard Walkthrough, Debugger Profiling Report Walkthrough, and Analyze Data Using the SMDebug Client Library.

Topics

- Defining Training Metrics
- Monitoring Training Job Metrics (CloudWatch Console)
- Monitoring Training Job Metrics (SageMaker Console)
- Example: Viewing a Training and Validation Curve

Defining Training Metrics

SageMaker automatically parses training job logs and sends training metrics to CloudWatch. By default, SageMaker sends system resource utilization metrics listed in SageMaker Jobs and Endpoint Metrics. If you want SageMaker to parse logs and send custom metrics from a training job of your own algorithm to CloudWatch, you need to specify metrics definitions by passing the name of metrics and regular expressions when you configure a SageMaker training job request.
You can specify the metrics that you want to track using the SageMaker console, the SageMaker Python SDK, or the low-level SageMaker API.

If you are using your own algorithm, do the following:

- Make sure that the algorithm writes the metrics that you want to capture to logs.
- Define a regular expression that accurately searches the logs to capture the values of the metrics that you want to send to CloudWatch.

For example, suppose your algorithm emits the following metrics for training error and validation error:

```
Train_error=0.138318;  Valid_error=0.324557;
```

If you want to monitor both of those metrics in CloudWatch, the dictionary for the metric definitions should look like the following example:

```
[
  {
    "Name": "train:error",
    "Regex": "Train_error=(.*?);"
  },
  {
    "Name": "validation:error",
    "Regex": "Valid_error=(.*?);"
  }
]
```

In the regex for the `train:error` metric defined in the preceding example, the first part of the regex finds the exact text "Train_error=", and the expression `(.*?);` captures any characters until the first semicolon character appears. In this expression, the parenthesis tell the regex to capture what is inside them, \* means any character, \*? means zero or more, and ? means capture only until the first instance of the ; character.

**Define Metrics Using the SageMaker Python SDK**

Define the metrics that you want to send to CloudWatch by specifying a list of metric names and regular expressions as the `metric_definitions` argument when you initialize an `Estimator`.
object. For example, if you want to monitor both the `train:error` and `validation:error` metrics in CloudWatch, your `Estimator` initialization would look like the following example:

```python
import sagemaker
from sagemaker.estimator import Estimator

estimator = Estimator(
    image_uri="your-own-image-uri",
    role=sagemaker.get_execution_role(),
    sagemaker_session=sagemaker.Session(),
    instance_count=1,
    instance_type='ml.c4.xlarge',
    metric_definitions=[
        {'Name': 'train:error', 'Regex': 'Train_error=(.*?);'},
        {'Name': 'validation:error', 'Regex': 'Valid_error=(.*?);'}
    ]
)
```

For more information about training by using Amazon SageMaker Python SDK estimators, see Sagemaker Python SDK on GitHub.

**Define Metrics Using the SageMaker Console**

If you choose the **Your own algorithm container in ECR** option as your algorithm source in the SageMaker console when you create a training job, add the metric definitions in the Metrics section. The following screenshot shows how it should look after you add the example metric names and the corresponding regular expressions.
Algorithm options
Use an Amazon SageMaker built-in algorithm, your own algorithm, or a third-party algorithm from AWS Marketplace.

▼ Algorithm source

- Amazon SageMaker built-in algorithm Learn more
- Your own algorithm resource
- Your own algorithm container in ECR Learn more
- An algorithm subscription from AWS Marketplace

▼ Provide container ECR path

Container
The registry path where the training image is stored in Amazon ECR. Learn more
`accountId.dkr.ecr.Region.amazonaws.com/repository[:tag] or [@digest]`

Input mode
You can provide your training data as a file or pipe.
File

Metrics
Define the metrics you want to emit to CloudWatch metrics.

<table>
<thead>
<tr>
<th>Metric name</th>
<th>Regex</th>
<th>Remove</th>
</tr>
</thead>
<tbody>
<tr>
<td>train:error</td>
<td>Train_error=(.*?);</td>
<td>Remove</td>
</tr>
<tr>
<td>validation:error</td>
<td>Valid_error=(.*?);</td>
<td>Remove</td>
</tr>
</tbody>
</table>

Add metric

Define Metrics Using the Low-level SageMaker API

Define the metrics that you want to send to CloudWatch by specifying a list of metric names and regular expressions in the MetricDefinitions field of the AlgorithmSpecification input parameter that you pass to the CreateTrainingJob operation. For example, if you want to monitor both the train:error and validation:error metrics in CloudWatch, your AlgorithmSpecification would look like the following example:
"AlgorithmSpecification": {
   "TrainingImage": "your-own-image-uri",
   "TrainingInputMode": "File",
   "MetricDefinitions": [
      {
         "Name": "train:error",
         "Regex": "Train_error=(.*?);"
      },
      {
         "Name": "validation:error",
         "Regex": "Valid_error=(.*?);"
      }
   ]
}

For more information about defining and running a training job by using the low-level SageMaker API, see CreateTrainingJob.

**Monitoring Training Job Metrics (CloudWatch Console)**

You can monitor the metrics that a training job emits in real time in the CloudWatch console.

**To monitor training job metrics (CloudWatch console)**

2. Choose **Metrics**, then choose `/aws/sagemaker/TrainingJobs`.
3. Choose **TrainingJobName**.
4. On the **All metrics** tab, choose the names of the training metrics that you want to monitor.
5. On the **Graphed metrics** tab, configure the graph options. For more information about using CloudWatch graphs, see Graph Metrics in the Amazon CloudWatch User Guide.

**Monitoring Training Job Metrics (SageMaker Console)**

You can monitor the metrics that a training job emits in real time by using the SageMaker console.

**To monitor training job metrics (SageMaker console)**

2. Choose **Training jobs**, then choose the training job whose metrics you want to see.
3. Choose **TrainingJobName**.

4. In the **Monitor** section, you can review the graphs of instance utilization and algorithm metrics.
Example: Viewing a Training and Validation Curve

Typically, you split the data on which you train your model into training and validation datasets. You use the training set to train the model parameters that are used to make predictions on the training dataset. Then you test how well the model makes predictions by calculating predictions for the validation set. To analyze the performance of a training job, you commonly plot a training curve against a validation curve.

Viewing a graph that shows the accuracy for both the training and validation sets over time can help you to improve the performance of your model. For example, if training accuracy continues to increase over time, but, at some point, validation accuracy starts to decrease, you are likely overfitting your model. To address this, you can make adjustments to your model, such as increasing regularization.

For this example, you can use the Image-classification-full-training example in the Example notebooks section of your SageMaker notebook instance. If you don't have a SageMaker notebook instance, create one by following the instructions at Step 1: Create an Amazon SageMaker Notebook Instance. If you prefer, you can follow along with the End-to-End Multiclass Image Classification Example in the example notebook on GitHub. You also need an Amazon S3 bucket to store the training data and for the model output.

To view training and validation error curves

2. Choose Notebooks, and then choose Notebook instances.
3. Choose the notebook instance that you want to use, and then choose Open.
4. On the dashboard for your notebook instance, choose SageMaker Examples.
5. Expand the Introduction to Amazon Algorithms section, and then choose Use next to Image-classification-fulltraining.ipynb.
7. Run all of the cells in the notebook up to the Inference section. You don't need to deploy an endpoint or get inference for this example.
8. After the training job starts, open the CloudWatch console at https://console.aws.amazon.com/cloudwatch.
9. Choose Metrics, then choose /aws/sagemaker/TrainingJobs.
10. Choose **TrainingJobName**.

11. On the **All metrics** tab, choose the **train:accuracy** and **validation:accuracy** metrics for the training job that you created in the notebook.

12. On the graph, choose an area that the metric's values to zoom in. You should see something like the following example.

---

**Use Amazon SageMaker Training Storage Paths for Training Datasets, Checkpoints, Model Artifacts, and Outputs**

This page provides a high-level summary of how the SageMaker training platform manages storage paths for training datasets, model artifacts, checkpoints, and outputs between AWS cloud storage and training jobs in SageMaker. Throughout this guide, you learn to identify the default paths set by the SageMaker platform and how the data channels can be streamlined with your data sources in Amazon Simple Storage Service (Amazon S3), FSx for Lustre, and Amazon EFS. For more information about various data channel input modes and storage options, see [Access Training Data](#).

**Topics**

- [Overview](#)
- [Uncompressed model output](#)
- [Tips and Considerations for Setting Up Storage Paths](#)
- [SageMaker Environment Variables and Default Paths for Training Storage Locations](#)
Overview

The following diagram shows the simplest example of how SageMaker manages input and output paths when you run a training job using the SageMaker Python SDK `Estimator` class and its `fit` method. It's based on using file mode as the data access strategy and Amazon S3 as the data source for the training input channels.

```python
estimator = Estimator(
    checkpoint_s3_uri='s3://checkpoint-dest/',
    output_path='s3://output-path/',
    base_job_name='job-name',
    input_mode='File'
    ...
)

estimator.fit(inputs={
    'channel1': 's3://bucket-data1/',
    ...
    'channel20': 's3://bucket-data20/'}
)
```

![Diagram showing SageMaker data paths](image)

This figure shows an overview of how SageMaker pairs storage paths between an Amazon S3 bucket as the data source and the SageMaker training instance based on how the paths are specified in a SageMaker estimator class. More information about the paths, how they read from or write to the paths, and purposes of the paths are described in the following section "SageMaker Environment Variables and Default Paths for Training Storage Locations".

You can use `OutputDataConfig` in the `CreateTrainingJob` API to find where your S3 bucket is located. Use the `ModelArtifacts` API to find the S3 location that contains your model artifacts. See the `abalone_build_train_deploy` notebook for an example of output paths and how they are used in API calls.

For more information and examples of how SageMaker manages data source, input modes, and local paths in SageMaker training instances, see `Access Training Data`. 
Uncompressed model output

SageMaker stores your model in /opt/ml/model and your data in /opt/ml/output/data. After the model and data are written to those locations, they're uploaded to your Amazon S3 bucket as compressed files by default.

You can save time on large data file compression by uploading model and data outputs to your S3 bucket as uncompressed files. To do this, create a training job in uncompressed upload mode by using either the AWS Command Line Interface (AWS CLI) or the SageMaker Python SDK.

The following code example shows how to create a training job in uncompressed upload mode when using the AWS CLI. To enable uncompressed upload mode, set CompressionType field in the OutputDataConfig API to NONE.

```json
{
    "TrainingJobName": "uncompressed_model_upload",
    ...  
    "OutputDataConfig": {
        "S3OutputPath": "s3://DOC-EXAMPLE-BUCKET/uncompressed_upload/output",
        "CompressionType": "NONE"
    },
    ...
}
```

The following code example shows you how to create a training job in uncompressed upload mode using the SageMaker Python SDK.

```python
import sagemaker
from sagemaker.estimator import Estimator

estimator = Estimator(
    image_uri="your-own-image-uri",
    role=sagemaker.get_execution_role(),
    sagemaker_session=sagemaker.Session(),
    instance_count=1,
    instance_type='ml.c4.xlarge',
    disable_output_compression=True
)
```
Tips and Considerations for Setting Up Storage Paths

Consider the following items when setting up storage paths for training jobs in SageMaker.

- If you want to store training artifacts for distributed training in the `/opt/ml/output/data` directory, you must properly append subdirectories or use unique file names for the artifacts through your model definition or training script. If the subdirectories and file names are not properly configured, all of the distributed training workers might write outputs to the same file name in the same output path in Amazon S3.

- If you use a custom training container, make sure you install the SageMaker Training Toolkit that helps set up the environment for SageMaker training jobs. Otherwise, you must specify the environment variables explicitly in your Dockerfile. For more information, see Create a container with your own algorithms and models.

- When using an ML instance with NVMe SSD volumes, SageMaker doesn't provision Amazon EBS gp2 storage. Available storage is fixed to the NVMe-type instance's storage capacity. SageMaker configures storage paths for training datasets, checkpoints, model artifacts, and outputs to use the entire capacity of the instance storage. For example, ML instance families with the NVMe-type instance storage include ml.p4d, ml.g4dn, and ml.g5. When using an ML instance with the EBS-only storage option and without instance storage, you must define the size of EBS volume through the `volume_size` parameter in the SageMaker estimator class (or `VolumeSizeInGB` if you are using the ResourceConfig API). For example, ML instance families that use EBS volumes include ml.c5 and ml.p2. To look up instance types and their instance storage types and volumes, see Amazon EC2 Instance Types.

- The default paths for SageMaker training jobs are mounted to Amazon EBS volumes or NVMe SSD volumes of the ML instance. When you adapt your training script to SageMaker, make sure that you use the default paths listed in the previous topic about the section called “SageMaker Environment Variables and Default Paths for Training Storage Locations”. We recommend that you use the `/tmp` directory as a scratch space for temporarily storing any large objects during training. This means that you must not use directories that are mounted to small disk space allocated for system, such as `/user` and `/home`, to avoid out-of-space errors.

To learn more, see the AWS machine learning blog Choose the best data source for your Amazon SageMaker training job that further discusses case studies and performance benchmarks of data sources and input modes.
SageMaker Environment Variables and Default Paths for Training Storage Locations

The following table summarizes input and output paths for training datasets, checkpoints, model artifacts, and outputs, managed by the SageMaker training platform.

<table>
<thead>
<tr>
<th>Local path in SageMaker training instance</th>
<th>SageMaker environment variable</th>
<th>Purpose</th>
<th>Read from S3 during start</th>
<th>Read from S3 during Spot-restart</th>
<th>Writes to S3 during training</th>
<th>Writes to S3 when job is terminated</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>/opt/ml/input/data</code> _<code>channel_name</code></td>
<td>SM_CHANNEL_<code>channel_name</code></td>
<td>Reading training data from the input channels specified through the SageMaker Python SDK <code>Estimator</code> class or the <code>CreateTrainingJob</code> API operation. For more information about how to specify it in your training script using the SageMaker Python SDK, see <code>Prepare a Training script</code>.</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td><code>/opt/ml/output/data</code> _<code>output_name</code></td>
<td>SM_OUTPUT_DIR</td>
<td>Saving outputs such as loss, accuracy, intermediate layers, weights, gradients, bias, and TensorBoard-compatible outputs. You can also save any arbitrary</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Local path in SageMaker training instance</td>
<td>SageMaker environment variable</td>
<td>Purpose</td>
<td>Read from S3 during start</td>
<td>Read from S3 during Spot-restart</td>
<td>Writes to S3 during training</td>
<td>Writes to S3 when job is terminated</td>
</tr>
<tr>
<td>------------------------------------------</td>
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<td>---------------------------</td>
<td>----------------------------------</td>
<td>-----------------------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>output you’d like using this path. Note that this is a different path from the one for storing the final model artifact /opt/ml/model/</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>/opt/ml/model ³</td>
<td>SM_MODEL_DIR</td>
<td>Storing the final model artifact. This is also the path from where the model artifact is deployed for Real-time inference in SageMaker Hosting.</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>/opt/ml/checkpoints ⁴</td>
<td>-</td>
<td>Saving model checkpoints (the state of model) to resume training from a certain point, and recover from unexpected or Managed Spot Training interruptions.</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Local path in SageMaker training instance</td>
<td>SageMaker environment variable</td>
<td>Purpose</td>
<td>Read from S3 during start</td>
<td>Read from S3 during Spot-restart</td>
<td>Writes to S3 during training</td>
<td>Writes to S3 when job is terminated</td>
</tr>
<tr>
<td>------------------------------------------</td>
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<td>---------</td>
<td>--------------------------</td>
<td>-------------------------------</td>
<td>-----------------------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>/opt/ml/code</td>
<td>SAGEMAKER_SUBMIT_DIRECTORY</td>
<td>Copying training scripts, additional libraries, and dependencies.</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>/tmp</td>
<td>-</td>
<td>Reading or writing to /tmp as a scratch space.</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

1 channel_name is the place to specify user-defined channel names for training data inputs. Each training job can contain several data input channels. You can specify up to 20 training input channels per training job. Note that the data downloading time from the data channels is counted to the billable time. For more information about data input paths, see [How Amazon SageMaker Provides Training Information](#). Also, there are three types of data input modes that SageMaker supports: file, FastFile, and pipe mode. To learn more about the data input modes for training in SageMaker, see [Access Training Data](#).

2 SageMaker compresses and writes training artifacts to TAR files (.tar.gz). Compression and uploading time is counted to the billable time. For more information, see [How Amazon SageMaker Processes Training Output](#).

3 SageMaker compresses and writes the final model artifact to a TAR file (.tar.gz). Compression and uploading time is counted to the billable time. For more information, see [How Amazon SageMaker Processes Training Output](#).

4 Sync with Amazon S3 during training. Write as is without compressing to TAR files. For more information, see [Use Checkpoints in Amazon SageMaker](#).
Provide Dataset Metadata to Training Jobs with an Augmented Manifest File

To include metadata with your dataset in a training job, use an augmented manifest file. When using an augmented manifest file, your dataset must be stored in Amazon Simple Storage Service (Amazon S3), and you must configure your training job to use the dataset stored there. You specify the location and format of this dataset for one or more Channel. Augmented manifests can only support Pipe input mode. See the section, InputMode in Channel to learn more about pipe input mode.

When specifying a channel's parameters, you specify a path to the file, called a S3Uri. Amazon SageMaker interprets this URI based on the specified S3DataType in S3DataSource. The AugmentedManifestFile option defines a manifest format that includes metadata with the input data. Using an augmented manifest file is an alternative to preprocessing when you have labeled data. For training jobs using labeled data, you typically need to preprocess the dataset to combine input data with metadata before training. If your training dataset is large, preprocessing can be time consuming and expensive.

Augmented Manifest File Format

An augmented manifest file must be formatted in JSON Lines format. In JSON Lines format, each line in the file is a complete JSON object followed by a newline separator.

During training, SageMaker parses each JSON line and sends some or all of its attributes on to the training algorithm. You specify which attribute contents to pass and the order in which to pass them with the AttributeNames parameter of the CreateTrainingJob API. The AttributeNames parameter is an ordered list of attribute names that SageMaker looks for in the JSON object to use as training input.

For example, if you list ["line", "book"] for AttributeNames, the input data must include the attribute names of line and book in the specified order. For this example, the following augmented manifest file content is valid:

```
{"author": "Herman Melville", "line": "Call me Ishmael", "book": "Moby Dick"}
{"line": "It was love at first sight.", "author": "Joseph Heller", "book": "Catch-22"}
```

SageMaker ignores unlisted attribute names even if they precede, follow, or are in between listed attributes.
When using augmented manifest files, observe the following guidelines:

- The order of the attributes listed in the AttributeNames parameter determines the order of the attributes passed to the algorithm in the training job.
- The listed AttributeNames can be a subset of all of the attributes in the JSON line. SageMaker ignores unlisted attributes in the file.
- You can specify any type of data allowed by the JSON format in AttributeNames, including text, numerical, data arrays, or objects.
- To include an S3 URI as an attribute name, add the suffix -ref to it.

If an attribute name contains the suffix -ref, the attribute's value must be an S3 URI to a data file that is accessible to the training job. For example, if AttributeNames contains ["image-ref", "is-a-cat"], the following example shows a valid augmented manifest file:

```json
{"image-ref": "s3://mybucket/sample01/image1.jpg", "is-a-cat": 1}
{"image-ref": "s3://mybucket/sample02/image2.jpg", "is-a-cat": 0}
```

In case of the first JSON line of this manifest file, SageMaker retrieves the image1.jpg file from s3://mybucket/sample01/ and the string representation of the is-a-cat attribute "1" for image classification.

Tip

To create an augmented manifest file, use Amazon SageMaker Ground Truth and create a labeling job. For more information about the output from a labeling job, see Output Data.

Stream Augmented Manifest File Data

Augmented manifest format enables you to do training in Pipe mode using files without needing to create RecordIO files. You need to specify both train and validation channels as values for the InputDataConfig parameter of the CreateTrainingJob request. Augmented manifest files are supported only for channels using Pipe input mode. For each channel, the data is extracted from its augmented manifest file and streamed (in order) to the algorithm through the channel's named pipe. Pipe mode uses the first in first out (FIFO) method, so records are processed in the order in which they are queued. For information about Pipe input mode, see Input Mode.
Attribute names with a "-ref" suffix point to preformatted binary data. In some cases, the algorithm knows how to parse the data. In other cases, you might need to wrap the data so that records are delimited for the algorithm. If the algorithm is compatible with RecordIO-formatted data, specifying RecordIO for RecordWrapperType solves this issue. If the algorithm is not compatible with RecordIO format, specify None for RecordWrapperType and make sure that your data is parsed correctly for your algorithm.

Using the ['image-ref', 'is-a-cat'] example, if you use RecordIO wrapping, the following stream of data is sent to the queue:

recordio_formatted(s3://mybucket/foo/image1.jpg)recordio_formatted("1")recordio_formatted(s3://mybucket/bar/image2.jpg)recordio_formatted("0")

Images that are not wrapped with RecordIO format, are streamed with the corresponding is-a-cat attribute value as one record. This can cause a problem because the algorithm might not delimit the images and attributes correctly. For more information about using augmented manifest files for image classification, see Train with Augmented Manifest Image Format.

With augmented manifest files and Pipe mode in general, size limits of the EBS volume do not apply. This includes settings that otherwise must be within the EBS volume size limit such as S3DataDistributionType. For more information about Pipe mode and how to use it, see Using Your Own Training Algorithms - Input Data Configuration.

Use an Augmented Manifest File (Console)

To complete this procedure, you need:

- The URL of the S3 bucket where you've stored the augmented manifest file.
- To store the data that is listed in the augmented manifest file in an S3 bucket.
- The URL of the S3 bucket where you want to store the output of the job.

To use an augmented manifest file in a training job (console)

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the navigation pane, choose Training, then choose Training jobs.
3. Choose Create training job.
4. Provide a name for the training job. The name must be unique within an AWS Region in an AWS account. It can have 1 to 63 characters. Valid characters: a-z, A-Z, 0-9, and . : + = @ _ % - (hyphen).

5. Choose the algorithm that you want to use. For information about supported built-in algorithms, see Use Amazon SageMaker Built-in Algorithms or Pre-trained Models. If you want to use a custom algorithm, make sure that it is compatible with Pipe mode.

6. (Optional) For **Resource configuration**, either accept the default values or, to reduce computation time, increase the resource consumption.
   a. (Optional) For **Instance type**, choose the ML compute instance type that you want to use. In most cases, `ml.m4.xlarge` is sufficient.
   b. For **Instance count**, use the default, 1.
   c. (Optional) For **Additional volume per instance (GB)**, choose the size of the ML storage volume that you want to provision. In most cases, you can use the default, 1. If you are using a large dataset, use a larger size.

7. Provide information about the input data for the training dataset.
   a. For **Channel name**, either accept the default (`train`) or enter a more meaningful name, such as `training-augmented-manifest-file`.
   b. For **InputMode**, choose **Pipe**.
   c. For **S3 data distribution type**, choose **FullyReplicated**. When training incrementally, fully replicating causes each ML compute instance to use a complete copy of the expanded dataset. For neural-based algorithms, such as Neural Topic Model (NTM) Algorithm, choose ShardedByS3Key.
   d. If the data specified in the augmented manifest file is uncompressed, set the **Compression type** to **None**. If the data is compressed using gzip, set it to **Gzip**.
   e. (Optional) For **Content type**, specify the appropriate MIME type. Content type is the multipurpose internet mail extension (MIME) type of the data.
   f. For **Record wrapper**, if the dataset specified in the augmented manifest file is saved in RecordIO format, choose **RecordIO**. If your dataset is not saved as a RecordIO-formatted file, choose **None**.
   g. For **S3 data type**, choose **AugmentedManifestFile**.
   h. For **S3 location**, provide the path to the bucket where you stored the augmented manifest file.
i. For **AugmentedManifestFile attribute names**, specify the name of an attribute that you want to use. The attribute name must be present within the augmented manifest file, and is case-sensitive.

j. (Optional) To add more attribute names, choose **Add row** and specify another attribute name for each attribute.

k. (Optional) To adjust the order of attribute names, choose the up or down buttons next to the names. When using an augmented manifest file, the order of the specified attribute names is important.

l. Choose **Done**.

8. For **Output data configuration**, provide the following information:

   a. For **S3 location**, type the path to the S3 bucket where you want to store the output data.

   b. (Optional) You can use your AWS Key Management Service (AWS KMS) encryption key to encrypt the output data at rest. For **Encryption key**, provide the key ID or its Amazon Resource Number (ARN). For more information, see [KMS-Managed Encryption Keys](#).

9. (Optional) For **Tags**, add one or more tags to the training job. A tag is metadata that you can define and assign to AWS resources. In this case, you can use tags to help you manage your training jobs. A tag consists of a key and a value, which you define. For example, you might want to create a tag with **Project** as a key and a value that refers to a project that is related to the training job, such as **Home value forecasts**.

10. Choose **Create training job**. SageMaker creates and runs the training job.

After the training job has finished, SageMaker stores the model artifacts in the bucket whose path you provided for **S3 output path** in the **Output data configuration** field. To deploy the model to get predictions, see [Step 5: Deploy the Model to Amazon EC2](#).

### Use an Augmented Manifest File (API)

The following shows how to train a model with an augmented manifest file using the SageMaker high-level Python library:

```python
import sagemaker

# Create a model object set to using "Pipe" mode.
model = sagemaker.estimator.Estimator(
    training_image,    
```
role,
instance_count=1,
instance_type='ml.p3.2xlarge',
volume_size = 50,
max_run = 360000,
input_mode = 'Pipe',
output_path=s3_output_location,
sagemaker_session=session
)

# Create a train data channel with S3_data_type as 'AugmentedManifestFile' and attribute names.
train_data = sagemaker.inputs.TrainingInput(
    your_augmented_manifest_file,
    distribution='FullyReplicated',
    content_type='application/x-recordio',
    s3_data_type='AugmentedManifestFile',
    attribute_names=['source-ref', 'annotations'],
    input_mode='Pipe',
    record_wrapping='RecordIO'
)

data_channels = {'train': train_data}

# Train a model.
model.fit(inputs=data_channels, logs=True)

After the training job has finished, SageMaker stores the model artifacts in the bucket whose path you provided for S3 output path in the Output data configuration field. To deploy the model to get predictions, see Step 5: Deploy the Model to Amazon EC2.

**Use checkpoints in Amazon SageMaker**

Use checkpoints in Amazon SageMaker to save the state of machine learning (ML) models during training. Checkpoints are snapshots of the model and can be configured by the callback functions of ML frameworks. You can use the saved checkpoints to restart a training job from the last saved checkpoint.

Using checkpoints, you can do the following:

- Save your model snapshots under training due to an unexpected interruption to the training job or instance.
Resume training the model in the future from a checkpoint.

Analyze the model at intermediate stages of training.

Use checkpoints with S3 Express One Zone for increased access speeds.

Use checkpoints with SageMaker managed spot training to save on training costs.

The SageMaker training mechanism uses training containers on Amazon EC2 instances, and the checkpoint files are saved under a local directory of the containers (the default is `/opt/ml/checkpoints`). SageMaker provides the functionality to copy the checkpoints from the local path to Amazon S3 and automatically syncs the checkpoints in that directory with S3. Existing checkpoints in S3 are written to the SageMaker container at the start of the job, enabling jobs to resume from a checkpoint. Checkpoints added to the S3 folder after the job has started are not copied to the training container. SageMaker also writes new checkpoints from the container to S3 during training. If a checkpoint is deleted in the SageMaker container, it will also be deleted in the S3 folder.

You can use checkpoints in Amazon SageMaker with the Amazon S3 Express One Zone storage class (S3 Express One Zone) for faster access to checkpoints. When you enable checkpointing and specify the S3 URI for your checkpoint storage destination, you can provide an S3 URI for a folder in either an S3 general purpose bucket or an S3 directory bucket. For more information on S3 Express One Zone and S3 directory buckets, see What is S3 Express One Zone.

If you are using checkpoints with SageMaker managed spot training, SageMaker manages checkpointing your model training on a spot instance and resuming the training job on the next spot instance. With SageMaker managed spot training, you can significantly reduce the billable time for training ML models. For more information, see Use Managed Spot Training in Amazon SageMaker.

Topics

- Checkpoints for frameworks and algorithms in SageMaker
- Enable checkpointing
- Browse checkpoint files
- Resume training from a checkpoint
- Cluster repairs for GPU errors
- Considerations for checkpointing
Checkpoints for frameworks and algorithms in SageMaker

Use checkpoints to save snapshots of ML models built on your preferred frameworks within SageMaker.

SageMaker frameworks and algorithms that support checkpointing

SageMaker supports checkpointing for AWS Deep Learning Containers and a subset of built-in algorithms without requiring training script changes. SageMaker saves the checkpoints to the default local path '/opt/ml/checkpoints' and copies them to Amazon S3.

- Deep Learning Containers: TensorFlow, PyTorch, MXNet, and HuggingFace

  ⚠️ Note
  If you are using the HuggingFace framework estimator, you need to specify a checkpoint output path through hyperparameters. For more information, see Run training on Amazon SageMaker in the HuggingFace documentation.

- Built-in algorithms: Image Classification, Object Detection, Semantic Segmentation, and XGBoost (0.90-1 or later)

  ⚠️ Note
  If you are using the XGBoost algorithm in framework mode (script mode), you need to bring an XGBoost training script with checkpointing that's manually configured. For more information about the XGBoost training methods to save model snapshots, see Training XGBoost in the XGBoost Python SDK documentation.

If a pre-built algorithm that does not support checkpointing is used in a managed spot training job, SageMaker does not allow a maximum wait time greater than an hour for the job in order to limit wasted training time from interrupts.

For custom training containers and other frameworks

If you are using your own training containers, training scripts, or other frameworks not listed in the previous section, you must properly set up your training script using callbacks or training APIs to save checkpoints to the local path ('/opt/ml/checkpoints') and load from the local
path in your training script. SageMaker estimators can sync up with the local path and save the checkpoints to Amazon S3.

**Enable checkpointing**

After you enable checkpointing, SageMaker saves checkpoints to Amazon S3 and syncs your training job with the checkpoint S3 bucket. You can use either S3 general purpose or S3 directory buckets for your checkpoint S3 bucket.

The following example shows how to configure checkpoint paths when you construct a SageMaker estimator. To enable checkpointing, add the `checkpoint_s3_uri` and `checkpoint_local_path` parameters to your estimator.

The following example template shows how to create a generic SageMaker estimator and enable checkpointing. You can use this template for the supported algorithms by specifying the `image_uri` parameter. To find Docker image URIs for algorithms with checkpointing supported by SageMaker, see [Docker Registry Paths and Example Code](#). You can also replace `estimator` and `Estimator` with other SageMaker frameworks' estimator parent classes and estimator classes, such as [TensorFlow](#), [PyTorch](#), [MXNet](#), [HuggingFace](#) and [XGBoost](#).

```python
import sagemaker
from sagemaker.estimator import Estimator

bucket=sagemaker.Session().default_bucket()
base_job_name="sagemaker-checkpoint-test"
checkpoint_in_bucket="checkpoints"
```
# The S3 URI to store the checkpoints
checkpoint_s3_bucket="s3:///{}/{}/{}".format(bucket, base_job_name,
    checkpoint_in_bucket)

# The local path where the model will save its checkpoints in the training container
checkpoint_local_path="/opt/ml/checkpoints"

estimator = Estimator(
    ...
    image_uri="<ecr_path>/{}:<tag>" # Specify to use built-in algorithms
    output_path=bucket,
    base_job_name=base_job_name,

    # Parameters required to enable checkpointing
    checkpoint_s3_uri=checkpoint_s3_bucket,
    checkpoint_local_path=checkpoint_local_path
)

The following two parameters specify paths for checkpointing:

- **checkpoint_local_path** – Specify the local path where the model saves the checkpoints periodically in a training container. The default path is set to '/opt/ml/checkpoints'. If you are using other frameworks or bringing your own training container, ensure that your training script's checkpoint configuration specifies the path to '/opt/ml/checkpoints'.

  ![Note]

  We recommend specifying the local paths as '/opt/ml/checkpoints' to be consistent with the default SageMaker checkpoint settings. If you prefer to specify your own local path, make sure you match the checkpoint saving path in your training script and the checkpoint_local_path parameter of the SageMaker estimators.

- **checkpoint_s3_uri** – The URI to an S3 bucket where the checkpoints are stored in real time. You can specify either an S3 general purpose or S3 directory bucket to store your checkpoints. For more information on S3 directory buckets, see Directory buckets in the Amazon Simple Storage Service User Guide.

To find a complete list of SageMaker estimator parameters, see the Estimator API in the Amazon SageMaker Python SDK documentation.
Browse checkpoint files

Locate checkpoint files using the SageMaker Python SDK and the Amazon S3 console.

To find the checkpoint files programmatically

To retrieve the S3 bucket URI where the checkpoints are saved, check the following estimator attribute:

```
estimator.checkpoint_s3_uri
```

This returns the S3 output path for checkpoints configured while requesting the CreateTrainingJob request. To find the saved checkpoint files using the S3 console, use the following procedure.

To find the checkpoint files from the S3 console

2. In the left navigation pane, choose Training jobs.
3. Choose the link to the training job with checkpointing enabled to open Job settings.
4. On the Job settings page of the training job, locate the Checkpoint configuration section.

<table>
<thead>
<tr>
<th>Checkpoint configuration</th>
</tr>
</thead>
<tbody>
<tr>
<td>S3 output path</td>
</tr>
<tr>
<td>s3://path-to-your-checkpoint</td>
</tr>
<tr>
<td>Local path</td>
</tr>
<tr>
<td>/opt/ml/checkpoints/</td>
</tr>
</tbody>
</table>

5. Use the link to the S3 bucket to access the checkpoint files.

Resume training from a checkpoint

To resume a training job from a checkpoint, run a new estimator with the same checkpoint_s3_uri that you created in the Enable checkpointing section. Once the training has
resumed, the checkpoints from this S3 bucket are restored to checkpoint_local_path in each instance of the new training job. Ensure that the S3 bucket is in the same Region as that of the current SageMaker session.

Cluster repairs for GPU errors

If you are running a training job that fails on a GPU, SageMaker will run a GPU health check to see whether the failure is related to a GPU issue. SageMaker takes the following actions based on the health check results:

- If the error is recoverable, and can be fixed by rebooting the instance or resetting the GPU, SageMaker will reboot the instance.
- If the error is not recoverable, and caused by a GPU that needs to be replaced, SageMaker will replace the instance.

The instance is either replaced or rebooted as part of a SageMaker cluster repair process. During this process, you will see the following message in your training job status:

Repairing training cluster due to hardware failure

SageMaker will attempt to repair the cluster up to 10 times. If the cluster repair is successful, SageMaker will automatically restart the training job from the previous checkpoint. If the cluster repair fails, the training job will also fail. You are not billed for the cluster repair process. Cluster repairs will not initiate unless your training job fails. If a GPU issue is detected for a warmpool cluster, the cluster will enter into repair mode to either reboot or replace the faulty instance. After repair, the cluster can still be used as a warmpool cluster.
The previously described cluster and instance repair process is depicted in the following diagram:

Considerations for checkpointing

Consider the following when using checkpoints in SageMaker.

- To avoid overwrites in distributed training with multiple instances, you must manually configure the checkpoint file names and paths in your training script. The high-level SageMaker checkpoint configuration specifies a single Amazon S3 location without additional suffixes or prefixes to tag checkpoints from multiple instances.
• The SageMaker Python SDK does not support high-level configuration for checkpointing frequency. To control the checkpointing frequency, modify your training script using the framework's model save functions or checkpoint callbacks.

• If you use SageMaker checkpoints with SageMaker Debugger and SageMaker distributed and are facing issues, see the following pages for troubleshooting and considerations.
  - Considerations for Amazon SageMaker Debugger
  - Troubleshooting for distributed training in Amazon SageMaker
  - Model Parallel Troubleshooting
Deploy models for inference

With Amazon SageMaker, you can deploy your machine learning (ML) models to make predictions, also known as inference. SageMaker provides a broad selection of ML infrastructure and model deployment options to help meet all your ML inference needs. It is a fully managed service and integrates with MLOps tools, so you can scale your model deployment, reduce inference costs, manage models more effectively in production, and reduce operational burden.

After you've built and trained a machine learning model, you can use SageMaker Inference to start getting predictions, or inferences, from your model. With SageMaker Inference, you can either set up an endpoint that returns inferences or run batch inferences from your model.

To get started with SageMaker Inference, see the following sections and review the Inference options to determine which feature best fits your use case.

You can refer to the Resources section for more troubleshooting and reference information, blogs and examples to help you get started, and common FAQs.

Topics

- Before you begin
- Steps for model deployment
- Inference options
- Advanced endpoint options
- Bring your own model
- Next steps

Before you begin

These topics assume that you have built and trained one or more machine learning models and are ready to deploy them. You don't need to train your model in SageMaker in order to deploy your model in SageMaker and get inferences. If you don't have your own model, you can also use SageMaker's built-in algorithms or pre-trained models.

If you are new to SageMaker and haven't picked out a model to deploy, work through the steps in the Get Started with Amazon SageMaker tutorial to familiarize yourself with an example of how
Steps for model deployment

For inference endpoints, the general workflow consists of the following:

- Create a model in SageMaker Inference by pointing to model artifacts stored in Amazon S3 and a container image.
- Select an inference option. For more information, see Inference options.
- Create a SageMaker Inference endpoint configuration by choosing the instance type and number of instances you need behind the endpoint. You can use Amazon SageMaker Inference Recommender to get recommendations for instance types. For Serverless Inference, you only need to provide the memory configuration you need based on your model size.
- Create a SageMaker Inference endpoint.
- Invoke your endpoint to receive an inference as a response.

The following diagram shows the preceding workflow.
You can perform these actions using the AWS console, the AWS SDKs, the SageMaker Python SDK, AWS CloudFormation or the AWS CLI.

For batch inference with batch transform, point to your model artifacts and input data and create a batch inference job. Instead of hosting an endpoint for inference, SageMaker outputs your inferences to an Amazon S3 location of your choice.

**Inference options**

SageMaker provides multiple inference options so that you can pick the option that best suits your workload:

- **Real-Time Inference**: Real-time inference is ideal for online inferences that have low latency or high throughput requirements. Use real-time inference for a persistent and fully managed endpoint (REST API) that can handle sustained traffic, backed by the instance type of your choice. Real-time inference can support payload sizes up to 6 MB and processing times of 60 seconds.

- **Serverless Inference**: Serverless inference is ideal when you have intermittent or unpredictable traffic patterns. SageMaker manages all of the underlying infrastructure, so there’s no need to manage instances or scaling policies. You pay only for what you use and not for idle time. It can support payload sizes up to 4 MB and processing times up to 60 seconds.

- **Batch Transform**: Batch transform is suitable for offline processing when large amounts of data are available upfront and you don’t need a persistent endpoint. You can also use batch transform for pre-processing datasets. It can support large datasets that are GBs in size and processing times of days.

- **Asynchronous Inference**: Asynchronous inference is ideal when you want to queue requests and have large payloads with long processing times. Asynchronous Inference can support payloads up to 1 GB and long processing times up to one hour. You can also scale down your endpoint to 0 when there are no requests to process.

The following diagram shows the preceding information in a flowchart and can help you choose the option that best fits your use case.
### Choosing Model Deployment Options

<table>
<thead>
<tr>
<th>Does your workload need to return an inference for each request to your model?</th>
<th>Would it be helpful to queue requests due to longer processing times or larger payloads?</th>
<th>Does your workload have intermittent traffic patterns or periods of no traffic?</th>
<th>Does your workload have sustained traffic and need lower and consistent latency?</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Batch Transform</td>
<td>Asynchronous Inference</td>
<td>Serverless Inference</td>
<td>Real-Time Inference</td>
</tr>
<tr>
<td>Payload size: GBs</td>
<td>Payload size: 1 GB</td>
<td>Payload size: 4 MB</td>
<td>Payload size: 6 MB</td>
</tr>
<tr>
<td>Runtime: days</td>
<td>Runtime: 1 hour</td>
<td>Runtime: 60 sec</td>
<td>Runtime: 60 sec</td>
</tr>
</tbody>
</table>

### Advanced endpoint options

With real-time inference, you can further optimize for performance and cost with the following advanced inference options:

- If you have multiple models that use the same framework and can share a container, then use **Host multiple models in one container behind one endpoint**. This option helps you optimize costs by improving endpoint utilization and reducing deployment overhead.

- If you have multiple models that use different frameworks and require their own containers, then use **Host multiple models which use different containers behind one endpoint**. With this option, you get many of the benefits of Multi-Model Endpoints and can deploy a variety of frameworks and models.

- If you want to host models with pre-processing and post-processing logic behind an endpoint, then use **Serial Inference Pipelines**. Inference pipelines are fully managed by SageMaker and provide lower latency because all of the containers are hosted on the same Amazon EC2 instances.
Bring your own model

To use an existing Docker container in SageMaker, see Adapting your own Docker container to work with SageMaker.

To create a new Docker container and receive more advanced guidance on how to run your own inference code, see the following links.

- To run your own inference code hosting services, see Use Your Own Inference Code with Hosting Services.
- To run your own inference code for batch inference, see Use Your Own Inference Code with Batch Transform.

Next steps

After you have an endpoint and understand the general inference workflow, you can use the following features within SageMaker Inference to improve your inference workflow.

Monitoring

To track your model over time through metrics such as model accuracy and drift, you can use Model Monitor. With Model Monitor, you can set alerts that notify you when there are deviations in your model's quality. To learn more, see the Model Monitor documentation. To learn more about tools that can be used to monitor model deployments and events that change your endpoint, see Monitor Amazon SageMaker. For example, you can monitor your endpoint’s health through metrics such as invocation errors and model latency using Amazon CloudWatch metrics. The SageMaker endpoint invocation metrics can provide you with valuable information about your endpoint's performance.

CI/CD for model deployment

To put together machine learning solutions in SageMaker, you can use SageMaker MLOps. You can use this feature to automate the steps in your machine learning workflow and practice CI/CD. You can use MLOps Project Templates to help with the setup and implementation of SageMaker MLOps projects. SageMaker also supports using your own third-party Git repo for creating a CI/CD system.

For your ML pipelines, use Model Registry to manage your model versions and the deployment and automation of your models.
Deployment guardrails

If you want to update your model while it’s in production without impacting production, you can use deployment guardrails. Deployment guardrails are a set of model deployment options in SageMaker Inference to update your machine learning models in production. Using the fully managed deployment options, you can control the switch from the current model in production to a new one. Traffic shifting modes give you granular control over the traffic shifting process, and built-in safeguards like auto-rollback help you catch issues early on. To learn more about deployment guardrails, see the [deployment guardrails documentation](#).

Inferentia

If you need to run large scale machine learning and deep learning applications for use cases such as image or speech recognition, natural language processing (NLP), personalization, forecasting, or fraud detection, you can use an Inf1 instance with a real-time endpoint.

Inf1 instances are built to support machine learning inference applications and feature the AWS Inferentia chips. Inf1 instances provide higher throughput and lower cost per inference than GPU-based instances.

To deploy a model on Inf1 instances, compile your model with SageMaker Neo and choose an Inf1 instance for your deployment option. To learn more, see [Optimize model performance using SageMaker Neo](#).

Optimize model performance

SageMaker provides features to manage resources and optimize inference performance when deploying machine learning models. You can use SageMaker’s [built-in algorithms and pre-built models](#), as well as [prebuilt Docker images](#), which are developed for machine learning. To train TensorFlow, Apache MXNet, PyTorch, ONNX, and XGBoost models once and optimize them to deploy on ARM, Intel, and Nvidia processors, see [Optimize model performance using SageMaker Neo](#).

Autoscaling

If you have varying amounts of traffic to your endpoints, you might want to try autoscaling. For example, during peak hours, you might require more instances to process requests, but during periods of low traffic, you might want to reduce your use of computing resources. To
dynamically adjust the number of instances provisioned in response to changes in your workload, see [Automatically Scale Amazon SageMaker Models](#).

If you have unpredictable traffic patterns or don’t want to set up scaling policies, you can also use Serverless Inference for an endpoint where SageMaker manages autoscaling for you. During periods of low traffic, SageMaker scales down your endpoint, and if traffic increases, then SageMaker scales your endpoint up. For more information, see the [Serverless Inference documentation](#).

## Deploy a Model in Amazon SageMaker

After you train your machine learning model, you can deploy it using Amazon SageMaker to get predictions in any of the following ways, depending on your use case:

- For persistent, real-time endpoints that make one prediction at a time, use SageMaker real-time hosting services. See [Real-time inference](#).
- Workloads that have idle periods between traffic spurts and can tolerate cold starts, use Serverless Inference. See [Serverless Inference](#).
- Requests with large payload sizes up to 1GB, long processing times, and near real-time latency requirements, use Amazon SageMaker Asynchronous Inference. See [Asynchronous inference](#).
- To get predictions for an entire dataset, use SageMaker batch transform. See [Use Batch Transform](#).

SageMaker also provides features to manage resources and optimize inference performance when deploying machine learning models:

- To manage models on edge devices so that you can optimize, secure, monitor, and maintain machine learning models on fleets of edge devices such as smart cameras, robots, personal computers, and mobile devices, see [Deploy models at the edge with SageMaker Edge Manager](#).
- To optimize Gluon, Keras, MXNet, PyTorch, TensorFlow, TensorFlow-Lite, and ONNX models for inference on Android, Linux, and Windows machines based on processors from Ambarella, ARM, Intel, Nvidia, NXP, Qualcomm, Texas Instruments, and Xilinx, see [Optimize model performance using Neo](#).

For more information about all deployment options, see [Deploy models for inference](#).
Create a model in Amazon SageMaker with ModelBuilder

Preparing your model for deployment on a SageMaker endpoint requires multiple steps, including choosing a model image, setting up the endpoint configuration, coding your serialization and deserialization functions to transfer data to and from server and client, identifying model dependencies, and uploading them to Amazon S3. ModelBuilder can reduce the complexity of initial setup and deployment to help you create a deployable model in a single step.

ModelBuilder performs the following tasks for you:

- Converts machine learning models trained using various frameworks like XGBoost or PyTorch into deployable models in one step.
- Performs automatic container selection based on the model framework so you don’t have to manually specify your container. You can still bring your own container by passing your own URI to ModelBuilder.
- Handles the serialization of data on the client side before sending it to the server for inference and deserialization of the results returned by the server. Data is correctly formatted without manual processing.
- Enables automatic capture of dependencies and packages the model according to model server expectations. ModelBuilder’s automatic capture of dependencies is a best-effort approach to dynamically load dependencies. (We recommend that you test the automated capture locally and update the dependencies to meet your needs.)
- For large language model (LLM) use cases, optionally performs local parameter tuning of serving properties that can be deployed for better performance when hosting on a SageMaker endpoint.
- Supports most of the popular model servers and containers like TorchServe, Triton, DJLServing and TGI container.

Build your model with ModelBuilder

ModelBuilder is a Python class that takes a framework model, such as XGBoost or PyTorch, or a user-specified inference specification and converts it into a deployable model. ModelBuilder provides a build function that generates the artifacts for deployment. The model artifact generated is specific to the model server, which you can also specify as one of the inputs. For more details about the ModelBuilder class, see ModelBuilder.
The following diagram illustrates the overall model creation workflow when you use `ModelBuilder`. `ModelBuilder` accepts a model or inference specification along with your schema to create a deployable model that you can test locally before deployment.

ModelBuilder can handle any customization you want to apply. However, to deploy a framework model, the model builder expects at minimum a model, sample input and output, and the role. In the following code example, `ModelBuilder` is called with a framework model and an instance of `SchemaBuilder` with minimum arguments (to infer the corresponding functions for serializing and deserializing the endpoint input and output). No container is specified and no packaged dependencies are passed—SageMaker automatically infers these resources when you build your model.

```python
from sagemaker.serve.builder.model_builder import ModelBuilder
from sagemaker.serve.builder.schema_builder import SchemaBuilder

model_builder = ModelBuilder(
    model=model,
    schema_builder=SchemaBuilder(input, output),
    role_arn="execution-role",
)
```

The following code sample invokes `ModelBuilder` with an inference specification (as an `InferenceSpec` instance) instead of a model, with additional customization. In this case, the call to model builder includes a path to store model artifacts and also turns on autocapture of all available dependencies. For additional details about `InferenceSpec`, see [Customize model loading and handling of requests](#).

```python
model_builder = ModelBuilder(
    mode=Mode.LOCAL_CONTAINER,
    model_path="model-artifact-directory",
    inference_spec="your-inference-spec",
)
```
Define serialization and deserialization methods

When invoking a SageMaker endpoint, the data is sent through HTTP payloads with different MIME types. For example, an image sent to the endpoint for inference needs to be converted to bytes at the client side and sent through an HTTP payload to the endpoint. When the endpoint receives the payload, it needs to deserialize the byte string back to the data type that is expected by the model (also known as server-side deserialization). After the model finishes prediction, the results also need to be serialized to bytes that can be sent back through the HTTP payload to the user or the client. Once the client receives the response byte data, it needs to perform client-side deserialization to convert the bytes data back to the expected data format, such as JSON. At minimum, you need to convert data for the following tasks:

1. Inference request serialization (handled by the client)
2. Inference request deserialization (handled by the server or algorithm)
3. Invoking the model against the payload and send response payload back
4. Inference response serialization (handled by the server or algorithm)
5. Inference response deserialization (handled by the client)

The following diagram shows the serialization and deserialization processes that occur when you invoke the endpoint.
When you supply sample input and output to SchemaBuilder, the schema builder generates the corresponding marshalling functions for serializing and deserializing the input and output. You can further customize your serialization functions with CustomPayloadTranslator. But for most cases, a simple serializer such as the following would work:

```python
input = "How is the demo going?"
output = "Comment la démo va-t-elle?"
schema = SchemaBuilder(input, output)
```

For further details about SchemaBuilder, see [SchemaBuilder](#).

The following code snippet outlines an example where you want to customize both serialization and deserialization functions at the client and server sides. You can define your own request and response translators with CustomPayloadTranslator and pass these translators to SchemaBuilder.

By including the inputs and outputs with the translators, the model builder can extract the data format the model expects. For example, suppose the sample input is a raw image, and your custom translators crop the image and send the cropped image to the server as a tensor. ModelBuilder needs both the raw input and any custom preprocessing or postprocessing code to derive a method to convert data on both the client and server sides.

```python
from sagemaker.serve import CustomPayloadTranslator

# request translator
class MyRequestTranslator(CustomPayloadTranslator):
    # This function converts the payload to bytes - happens on client side
def serialize_payload_to_bytes(self, payload: object) -> bytes:
        # converts the input payload to bytes
        # ... ...
        return  //return object as bytes

    # This function converts the bytes to payload - happens on server side
def deserialize_payload_from_stream(self, stream) -> object:
        # convert bytes to in-memory object
        # ... ...
        return //return in-memory object

# response translator
class MyResponseTranslator(CustomPayloadTranslator):
    # This function converts the payload to bytes - happens on server side
```
def serialize_payload_to_bytes(self, payload: object) -> bytes:
    # converts the response payload to bytes
    return //return object as bytes

# This function converts the bytes to payload - happens on client side
def deserialize_payload_from_stream(self, stream) -> object:
    # convert bytes to in-memory object
    return //return in-memory object

You pass in the sample input and output along with the previously-defined custom translators when you create the SchemaBuilder object, as shown in the following example:

my_schema = SchemaBuilder(
    sample_input=image,
    sample_output=output,
    input_translator=MyRequestTranslator(),
    output_translator=MyResponseTranslator()
)

Then you pass in the sample input and output, along with the custom translators defined previously, to the SchemaBuilder object.

my_schema = SchemaBuilder(
    sample_input=image,
    sample_output=output,
    input_translator=MyRequestTranslator(),
    output_translator=MyResponseTranslator()
)

The following sections explain in detail how to build your model with ModelBuilder and use its supporting classes to customize the experience for your use case.

**Topics**

- [Customize model loading and handling of requests](#)
- [Build your model and deploy](#)
- [Bring your own container (BYOC)](#)
- [Using ModelBuilder in local mode](#)
- [ModelBuilder examples](#)
Customize model loading and handling of requests

Providing your own inference code through `InferenceSpec` offers an additional layer of customization. With `InferenceSpec`, you can customize how the model is loaded and how it handles incoming inference requests, bypassing its default loading and inference handling mechanisms. This flexibility is particularly beneficial when working with non-standard models or custom inference pipelines. You can customize the `invoke` method to control how the model preprocesses and postprocesses incoming requests. The `invoke` method ensures that the model handles inference requests correctly. The following example uses `InferenceSpec` to generate a model with the HuggingFace pipeline. For further details about `InferenceSpec`, refer to the `InferenceSpec`.

```python
from sagemaker.serve.spec.inference_spec import InferenceSpec
from transformers import pipeline

class MyInferenceSpec(InferenceSpec):
    def load(self, model_dir: str):
        return pipeline("translation_en_to_fr", model="t5-small")

    def invoke(self, input, model):
        return model(input)

inf_spec = MyInferenceSpec()

model_builder = ModelBuilder(
    inference_spec=your_inference_spec,
    schema_builder=SchemaBuilder(X_test, y_pred)
)
```

The following example illustrates a more customized variation of a previous example. A model is defined with an inference specification that has dependencies. In this case, the code in the inference specification is dependent on the `lang-segment` package. The argument for dependencies contains a statement that directs the builder to install `lang-segment` using Git. Since the model builder is directed by the user to custom install a dependency, the auto key is False to turn off autocapture of dependencies.

```python
model_builder = ModelBuilder(
    mode=Mode.LOCAL_CONTAINER,
    model_path=model_artifact_directory,
    inference_spec=your_inference-spec,
)```
Build your model and deploy

Call the build function to create your deployable model. This step creates inference code (as `inference.py`) in your working directory with the code necessary to create your schema, run serialization and deserialization of inputs and outputs, and run other user-specified custom logic.

As an integrity check, SageMaker packages and pickles the necessary files for deployment as part of the ModelBuilder build function. During this process, SageMaker also creates HMAC signing for the pickle file and adds the secret key in the CreateModel API as an environment variable during deploy (or create). The endpoint launch uses the environment variable to validate the integrity of the pickle file.

```python
# Build the model according to the model server specification and save it as files in the working directory
model = model_builder.build()
```

Deploy your model with the model's existing deploy method. In this step, SageMaker sets up an endpoint to host your model as it starts making predictions on incoming requests. Although the ModelBuilder infers the endpoint resources needed to deploy your model, you can override those estimates with your own parameter values. The following example directs SageMaker to deploy the model on a single `ml.c6i.xlarge` instance. A model constructed from ModelBuilder enables live logging during deployment as an added feature.

```python
predictor = model.deploy(
    initial_instance_count=1,
    instance_type="ml.c6i.xlarge"
)
```

If you want more fine-grained control over the endpoint resources assigned to your model, you can use a ResourceRequirements object. With the ResourceRequirements object, you can request a minimum number of CPUs, accelerators, and copies of models you want to deploy. You can also request a minimum and maximum bound of memory (in MB). To use this feature, you need to specify your endpoint type as EndpointType.INFERENCE_COMPONENT_BASED. The following
example requests four accelerators, a minimum memory size of 1024 MB, and one copy of your model to be deployed to an endpoint of type `EndpointType.INFERENCE_COMPONENT_BASED`.

```python
resource_requirements = ResourceRequirements(
    requests={
        "num_accelerators": 4,
        "memory": 1024,
        "copies": 1,
    },
    limits={},
)
predictor = model.deploy(
    mode=Mode.SAGEMAKER_ENDPOINT,
    endpoint_type=EndpointType.INFERENCE_COMPONENT_BASED,
    resources=resource_requirements,
    role="role"
)
```

### Bring your own container (BYOC)

If you want to bring your own container (extended from a SageMaker container), you can also specify the image URI as shown in the following example. You also need to identify the model server that corresponds to the image for `ModelBuilder` to generate artifacts specific to the model server.

```python
model_builder = ModelBuilder(
    model=model,
    model_server=ModelServer.TORCHSERVE,
    schema_builder=SchemaBuilder(X_test, y_pred),
    image_uri="123123123123.dkr.ecr.ap-southeast-2.amazonaws.com/byoc-image:xgb-1.7-1")
```

### Using ModelBuilder in local mode

You can deploy your model locally by using the mode argument to switch between local testing and deployment to an endpoint. You need to store the model artifacts in the working directory, as shown in the following snippet:

```python
model = XGBClassifier()
model.fit(X_train, y_train)
```
Pass the model object, a SchemaBuilder instance, and set mode to Mode.LOCAL_CONTAINER. When you call the build function, ModelBuilder automatically identifies the supported framework container and scans for dependencies. The following example demonstrates model creation with an XGBoost model in local mode.

```python
model.save_model(model_dir + "\my_model.xgb")
```

```python
model_builder_local = ModelBuilder(
    model=model,
    schema_builder=SchemaBuilder(X_test, y_pred),
    role_arn=execution_role,
    mode=Mode.LOCAL_CONTAINER
)

xgb_local_builder = model_builder_local.build()
```

Call the deploy function to deploy locally, as shown in the following snippet. If you specify parameters for instance type or count, these arguments are ignored.

```python
predictor_local = xgb_local_builder.deploy()
```

**Troubleshooting local mode**

Depending on your individual local setup, you may encounter difficulties running ModelBuilder smoothly in your environment. See the following list for some issues you may face and how to resolve them.

- **Already already in use**: You may encounter an Address already in use error. In this case, it is possible that a Docker container is running on that port or another process is utilizing it. You can follow the approach outlined in [Linux documentation](https://www.linuxdocumentation.org) to identify the process and gracefully redirect your local process from port 8080 to another port or clean up the Docker instance.

- **IAM Permission Issue**: You might encounter a permission issue when trying to pull an Amazon ECR image or access Amazon S3. In this case, navigate to the execution role of the notebook or Studio Classic instance to verify the policy for SageMakerFullAccess or the respective API permissions.

- **EBS volume capacity issue**: If you deploy a large language model (LLM), you might run out of space while running Docker in local mode or experience space limitations for the Docker cache. In this case, you can try to move your Docker volume to a filesystem that has enough space. To move your Docker volume, complete the following steps:
1. Open a terminal and run `df` to display disk usage, as shown in the following output:

```bash
(python3) sh-4.2$ df
Filesystem     1K-blocks      Used  Available  Use%  Mounted on
devtmpfs       195928700       0   195928700  0%   /dev
tmpfs          195939296       0   195939296  0%   /dev/shm
tmpfs          195939296  1048  195938248   1%   /run
tmpfs          195939296       0   195939296  0%   /sys/fs/cgroup
/dev/nvme0n1p1 141545452 135242112    6303340  96%   /
tmpfs          39187860       0   39187860  0%   /run/user/0
/dev/nvme2n1   264055236  76594068 176644712  31%   /home/ec2-user/SageMaker
tmpfs          39187860       0   39187860  0%   /run/user/1002
tmpfs          39187860       0   39187860  0%   /run/user/1001
tmpfs          39187860       0   39187860  0%   /run/user/1000
```

2. Move the default Docker directory from `/dev/nvme0n1p1` to `/dev/nvme2n1` so you can fully utilize the 256 GB SageMaker volume. For more details, see documentation about how to move your Docker directory.

3. Stop Docker with the following command:

```
sudo service docker stop
```

4. Add a `daemon.json` to `/etc/docker` or append the following JSON blob to the existing one.

```json
{
    "data-root": "/home/ec2-user/SageMaker/{created_docker_folder}"
}
```

5. Move the Docker directory in `/var/lib/docker` to `/home/ec2-user/SageMaker` with the following command:

```
sudo rsync -aP /var/lib/docker/ /home/ec2-user/SageMaker/{created_docker_folder}
```

6. Start Docker with the following command:

```
sudo service docker start
```

7. Clean trash with the following command:
8. If you are using a SageMaker notebook instance, you can follow the steps in the Docker prep file to prepare Docker for local mode.

**ModelBuilder examples**

For more examples of using ModelBuilder to build your models, see ModelBuilder sample notebooks.

**Validate a Machine Learning Model**

After training a model, evaluate it to determine whether its performance and accuracy enable you to achieve your business goals. You might generate multiple models using different methods and evaluate each. For example, you could apply different business rules for each model, and then apply various measures to determine each model's suitability. You might consider whether your model needs to be more sensitive than specific (or vice versa).

You can evaluate your model using historical data (offline) or live data:

- **Offline testing**—Use historical, not live, data to send requests to the model for inferences.

  Deploy your trained model to an alpha endpoint, and use historical data to send inference requests to it. To send the requests, use a Jupyter notebook in your Amazon SageMaker notebook instance and either the AWS SDK for Python (Boto) or the high-level Python library provided by SageMaker.

- **Online testing with live data**—SageMaker supports A/B testing for models in production by using production variants. Production variants are models that use the same inference code and are deployed on the same SageMaker endpoint. You configure the production variants so that a small portion of the live traffic goes to the model that you want to validate. For example, you might choose to send 10% of the traffic to a model variant for evaluation. After you are satisfied with the model's performance, you can route 100% traffic to the updated model. For an example of testing models in production, see Production variants.

For more information, see articles and books about how to evaluate models, for example, Evaluating Machine Learning Models.
Options for offline model evaluation include:

- **Validating using a holdout set**—Machine learning practitioners often set aside a part of the data as a "holdout set." They don’t use this data for model training.

  With this approach, you evaluate how well your model provides inferences on the holdout set. You then assess how effectively the model generalizes what it learned in the initial training, as opposed to using model memory. This approach to validation gives you an idea of how often the model is able to infer the correct answer.

  In some ways, this approach is similar to teaching elementary school students. First, you provide them with a set of examples to learn, and then test their ability to generalize from their learning. With homework and tests, you pose problems that were not included in the initial learning and determine whether they are able to generalize effectively. Students with perfect memories could memorize the problems, instead of learning the rules.

  Typically, the holdout dataset is of 20-30% of the training data.

- **k-fold validation**—In this validation approach, you split the example dataset into $k$ parts. You treat each of these parts as a holdout set for $k$ training runs, and use the other $k-1$ parts as the training set for that run. You produce $k$ models using a similar process, and aggregate the models to generate your final model. The value $k$ is typically in the range of 5-10.

### Amazon SageMaker Inference Recommender

Amazon SageMaker Inference Recommender is a capability of Amazon SageMaker that reduces the time required to get machine learning (ML) models in production by automating load testing and model tuning across SageMaker ML instances. You can use Inference Recommender to deploy your model to a real-time or serverless inference endpoint that delivers the best performance at the lowest cost. Inference Recommender helps you select the best instance type and configuration (such as instance count, container parameters, and model optimizations) or serverless configuration (such as max concurrency and memory size) for your ML models and workloads.
Amazon SageMaker Inference Recommender only charges you for the instances used while your jobs are executing.

**How it Works**

To use Amazon SageMaker Inference Recommender, you can either create a SageMaker model or register a model to the SageMaker model registry with your model artifacts. Use the AWS SDK for Python (Boto3) or the SageMaker console to run benchmarking jobs for different SageMaker endpoint configurations. Inference Recommender jobs help you collect and visualize metrics across performance and resource utilization to help you decide on which endpoint type and configuration to choose.

**How to Get Started**

If you are a first-time user of Amazon SageMaker Inference Recommender, we recommend that you do the following:

1. Read through the Prerequisites section to make sure you have satisfied the requirements to use Amazon SageMaker Inference Recommender.
2. Read through the Recommendation jobs section to launch your first Inference Recommender recommendation jobs.
3. Explore the introductory Amazon SageMaker Inference Recommender Jupyter notebook example, or review the example notebooks in the following section.

**Example notebooks**

The following example Jupyter notebooks can help you with the workflows for multiple use cases in Inference Recommender:

- If you want an introductory notebook that benchmarks a TensorFlow model, see the [SageMaker Inference Recommender TensorFlow](#) notebook.
- If you want to benchmark a HuggingFace model, see the [SageMaker Inference Recommender for HuggingFace](#) notebook.
- If you want to benchmark an XGBoost model, see the [SageMaker Inference Recommender XGBoost](#) notebook.
- If you want to review CloudWatch metrics for your Inference Recommender jobs, see the [SageMaker Inference Recommender CloudWatch metrics](#) notebook.
Prerequisites

To use Amazon SageMaker Inference Recommender, first make sure you have met the prerequisites in the following list. As an example, we show how to use a PyTorch (v1.7.1) ResNet-18 pre-trained model for both types of Amazon SageMaker Inference Recommender recommendation jobs. The examples shown use the AWS SDK for Python (Boto3).

Note

- The following code examples use Python. Remove the ! prefix character if you run any of the following code samples in your terminal or AWS CLI.
- You can run the following examples with the Python 3 (TensorFlow 2.6 Python 3.8 CPU Optimized) kernel in an Amazon SageMaker Studio notebook. For more information about Studio, see Amazon SageMaker Studio.

1. Create an IAM role for Amazon SageMaker.

Create an IAM role for Amazon SageMaker that has the AmazonSageMakerFullAccess IAM managed policy attached.

2. Set up your environment.

Import dependencies and create variables for your AWS Region, your SageMaker IAM role (from Step 1), and the SageMaker client.

```python
!pip install --upgrade pip awscli botocore boto3 --quiet
from sagemaker import get_execution_role, Session, image_uris
import boto3
region = boto3.Session().region_name
role = get_execution_role()
sagemaker_client = boto3.client("sagemaker", region_name=region)
sagemaker_session = Session()
```

3. (Optional) Review existing models benchmarked by Inference Recommender.

Inference Recommender benchmarks models from popular model zoos. Inference Recommender supports your model even if it is not already benchmarked.
Use ListModelMetaData to get a response object that lists the domain, framework, task, and model name of machine learning models found in common model zoos.

You use the domain, framework, framework version, task, and model name in later steps to both select an inference Docker image and register your model with SageMaker Model Registry. The following demonstrates how to list model metadata with SDK for Python (Boto3):

```python
list_model_metadata_response=sagemaker_client.list_model_metadata()
```

The output includes model summaries (ModelMetadataSummaries) and response metadata (ResponseMetadata) similar to the following example:

```json
{
    'ModelMetadataSummaries': [{
        'Domain': 'NATURAL_LANGUAGE_PROCESSING',
        'Framework': 'PYTORCH:1.6.0',
        'Model': 'bert-base-cased',
        'Task': 'FILL_MASK'
    },
    {
        'Domain': 'NATURAL_LANGUAGE_PROCESSING',
        'Framework': 'PYTORCH:1.6.0',
        'Model': 'bert-base-uncased',
        'Task': 'FILL_MASK'
    },
    {
        'Domain': 'COMPUTER_VISION',
        'Framework': 'MXNET:1.8.0',
        'Model': 'resnet18v2-gluon',
        'Task': 'IMAGECLASSIFICATION'
    },
    {
        'Domain': 'COMPUTER_VISION',
        'Framework': 'PYTORCH:1.6.0',
        'Model': 'resnet152',
        'Task': 'IMAGECLASSIFICATION'
    }],
    'ResponseMetadata': {
        'HTTPHeaders': {
            'content-length': '2345',
```
For this demo, we use a PyTorch (v1.7.1) ResNet-18 model to perform image classification. The following Python code sample stores the framework, framework version, domain, and task into variables for later use:

```python
# ML framework details
framework = 'pytorch'
framework_version = '1.7.1'

# ML model details
ml_domain = 'COMPUTER_VISION'
ml_task = 'IMAGE_CLASSIFICATION'
```

4. **Upload your machine learning model to Amazon S3.**

Use this PyTorch (v1.7.1) ResNet-18 model if you do not have a pre-trained machine learning model:

```python
# Optional: Download a sample PyTorch model
import torch
from torchvision import models, transforms, datasets

# Create an example input for tracing
image = torch.zeros([1, 3, 256, 256], dtype=torch.float32)

# Load a pretrained resnet18 model from TorchHub
model = models.resnet18(pretrained=True)

# Tell the model we are using it for evaluation (not training). Note this is required for Inferentia compilation.
model.eval()
model_trace = torch.jit.trace(model, image)
```
# Save your traced model
model_trace.save('model.pth')

Download a sample inference script `inference.py`. Create a code directory and move the inference script to the code directory.

```bash
# Download the inference script

# move it into a code/ directory
!mkdir code
!mv inference.py code/
```

Amazon SageMaker requires pre-trained machine learning models to be packaged as a compressed TAR file (*tar.gz*). Compress your model and inference script to satisfy this requirement:

```bash
!tar -czf test.tar.gz model.pth code/inference.py
```

When your endpoint is provisioned, the files in the archive are extracted to `/opt/ml/model/` on the endpoint.

After you compress your model and model artifacts as a `.tar.gz` file, upload them to your Amazon S3 bucket. The following example demonstrates how to upload your model to Amazon S3 using the AWS CLI:

```bash
!aws s3 cp test.tar.gz s3://{your-bucket}/models/
```

5. **Select a prebuilt Docker inference image or create your own Inference Docker Image.**

SageMaker provides containers for its built-in algorithms and prebuilt Docker images for some of the most common machine learning frameworks, such as Apache MXNet, TensorFlow, PyTorch, and Chainer. For a full list of the available SageMaker images, see [Available Deep Learning Containers Images](#).

If none of the existing SageMaker containers meet your needs and you don't have an existing container of your own, create a new Docker image. See [Use your own inference code](#) for information about how to create your Docker image. 
The following demonstrates how to retrieve a PyTorch version 1.7.1 inference image using the SageMaker Python SDK:

```python
from sagemaker import image_uris

## Uncomment and replace with your own values if you did not define these variables a previous step.
# framework = 'pytorch'
# framework_version = '1.7.1'

# Note: you can use any CPU-based instance here,
# this is just to set the arch as CPU for the Docker image
instance_type = 'ml.m5.2xlarge'

image_uri = image_uris.retrieve(framework,
                               region,
                               version=framework_version,
                               py_version='py3',
                               instance_type=instance_type,
                               image_scope='inference')
```

For a list of available SageMaker Instances, see Amazon SageMaker Pricing.

6. Create a sample payload archive.

Create an archive that contains individual files that the load testing tool can send to your SageMaker endpoints. Your inference code must be able to read the file formats from the sample payload.

The following downloads a .jpg image that this example uses in a later step for the ResNet-18 model.

```bash
!wget https://cdn.pixabay.com/photo/2020/12/18/05/56/flowers-5841251_1280.jpg
```

Compress the sample payload as a tarball:

```bash
!tar -cvzf payload.tar.gz flowers-5841251_1280.jpg
```

Upload the sample payload to Amazon S3 and note the Amazon S3 URI:
You need the Amazon S3 URI in a later step, so store it in a variable:

```python
bucket_prefix='models'
bucket = '<your-bucket-name>'  # Provide the name of your S3 bucket
payload_s3_key = f'{bucket_prefix}/payload.tar.gz'
sample_payload_url= f's3://{bucket}/{payload_s3_key}'
```

7. **Prepare your model input for the recommendations job**

For the last prerequisite, you have two options to prepare your model input. You can either register your model with SageMaker Model Registry, which you can use to catalog models for production, or you can create a SageMaker model and specify it in the ContainerConfig field when creating a recommendations job. The first option is best if you want to take advantage of the features that Model Registry provides, such as managing model versions and automating model deployment. The second option is ideal if you want to get started quickly. For the first option, go to step 7. For the second option, skip step 7 and go to step 8.

8. **Option 1: Register your model in the model registry**

With SageMaker Model Registry, you can catalog models for production, manage model versions, associate metadata (such as training metrics) with a model, manage the approval status of a model, deploy models to production, and automate model deployment with CI/CD.

When you use SageMaker Model Registry to track and manage your models, they are represented as a versioned model package within model package groups. Unversioned model packages are not part of a model group. Model package groups hold multiple versions or iterations of a model. Though it is not required to create them for every model in the registry, they help organize various models that all have the same purpose and provide automatic versioning.

To use Amazon SageMaker Inference Recommender, you must have a versioned model package. You can create a versioned model package programmatically with the AWS SDK for Python (Boto3) or with Amazon SageMaker Studio Classic. To create a versioned model package programmatically, first create a model package group with the `CreateModelPackageGroup` API. Next, create a model package using the `CreateModelPackage` API. Calling this method makes a versioned model package.
See Create a Model Group and Register a Model Version for detailed instructions about how to programmatically and interactively create a model package group and how to create a versioned model package, respectively, with the AWS SDK for Python (Boto3) and Amazon SageMaker Studio Classic.

The following code sample demonstrates how to create a versioned model package using the AWS SDK for Python (Boto3).

### Note
You do not need to approve the model package to create an Inference Recommender job.

#### a. Create a model package group

Create a model package group with the CreateModelPackageGroup API. Provide a name to the model package group for the ModelPackageGroupName and optionally provide a description of the model package in the ModelPackageGroupDescription field.

```python
model_package_group_name = '<INSERT>'
model_package_group_description = '<INSERT>'

model_package_group_input_dict = {
    'ModelPackageGroupName' : model_package_group_name,
    'ModelPackageGroupDescription' : model_package_group_description,
}

model_package_group_response =
sagemaker_client.create_model_package_group(**model_package_group_input_dict)
```

See the Amazon SageMaker API Reference Guide for a full list of optional and required arguments you can pass to CreateModelPackageGroup.

Create a model package by specifying a Docker image that runs your inference code and the Amazon S3 location of your model artifacts and provide values for InferenceSpecification. InferenceSpecification should contain information...
about inference jobs that can be run with models based on this model package, including the following:

- The Amazon ECR paths of images that run your inference code.
- (Optional) The instance types that the model package supports for transform jobs and real-time endpoints used for inference.
- The input and output content formats that the model package supports for inference.

In addition, you must specify the following parameters when you create a model package:

- **Domain**: The machine learning domain of your model package and its components. Common machine learning domains include computer vision and natural language processing.
- **Task**: The machine learning task your model package accomplishes. Common machine learning tasks include object detection and image classification. Specify "OTHER" if none of the tasks listed in the [API Reference Guide](#) satisfy your use case. See the **Task** API field descriptions for a list of supported machine learning tasks.
- **SamplePayloadUrl**: The Amazon Simple Storage Service (Amazon S3) path where the sample payload are stored. This path must point to a single GZIP compressed TAR archive (.tar.gz suffix).
- **Framework**: The machine learning framework of the model package container image.
- **FrameworkVersion**: The framework version of the model package container image.

If you provide an allow list of instance types to use to generate inferences in real-time for the [SupportedRealtimeInferenceInstanceTypes](#), Inference Recommender limits the search space for instance types during a Default job. Use this parameter if you have budget constraints or know there's a specific set of instance types that can support your model and container image.

In a previous step, we downloaded a pre-trained ResNet18 model and stored it in an Amazon S3 bucket in a directory called `models`. We retrieved a PyTorch (v1.7.1) Deep Learning Container inference image and stored the URI in a variable called `image_uri`. Use those variables in the following code sample to define a dictionary used as input to the [CreateModelPackage](#) API.

```python
# Provide the Amazon S3 URI of your compressed tarfile
```
# so that Model Registry knows where to find your model artifacts
bucket_prefix='models'
bucket = '<your-bucket-name>'  # Provide the name of your S3 bucket
model_s3_key = f"{bucket_prefix}/test.tar.gz"
model_url= f"s3://{bucket}/{model_s3_key}"

# Similar open source model to the packaged model
# The name of the ML model as standardized by common model zoos
nearest_model_name = 'resnet18'

# The supported MIME types for input and output data. In this example,
# we are using images as input.
input_content_type='image/jpeg'

# Optional - provide a description of your model.
model_package_description = '<INSERT>'

## Uncomment if you did not store the domain and task in an earlier
## step
#ml_domain = 'COMPUTER_VISION'
#ml_task = 'IMAGE_CLASSIFICATION'

## Uncomment if you did not store the framework and framework version
## in a previous step.
#framework = 'PYTORCH'
#framework_version = '1.7.1'

# Optional: Used for optimizing your model using SageMaker Neo
# PyTorch uses NCHW format for images
data_input_configuration = "[[1,3,256,256]]"

# Create a dictionary to use as input for creating a model package group
model_package_input_dict = {
    "ModelPackageName": model_package_group_name,
    "ModelPackageDescription": model_package_description,
    "Domain": ml_domain,
    "Task": ml_task,
    "SamplePayloadUrl": sample_payload_url,
    "InferenceSpecification": {
        "Containers": [  
            {
                "Image": image_uri,
                "ModelDataUrl": model_url,
            }
        ]
    }
}
b. **Create a model package**

Use the CreateModelPackage API to create a model package. Pass the input dictionary defined in the previous step:

```python
model_package_response = sagemaker_client.create_model_package(**model_package_input_dict)
```

You need the model package ARN to use Amazon SageMaker Inference Recommender. Note the ARN of the model package or store it in a variable:

```python
model_package_arn = model_package_response['ModelPackageArn']
print('ModelPackage Version ARN : {}'.format(model_package_arn))
```

9. **Option 2: Create a model and configure the ContainerConfig field**

Use this option if you want to start an inference recommendations job and don't need to register your model in the Model Registry. In the following steps, you create a model in SageMaker and configure the ContainerConfig field as input for the recommendations job.

a. **Create a model**

Create a model with the CreateModel API. For an example that calls this method when deploying a model to SageMaker Hosting, see [Create a Model (AWS SDK for Python (Boto3))](https://docs.aws.amazon.com/boto3/guide/quickstart-create-model.html).

In a previous step, we downloaded a pre-trained ResNet18 model and stored it in an Amazon S3 bucket in a directory called `models`. We retrieved a PyTorch (v1.7.1) Deep Learning Container inference image and stored the URI in a variable called `image_uri`. 

```python
"Framework": framework.upper(),
"FrameworkVersion": framework_version,
"NearestModelName": nearest_model_name,
"ModelInput": {"DataInputConfig":
    data_input_configuration}
   ],
   "SupportedContentTypes": [input_content_type]}
}

```
We use those variables in the following code example where we define a dictionary used as input to the `CreateModel` API.

```python
model_name = '<name_of_the_model>'
# Role to give SageMaker permission to access AWS services.
sagemaker_role = 'arn:aws:iam::<region>:<account>:role/*'

# Provide the Amazon S3 URI of your compressed tarfile
# so that Model Registry knows where to find your model artifacts
bucket_prefix = 'models'
bucket = '<your-bucket-name>'
model_s3_key = f'{bucket_prefix}/test.tar.gz'
model_url = f's3://{bucket}/{model_s3_key}'

# Create model
create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    PrimaryContainer = {
        'Image': image_uri,
        'ModelDataUrl': model_url,
    })
```

b. **Configure the ContainerConfig field**

Next, you must configure the `ContainerConfig` field with the model you just created and specify the following parameters in it:

- **Domain**: The machine learning domain of the model and its components, such as computer vision or natural language processing.
- **Task**: The machine learning task that the model accomplishes, such as image classification or object detection.
- **PayloadConfig**: The configuration for the payload for a recommendation job. For more information about the subfields, see `RecommendationJobPayloadConfig`.
- **Framework**: The machine learning framework of the container image, such as PyTorch.
- **FrameworkVersion**: The framework version of the container image.
- **(Optional) SupportedInstanceTypes**: A list of the instance types that are used to generate inferences in real-time.
If you use the `SupportedInstanceTypes` parameter, Inference Recommender limits the search space for instance types during a Default job. Use this parameter if you have budget constraints or know there's a specific set of instance types that can support your model and container image.

In the following code example, we use the previously defined parameters, along with `NearestModelName`, to define a dictionary used as input to the `CreateInferenceRecommendationsJob` API.

```python
## Uncomment if you did not store the domain and task in a previous step
# ml_domain = 'COMPUTER_VISION'
# ml_task = 'IMAGE_CLASSIFICATION'

## Uncomment if you did not store the framework and framework version in a previous step
# framework = 'PYTORCH'
# framework_version = '1.7.1'

# The name of the ML model as standardized by common model zoos
nearest_model_name = 'resnet18'

# The supported MIME types for input and output data. In this example, # we are using images as input
input_content_type='image/jpeg'

# Optional: Used for optimizing your model using SageMaker Neo
# PyTorch uses NCHW format for images
data_input_configuration = "[[1,3,256,256]]"

# Create a dictionary to use as input for creating an inference recommendation job
container_config = {
    "Domain": ml_domain,
    "Framework": framework.upper(),
    "FrameworkVersion": framework_version,
    "NearestModelName": nearest_model_name,
    "PayloadConfig": {  
        "SamplePayloadUrl": sample_payload_url,
        "SupportedContentTypes": [ input_content_type ]  
    },
    "DataInputConfig": data_input_configuration
}
Recommendation jobs

Amazon SageMaker Inference Recommender can make two types of recommendations:

1. Inference recommendations (Default job type) run a set of load tests on the recommended instance types. You can also load test for a serverless endpoint. You only need to provide a model package Amazon Resource Name (ARN) to launch this type of recommendation job. Inference recommendation jobs complete within 45 minutes.

2. Endpoint recommendations (Advanced job type) are based on a custom load test where you select your desired ML instances or a serverless endpoint, provide a custom traffic pattern, and provide requirements for latency and throughput based on your production requirements. This job takes an average of 2 hours to complete depending on the job duration set and the total number of inference configurations tested.

Both types of recommendations use the same APIs to create, describe, and stop jobs. The output is a list of instance configuration recommendations with associated environment variables, cost, throughput, and latency metrics. Recommendation jobs also provide an initial instance count, which you can use to configure an autoscaling policy. To differentiate between the two types of jobs, when you’re creating a job through either the SageMaker console or the APIs, specify Default to create preliminary endpoint recommendations and Advanced for custom load testing and endpoint recommendations.

Note
You do not need to do both types of recommendation jobs in your own workflow. You can do either independently of the other.

Inference Recommender can also provide you with a list of prospective instances, or the top five instance types that are optimized for cost, throughput and latency for model deployment, along with a confidence score. You can choose these instances when deploying your model. Inference Recommender automatically performs benchmarking against your model for you to provide the prospective instances. Since these are preliminary recommendations, we recommend that you run further instance recommendation jobs to get more accurate results. To view the prospective
instances, go to your SageMaker model details page. For more information, see Get instant prospective instances.

Topics

- Get instant prospective instances
- Get an inference recommendation
- Get an inference recommendation for an existing endpoint
- Get compiled recommendations with Neo
- Interpret recommendation results
- Get autoscaling policy recommendations
- Run a custom load test
- Troubleshoot Inference Recommender errors

Get instant prospective instances

Inference Recommender can also provide you with a list of prospective instances, or instance types that might be suitable for your model, on your SageMaker model details page. Inference Recommender automatically performs preliminary benchmarking against your model for you to provide the top five prospective instances. Since these are preliminary recommendations, we recommend that you run further instance recommendation jobs to get more accurate results.

You can view a list of prospective instances for your model either programmatically by using the DescribeModel API, the SageMaker Python SDK, or the SageMaker console.

Note

You won't get prospective instances for models that you created in SageMaker before this feature became available.

To view the prospective instances for your model through the console, do the following:

2. In the left navigation pane, choose Inference, and then choose Models.
3. From the list of models, choose your model.
On the details page for your model, go to the **Prospective instances to deploy model** section. The following screenshot shows this section.

![Prospective instances to deploy model](image)

In this section, you can view the prospective instances that are optimized for cost, throughput, and latency for model deployment, along with additional information for each instance type such as the memory size, CPU and GPU count, and cost per hour.

If you decide that you want to benchmark a sample payload and run a full inference recommendation job for your model, you can start a default inference recommendation job from this page. To start a default job through the console, do the following:

1. On your model details page in the **Prospective instances to deploy model section**, choose **Run Inference recommender job**.
2. In the dialog box that pops up, for **S3 bucket for benchmarking payload**, enter the Amazon S3 location where you’ve stored a sample payload for your model.
3. For **Payload content type**, enter the MIME types for your payload data.
4. (Optional) In the **Model compilation using SageMaker Neo** section, for the **Data input configuration**, enter a data shape in dictionary format.
5. Choose **Run job**.

Inference Recommender starts the job, and you can view the job and its results from the **Inference recommender** list page in the SageMaker console.

If you want to run an advanced job and perform custom load tests, or if you want to configure additional settings and parameters for your job, see **Run a custom load test**.
Get an inference recommendation

Inference recommendation jobs run a set of load tests on recommended instance types or a serverless endpoint. Inference recommendation jobs use performance metrics that are based on load tests using the sample data you provided during model version registration.

ℹ️ Note
Before you create an Inference Recommender recommendation job, make sure you have satisfied the Prerequisites.

The following demonstrates how to use Amazon SageMaker Inference Recommender to create an inference recommendation based on your model type using the AWS SDK for Python (Boto3), AWS CLI, and Amazon SageMaker Studio Classic, and the SageMaker console.

Create an inference recommendation

Create an inference recommendation programmatically using the AWS SDK for Python (Boto3) or the AWS CLI, or interactively using Studio Classic or the SageMaker console. Specify a job name for your inference recommendation, an AWS IAM role ARN, an input configuration, and either a model package ARN when you registered your model with the model registry, or your model name and a ContainerConfig dictionary from when you created your model in the Prerequisites section.

AWS SDK for Python (Boto3)

Use the CreateInferenceRecommendationsJob API to start an inference recommendation job. Set the JobType field to 'Default' for inference recommendation jobs. In addition, provide the following:

- The Amazon Resource Name (ARN) of an IAM role that enables Inference Recommender to perform tasks on your behalf. Define this for the RoleArn field.
- A model package ARN or model name. Inference Recommender supports either one model package ARN or a model name as input. Specify one of the following:
  - The ARN of the versioned model package you created when you registered your model with SageMaker model registry. Define this for ModelPackageVersionArn in the InputConfig field.
  - The name of the model you created. Define this for ModelName in the InputConfig field. Also, provide the ContainerConfig dictionary, which includes the required fields.
that need to be provided with the model name. Define this for ContainerConfig in the InputConfig field. In the ContainerConfig, you can also optionally specify the SupportedEndpointType field as either RealTime or Serverless. If you specify this field, Inference Recommender returns recommendations for only that endpoint type. If you don't specify this field, Inference Recommender returns recommendations for both endpoint types.

- A name for your Inference Recommender recommendation job for the JobName field. The Inference Recommender job name must be unique within the AWS Region and within your AWS account.

Import the AWS SDK for Python (Boto3) package and create a SageMaker client object using the client class. If you followed the steps in the Prerequisites section, only specify one of the following:

- Option 1: If you would like to create an inference recommendations job with a model package ARN, then store the model package group ARN in a variable named model_package_arn.
- Option 2: If you would like to create an inference recommendations job with a model name and ContainerConfig, store the model name in a variable named model_name and the ContainerConfig dictionary in a variable named container_config.

```python
# Create a low-level SageMaker service client.
import boto3
aws_region = '<INSERT>'
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Provide only one of model package ARN or model name, not both.
# Provide your model package ARN that was created when you registered your
# model with Model Registry
model_package_arn = '<INSERT>'
## Uncomment if you would like to create an inference recommendations job with a
## model name instead of a model package ARN, and comment out model_package_arn
## above
## Provide your model name
# model_name = '<INSERT>'
## Provide your container config
# container_config = '<INSERT>'

# Provide a unique job name for SageMaker Inference Recommender job
```
job_name = '<INSERT>'

# Inference Recommender job type. Set to Default to get an initial recommendation
job_type = 'Default'

# Provide an IAM Role that gives SageMaker Inference Recommender permission to
# access AWS services
role_arn = 'arn:aws:iam::<account>:role/*'

sagemaker_client.create_inference_recommendations_job(
    JobName = job_name,
    JobType = job_type,
    RoleArn = role_arn,
    # Provide only one of model package ARN or model name, not both.
    # If you would like to create an inference recommendations job with a model
    # name,
    # uncomment ModelName and ContainerConfig, and comment out
    ModelPackageVersionArn.
    InputConfig = {
        'ModelPackageVersionArn': model_package_arn
        # 'ModelName': model_name,
        # 'ContainerConfig': container_config
    }
)

See the Amazon SageMaker API Reference Guide for a full list of optional and required arguments you can pass to CreateInferenceRecommendationsJob.

AWS CLI

Use the create-inference-recommendations-job API to start an inference recommendation job. Set the job-type field to 'Default' for inference recommendation jobs. In addition, provide the following:

- The Amazon Resource Name (ARN) of an IAM role that enables Amazon SageMaker Inference Recommender to perform tasks on your behalf. Define this for the role-arn field.
- A model package ARN or model name. Inference Recommender supports either one model package ARN or a model name as input. Specify one of the following
  - The ARN of the versioned model package you created when you registered your model with Model Registry. Define this for ModelPackageVersionArn in the input-config field.
• The name of the model you created. Define this for `ModelName` in the `input-config` field. Also, provide the `ContainerConfig` dictionary which includes the required fields that need to be provided with the model name. Define this for `ContainerConfig` in the `input-config` field. In the `ContainerConfig`, you can also optionally specify the `SupportedEndpointType` field as either `RealTime` or `Serverless`. If you specify this field, Inference Recommender returns recommendations for only that endpoint type. If you don't specify this field, Inference Recommender returns recommendations for both endpoint types.

• A name for your Inference Recommender recommendation job for the `job-name` field. The Inference Recommender job name must be unique within the AWS Region and within your AWS account.

To create an inference recommendation jobs with a model package ARN, use the following example:

```bash
aws sagemaker create-inference-recommendations-job
    --region <region>
    --job-name <job_name>
    --job-type Default
    --role-arn arn:aws:iam::<account:role/*/>
    --input-config "{"ModelPackageVersionArn": "arn:aws:sagemaker::<region:account:role/*/",}
                     "{"
                     "}"
```

To create an inference recommendation jobs with a model name and `ContainerConfig`, use the following example. The example uses the `SupportedEndpointType` field to specify that we only want to return real-time inference recommendations:

```bash
aws sagemaker create-inference-recommendations-job
    --region <region>
    --job-name <job_name>
    --job-type Default
    --role-arn arn:aws:iam::<account:role/*/>
    --input-config "{"ModelName": "model-name",
                     "ContainerConfig": {
                     "Domain": "COMPUTER_VISION",
                     "Framework": "PYTORCH",
                     "FrameworkVersion": "1.7.1",
```
```json
"NearestModelName": "resnet18",
"PayloadConfig": {
    "SamplePayloadUrl": "s3://{bucket}/{payload_s3_key}",
    "SupportedContentTypes": ["image/jpeg"]
},
"SupportedEndpointType": "RealTime",
"DataInputConfig": "[[1,3,256,256]]",
"Task": "IMAGE_CLASSIFICATION",
}
```

Amazon SageMaker Studio Classic

Create an inference recommendation job in Studio Classic.

1. In your Studio Classic application, choose the home icon.
2. In the left sidebar of Studio Classic, choose **Models**.
3. Choose **Model Registry** from the dropdown list to display models you have registered with the model registry.

   The left panel displays a list of model groups. The list includes all the model groups registered with the model registry in your account, including models registered outside of Studio Classic.

4. Select the name of your model group. When you select your model group, the right pane of Studio Classic displays column heads such as **Versions** and **Setting**.

   If you have one or more model packages within your model group, you see a list of those model packages within the **Versions** column.

5. Choose the **Inference recommender** column.
6. Choose an IAM role that grants Inference Recommender permission to access AWS services. You can create a role and attach the AmazonSageMakerFullAccess IAM managed policy to accomplish this. Or you can let Studio Classic create a role for you.
7. Choose **Get recommendations**.

   The inference recommendation can take up to 45 minutes.
Warning

Do not close this tab. If you close this tab, you cancel the instance recommendation job.

SageMaker console

Create an instance recommendation job through the SageMaker console by doing the following:


2. In the left navigation pane, choose Inference, and then choose Inference recommender.

3. On the Inference recommender jobs page, choose Create job.

4. For Step 1: Model configuration, do the following:
   a. For Job type, choose Default recommender job.
   b. If you're using a model registered in the SageMaker model registry, then turn on the Choose a model from the model registry toggle and do the following:
      i. From the Model group dropdown list, choose the model group in SageMaker model registry where your model is located.
      ii. From the Model version dropdown list, choose the desired version of your model.
   c. If you’re using a model that you’ve created in SageMaker, then turn off the Choose a model from the model registry toggle and do the following:
      • For the Model name field, enter the name of your SageMaker model.
   d. From the IAM role dropdown list, you can select an existing AWS IAM role that has the necessary permissions to create an instance recommendation job. Alternatively, if you don’t have an existing role, you can choose Create a new role to open the role creation pop-up, and SageMaker adds the necessary permissions to the new role that you create.
   e. For S3 bucket for benchmarking payload, enter the Amazon S3 path to your sample payload archive, which should contain sample payload files that Inference Recommender uses to benchmark your model on different instance types.
   f. For Payload content type, enter the MIME types of your sample payload data.
g. **(Optional)** If you turned off the **Choose a model from the model registry toggle** and specified a SageMaker model, then for **Container configuration**, do the following:

i. For the **Domain** dropdown list, select the machine learning domain of the model, such as computer vision, natural language processing, or machine learning.

ii. For the **Framework** dropdown list, select the framework of your container, such as TensorFlow or XGBoost.

iii. For **Framework version**, enter the framework version of your container image.

iv. For the **Nearest model name** dropdown list, select the pre-trained model that mostly closely matches your own.

v. For the **Task** dropdown list, select the machine learning task that the model accomplishes, such as image classification or regression.

h. **(Optional)** For **Model compilation using SageMaker Neo**, you can configure the recommendation job for a model that you've compiled using SageMaker Neo. For **Data input configuration**, enter the correct input data shape for your model in a format similar to `{ 'input': [1, 1024, 1024, 3] }.

   i. Choose **Next**.

5. For **Step 2: Instances and environment parameters**, do the following:

   a. **(Optional)** For **Select instances for benchmarking**, you can select up to 8 instance types that you want to benchmark. If you don’t select any instances, Inference Recommender considers all instance types.

   b. Choose **Next**.

6. For **Step 3: Job parameters**, do the following:

   a. **(Optional)** For the **Job name** field, enter a name for your instance recommendation job. When you create the job, SageMaker appends a timestamp to the end of this name.

   b. **(Optional)** For the **Job description** field, enter a description for the job.

   c. **(Optional)** For the **Encryption key** dropdown list, choose an AWS KMS key by name or enter its ARN to encrypt your data.

   d. **(Optional)** For **Max test duration (s)**, enter the maximum number of seconds you want each test to run for.
e. (Optional) For **Max invocations per minute**, enter the maximum number of requests per minute the endpoint can reach before stopping the recommendation job. After reaching this limit, SageMaker ends the job.

f. (Optional) For **P99 Model latency threshold (ms)**, enter the model latency percentile in milliseconds.

g. Choose **Next**.

7. For **Step 4: Review job**, review your configurations and then choose **Submit**.

### Get your inference recommendation job results

Collect the results of your inference recommendation job programmatically with AWS SDK for Python (Boto3), the AWS CLI, Studio Classic, or the SageMaker console.

**AWS SDK for Python (Boto3)**

Once an inference recommendation is complete, you can use `DescribeInferenceRecommendationsJob` to get the job details and recommendations. Provide the job name that you used when you created the inference recommendation job.

```python
job_name='<INSERT>'
response = sagemaker_client.describe_inference_recommendations_job(
    JobName=job_name)
```

Print the response object. The previous code sample stored the response in a variable named `response`.

```python
print(response['Status'])
```

This returns a JSON response similar to the following example. Note that this example shows the recommended instance types for real-time inference (for an example showing serverless inference recommendations, see the example after this one).

```json
{
    'JobName': 'job-name',
    'JobDescription': 'job-description',
    'JobType': 'Default',
    'Status': 'COMPLETED',
}
```
Recommendation jobs 4012
The first few lines provide information about the inference recommendation job itself. This includes the job name, role ARN, and creation and deletion times.

The InferenceRecommendations dictionary contains a list of Inference Recommender inference recommendations.

The EndpointConfiguration nested dictionary contains the instance type (InstanceType) recommendation along with the endpoint and variant name (a deployed AWS machine learning model) that was used during the recommendation job. You can use the endpoint and variant name for monitoring in Amazon CloudWatch Events. See Monitor Amazon SageMaker with Amazon CloudWatch for more information.
The Metrics nested dictionary contains information about the estimated cost per hour (CostPerHour) for your real-time endpoint in US dollars, the estimated cost per inference (CostPerInference) in US dollars for your real-time endpoint, the expected maximum number of InvokeEndpoint requests per minute sent to the endpoint (MaxInvocations), and the model latency (ModelLatency), which is the interval of time (in microseconds) that your model took to respond to SageMaker. The model latency includes the local communication times taken to send the request and to fetch the response from the container of a model and the time taken to complete the inference in the container.

The following example shows the InferenceRecommendations part of the response for an inference recommendations job configured to return serverless inference recommendations:

```json
"InferenceRecommendations": [ {
   "EndpointConfiguration": {  "EndpointName": "value",  "InitialInstanceCount": "value",  "InstanceType": "value",  "VariantName": "value",  "ServerlessConfig": {  "MaxConcurrency": "value",  "MemorySizeInMb": "value"  }
},
   "InvocationEndTime": "value",
   "InvocationStartTime": "value",
   "Metrics": {  "CostPerHour": "value",  "CostPerInference": "value",  "CpuUtilization": "value",  "MaxInvocations": "value",  "MemoryUtilization": "value",  "ModelLatency": "value",  "ModelSetupTime": "value"
   },
   "ModelConfiguration": {  "Compiled": "False",  "EnvironmentParameters": [],  "InferenceSpecificationName": "value"
   },
   "RecommendationId": "value"
}]
```
You can interpret the recommendations for serverless inference similarly to the results for real-time inference, with the exception of the ServerlessConfig, which tells you the metrics returned for a serverless endpoint with the given MemorySizeInMB and when MaxConcurrency = 1. To increase the throughput possible on the endpoint, increase the value of MaxConcurrency linearly. For example, if the inference recommendation shows MaxInvocations as 1000, then increasing MaxConcurrency to 2 would support 2000 MaxInvocations. Note that this is true only up to a certain point, which can vary based on your model and code. Serverless recommendations also measure the metric ModelSetupTime, which measures (in microseconds) the time it takes to launch computer resources on a serverless endpoint. For more information about setting up serverless endpoints, see the Serverless Inference documentation.

AWS CLI

Once an inference recommendation is complete, you can use describe-inference-recommendations-job to get the job details and recommended instance types. Provide the job name that you used when you created the inference recommendation job.

```
aws sagemaker describe-inference-recommendations-job
  --job-name <job-name>
  --region <aws-region>
```

The JSON response similar should resemble the following example. Note that this example shows the recommended instance types for real-time inference (for an example showing serverless inference recommendations, see the example after this one).

```json
{
  'JobName': 'job-name',
  'JobDescription': 'job-description',
  'JobType': 'Default',
  'Status': 'COMPLETED',
  'CreationTime': datetime.datetime(2021, 10, 26, 20, 4, 57, 627000, tzinfo=tzlocal()),
  'LastModifiedTime': datetime.datetime(2021, 10, 26, 20, 25, 1, 997000, tzinfo=tzlocal()),
  'InputConfig': {
```
  'JobDurationInSeconds': 0
],
'InferenceRecommendations': [{
  'Metrics': {
    'CostPerHour': 0.20399999618530273,
    'CostPerInference': 5.246913588052848e-06,
    'MaximumInvocations': 648,
    'ModelLatency': 263596
  },
  'EndpointConfiguration': {
    'EndpointName': 'endpoint-name',
    'VariantName': 'variant-name',
    'InstanceType': 'ml.c5.xlarge',
    'InitialInstanceCount': 1
  },
  'ModelConfiguration': {
    'Compiled': False,
    'EnvironmentParameters': []
  }
},
{
  'Metrics': {
    'CostPerHour': 0.11500000208616257,
    'CostPerInference': 2.92620870823157e-06,
    'MaximumInvocations': 655,
    'ModelLatency': 826019
  },
  'EndpointConfiguration': {
    'EndpointName': 'endpoint-name',
    'VariantName': 'variant-name',
    'InstanceType': 'ml.c5d.large',
    'InitialInstanceCount': 1
  },
  'ModelConfiguration': {
    'Compiled': False,
    'EnvironmentParameters': []
  }
},
{
  'Metrics': {
    'CostPerHour': 0.11500000208616257,
    'CostPerInference': 3.3625731248321244e-06,
    'MaximumInvocations': 655,
    'ModelLatency': 826019
  },
  'EndpointConfiguration': {
    'EndpointName': 'endpoint-name',
    'VariantName': 'variant-name',
    'InstanceType': 'ml.c5d.large',
    'InitialInstanceCount': 1
  },
  'ModelConfiguration': {
    'Compiled': False,
    'EnvironmentParameters': []
  }
},
{
  'Metrics': {
    'CostPerHour': 0.11500000208616257,
    'CostPerInference': 3.3625731248321244e-06,
    'MaximumInvocations': 655,
    'ModelLatency': 826019
  },
  'EndpointConfiguration': {
    'EndpointName': 'endpoint-name',
    'VariantName': 'variant-name',
    'InstanceType': 'ml.c5d.large',
    'InitialInstanceCount': 1
  },
  'ModelConfiguration': {
    'Compiled': False,
    'EnvironmentParameters': []
  }
},
{
  'Metrics': {
    'CostPerHour': 0.11500000208616257,
    'CostPerInference': 3.3625731248321244e-06,
    'MaximumInvocations': 655,
    'ModelLatency': 826019
  },
  'EndpointConfiguration': {
    'EndpointName': 'endpoint-name',
    'VariantName': 'variant-name',
    'InstanceType': 'ml.c5d.large',
    'InitialInstanceCount': 1
  },
  'ModelConfiguration': {
    'Compiled': False,
    'EnvironmentParameters': []
  }
}]}
The first few lines provide information about the inference recommendation job itself. This includes the job name, role ARN, creation, and deletion time.

The InferenceRecommendations dictionary contains a list of Inference Recommender inference recommendations.

The EndpointConfiguration nested dictionary contains the instance type (InstanceType) recommendation along with the endpoint and variant name (a deployed AWS machine learning model) used during the recommendation job. You can use the endpoint and variant name for monitoring in Amazon CloudWatch Events. See Monitor Amazon SageMaker with Amazon CloudWatch for more information.

The Metrics nested dictionary contains information about the estimated cost per hour (CostPerHour) for your real-time endpoint in US dollars, the estimated cost per inference (CostPerInference) in US dollars for your real-time endpoint, the expected maximum
number of InvokeEndpoint requests per minute sent to the endpoint (MaxInvocations), and the model latency (ModelLatency), which is the interval of time (in milliseconds) that your model took to respond to SageMaker. The model latency includes the local communication times taken to send the request and to fetch the response from the container of a model and the time taken to complete the inference in the container.

The following example shows the InferenceRecommendations part of the response for an inference recommendations job configured to return serverless inference recommendations:

```json
"InferenceRecommendations": [
  {
    "EndpointConfiguration": {
      "EndpointName": "value",
      "InitialInstanceCount": "value",
      "InstanceType": "value",
      "VariantName": "value",
      "ServerlessConfig": {
        "MaxConcurrency": "value",
        "MemorySizeInMb": "value"
      }
    },
    "InvocationEndTime": "value",
    "InvocationStartTime": "value",
    "Metrics": {
      "CostPerHour": "value",
      "CostPerInference": "value",
      "CpuUtilization": "value",
      "MaxInvocations": "value",
      "MemoryUtilization": "value",
      "ModelLatency": "value",
      "ModelSetupTime": "value"
    },
    "ModelConfiguration": {
      "Compiled": "False",
      "EnvironmentParameters": [],
      "InferenceSpecificationName": "value"
    },
    "RecommendationId": "value"
  }
]
```
You can interpret the recommendations for serverless inference similarly to the results for real-time inference, with the exception of the ServerlessConfig, which tells you the metrics returned for a serverless endpoint with the given MemorySizeInMB and when MaxConcurrency = 1. To increase the throughput possible on the endpoint, increase the value of MaxConcurrency linearly. For example, if the inference recommendation shows MaxInvocations as 1000, then increasing MaxConcurrency to 2 would support 2000 MaxInvocations. Note that this is true only up to a certain point, which can vary based on your model and code. Serverless recommendations also measure the metric ModelSetupTime, which measures (in microseconds) the time it takes to launch computer resources on a serverless endpoint. For more information about setting up serverless endpoints, see the Serverless Inference documentation.

Amazon SageMaker Studio Classic

The inference recommendations populate in a new Inference recommendations tab within Studio Classic. It can take up to 45 minutes for the results to show up. This tab contains Results and Details column headings.

The Details column provides information about the inference recommendation job, such as the name of the inference recommendation, when the job was created (Creation time), and more. It also provides Settings information, such as the maximum number of invocations that occurred per minute and information about the Amazon Resource Names used.

The Results column provides a Deployment goals and SageMaker recommendations window in which you can adjust the order that the results are displayed based on deployment importance. There are three dropdown menus that you can use to provide the level of importance of the Cost, Latency, and Throughput for your use case. For each goal (cost, latency, and throughput), you can set the level of importance: Lowest Importance, Low Importance, Moderate importance, High importance, or Highest importance.

Based on your selections of importance for each goal, Inference Recommender displays its top recommendation in the SageMaker recommendation field on the right of the panel, along with the estimated cost per hour and inference request. It also provides information about the expected model latency, maximum number of invocations, and the number of instances. For serverless recommendations, you can see the ideal values for the maximum concurrency and endpoint memory size.

In addition to the top recommendation displayed, you can also see the same information displayed for all instances that Inference Recommender tested in the All runs section.
SageMaker console

You can view your instance recommendation jobs in the SageMaker console by doing the following:

2. In the left navigation pane, choose Inference, and then choose Inference recommender.
3. On the Inference recommender jobs page, choose the name of your inference recommendation job.

On the details page for your job, you can view the Inference recommendations, which are the instance types SageMaker recommends for your model, as shown in the following screenshot.

<table>
<thead>
<tr>
<th>Instance</th>
<th>Status</th>
<th>Model latency</th>
<th>Cost per hour</th>
<th>Cost per inference</th>
<th>Invocations per minute</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.in1.xlarge</td>
<td>In progress</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ml.m5.8xlarge</td>
<td>Success</td>
<td>11ms</td>
<td>$12.12</td>
<td>$12.12</td>
<td>14</td>
</tr>
<tr>
<td>ml.g4dn.8xlarge</td>
<td>Success</td>
<td>12ms</td>
<td>$12.12</td>
<td>$12.12</td>
<td>21</td>
</tr>
<tr>
<td>ml.g4dn.xlarge</td>
<td>Error</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

In this section, you can compare the instance types by various factors such as **Model latency**, **Cost per hour**, **Cost per inference**, and **Invocations per minute**.

On this page, you can also view the configurations you specified for your job. In the Monitor section, you can view the Amazon CloudWatch metrics that were logged for each instance type. To learn more about interpreting these metrics, see [Interpret results](#).

For more information about interpreting the results of your recommendation job, see [Interpret recommendation results](#).

**Stop your inference recommendation**

You might want to stop a job that is currently running if you began a job by mistake or no longer need to run the job. Stop your Inference Recommender inference recommendation jobs programmatically with the StopInferenceRecommendationsJob API or with Studio Classic.
**AWS SDK for Python (Boto3)**

Specify the name of the inference recommendation job for the `JobName` field:

```python
sagemaker_client.stop_inference_recommendations_job(
    JobName='<INSERT>'
)
```

**AWS CLI**

Specify the job name of the inference recommendation job for the `job-name` flag:

```bash
aws sagemaker stop-inference-recommendations-job --job-name <job-name>
```

**Amazon SageMaker Studio Classic**

Close the tab in which you initiated the inference recommendation to stop your Inference Recommender inference recommendation.

**SageMaker console**

To stop your instance recommendation job through the SageMaker console, do the following:

2. In the left navigation pane, choose **Inference**, and then choose **Inference recommender**.
3. On the **Inference recommender jobs** page, select your instance recommendation job.
4. Choose **Stop job**.
5. In the dialog box that pops up, choose **Confirm**.

After stopping your job, the job's **Status** should change to **Stopping**.

**Get an inference recommendation for an existing endpoint**

Inference recommendation jobs run a set of load tests on recommended instance types and an existing endpoint. Inference recommendation jobs use performance metrics that are based on load tests using the sample data you provided during model version registration.

You can benchmark and get inference recommendations for an existing SageMaker Inference endpoint to help you improve the performance of your endpoint. The procedure of getting
recommendations for an existing SageMaker Inference endpoint is similar to the procedure for getting inference recommendations without an endpoint. There are several feature exclusions to take note of when benchmarking an existing endpoint:

- You can only use one existing endpoint per Inference Recommender job.
- You can only have one variant on your endpoint.
- You can’t use an endpoint that enables autoscaling.
- This functionality is only supported for Real-Time Inference.
- This functionality doesn't support Real-Time Multi-Model Endpoints.

⚠️ Warning

We strongly recommend that you don't run an Inference Recommender job on a production endpoint that handles live traffic. The synthetic load during benchmarking can affect your production endpoint and cause throttling or provide inaccurate benchmark results. We recommend that you use a non-production or developer endpoint for comparison purposes.

The following sections demonstrate how to use Amazon SageMaker Inference Recommender to create an inference recommendation for an existing endpoint based on your model type using the AWS SDK for Python (Boto3) and the AWS CLI.

⚠️ Note

Before you create an Inference Recommender recommendation job, make sure you have satisfied the Prerequisites.

Prerequisites

If you don't already have a SageMaker Inference endpoint, you can either get an inference recommendation without an endpoint, or you can create a Real-Time Inference endpoint by following the instructions in Create your endpoint and deploy your model.
Create an inference recommendation job for an existing endpoint

Create an inference recommendation programatically using AWS SDK for Python (Boto3), or the AWS CLI. Specify a job name for your inference recommendation, the name of an existing SageMaker Inference endpoint, an AWS IAM role ARN, an input configuration, and your model package ARN from when you registered your model with the model registry.

AWS SDK for Python (Boto3)

Use the CreateInferenceRecommendationsJob API to get an inference recommendation. Set the JobType field to 'Default' for inference recommendation jobs. In addition, provide the following:

- Provide a name for your Inference Recommender recommendation job for the JobName field. The Inference Recommender job name must be unique within the AWS Region and within your AWS account.
- The Amazon Resource Name (ARN) of an IAM role that enables Inference Recommender to perform tasks on your behalf. Define this for the RoleArn field.
- The ARN of the versioned model package you created when you registered your model with the model registry. Define this for ModelPackageVersionArn in the InputConfig field.
- Provide the name of an existing SageMaker Inference endpoint that you want to benchmark in Inference Recommender for Endpoints in the InputConfig field.

Import the AWS SDK for Python (Boto3) package and create a SageMaker client object using the client class. If you followed the steps in the Prerequisites section, the model package group ARN was stored in a variable named model_package_arn.

```python
# Create a low-level SageMaker service client.
import boto3
aws_region = '<region>'
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Provide your model package ARN that was created when you registered your
# model with Model Registry
model_package_arn = '<model-package-arn>'

# Provide a unique job name for SageMaker Inference Recommender job
job_name = '<job-name>'
```
Inference Recommender job type. Set to Default to get an initial recommendation:

```python
job_type = 'Default'
```

Provide an IAM Role that gives SageMaker Inference Recommender permission to access AWS services:

```python
role_arn = '<arn:aws:iam::<account>:role/*>'
```

Provide endpoint name for your endpoint that want to benchmark in Inference Recommender:

```python
endpoint_name = '<existing-endpoint-name>'
```

Use the `create-inference-recommendations-job` API to get an instance endpoint recommendation. Set the `job-type` field to 'Default' for instance endpoint recommendation jobs. In addition, provide the following:

- Provide a name for your Inference Recommender recommendation job for the `job-name` field. The Inference Recommender job name must be unique within the AWS Region and within your AWS account.
- The Amazon Resource Name (ARN) of an IAM role that enables Amazon SageMaker Inference Recommender to perform tasks on your behalf. Define this for the `role-arn` field.
- The ARN of the versioned model package you created when you registered your model with Model Registry. Define this for `ModelPackageVersionArn` in the `input-config` field.
- Provide the name of an existing SageMaker Inference endpoint that you want to benchmark in Inference Recommender for Endpoints in the `input-config` field.

See the [Amazon SageMaker API Reference Guide](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateInferenceRecommendationsJob.html) for a full list of optional and required arguments you can pass to `CreateInferenceRecommendationsJob`.

**AWS CLI**

Use the `create-inference-recommendations-job` API to get an instance endpoint recommendation.
aws sagemaker create-inference-recommendations-job
   --region <region>\n   --job-name <job_name>\n   --job-type Default\n   --role-arn arn:aws:iam::<account:role/*>\n   --input-config "{
      "Endpoints": [{"EndpointName": <endpoint_name>}]}
"

Get your inference recommendation job results

You can collect the results of your inference recommendation job programmatically with the same procedure for standard inference recommendation jobs. For more information, see Get your inference recommendation job results.

When you get inference recommendation job results for an existing endpoint, you should receive a JSON response similar to the following:

```json
{
   "JobName": "job-name",
   "JobType": "Default",
   "RoleArn": "iam-role-arn",
   "Status": "COMPLETED",
   "CreationTime": 1664922919.2,
   "LastModifiedTime": 1664924208.291,
   "InputConfig": {
      "Endpoints": [
         {
            "EndpointName": "endpoint-name"
         }
      ],
   },
   "InferenceRecommendations": [
      {
         "Metrics": {
            "CostPerHour": 0.7360000014305115,
            "CostPerInference": 7.456940238625975e-06,
```
The first few lines provide information about the inference recommendation job itself. This includes the job name, role ARN, and creation and latest modification times.

The `InferenceRecommendations` dictionary contains a list of Inference Recommender inference recommendations.

The `EndpointConfiguration` nested dictionary contains the instance type (InstanceType) recommendation along with the endpoint and variant name (a deployed AWS machine learning model) that was used during the recommendation job.
The Metrics nested dictionary contains information about the estimated cost per hour (CostPerHour) for your real-time endpoint in US dollars, the estimated cost per inference (CostPerInference) in US dollars for your real-time endpoint, the expected maximum number of InvokeEndpoint requests per minute sent to the endpoint (MaxInvocations), and the model latency (ModelLatency), which is the interval of time (in milliseconds) that your model took to respond to SageMaker. The model latency includes the local communication times taken to send the request and to fetch the response from the container of a model and the time taken to complete the inference in the container.

The EndpointPerformances nested dictionary contains the name of your existing endpoint on which the recommendation job was run (EndpointName) and the performance metrics for your endpoint (MaxInvocations and ModelLatency).

**Stop your instance endpoint recommendation**

You might want to stop a job that is currently running if you began a job by mistake or no longer need to run the job. You can stop your Inference Recommender recommendation job programmatically with the same procedure for standard inference recommendation jobs. For more information, see [Stop your inference recommendation](#).

**Get compiled recommendations with Neo**

In Inference Recommender, you can compile your model with Neo and get endpoint recommendations for your compiled model. SageMaker Neo is a service that can optimize your model for a target hardware platform (that is, a specific instance type or environment). Optimizing a model with Neo might improve the performance of your hosted model.

For Neo-supported frameworks and containers, Inference Recommender automatically suggests Neo-optimized recommendations. To be eligible for Neo compilation, your input must meet the following prerequisites:

- You are using a SageMaker owned [DLC](#) or XGBoost container.
- You are using a framework version supported by Neo. For the framework versions supported by Neo, see [Cloud Instances](#) in the SageMaker Neo documentation.
- Neo requires that you provide a correct input data shape for your model. You can specify this data shape as the [DataInputConfig](#) in the [InferenceSpecification](#) when you create a model package. For information about the correct data shapes for each framework, see [Prepare Model for Compilation](#) in the SageMaker Neo documentation.
The following example shows how to specify the `DataInputConfig` field in the `InferenceSpecification`, where `data_input_configuration` is a variable that contains the data shape in dictionary format (for example, `{'input':[1,1024,1024,3]}`).

```json
"InferenceSpecification": {
    "Containers": [
        {
            "Image": dlc_uri,
            "Framework": framework.upper(),
            "FrameworkVersion": framework_version,
            "NearestModelName": model_name,
            "ModelInput": {"DataInputConfig": data_input_configuration},
        }
    ],
    "SupportedContentTypes": input_mime_types,  # required, must be non-null
    "SupportedResponseMIMETypes": [],
    "SupportedRealtimeInferenceInstanceTypes": supported_realtime_inference_types,  # optional
}
```

If these conditions are met in your request, then Inference Recommender runs scenarios for both compiled and uncompiled versions of your model, giving you multiple recommendation combinations to choose from. You can compare the configurations for compiled and uncompiled versions of the same inference recommendation and determine which one best suits your use case. The recommendations are ranked by cost per inference.

To get the Neo compilation recommendations, you don’t have to do any additional configuration besides making sure that your input meets the preceding requirements. Inference Recommender automatically runs Neo compilation on your model if your input meets the requirements, and you receive a response that includes Neo recommendations.

If you run into errors during your Neo compilation, see [Troubleshoot Neo Compilation Errors](#).

The following table is an example of a response you might get from an Inference Recommender job that includes recommendations for compiled models. If the `InferenceSpecificationName` field is None, then the recommendation is an uncompiled model. The last row, in which the value for the `InferenceSpecificationName` field is neo-00011122-2333-4445-5566-677788899900, is for a model compiled with Neo. The value in the field is the name of the Neo job used to compile and optimize your model.
<table>
<thead>
<tr>
<th>EndpointName</th>
<th>InstanceType</th>
<th>InitialInstanceCount</th>
<th>EnvironmentParameters</th>
<th>CostPerHour</th>
<th>CostPerInference</th>
<th>MaxInvocations</th>
<th>ModelLatency</th>
<th>InferenceSpecificationName</th>
</tr>
</thead>
<tbody>
<tr>
<td>sm-epc-example-00111222</td>
<td>ml.c5.9xlarge</td>
<td>1</td>
<td>[]</td>
<td>1.836</td>
<td>9.15E-07</td>
<td>33456</td>
<td>7</td>
<td>None</td>
</tr>
<tr>
<td>sm-epc-example-11222333</td>
<td>ml.c5.2xlarge</td>
<td>1</td>
<td>[]</td>
<td>0.408</td>
<td>2.11E-07</td>
<td>32211</td>
<td>21</td>
<td>None</td>
</tr>
<tr>
<td>sm-epc-example-22333444</td>
<td>ml.c5.xlarge</td>
<td>1</td>
<td>[]</td>
<td>0.204</td>
<td>1.86E-07</td>
<td>18276</td>
<td>92</td>
<td>None</td>
</tr>
<tr>
<td>sm-epc-example-33444555</td>
<td>ml.c5.xlarge</td>
<td>1</td>
<td>[]</td>
<td>0.204</td>
<td>1.60E-07</td>
<td>21286</td>
<td>42</td>
<td>neo-00011122-2333-4445-5566-677788899900</td>
</tr>
</tbody>
</table>

**Get started**

The general steps for creating an Inference Recommender job that includes Neo-optimized recommendations are as follows:

- Prepare your ML model for compilation. For more information, see [Prepare Model for Compilation](#) in the Neo documentation.
- Package your model in a model archive (.tar.gz file).
- Create a sample payload archive.
- Register your model in SageMaker Model Registry.
- Create an Inference Recommender job.
• View the results of the Inference Recommender job and choose a configuration.

• Debug compilation failures, if any. For more information, see Troubleshoot Neo Compilation Errors.

For an example that demonstrates the previous workflow and how to get Neo-optimized recommendations using XGBoost, see the following example notebook. For an example that show how to get Neo-optimized recommendations using TensorFlow, see the following example notebook.

Interpret recommendation results

Each Inference Recommender job result includes InstanceType, InitialInstanceCount, and EnvironmentParameters, which are tuned environment variable parameters for your container to improve its latency and throughput. The results also include performance and cost metrics such as MaxInvocations, ModelLatency, CostPerHour, CostPerInference, CpuUtilization, and MemoryUtilization.

In the table below we provide a description of these metrics. These metrics can help you narrow down your search for the best endpoint configuration that suits your use case. For example, if your motivation is overall price performance with an emphasis on throughput, then you should focus on CostPerInference.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
<th>Use case</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelLatency</td>
<td>The interval of time taken by a model to respond as viewed from SageMaker. This interval includes the local communication times taken to send the request and to fetch the response from the container of a model and the time taken to complete the inference in the container. Units: Milliseconds</td>
<td>Latency sensitive workloads such as ad serving and medical diagnosis</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
<td>Use case</td>
</tr>
<tr>
<td>----------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>MaximumInvocations</td>
<td>The maximum number of InvokeEndpoint requests sent to a model endpoint in a minute.</td>
<td>Throughput-focused workloads such as video processing or batch inference</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
<td></td>
</tr>
<tr>
<td>CostPerHour</td>
<td>The estimated cost per hour for your real-time endpoint.</td>
<td>Cost sensitive workloads with no latency deadlines</td>
</tr>
<tr>
<td></td>
<td>Units: US Dollars</td>
<td></td>
</tr>
<tr>
<td>CostPerInference</td>
<td>The estimated cost per inference call for your real-time endpoint.</td>
<td>Maximize overall price performance with a focus on throughput</td>
</tr>
<tr>
<td></td>
<td>Units: US Dollars</td>
<td></td>
</tr>
<tr>
<td>CpuUtilization</td>
<td>The expected CPU utilization at maximum invocations per minute for the endpoint instance.</td>
<td>Understand instance health during benchmarking by having visibility into core CPU utilization of the instance</td>
</tr>
<tr>
<td></td>
<td>Units: Percent</td>
<td></td>
</tr>
<tr>
<td>MemoryUtilization</td>
<td>The expected memory utilization at maximum invocations per minute for the endpoint instance.</td>
<td>Understand instance health during benchmarking by having visibility into core memory utilization of the instance</td>
</tr>
<tr>
<td></td>
<td>Units: Percent</td>
<td></td>
</tr>
</tbody>
</table>

In some cases you might want to explore other SageMaker Endpoint Invocation metrics such as CpuUtilization. Every Inference Recommender job result includes the names of endpoints spun up during the load test. You can use CloudWatch to review the logs for these endpoints even after they've been deleted.
The following image is an example of CloudWatch metrics and charts you can review for a single endpoint from your recommendation result. This recommendation result is from a Default job. The way to interpret the scalar values from the recommendation results is that they are based on the time point when the Invocations graph first begins to level out. For example, the ModelLatency value reported is at the beginning of the plateau around 03:00:31.

For full descriptions of the CloudWatch metrics used in the preceding charts, see SageMaker Endpoint Invocation metrics.

You can also see performance metrics like ClientInvocations and NumberOfUsers published by Inference Recommender in the /aws/sagemaker/InferenceRecommendationsJobs namespace. For a full list of metrics and descriptions published by Inference Recommender, see SageMaker Inference Recommender Jobs Metrics.
See the Amazon SageMaker Inference Recommender - CloudWatch Metrics Jupyter notebook in the amazon-sagemaker-examples Github repository for an example of how to use the AWS SDK for Python (Boto3) to explore CloudWatch metrics for your endpoints.

Get autoscaling policy recommendations

With Amazon SageMaker Inference Recommender, you can get recommendations for autoscaling policies for your SageMaker endpoint based on your anticipated traffic pattern. If you've already completed an inference recommendation job, you can provide the details of the job to get a recommendation for an autoscaling policy that you can apply to your endpoint.

Inference Recommender benchmarks different values for each metric to determine the ideal autoscaling configuration for your endpoint. The autoscaling recommendation returns a recommended autoscaling policy for each metric that was defined in your inference recommendation job. You can save the policies and apply them to your endpoint with the PutScalingPolicy API.

To get started, review the following prerequisites.

Prerequisites

Before you begin, you must have completed a successful inference recommendation job. In the following section, you can provide either an inference recommendation ID or the name of a SageMaker endpoint that was benchmarked during an inference recommendation job.

To retrieve your recommendation job ID or endpoint name, you can either view the details of your inference recommendation job in the SageMaker console, or you can use the RecommendationId or EndpointName fields returned by the DescribeInferenceRecommendationsJob API.

Create an autoscaling configuration recommendation

To create an autoscaling recommendation policy, you can use the AWS SDK for Python (Boto3).

The following example shows the fields for the GetScalingConfigurationRecommendation API. Use the following fields when you call the API:

- InferenceRecommendationsJobName – Enter the name of your inference recommendation job.
- RecommendationId – Enter the ID of an inference recommendation from a recommendation job. This is optional if you've specified the EndpointName field.
• EndpointName – Enter the name of an endpoint that was benchmarked during an inference recommendation job. This is optional if you’ve specified the RecommendationId field.

• TargetCpuUtilizationPerCore – (Optional) Enter a percentage value of how much utilization you want an instance on your endpoint to use before autoscaling. The default value if you don’t specify this field is 50%.

• ScalingPolicyObjective – (Optional) An object where you specify your anticipated traffic pattern.
  • MinInvocationsPerMinute – (Optional) The minimum number of expected requests to your endpoint per minute.
  • MaxInvocationsPerMinute – (Optional) The maximum number of expected requests to your endpoint per minute.

```json
{
  "InferenceRecommendationsJobName": "string", // Required
  "RecommendationId": "string", // Optional, provide one of RecommendationId or EndpointName
  "EndpointName": "string", // Optional, provide one of RecommendationId or EndpointName
  "TargetCpuUtilizationPerCore": number, // Optional
  "ScalingPolicyObjective": { // Optional
    "MinInvocationsPerMinute": number,
    "MaxInvocationsPerMinute": number
  }
}
```

After submitting your request, you’ll receive a response with autoscaling policies defined for each metric. See the following section for information about interpreting the response.

**Review your autoscaling configuration recommendation results**

The following example shows the response from the [GetScalingConfigurationRecommendation](https://docs.aws.amazon.com/sagemaker/latest/dg/API_GetScalingConfigurationRecommendation.html) API:

```json
{
  "InferenceRecommendationsJobName": "string",
  "RecommendationId": "string", // One of RecommendationId or EndpointName is shown
  "EndpointName": "string",
  "TargetUtilizationPercentage": Integer,
  "ScalingPolicyObjective": {
```

**Recommendation jobs**

---

*Page 4034*
The InferenceRecommendationsJobName, RecommendationID or EndpointName, TargetCpuUtilizationPerCore, and the ScalingPolicyObjective object fields are copied from your initial request.

The Metric object lists the metrics that were benchmarked in your inference recommendation job, along with a calculation of the values for each metric when the instance utilization would be the same as the TargetCpuUtilizationPerCore value. This is useful for anticipating the performance metrics on your endpoint when it scales in and out with the recommended autoscaling policy. For example, consider if your instance utilization was 50% in your inference recommendation job and your InvocationsPerInstance value was originally 4. If you specify
the TargetCpuUtilizationPerCore value to be 100% in your autoscaling recommendation request, then the InvocationsPerInstance metric value returned in the response is 2 because you anticipated allocating twice as much instance utilization.

The DynamicScalingConfiguration object returns the values that you should specify for the TargetTrackingScalingPolicyConfiguration when you call the PutScalingPolicy API. This includes the recommended minimum and maximum capacity values, the recommended scale in and scale out cooldown times, and the ScalingPolicies object, which contains the recommended TargetValue you should specify for each metric.

Run a custom load test

Amazon SageMaker Inference Recommender load tests conduct extensive benchmarks based on production requirements for latency and throughput, custom traffic patterns, and either serverless endpoints or real-time instances (up to 10) that you select.

The following sections demonstrate how to create, describe, and stop a load test programmatically using the AWS SDK for Python (Boto3) and the AWS CLI, or interactively using Amazon SageMaker Studio Classic or the SageMaker console.

Create a load test job

Create a load test programmatically using the AWS SDK for Python (Boto3), with the AWS CLI, or interactively using Studio Classic or the SageMaker console. As with Inference Recommender inference recommendations, specify a job name for your load test, an AWS IAM role ARN, an input configuration, and your model package ARN from when you registered your model with the model registry. Load tests require that you also specify a traffic pattern and stopping conditions.

AWS SDK for Python (Boto3)

Use the CreateInferenceRecommendationsJob API to create an Inference Recommender load test. Specify Advanced for the JobType field and provide:

- A job name for your load test (JobName). The job name must be unique within your AWS Region and within your AWS account.
- The Amazon Resource Name (ARN) of an IAM role that enables Inference Recommender to perform tasks on your behalf. Define this for the RoleArn field.
- An endpoint configuration dictionary (InputConfig) where you specify the following:
  - For TrafficPattern, specify either the phases or stairs traffic pattern. With the phases traffic pattern, new users spawn every minute at the rate you specify. With the stairs traffic pattern,
pattern, new users spawn at timed intervals (or steps) at a rate you specify. Choose one of the following:

- For **TrafficType**, specify **PHASES**. Then, for the **Phases** array, specify the **InitialNumberOfUsers** (how many concurrent users to start with, with a minimum of 1 and a maximum of 3), **SpawnRate** (the number of users to be spawned in a minute for a specific phase of load testing, with a minimum of 0 and maximum of 3), and **DurationInSeconds** (how long the traffic phase should be, with a minimum of 120 and maximum of 3600).

- For **TrafficType**, specify **STAIRS**. Then, for the **Stairs** array, specify the **DurationInSeconds** (how long the traffic phase should be, with a minimum of 120 and maximum of 3600), **NumberOfSteps** (how many intervals are used during the phase), and **UsersPerStep** (how many users are added during each interval). Note that the length of each step is the value of DurationInSeconds / NumberOfSteps. For example, if your DurationInSeconds is 600 and you specify 5 steps, then each step is 120 seconds long.

### Note

A user is defined as a system-generated actor that runs in a loop and invokes requests to an endpoint as part of Inference Recommender. For a typical XGBoost container running on an `ml.c5.large` instance, endpoints can reach 30,000 invocations per minute (500 tps) with just 15-20 users.

- For **ResourceLimit**, specify **MaxNumberOfTests** (the maximum number of benchmarking load tests for an Inference Recommender job, with a minimum of 1 and a maximum of 10) and **MaxParallelOfTests** (the maximum number of parallel benchmarking load tests for an Inference Recommender job, with a minimum of 1 and a maximum of 10).

- For **EndpointConfigurations**, you can specify one of the following:
  - The **InstanceType** field, where you specify the instance type on which you want to run your load tests.
  - The **ServerlessConfig**, in which you specify your ideal values for **MaxConcurrency** and **MemorySizeInMB** for a serverless endpoint. For more information, see the [Serverless Inference documentation](#).
- A stopping conditions dictionary (StoppingConditions), where if any of the conditions are met, the Inference Recommender job stops. For this example, specify the following fields in the dictionary:
  - For MaxInvocations, specify the maximum number of requests per minute expected for the endpoint, with a minimum of 1 and a maximum of 30,000.
  - For ModelLatencyThresholds, specify Percentile (the model latency percentile threshold) and ValueInMilliseconds (the model latency percentile value in milliseconds).
  - (Optional) For FlatInvocations, you can specify whether to continue the load test when the TPS (invocations per minute) rate flattens. A flattened TPS rate usually means that the endpoint has reached capacity. However, you might want to continue monitoring the endpoint under full capacity conditions. To continue the load test when this happens, specify this value as Continue. Otherwise, the default value is Stop.

```python
# Create a low-level SageMaker service client.
import boto3
aws_region=<INSERT>
sagemaker_client=boto3.client('sagemaker', region=aws_region)

# Provide a name to your recommendation based on load testing
load_test_job_name=<INSERT>

# Provide the name of the sagemaker instance type
instance_type=<INSERT>

# Provide the IAM Role that gives SageMaker permission to access AWS services
role_arn='arn:aws:iam::<account>:role/*'

# Provide your model package ARN that was created when you registered your
# model with Model Registry
model_package_arn='arn:aws:sagemaker:<region>:<account>:role/*'

sagemaker_client.create_inference_recommendations_job(
    JobName=load_test_job_name,
    JobType="Advanced",
    RoleArn=role_arn,
    InputConfig={
        'ModelPackageVersionArn': model_package_arn,
        "JobDurationInSeconds": 7200,
    })
```
'TrafficPattern': {
  # Replace PHASES with STAIRS to use the stairs traffic pattern
  'TrafficType': 'PHASES',
  'Phases': [
    {
      'InitialNumberOfUsers': 1,
      'SpawnRate': 1,
      'DurationInSeconds': 120
    },
    {
      'InitialNumberOfUsers': 1,
      'SpawnRate': 1,
      'DurationInSeconds': 120
    }
  ]
},
# Uncomment this section and comment out the Phases object above to use the stairs traffic pattern
# 'Stairs': {
#   'DurationInSeconds': 240,
#   'NumberOfSteps': 2,
#   'UsersPerStep': 2
# },
'ResourceLimit': {
  'MaxNumberOfTests': 10,
  'MaxParallelOfTests': 3
},
"EndpointConfigurations": [
  {
    'InstanceType': 'ml.c5.xlarge'
  },
  {
    'InstanceType': 'ml.m5.xlarge'
  },
  {
    'InstanceType': 'ml.r5.xlarge'
  }
]
# Uncomment the ServerlessConfig and comment out the InstanceType field if you want recommendations for a serverless endpoint
# "ServerlessConfig": {
#   "MaxConcurreny": value,
#   "MemorySizeInMB": value
# }
}
StoppingConditions={
    'MaxInvocations': 1000,
    'ModelLatencyThresholds': [{
        'Percentile': 'P95',
        'ValueInMilliseconds': 100
    }],
    # Change 'Stop' to 'Continue' to let the load test continue if invocations flatten
    'FlatInvocations': 'Stop'
}

See the Amazon SageMaker API Reference Guide for a full list of optional and required arguments you can pass to CreateInferenceRecommendationsJob.

AWS CLI

Use the create-inference-recommendations-job API to create an Inference Recommender load test. Specify Advanced for the JobType field and provide:

- A job name for your load test (job-name). The job name must be unique within your AWS Region and within your AWS account.
- The Amazon Resource Name (ARN) of an IAM role that enables Inference Recommender to perform tasks on your behalf. Define this for the role-arn field.
- An endpoint configuration dictionary (input-config) where you specify the following:
  - For TrafficPattern, specify either the phases or stairs traffic pattern. With the phases traffic pattern, new users spawn every minute at the rate you specify. With the stairs traffic pattern, new users spawn at timed intervals (or steps) at a rate you specify. Choose one of the following:
    - For TrafficType, specify PHASES. Then, for the Phases array, specify the InitialNumberOfUsers (how many concurrent users to start with, with a minimum of 1 and a maximum of 3), SpawnRate (the number of users to be spawned in a minute for a specific phase of load testing, with a minimum of 0 and maximum of 3), and DurationInSeconds (how long the traffic phase should be, with a minimum of 120 and maximum of 3600).
    - For TrafficType, specify STAIRS. Then, for the Stairs array, specify the DurationInSeconds (how long the traffic phase should be, with a minimum of 120 and maximum of 3600), NumberOfSteps (how many intervals are used during the phase), and UsersPerStep (how many users are added during each interval). Note that
the length of each step is the value of DurationInSeconds / NumberOfSteps. For example, if your DurationInSeconds is 600 and you specify 5 steps, then each step is 120 seconds long.

Note

A user is defined as a system-generated actor that runs in a loop and invokes requests to an endpoint as part of Inference Recommender. For a typical XGBoost container running on an ml.c5.large instance, endpoints can reach 30,000 invocations per minute (500 tps) with just 15-20 users.

- For ResourceLimit, specify MaxNumberOfTests (the maximum number of benchmarking load tests for an Inference Recommender job, with a minimum of 1 and a maximum of 10) and MaxParallelNumberOfTests (the maximum number of parallel benchmarking load tests for an Inference Recommender job, with a minimum of 1 and a maximum of 10).

- For EndpointConfigurations, you can specify one of the following:
  - The InstanceType field, where you specify the instance type on which you want to run your load tests.
  - The ServerlessConfig, in which you specify your ideal values for MaxConcurrency and MemorySizeInMB for a serverless endpoint.

- A stopping conditions dictionary (stopping-conditions), where if any of the conditions are met, the Inference Recommender job stops. For this example, specify the following fields in the dictionary:
  - For MaxInvocations, specify the maximum number of requests per minute expected for the endpoint, with a minimum of 1 and a maximum of 30,000.
  - For ModelLatencyThresholds, specify Percentile (the model latency percentile threshold) and ValueInMilliseconds (the model latency percentile value in milliseconds).

- (Optional) For FlatInvocations, you can specify whether to continue the load test when the TPS (invocations per minute) rate flattens. A flattened TPS rate usually means that the endpoint has reached capacity. However, you might want to continue monitoring the endpoint under full capacity conditions. To continue the load test when this happens, specify this value as Continue. Otherwise, the default value is Stop.
aws sagemaker create-inference-recommendations-job
   --region <region>\n   --job-name <job-name>\n   --job-type ADVANCED\n   --role-arn arn:aws:iam::<account>:role/*\n   --input-config "{
        "JobDurationInSeconds": 7200,
        "TrafficPattern": {
            # Replace PHASES with STAIRS to use the stairs traffic pattern
            "TrafficType": "PHASES",
            "Phases": [
                {
                    "InitialNumberOfUsers": 1,
                    "SpawnRate": 60,
                    "DurationInSeconds": 300
                }
            ]
        # Uncomment this section and comment out the Phases object above to use the stairs traffic pattern
        # 'Stairs': {
        # 'DurationInSeconds': 240,
        # 'NumberOfSteps': 2,
        # 'UsersPerStep': 2
        # }
        },
        "ResourceLimit": {
            "MaxNumberOfTests": 10,
            "MaxParallelOfTests": 3
        },
        "EndpointConfigurations": [
            {
                "InstanceType": "ml.c5.xlarge"
            },
            {
                "InstanceType": "ml.m5.xlarge"
            },
            {
                "InstanceType": "ml.r5.xlarge"
            }
        ]
        # Use the ServerlessConfig and leave out the InstanceType fields if you want recommendations for a serverless endpoint
        # "ServerlessConfig": {
    }
Amazon SageMaker Studio Classic

Create a load test with Studio Classic.

1. In your Studio Classic application, choose the home icon
   ![home icon]
2. In the left sidebar of Studio Classic, choose Deployments.
3. Choose Inference recommender from the dropdown list.
5. Select the name of your model group from the dropdown Model group field. The list includes all the model groups registered with the model registry in your account, including models registered outside of Studio Classic.
6. Select a model version from the dropdown Model version field.
7. Choose Continue.
8. Provide a name for the job in the Name field.
9. (Optional) Provide a description of your job in the Description field.
10. Choose an IAM role that grants Inference Recommender permission to access AWS services. You can create a role and attach the AmazonSageMakerFullAccess IAM managed policy to accomplish this, or you can let Studio Classic create a role for you.
11. Choose **Stopping Conditions** to expand the available input fields. Provide a set of conditions for stopping a deployment recommendation.

   a. Specify the maximum number of requests per minute expected for the endpoint in the **Max Invocations Per Minute** field.

   b. Specify the model latency threshold in microseconds in the **Model Latency Threshold** field. The **Model Latency Threshold** depicts the interval of time taken by a model to respond as viewed from Inference Recommender. The interval includes the local communication time taken to send the request and to fetch the response from the model container and the time taken to complete the inference in the container.

12. Choose **Traffic Pattern** to expand the available input fields.

   a. Set the initial number of virtual users by specifying an integer in the **Initial Number of Users** field.

   b. Provide an integer number for the **Spawn Rate** field. The spawn rate sets the number of users created per second.

   c. Set the duration for the phase in seconds by specifying an integer in the **Duration** field.

   d. (Optional) Add additional traffic patterns. To do so, choose **Add**.

13. Choose the **Additional** setting to reveal the **Max test duration** field. Specify, in seconds, the maximum time a test can take during a job. New jobs are not scheduled after the defined duration. This helps ensure jobs that are in progress are not stopped and that you only view completed jobs.

14. Choose **Continue**.

15. Choose **Selected Instances**.

16. In the **Instances for benchmarking** field, choose **Add instances to test**. Select up to 10 instances for Inference Recommender to use for load testing.

17. Choose **Additional settings**.

   a. Provide an integer that sets an upper limit on the number of tests a job can make for the **Max number of tests field**. Note that each endpoint configuration results in a new load test.

   b. Provide an integer for the **Max parallel** test field. This setting defines an upper limit on the number of load tests that can run in parallel.

18. Choose **Submit**.
The load test can take up to 2 hours.

⚠️ Warning
Do not close this tab. If you close this tab, you cancel the Inference Recommender load test job.

SageMaker console

Create a custom load test through the SageMaker console by doing the following:

2. In the left navigation pane, choose Inference, and then choose Inference recommender.
3. On the Inference recommender jobs page, choose Create job.
4. For Step 1: Model configuration, do the following:
   a. For Job type, choose Advanced recommender job.
   b. If you're using a model registered in the SageMaker model registry, then turn on the Choose a model from the model registry toggle and do the following:
      i. For the Model group dropdown list, choose the model group in SageMaker model registry where your model is.
      ii. For the Model version dropdown list, choose the desired version of your model.
   c. If you're using a model that you've created in SageMaker, then turn off the Choose a model from the model registry toggle and do the following:
      • For the Model name field, enter the name of your SageMaker model.
   d. For IAM role, you can select an existing AWS IAM role that has the necessary permissions to create an instance recommendation job. Alternatively, if you don't have an existing role, you can choose Create a new role to open the role creation pop-up, and SageMaker adds the necessary permissions to the new role that you create.
   e. For S3 bucket for benchmarking payload, enter the Amazon S3 path to your sample payload archive, which should contain sample payload files that Inference Recommender uses to benchmark your model on different instance types.
   f. For Payload content type, enter the MIME types of your sample payload data.
g. For Traffic pattern, configure phases for the load test by doing the following:

i. For Initial number of users, specify how many concurrent users you want to start with (with a minimum of 1 and a maximum of 3).

ii. For Spawn rate, specify the number of users to be spawned in a minute for the phase (with a minimum of 0 and a maximum of 3).

iii. For Duration (seconds), specify how low the traffic phase should be in seconds (with a minimum of 120 and a maximum of 3600).

h. (Optional) If you turned off the Choose a model from the model registry toggle and specified a SageMaker model, then for Container configuration, do the following:

i. For the Domain dropdown list, select the machine learning domain of the model, such as computer vision, natural language processing, or machine learning.

ii. For the Framework dropdown list, select the framework of your container, such as TensorFlow or XGBoost.

iii. For Framework version, enter the framework version of your container image.

iv. For the Nearest model name dropdown list, select the pre-trained model that mostly closely matches your own.

v. For the Task dropdown list, select the machine learning task that the model accomplishes, such as image classification or regression.

i. (Optional) For Model compilation using SageMaker Neo, you can configure the recommendation job for a model that you've compiled using SageMaker Neo. For Data input configuration, enter the correct input data shape for your model in a format similar to {'input': [1, 1024, 1024, 3]}.

j. Choose Next.

5. For Step 2: Instances and environment parameters, do the following:

a. For Select instances for benchmarking, select up to 8 instance types that you want to benchmark against.

b. (Optional) For Environment parameter ranges, you can specify environment parameters that help optimize your model. Specify the parameters as Key and Value pairs.

c. Choose Next.

6. For Step 3: Job parameters, do the following:
a. (Optional) For the **Job name** field, enter a name for your instance recommendation job. When you create the job, SageMaker appends a timestamp to the end of this name.

b. (Optional) For the **Job description** field, enter a description for the job.

c. (Optional) For the **Encryption key** dropdown list, choose an AWS KMS key by name or enter its ARN to encrypt your data.

d. (Optional) For **Max number of tests**, enter the number of test that you want to run during the recommendation job.

e. (Optional) For **Max parallel tests**, enter the maximum number of parallel tests that you want to run during the recommendation job.

f. For **Max test duration (s)**, enter the maximum number of seconds you want each test to run for.

g. For **Max invocations per minute**, enter the maximum number of requests per minute the endpoint can reach before stopping the recommendation job. After reaching this limit, SageMaker ends the job.

h. For **P99 Model latency threshold (ms)**, enter the model latency percentile in milliseconds.

i. Choose **Next**.

7. For **Step 4: Review job**, review your configurations and then choose **Submit**.

**Get your load test results**

You can programmatically collect metrics across all load tests once the load tests are done with AWS SDK for Python (Boto3), the AWS CLI, Studio Classic, or the SageMaker console.

**AWS SDK for Python (Boto3)**

Collect metrics with the `DescribeInferenceRecommendationsJob` API. Specify the job name of the load test for the `JobName` field:

```python
load_test_response = sagemaker_client.describe_inference_recommendations_job(
    JobName=load_test_job_name
)
```

Print the response object.
This returns a JSON response similar to the following example. Note that this example shows the recommended instance types for real-time inference (for an example showing serverless inference recommendations, see the example after this one).

```json
{
    'JobName': 'job-name',
    'JobDescription': 'job-description',
    'JobType': 'Advanced',
    'Status': 'COMPLETED',
    'CreationTime': datetime.datetime(2021, 10, 26, 19, 38, 30, 957000, tzinfo=tzlocal()),
    'LastModifiedTime': datetime.datetime(2021, 10, 26, 19, 46, 31, 399000, tzinfo=tzlocal()),
    'InputConfig': {
        'JobDurationInSeconds': 7200,
        'TrafficPattern': {
            'TrafficType': 'PHASES'
        },
        'ResourceLimit': {
            'MaxNumberOfTests': 100,
            'MaxParallelOfTests': 100
        },
        'EndpointConfigurations': [{
            'InstanceType': 'ml.c5d.xlarge'
        }]
    },
    'StoppingConditions': {
        'MaxInvocations': 1000,
        'ModelLatencyThresholds': [{
            'Percentile': 'P95',
            'ValueInMilliseconds': 100
        }],
        'InferenceRecommendations': [{
            'Metrics': {
                'CostPerHour': 0.6899999976158142,
                'CostPerInference': 1.0332434612791985e-05,
            }
        }]
    }
}
```
The first few lines provide information about the load test job itself. This includes the job name, role ARN, creation, and deletion time.

The InferenceRecommendations dictionary contains a list of Inference Recommender inference recommendations.

The EndpointConfiguration nested dictionary contains the instance type (InstanceType) recommendation along with the endpoint and variant name (a deployed AWS machine learning model) used during the recommendation job. You can use the endpoint and variant name for monitoring in Amazon CloudWatch Events. See Monitor Amazon SageMaker with Amazon CloudWatch for more information.

The EndpointConfiguration nested dictionary also contains the instance count (InitialInstanceCount) recommendation. This is the number of instances that you should provision in the endpoint to meet the MaxInvocations specified in the StoppingConditions. For example, if the InstanceType is ml.m5.large and the
InitialInstanceCount is 2, then you should provision 2 ml.m5.large instances for your endpoint so that it can handle the TPS specified in the MaxInvocations stopping condition.

The Metrics nested dictionary contains information about the estimated cost per hour (CostPerHour) for your real-time endpoint in US dollars, the estimated cost per inference (CostPerInference) for your real-time endpoint, the maximum number of InvokeEndpoint requests sent to the endpoint, and the model latency (ModelLatency), which is the interval of time (in microseconds) that your model took to respond to SageMaker. The model latency includes the local communication times taken to send the request and to fetch the response from the model container and the time taken to complete the inference in the container.

The following example shows the InferenceRecommendations part of the response for a load test job that was configured to return serverless inference recommendations:

```
"InferenceRecommendations": [ 
  {
    "EndpointConfiguration": {
      "EndpointName": "value",
      "InitialInstanceCount": "value",
      "InstanceType": "value",
      "VariantName": "value",
      "ServerlessConfig": {
        "MaxConcurrency": "value",
        "MemorySizeInMb": "value"
      }
    },
    "InvocationEndTime": "value",
    "InvocationStartTime": "value",
    "Metrics": {
      "CostPerHour": "value",
      "CostPerInference": "value",
      "CpuUtilization": "value",
      "MaxInvocations": "value",
      "MemoryUtilization": "value",
      "ModelLatency": "value",
      "ModelSetupTime": "value"
    },
    "ModelConfiguration": {
      "Compiled": "False",
      "EnvironmentParameters": [],
      "InferenceSpecificationName": "value"
    }
  }]
```

Recommendation jobs 4050
You can interpret the recommendations for serverless inference similarly to the results for real-time inference, with the exception of the ServerlessConfig, which tells you the values you specified for MaxConcurrency and MemorySizeInMB when setting up the load test. Serverless recommendations also measure the metric ModelSetupTime, which measures (in microseconds) the time it takes to launch compute resources on a serverless endpoint. For more information about setting up serverless endpoints, see the Serverless Inference documentation.

AWS CLI

Collect metrics with the describe-inference-recommendations-job API. Specify the job name of the load test for the job-name flag:

```
aws sagemaker describe-inference-recommendations-job --job-name <job-name>
```

This returns a response similar to the following example. Note that this example shows the recommended instance types for real-time inference (for an example showing Serverless Inference recommendations, see the example after this one).

```json
{
    'JobName': 'job-name',
    'JobDescription': 'job-description',
    'JobType': 'Advanced',
    'Status': 'COMPLETED',
    'CreationTime': datetime.datetime(2021, 10, 26, 19, 38, 30, 957000, tzinfo=tzlocal()),
    'LastModifiedTime': datetime.datetime(2021, 10, 26, 19, 46, 31, 399000, tzinfo=tzlocal()),
    'InputConfig': {
        'JobDurationInSeconds': 7200,
        'TrafficPattern': {
            'TrafficType': 'PHASES'
        },
        'ResourceLimit': {
            'MaxNumberOfTests': 100,
```
{'MaxParallelOfTests': 100},
'EndpointConfigurations': [{
    'InstanceType': 'ml.c5d.xlarge'}],
'StoppingConditions': {
    'MaxInvocations': 1000,
    'ModelLatencyThresholds': [{
        'Percentile': 'P95',
        'ValueInMilliseconds': 100}]
},
'InferenceRecommendations': [{
    'Metrics': {
        'CostPerHour': 0.6899999976158142,
        'CostPerInference': 1.0332434612791985e-05,
        'MaximumInvocations': 1113,
        'ModelLatency': 100000}
},
'EndpointConfiguration': {
    'EndpointName': 'endpoint-name',
    'VariantName': 'variant-name',
    'InstanceType': 'ml.c5d.xlarge',
    'InitialInstanceCount': 3}
},
'ModelConfiguration': {
    'Compiled': False,
    'EnvironmentParameters': []}
},
'ResponseMetadata': {
    'RequestId': 'request-id',
    'HttpStatus': 200,
    'HTTPHeaders': {
        'x-amzn-requestid': 'x-amzn-requestid',
        'content-type': 'content-type',
        'content-length': '1199',
        'date': 'Tue, 26 Oct 2021 19:57:42 GMT'},
    'RetryAttempts': 0}
The first few lines provide information about the load test job itself. This includes the job name, role ARN, creation, and deletion time.

The InferenceRecommendations dictionary contains a list of Inference Recommender inference recommendations.

The EndpointConfiguration nested dictionary contains the instance type (InstanceType) recommendation along with the endpoint and variant name (a deployed AWS machine learning model) used during the recommendation job. You can use the endpoint and variant name for monitoring in Amazon CloudWatch Events. See Monitor Amazon SageMaker with Amazon CloudWatch for more information.

The Metrics nested dictionary contains information about the estimated cost per hour (CostPerHour) for your real-time endpoint in US dollars, the estimated cost per inference (CostPerInference) for your real-time endpoint, the maximum number of InvokeEndpoint requests sent to the endpoint, and the model latency (ModelLatency), which is the interval of time (in microseconds) that your model took to respond to SageMaker. The model latency includes the local communication times taken to send the request and to fetch the response from the model container and the time taken to complete the inference in the container.

The following example shows the InferenceRecommendations part of the response for a load test job that was configured to return serverless inference recommendations:

```json
"InferenceRecommendations": [
  {
    "EndpointConfiguration": {
      "EndpointName": "value",
      "InitialInstanceCount": value,
      "InstanceType": "value",
      "VariantName": "value",
      "ServerlessConfig": {
        "MaxConcurrency": value,
        "MemorySizeInMb": value
      }
    },
    "InvocationEndTime": value,
    "InvocationStartTime": value,
    "Metrics": {
      "CostPerHour": value,
      "CostPerInference": value,
      "CpuUtilization": value,
      "MaxInvocations": value,
    }
  }]
```
You can interpret the recommendations for serverless inference similarly to the results for real-time inference, with the exception of the ServerlessConfig, which tells you the values you specified for MaxConcurrency and MemorySizeInMB when setting up the load test. Serverless recommendations also measure the metric ModelSetupTime, which measures (in microseconds) the time it takes to launch computer resources on a serverless endpoint. For more information about setting up serverless endpoints, see the Serverless Inference documentation.

Amazon SageMaker Studio Classic

The recommendations populate in a new tab called Inference recommendations within Studio Classic. It can take up to 2 hours for the results to show up. This tab contains Results and Details columns.

The Details column provides information about the load test job, such as the name given to the load test job, when the job was created (Creation time), and more. It also contains Settings information, such as the maximum number of invocation that occurred per minute and information about the Amazon Resource Names used.

The Results column provides Deployment goals and SageMaker recommendations windows in which you can adjust the order in which results are displayed based on deployment importance. There are three dropdown menus in which you can provide the level of importance of the Cost, Latency, and Throughput for your use case. For each goal (cost, latency, and throughput), you can set the level of importance: Lowest Importance, Low Importance, Moderate importance, High importance, or Highest importance.

Based on your selections of importance for each goal, Inference Recommender displays its top recommendation in the SageMaker recommendation field on the right of the panel, along
with the estimated cost per hour and inference request. It also provides information about the expected model latency, maximum number of invocations, and the number of instances.

In addition to the top recommendation displayed, you can also see the same information displayed for all instances that Inference Recommender tested in the All runs section.

SageMaker console

You can view your custom load test job results in the SageMaker console by doing the following:

2. In the left navigation pane, choose Inference, and then choose Inference recommender.
3. On the Inference recommender jobs page, choose the name of your inference recommendation job.

On the details page for your job, you can view the Inference recommendations, which are the instance types SageMaker recommends for your model, as shown in the following screenshot.

In this section, you can compare the instance types by various factors such as Model latency, Cost per hour, Cost per inference, and Invocations per minute.

On this page, you can also view the configurations you specified for your job. In the Monitor section, you can view the Amazon CloudWatch metrics that were logged for each instance type. To learn more about interpreting these metrics, see Interpret results.
Stop your load test

You might want to stop a job that is currently running if you began a job by mistake or no longer need to run the job. Stop your load test jobs programmatically with the StopInferenceRecommendationsJob API, or through Studio Classic or the SageMaker console.

AWS SDK for Python (Boto3)

Specify the job name of the load test for the JobName field:

```python
sagemaker_client.stop_inference_recommendations_job(
    JobName='<INSERT>'
)
```

AWS CLI

Specify the job name of the load test for the job-name flag:

```bash
aws sagemaker stop-inference-recommendations-job --job-name <job-name>
```

Amazon SageMaker Studio Classic

Close the tab where you initiated your custom load job to stop your Inference Recommender load test.

SageMaker console

To stop your load test job through the SageMaker console, do the following:

2. In the left navigation pane, choose **Inference**, and then choose **Inference recommender**.
3. On the **Inference recommender jobs** page, select your load test job.
4. Choose **Stop job**.
5. In the dialog box that pops up, choose **Confirm**.

After stopping your job, the job’s **Status** should change to **Stopping**.
Troubleshoot Inference Recommender errors

This section contains information about how to understand and prevent common errors, the error messages they generate, and guidance on how to resolve these errors.

How to troubleshoot

You can attempt to resolve your error by going through the following steps:

- Check if you've covered all the prerequisites to use Inference Recommender. See the Inference Recommender Prerequisites.
- Check that you are able to deploy your model from Model Registry to an endpoint and that it can process your payloads without errors. See Deploy a Model from the Registry.
- When you kick off an Inference Recommender job, you should see endpoints being created in the console and you can review the CloudWatch logs.

Common errors

Review the following table for common Inference Recommender errors and their solutions.

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specify Domain in the Model Package version 1. Domain is a mandatory parameter for the job.</td>
<td>Make sure you provide the ML domain or OTHER if unknown.</td>
</tr>
<tr>
<td>Provided role ARN cannot be assumed and an AWSSecurityTokenServiceException error occurred.</td>
<td>Make sure the execution role provided has the necessary permissions specified in the prerequisites.</td>
</tr>
<tr>
<td>Specify Framework in the Model Package version 1. Framework is a mandatory parameter for the job.</td>
<td>Make sure you provide the ML Framework or OTHER if unknown.</td>
</tr>
<tr>
<td>Users at the end of prev phase is 0 while initial users of current phase is 1.</td>
<td>Users here refers to virtual users or threads used to send requests. Each phase starts with A users and ends with B users such that B &gt; A.</td>
</tr>
<tr>
<td>Error</td>
<td>Solution</td>
</tr>
<tr>
<td>--------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Between sequential phases, x_1 and x_2, we require that ( \text{abs}(x_2.A - x_1.B) \leq 3 ) and ( \geq 0 ).</td>
<td></td>
</tr>
<tr>
<td>Total Traffic duration (across) should not be more than Job duration.</td>
<td>The total duration of all your Phases cannot exceed the Job duration.</td>
</tr>
<tr>
<td>Burstable instance type ml.t2.medium is not allowed.</td>
<td>Inference Recommender doesn't support load testing on t2 instance family because burstable instances do not provide consistent performance.</td>
</tr>
<tr>
<td>ResourceLimitExceeded when calling CreateEndpoint operation</td>
<td>You have exceeded a SageMaker resource limit. For example, Inference Recommender might be unable to provision endpoints for benchmarking if the account has reached the endpoint quota. For more information about SageMaker limits and quotas, see Amazon SageMaker endpoints and quotas.</td>
</tr>
<tr>
<td>ModelError when calling InvokeEndpoint operation</td>
<td>A model error can happen for the following reasons:</td>
</tr>
<tr>
<td></td>
<td>• The invocation timed out while waiting for a response from the model container.</td>
</tr>
<tr>
<td></td>
<td>• The model couldn't process the input payload.</td>
</tr>
<tr>
<td>Error</td>
<td>Solution</td>
</tr>
<tr>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>PayloadError when calling InvokeEndpoint operation</td>
<td>A payload error can happen for following reasons:</td>
</tr>
<tr>
<td></td>
<td>• The payload source isn't in the Amazon S3 bucket.</td>
</tr>
<tr>
<td></td>
<td>• The payload is in a non-file object format.</td>
</tr>
<tr>
<td></td>
<td>• The payload is in an invalid file type. For example, a model expects an image type payload but is passed a text file.</td>
</tr>
<tr>
<td></td>
<td>• The payload is empty.</td>
</tr>
</tbody>
</table>

**Check CloudWatch**

When you kick off an Inference Recommender job, you should see endpoints being created in the console. Select one of the endpoints and view the CloudWatch logs to monitor for any 4xx/5xx errors. If you have a successful Inference Recommender job, you will be able to see the endpoint names as part of the results. Even if your Inference Recommender job is unsuccessful, you can still check the CloudWatch logs for the deleted endpoints by following the steps below:

1. Open the Amazon CloudWatch console at [https://console.aws.amazon.com/cloudwatch/](https://console.aws.amazon.com/cloudwatch/).
2. Select the Region in which you created the Inference Recommender job from the Region dropdown list in the top right.
3. In the navigation pane of CloudWatch, choose Logs, and then select Log groups.
4. Search for the log group called /aws/sagemaker/Endpoints/sm-epc-*. Select the log group based on your most recent Inference Recommender job.

You can also troubleshoot your job by checking the Inference Recommender CloudWatch logs. The Inference Recommender logs, which are published in the /aws/sagemaker/InferenceRecommendationsJobs CloudWatch log group, give a high level view on the progress of the job in the <jobName>/execution log stream. You can find detailed information on each of the endpoint configurations being tested in the <jobName>/Endpoint/<endpointName> log stream.

**Overview of the Inference Recommender log streams**
• `<jobName>`/execution contains overall job information such as endpoint configurations scheduled for benchmarking, compilation job skip reason, and validation failure reason.

• `<jobName>`/Endpoint/<endpointName> contains information such as resource creation progress, test configuration, load test stop reason, and resource cleanup status.

• `<jobName>`/CompilationJob/<compilationJobName> contains information on compilation jobs created by Inference Recommender, such as the compilation job configuration and compilation job status.

Create an alarm for Inference Recommender error messages

Inference Recommender outputs log statements for errors that might be helpful while troubleshooting. With a CloudWatch log group and a metric filter, you can look for terms and patterns in this log data as the data is sent to CloudWatch. Then, you can create a CloudWatch alarm based on the log group-metric filter. For more information, see Create a CloudWatch alarm based on a log group-metric filter.

Check benchmarks

When you kick off an Inference Recommender job, Inference Recommender creates several benchmarks to evaluate the performance of your model on different instance types. You can use the ListInferenceRecommendationsJobSteps API to view the details for all the benchmarks. If you have a failed benchmark, you can see the failure reasons as part of the results.

To use the ListInferenceRecommendationsJobSteps API, provide the following values:

• For JobName, provide the name of the Inference Recommender job.

• For StepType, use BENCHMARK to return details about the job's benchmarks.

• For Status, use FAILED to return details about only the failed benchmarks. For a list of the other status types, see the Status field in the ListInferenceRecommendationsJobSteps API.

```python
# Create a low-level SageMaker service client.
import boto3
aws_region = '<region>'
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Provide the job name for the SageMaker Inference Recommender job
job_name = '<job-name>'
```
# Filter for benchmarks
step_type = 'BENCHMARK'

# Filter for benchmarks that have a FAILED status
status = 'FAILED'

response = sagemaker_client.list_inference_recommendations_job_steps(
    JobName = job_name,
    StepType = step_type,
    Status = status
)

You can print the response object to view the results. The preceding code example stored the response in a variable called response:

```
print(response)
```

## Real-time inference

Real-time inference is ideal for inference workloads where you have real-time, interactive, low latency requirements. You can deploy your model to SageMaker hosting services and get an endpoint that can be used for inference. These endpoints are fully managed and support autoscaling (see [Automatically Scale Amazon SageMaker Models](#)).

### Topics
- Deploy models for real-time inference
- Invoke models for real-time inference
- Manage your endpoints
- Hosting options
- Automatically Scale Amazon SageMaker Models
- Host instance storage volumes
- Safely validate models in production
- Online Explainability with SageMaker Clarify
Deploy models for real-time inference

There are several options to deploy a model using SageMaker hosting services. You can interactively deploy a model with SageMaker Studio. Or, you can programmatically deploy a model using an AWS SDK, such as the SageMaker Python SDK or the SDK for Python (Boto3). You can also deploy by using the AWS CLI.

Before you begin

Before you deploy a SageMaker model, locate and make note of the following:

- The AWS Region where your Amazon S3 bucket is located
- The Amazon S3 URI path where the model artifacts are stored
- The IAM role for SageMaker
- The Docker Amazon ECR URI registry path for the custom image that contains the inference code, or the framework and version of a built-in Docker image that is supported and by AWS

For a list of AWS services available in each AWS Region, see Region Maps and Edge Networks. See Creating IAM roles for information on how to create an IAM role.

⚠️ Important

The Amazon S3 bucket where the model artifacts are stored must be in the same AWS Region as the model that you are creating.

Shared resource utilization with multiple models

You can deploy one or more models to an endpoint with Amazon SageMaker. When multiple models share an endpoint, they jointly utilize the resources that are hosted there, such as the ML compute instances, CPUs, and accelerators. The most flexible way to deploy multiple models to an endpoint is to define each model as an *inference component*.

Inference components

An inference component is a SageMaker hosting object that you can use to deploy a model to an endpoint. In the inference component settings, you specify the model, the endpoint, and how
the model utilizes the resources that the endpoint hosts. To specify the model, you can specify a SageMaker Model object, or you can directly specify the model artifacts and image.

In the settings, you can optimize resource utilization by tailoring how the required CPU cores, accelerators, and memory are allocated to the model. You can deploy multiple inference components to an endpoint, where each inference component contains one model and the resource utilization needs for that model.

After you deploy an inference component, you can directly invoke the associated model when you use the InvokeEndpoint action in the SageMaker API.

Inference components provide the following benefits:

**Flexibility**

The inference component decouples the details of hosting the model from the endpoint itself. This provides more flexibility and control over how models are hosted and served with an endpoint. You can host multiple models on the same infrastructure, and you can add or remove models from an endpoint as needed. You can update each model independently.

**Scalability**

You can specify how many copies of each model to host, and you can set a minimum number of copies to ensure that the model loads in the quantity that you require to serve requests. You can scale any inference component copy down to zero, which makes room for another copy to scale up.

SageMaker packages your models as inference components when you deploy them by using:

- SageMaker Studio Classic.
- The SageMaker Python SDK to deploy a Model object (where you set the endpoint type to `EndpointType.INFERENCE_COMPONENT_BASED`).
- The AWS SDK for Python (Boto3) to define `InferenceComponent` objects that you deploy to an endpoint.

**Deploy models with SageMaker Studio**

Complete the following steps to create and deploy your model interactively through SageMaker Studio. For more information about Studio, see the [Studio](#) documentation. For more walkthroughs
of various deployment scenarios, see the blog Package and deploy classical ML models and LLMs easily with Amazon SageMaker – Part 2.

**Prepare your artifacts and permissions**

Complete this section before creating a model in SageMaker Studio.

You have two options for bringing your artifacts and creating a model in Studio:

1. You can bring a pre-packaged tar.gz archive, which should include your model artifacts, any custom inference code, and any dependencies listed in a requirements.txt file.

2. SageMaker can package your artifacts for you. You only have to bring your raw model artifacts and any dependencies in a requirements.txt file, and SageMaker can provide default inference code for you (or you can override the default code with your own custom inference code). SageMaker supports this option for the following frameworks: PyTorch, XGBoost.

In addition to bringing your model, your AWS Identity and Access Management (IAM) role, and a Docker container (or desired framework and version for which SageMaker has a pre-built container), you must also grant permissions to create and deploy models through SageMaker Studio.

You should have the AmazonSageMakerFullAccess policy attached to your IAM role so that you can access SageMaker and other relevant services. To see the prices of the instance types in Studio, you also must attach the AWSPriceListServiceFullAccess policy (or if you don’t want to attach the whole policy, more specifically, the pricing:GetProducts action).

If you choose to upload your model artifacts when creating a model (or upload a sample payload file for inference recommendations), then you must create an Amazon S3 bucket. The bucket name must be prefixed by the word SageMaker. Alternate capitalizations of SageMaker are also acceptable: Sagemaker or sagemaker.

We recommend that you use the bucket naming convention sagemaker-\{Region\}-\{accountID\}. This bucket is used to store the artifacts that you upload.

After creating the bucket, attach the following CORS (cross-origin resource sharing) policy to the bucket:
You can attach a CORS policy to an Amazon S3 bucket by using any of the following methods:

- Through the [Edit cross-origin resource sharing (CORS)](https://console.aws.amazon.com/s3) page in the Amazon S3 console
- Using the Amazon S3 API [PutBucketCors](https://docs.aws.amazon.com/AmazonS3/latest/API/REST-Bucket-CORS.html)
- Using the put-bucket-cors AWS CLI command:

  ```bash
  aws s3api put-bucket-cors --bucket="..." --cors-configuration="..."
  ```

### Create a deployable model

In this step, you create a deployable version of your model in SageMaker by providing your artifacts along with additional specifications, such as your desired container and framework, any custom inference code, and network settings.

Create a deployable model in SageMaker Studio by doing the following:

1. Open the SageMaker Studio application.
2. In the left navigation pane, choose **Models**.
3. Choose the **Deployable models** tab.
4. On the **Deployable models** page, choose **Create**.
5. On the **Create deployable model** page, for the **Model name** field, enter a name for the model.

There are several more sections for you to fill out on the **Create deployable model** page.

The **Container definition** section looks like the following screenshot:
For the Container definition section, do the following:

1. For Container type, select Pre-built container if you'd like to use a SageMaker managed container, or select Bring your own container if you have your own container.

2. If you selected Pre-built container, select the Container framework, Framework version, and Hardware type that you'd like to use.

3. If you selected Bring your own container, enter an Amazon ECR path for ECR path to container image.

Then, fill out the Artifacts section, which looks like the following screenshot:
For the Artifacts section, do the following:

1. If you're using one of the frameworks that SageMaker supports for packaging model artifacts (PyTorch or XGBoost), then for **Artifacts**, you can choose the **Upload artifacts** option. With this option, you can simply specify your raw model artifacts, any custom inference code you have, and your requirements.txt file, and SageMaker handles packaging the archive for you. Do the following:

   a. For **Artifacts**, select **Upload artifacts** to continue providing your files. Otherwise, if you already have a tar.gz archive that contains your model files, inference code, and requirements.txt file, then select **Input S3 URI to pre-packaged artifacts**.

   b. If you chose to upload your artifacts, then for **S3 bucket**, enter the Amazon S3 path to a bucket where you'd like SageMaker to store your artifacts after packaging them for you. Then, complete the following steps.

   c. For **Upload model artifacts**, upload your model files.
d. For **Inference code**, select **Use default inference code** if you'd like to use default code that SageMaker provides for serving inference. Otherwise, select **Upload customized inference code** to use your own inference code.

e. For **Upload requirements.txt**, upload a text file that lists any dependencies that you want to install at runtime.

2. If you're not using a framework that SageMaker supports for packaging model artifacts, then Studio shows you the **Pre-packaged artifacts** option, and you must provide all of your artifacts already packaged as a `.tar.gz` archive. Do the following:

   a. For **Pre-packaged artifacts**, select **Input S3 URI for pre-packaged model artifacts** if you have your `.tar.gz` archive already uploaded to Amazon S3. Select **Upload pre-packaged model artifacts** if you want to directly upload your archive to SageMaker.

   b. If you selected **Input S3 URI for pre-packaged model artifacts**, enter the Amazon S3 path to your archive for **S3 URI**. Otherwise, select and upload the archive from your local machine.

The next section is **Security**, which looks like the following screenshot:

![Security](image)

**For the Security section, do the following:**

1. For **IAM role**, enter the ARN for an IAM role.

2. (Optional) For **Virtual Private Cloud (VPC)**, you can select an Amazon VPC for storing your model configuration and artifacts.
3. (Optional) Turn on the **Network isolation** toggle if you want to restrict your container's internet access.

Finally, you can optionally fill out the **Advanced options** section, which looks like the following screenshot:

![Advanced options section](image)

(Optional) For the Advanced options section, do the following:

1. Turn on the **Customized instance recommendations** toggle if you want to run an Amazon SageMaker Inference Recommender job on your model after its creation. Inference Recommender is a feature that provides you with recommended instance types for optimizing inference performance and cost. You can view these instance recommendations when preparing to deploy your model.

2. For **Add environment variables**, enter an environment variables for your container as key-value pairs.

3. For **Tags**, enter any tags as key-value pairs.

4. After finishing your model and container configuration, choose **Create deployable model**.
You should now have a model in SageMaker Studio that is ready for deployment.

**Deploy your model**

Finally, you deploy the model you configured in the previous step to an HTTPS endpoint. You can deploy either a single model or multiple models to the endpoint.

One way to deploy a model is by doing the following in Studio:

1. Open the SageMaker Studio application.
2. In the left navigation pane, choose **Models**.
3. On the **Models** page, select one or more models from the list of SageMaker models.
4. Choose **Deploy**.
5. For **Endpoint name**, open the dropdown menu. You can either select an existing endpoint or you can create a new endpoint to which you deploy the model.
6. For **Instance type**, select the instance type that you want to use for the endpoint. If you previously ran an Inference Recommender job for the model, your recommended instance types appear in the list under the title **Recommended**. Otherwise, you'll see a few **Prospective instances** that might be suitable for your model.
7. For **Initial instance count**, enter the initial number of instances that you'd like to provision for your endpoint.
8. If the model you're deploying is one of the most used SageMaker JumpStart LLMs from the model hub, then the **Alternate configurations** option appears after the instance type and instance count fields.

For the most popular SageMaker JumpStart LLMs, AWS has pre-benchmarked instance types to optimize for either cost or performance. This data can help you decide which instance type to use for deploying your LLM. Choose **Alternate configurations** to open a dialog box that contains the pre-benchmarked data. The panel looks like the following screenshot:
In the **Alternate configurations** box, do the following:

a. Select an instance type. You can choose **Cost per hour** or **Best performance** to see instance types that optimize either cost or performance for the specified model. You can also choose **Other supported instances** to see a list of other instance types that are compatible with the SageMaker JumpStart model. Note that selecting an instance type here overwrites any previous instance selection specified in Step 6.

b. (Optional) Turn on the **Customize the selected configuration** toggle to specify **Max total tokens** (the maximum number of tokens that you want to allow, which is the sum of your input tokens and the model's generated output), **Max input token length** (the maximum number of tokens you want to allow for the input of each request), and **Max concurrent requests** (the maximum number of requests that the model can process at a time).

c. Choose **Select** to confirm your instance type and configuration settings.

9. The **Model** field should already be populated with the name of the model or models that you're deploying. You can choose **Add model** to add more models to the deployment. For each model that you add, fill out the following fields:

   a. For **Number of CPU cores**, enter the CPU cores that you'd like to dedicate for the model's usage.
b. For **Min number of copies**, enter the minimum number of model copies that you want to have hosted on the endpoint at any given time.

c. For **Min CPU memory (MB)**, enter the minimum amount of memory (in MB) that the model requires.

d. For **Max CPU memory (MB)**, enter the maximum amount of memory (in MB) that you'd like to allow the model to use.

10. (Optional) For the **Advanced options**, do the following:

   a. For **IAM role**, use either the default SageMaker IAM execution role, or specify your own role that has the permissions you need. Note that this IAM role must be the same as the role that you specified when creating the deployable model.

   b. For **Virtual Private Cloud (VPC)**, you can specify a VPC in which you want to host your endpoint.

   c. For **Encryption KMS key**, select an AWS KMS key to encrypt data on the storage volume attached to the ML compute instance that hosts the endpoint.

   d. Turn on the **Enable network isolation** toggle to restrict your container's internet access.

   e. For **Timeout configuration**, enter values for the **Model data download timeout (seconds)** and **Container startup health check timeout (seconds)** fields. These values determine the maximum amount of time that SageMaker allows for downloading the model to the container and starting up the container, respectively.

   f. For **Tags**, enter any tags as key-value pairs.

After configuring your options, the page should look like the following screenshot.
After configuring your deployment, choose **Deploy** to create the endpoint and deploy your model.

**Deploy models with the Python SDKs**

Using the SageMaker Python SDK, you can build your model in two ways. The first is to create a model object from the `Model` or `ModelBuilder` class. If you use the `Model` class to create your `Model` object, you need to specify the model package or inference code (depending on your model server), scripts to handle serialization and deserialization of data between the client and server, and any dependencies to be uploaded to Amazon S3 for consumption. The second way to build your model is to use `ModelBuilder` for which you provide model artifacts or inference code. `ModelBuilder` automatically captures your dependencies, infers the needed serialization and deserialization functions, and packages your dependencies to create your `Model` object. For more information about `ModelBuilder`, see [Create a model in Amazon SageMaker with ModelBuilder](#).

The following section describes both methods to create your model and deploy your model object.

**Set up**

The following examples prepare for the model deployment process. They import the necessary libraries and define the S3 URL that locates the model artifacts.
SageMaker Python SDK

**Example import statements**

The following example imports modules from the SageMaker Python SDK, the SDK for Python (Boto3), and the Python Standard Library. These modules provide useful methods that help you deploy models, and they're used by the remaining examples that follow.

```python
import boto3
from datetime import datetime
from sagemaker.compute_resource_requirements.resource_requirements import ResourceRequirements
from sagemaker.predictor import Predictor
from sagemaker.enums import EndpointType
from sagemaker.model import Model
from sagemaker.session import Session
```

**boto3 inference components**

**Example import statements**

The following example imports modules from the SDK for Python (Boto3) and the Python Standard Library. These modules provide useful methods that help you deploy models, and they're used by the remaining examples that follow.

```python
import boto3
import botocore
import sys
import time
```

**boto3 models (without inference components)**

**Example import statements**

The following example imports modules from the SDK for Python (Boto3) and the Python Standard Library. These modules provide useful methods that help you deploy models, and they're used by the remaining examples that follow.

```python
import boto3
import botocore
```
import datetime
from time import gmtime, strftime

Example model artifact URL

The following code builds an example Amazon S3 URL. The URL locates the model artifacts for a pre-trained model in an Amazon S3 bucket.

```python
# Create a variable w/ the model S3 URL

# The name of your S3 bucket:
s3_bucket = "DOC-EXAMPLE-BUCKET"
# The directory within your S3 bucket your model is stored in:
bucket_prefix = "sagemaker/model/path"
# The file name of your model artifact:
model_filename = "my-model-artifact.tar.gz"
# Relative S3 path:
model_s3_key = f"{bucket_prefix}/"+model_filename
# Combine bucket name, model file name, and relate S3 path to create S3 model URL:
model_url = f"s3://{s3_bucket}/{model_s3_key}"
```

The full Amazon S3 URL is stored in the variable `model_url`, which is used in the examples that follow.

Overview

There are multiple ways that you can deploy models with the SageMaker Python SDK or the SDK for Python (Boto3). The following sections summarize the steps that you complete for several possible approaches. These steps are demonstrated by the examples that follow.

SageMaker Python SDK

Using the SageMaker Python SDK, you can build your model in either of the following ways:

- **Create a model object from the Model class** – You must specify the model package or inference code (depending on your model server), scripts to handle serialization and deserialization of data between the client and server, and any dependencies to be uploaded to Amazon S3 for consumption.
- **Create a model object from the ModelBuilder class** – You provide model artifacts or inference code, and ModelBuilder automatically captures your dependencies, infers the
needed serialization and deserialization functions, and packages your dependencies to create your Model object.

For more information about ModelBuilder, see [Create a model in Amazon SageMaker with ModelBuilder](https://docs.aws.amazon.com/sagemaker/latest/dg/modelbuilder-create-model.html). You can also see the blog [Package and deploy classical ML models and LLMs easily with SageMaker – Part 1](https://aws.amazon.com/blogs/machine-learning/package-and-deploy-classical-ml-models-and-llms-easily-with-sagemaker-part-1/) for more information.

The examples that follow describe both methods to create your model and deploy your model object. To deploy a model in these ways, you complete the following steps:

1. Define the endpoint resources to allocate to the model with a ResourceRequirements object.
2. Create a model object from the Model or ModelBuilder classes. The ResourceRequirements object is specified in the model settings.
3. Deploy the model to an endpoint by using the deploy method of the Model object.

**boto3 inference components**

The examples that follow demonstrate how to assign a model to an inference component and then deploy the inference component to an endpoint. To deploy a model in this way, you complete the following steps:

1. (Optional) Create a SageMaker model object by using the create_model method.
2. Specify the settings for your endpoint by creating an endpoint configuration object. To create one, you use the create_endpoint_config method.
3. Create your endpoint by using the create_endpoint method, and in your request, provide the endpoint configuration that you created.
4. Create an inference component by using the create_inference_component method. In the settings, you specify a model by doing either of the following:
   - Specifying a SageMaker model object
   - Specifying the model image URI and S3 URL

You also allocate endpoint resources to the model. By creating the inference component, you deploy the model to the endpoint. You can deploy multiple models to an endpoint by creating multiple inference components — one for each model.
The examples that follow demonstrate how to create a model object and then deploy the model to an endpoint. To deploy a model in this way, you complete the following steps:

1. Create a SageMaker model by using the `create_model` method.

2. Specify the settings for your endpoint by creating an endpoint configuration object. To create one, you use the `create_endpoint_config` method. In the endpoint configuration, you assign the model object to a production variant.

3. Create your endpoint by using the `create_endpoint` method. In your request, provide the endpoint configuration that you created.

   When you create the endpoint, SageMaker provisions the endpoint resources, and it deploys the model to the endpoint.

**Configure**

The following examples configure the resources that you require to deploy a model to an endpoint.

**SageMaker Python SDK**

The following example assigns endpoint resources to a model with a `ResourceRequirements` object. These resources include CPU cores, accelerators, and memory. Then, the example creates a model object from the `Model` class. Alternatively you can create a model object by instantiating the `ModelBuilder` class and running `build`—this method is also shown in the example. `ModelBuilder` provides a unified interface for model packaging, and in this instance, prepares a model for a large model deployment. The example utilizes `ModelBuilder` to construct a Hugging Face model. (You can also pass a JumpStart model). Once you build the model, you can specify resource requirements in the model object. In the next step, you use this object to deploy the model to an endpoint.

```python
resources = ResourceRequirements(
    requests = {
        "num_cpus": 2,  # Number of CPU cores required:
        "num_accelerators": 1,  # Number of accelerators required
        "memory": 8192,  # Minimum memory required in Mb (required)
        "copies": 1,
    },
    limits = {},
```
```python
now = datetime.now()
dt_string = now.strftime("%d-%m-%Y-%H-%M-%S")
model_name = "my-sm-model"+dt_string

# build your model with Model class
model = Model(
    name = "model-name",
    image_uri = "image-uri",
    model_data = model_url,
    role = "arn:aws:iam::111122223333:role/service-role/role-name",
    resources = resources,
    predictor_cls = Predictor,
)

# Alternate mechanism using ModelBuilder
# uncomment the following section to use ModelBuilder
/*@ 
model_builder = ModelBuilder(
    model="<HuggingFace-ID>", # like "meta-llama/Llama-2-7b-hf"
    schema_builder=SchemaBuilder(sample_input,sample_output),
    env_vars={ "HUGGING_FACE_HUB_TOKEN": "<HuggingFace_token>" } 
)
*/

# build your Model object
model = model_builder.build()

# create a unique name from string 'mb-inference-component'
model.model_name = unique_name_from_base("mb-inference-component")

# assign resources to your model
model.resources = resources
*/
```

boto3 inference components

The following example configures an endpoint with the `create_endpoint_config` method. You assign this configuration to an endpoint when you create it. In the configuration, you define one or more production variants. For each variant, you can choose the instance type that you want Amazon SageMaker to provision, and you can enable managed instance scaling.

```python
endpoint_config_name = "endpoint-config-name"
```
```python
endpoint_name = "endpoint-name"
inference_component_name = "inference-component-name"
variant_name = "variant-name"

sagemaker_client.create_endpoint_config(
    EndpointConfigName = endpoint_config_name,
    ExecutionRoleArn = "arn:aws:iam::111122223333:role/service-role/role-name",
    ProductionVariants = [
        {
            "VariantName": variant_name,
            "InstanceType": "ml.p4d.24xlarge",
            "InitialInstanceCount": 1,
            "ManagedInstanceScaling": {
                "Status": "ENABLED",
                "MinInstanceCount": 1,
                "MaxInstanceCount": 2,
            },
        },
    ],
)
```

boto3 models (without inference components)

**Example model definition**

The following example defines a SageMaker model with the `create_model` method in the AWS SDK for Python (Boto3).

```python
model_name = "model-name"

create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = "arn:aws:iam::111122223333:role/service-role/role-name",
    PrimaryContainer = {
        "Image": "image-uri",
        "ModelDataUrl": model_url,
    }
)
```

This example specifies the following:

- ModelName: A name for your model (in this example it is stored as a string variable called model_name).
• **ExecutionRoleArn**: The Amazon Resource Name (ARN) of the IAM role that Amazon SageMaker can assume to access model artifacts and Docker images for deployment on ML compute instances or for batch transform jobs.

• **PrimaryContainer**: The location of the primary Docker image containing inference code, associated artifacts, and custom environment maps that the inference code uses when the model is deployed for predictions.

### Example endpoint configuration

The following example configures an endpoint with the `create_endpoint_config` method. Amazon SageMaker uses this configuration to deploy models. In the configuration, you identify one or more models, created with the `create_model` method, to deploy the resources that you want Amazon SageMaker to provision.

```python
endpoint_config_response = sagemaker_client.create_endpoint_config(
    EndpointConfigName = "endpoint-config-name",
    # List of ProductionVariant objects, one for each model that you want to host at this endpoint:
    ProductionVariants = [
        {
            "VariantName": "variant-name", # The name of the production variant.
            "ModelName": model_name,
            "InstanceType": "ml.p4d.24xlarge",
            "InitialInstanceCount": 1 # Number of instances to launch initially.
        }
    ]
)
```

This example specifies the following keys for the `ProductionVariants` field:

• **VariantName**: The name of the production variant.

• **ModelName**: The name of the model that you want to host. This is the name that you specified when creating the model.

• **InstanceType**: The compute instance type. See the `InstanceType` field in [https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_ProductionVariant.html](https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_ProductionVariant.html) and [SageMaker Pricing](https://docs.aws.amazon.com/sagemaker/latest/APIReference/API_ProductionVariant.html) for a list of supported compute instance types and pricing for each instance type.
Deploy

The following examples deploy a model to an endpoint.

SageMaker Python SDK

The following example deploys the model to a real-time, HTTPS endpoint with the deploy method of the model object. If you specify a value for the resources argument for both model creation and deployment, the resources you specify for deployment take precedence.

```python
predictor = model.deploy(
    initial_instance_count = 1,
    instance_type = "ml.p4d.24xlarge",
    endpoint_type = EndpointType.INFERENCE_COMPONENT_BASED,
    resources = resources,
)
```

For the `instance_type` field, the example specifies the name of the Amazon EC2 instance type for the model. For the `initial_instance_count` field, it specifies the initial number of instances to run the endpoint on.

The following code sample demonstrates another case where you deploy a model to an endpoint and then deploy another model to the same endpoint. In this case you must supply the same endpoint name to the deploy methods of both models.

```python
# Deploy the model to inference-component-based endpoint
falcon_predictor = falcon_model.deploy(
    initial_instance_count = 1,
    instance_type = "ml.p4d.24xlarge",
    endpoint_type = EndpointType.INFERENCE_COMPONENT_BASED,
    endpoint_name = "<endpoint_name>"
    resources = resources,
)

# Deploy another model to the same inference-component-based endpoint
llama2_predictor = llama2_model.deploy( # resources already set inside llama2_model
    endpoint_type = EndpointType.INFERENCE_COMPONENT_BASED,
    endpoint_name = "<endpoint_name>"  # same endpoint name as for falcon model
)
boto3 inference components

Once you have an endpoint configuration, use the create_endpoint method to create your endpoint. The endpoint name must be unique within an AWS Region in your AWS account.

The following example creates an endpoint using the endpoint configuration specified in the request. Amazon SageMaker uses the endpoint to provision resources.

```python
sagemaker_client.create_endpoint(  
    EndpointName = endpoint_name,  
    EndpointConfigName = endpoint_config_name,  
)
```

After you've created an endpoint, you can deploy one or models to it by creating inference components. The following example creates one with the create_inference_component method.

```python
sagemaker_client.create_inference_component(  
    InferenceComponentName = inference_component_name,  
    EndpointName = endpoint_name,  
    VariantName = variant_name,  
    Specification = {  
        "Container": {  
            "Image": "image-uri",  
            "ArtifactUrl": model_url,  
        },  
        "ComputeResourceRequirements": {  
            "NumberOfCpuCoresRequired": 1,  
            "MinMemoryRequiredInMb": 1024  
        }  
    },  
    RuntimeConfig = {"CopyCount": 2}  
)
```

boto3 models (without inference components)

**Example deployment**

Provide the endpoint configuration to SageMaker. The service launches the ML compute instances and deploys the model or models as specified in the configuration.
Once you have your model and endpoint configuration, use the `create_endpoint` method to create your endpoint. The endpoint name must be unique within an AWS Region in your AWS account.

The following example creates an endpoint using the endpoint configuration specified in the request. Amazon SageMaker uses the endpoint to provision resources and deploy models.

```python
create_endpoint_response = sagemaker_client.create_endpoint(
    # The endpoint name must be unique within an AWS Region in your AWS account:
    EndpointName = "endpoint-name",
    # The name of the endpoint configuration associated with this endpoint:
    EndpointConfigName = "endpoint-config-name"
)
```

**Deploy models with the AWS CLI**

You can deploy a model to an endpoint by using the AWS CLI.

**Overview**

When you deploy a model with the AWS CLI, you can deploy it with or without using an inference component. The following sections summarize the commands that you run for both approaches. These commands are demonstrated by the examples that follow.

**With inference components**

To deploy a model with an inference component, do the following:

1. (Optional) Create a model with the `create-model` command.
2. Specify the settings for your endpoint by creating an endpoint configuration. To create one, you run the `create-endpoint-config` command.
3. Create your endpoint by using the `create-endpoint` command. In the command body, specify the endpoint configuration that you created.
4. Create an inference component by using the `create-inference-component` command. In the settings, you specify a model by doing either of the following:
   - Specifying a SageMaker model object
   - Specifying the model image URI and S3 URL
You also allocate endpoint resources to the model. By creating the inference component, you deploy the model to the endpoint. You can deploy multiple models to an endpoint by creating multiple inference components — one for each model.

Without inference components

To deploy a model without using an inference component, do the following:

1. Create a SageMaker model by using the `create-model` command.
2. Specify the settings for your endpoint by creating an endpoint configuration object. To create one, you use the `create-endpoint-config` command. In the endpoint configuration, you assign the model object to a production variant.
3. Create your endpoint by using the `create-endpoint` command. In your command body, specify the endpoint configuration that you created.

When you create the endpoint, SageMaker provisions the endpoint resources, and it deploys the model to the endpoint.

Configure

The following examples configure the resources that you require to deploy a model to an endpoint.

With inference components

**Example create-endpoint-config command**

The following example creates an endpoint configuration with the `create-endpoint-config` command.

```bash
aws sagemaker create-endpoint-config \
--endpoint-config-name endpoint-config-name \
--execution-role-arn arn:aws:iam::111122223333:role/service-role/role-name\ 
--production-variants file://production-variants.json
```

In this example, the file `production-variants.json` defines a production variant with the following JSON:

```json
[ ]
```
If the command succeeds, the AWS CLI responds with the ARN for the resource you created.

```json
{
}
```

**Without inference components**

**Example create-model command**

The following example creates a model with the `create-model` command.

```
aws sagemaker create-model \
--model-name model-name \
--execution-role-arn arn:aws:iam::111122223333:role/service-role/role-name \
--primary-container "{ "Image": "image-uri", "ModelDataUrl": "model-s3-url" }"
```

If the command succeeds, the AWS CLI responds with the ARN for the resource you created.

```json
{
}
```

**Example create-endpoint-config command**

The following example creates an endpoint configuration with the `create-endpoint-config` command.

```
aws sagemaker create-endpoint-config \
--endpoint-config-name endpoint-config-name \
```
In this example, the file `production-variants.json` defines a production variant with the following JSON:

```
[
  {
    "VariantName": "variant-name",
    "ModelName": "model-name",
    "InstanceType": "ml.p4d.24xlarge",
    "InitialInstanceCount": 1
  }
]
```

If the command succeeds, the AWS CLI responds with the ARN for the resource you created.

```
{
}
```

## Deploy

The following examples deploy a model to an endpoint.

### With inference components

#### Example create-endpoint command

The following example creates an endpoint with the `create-endpoint` command.

```
aws sagemaker create-endpoint \
  --endpoint-name endpoint-name \
  --endpoint-config-name endpoint-config-name
```

If the command succeeds, the AWS CLI responds with the ARN for the resource you created.

```
{
  "EndpointArn": "arn:aws:sagemaker:us-west-2:111122223333:endpoint/endpoint-name"
}
```
Example create-inference-component command

The following example creates an inference component with the `create-inference-component` command.

```bash
aws sagemaker create-inference-component \
  --inference-component-name inference-component-name \
  --endpoint-name endpoint-name \
  --variant-name variant-name \
  --specification file://specification.json \
  --runtime-config "{"CopyCount": 2}"
```

In this example, the file `specification.json` defines the container and compute resources with the following JSON:

```json
{
  "Container": {
    "Image": "image-uri",
    "ArtifactUrl": "model-s3-url"
  },
  "ComputeResourceRequirements": {
    "NumberOfCpuCoresRequired": 1,
    "MinMemoryRequiredInMb": 1024
  }
}
```

If the command succeeds, the AWS CLI responds with the ARN for the resource you created.

```json
{
}
```

Without inference components

Example create-endpoint command

The following example creates an endpoint with the `create-endpoint` command.

```bash
aws sagemaker create-endpoint \
  --endpoint-name endpoint-name \
  --endpoint-config-name endpoint-config-name
```
If the command succeeds, the AWS CLI responds with the ARN for the resource you created.

```json
{
  "EndpointArn": "arn:aws:sagemaker:us-west-2:111122223333:endpoint/endpoint-name"
}
```

**Invoke models for real-time inference**

After you deploy your model using SageMaker hosting services, you can test your model on that endpoint by sending it test data. You can test your endpoints using Amazon SageMaker Studio, the AWS SDKs, or the AWS CLI.

**Invoke Your Endpoint Using Amazon SageMaker Studio**

After you deploy your model to an endpoint, you can view the endpoint through Amazon SageMaker Studio and test your endpoint by sending single inference requests.

**Note**

SageMaker only supports endpoint testing in Studio for real-time endpoints.

**To send a test inference request to your endpoint**

1. Launch Amazon SageMaker Studio.
2. In the navigation pane on the left, choose **Deployments**.
3. From the dropdown, choose **Endpoints**.
4. Find for your endpoint by name, and choose the name in the table. The endpoint names listed in the **Endpoints** panel are defined when you deploy a model. The Studio workspace opens the **Endpoint** page in a new tab.
5. Choose the **Test inference** tab.
6. For **Testing Options**, select one of the following:

   a. Select **Test the sample request** to immediately send a request to your endpoint. Use the **JSON editor** to provide sample data in JSON format, and choose **Send Request** to submit the request to your endpoint. After submitting your request, Studio shows the inference output in a card to the right of the JSON editor.
b. Select **Use Python SDK example code** to view the code for sending a request to the endpoint. Then, copy the code example from the **Example inference request** section and run the code from your testing environment.

The top of the card shows the type of request that was sent to the endpoint (only JSON is accepted). The card shows the following fields:

- **Status** – displays one of the following status types:
  - **Success** – The request succeeded.
  - **Failed** – The request failed. A response appears under **Failure Reason**.
  - **Pending** – While the inference request is pending, the status shows a spinning, circular icon.
- **Execution Length** – How long the invocation took (end time minus the start time) in milliseconds.
- **Request Time** – How many minutes have passed since the request was sent.
- **Result Time** – How many minutes have passed since the result was returned.

**Invoke Your Endpoint by Using the AWS SDK for Python (Boto3)**

After you deploy your model to an endpoint you can check your endpoint by using one of the AWS SDKs, including as the AWS SDK for Python (Boto3). To test your endpoint with this SDK, you use one of the following methods:

- **invoke_endpoint** – Sends an inference request to a model endpoint and returns the response that the model generates. This method returns the inference payload as one response after the model finishes generating it. For more information, see **invoke_endpoint** in the **AWS SDK for Python (Boto3) API Reference**.

- **invoke_endpoint_with_response_stream** – Sends an inference request to a model endpoint and streams the response in incremental parts while the model generates the inference. With this method, your client application immediately starts receiving parts of the response as the parts become available. Your client doesn't need to wait for the model to generate the whole response payload. You can implement streaming to support fast interactive experiences, such as chatbots, virtual assistants, and music generators.

  Use this method only to invoke models that support inference streaming.
Amazon SageMaker

Developer Guide

When a container handles a streaming inference request, it returns the model's inference
as a series of parts incrementally as the model generates them. Client applications start
receiving responses immediately when they're available. They don't need to wait for the model
to generate the entire response. You can implement streaming to support fast interactive
experiences, such as chatbots, virtual assistants, and music generators.
Before you can use these methods in your client code, you must create a SageMaker Runtime client,
and you must specify the name of your endpoint. The following example sets up the client and
endpoint for the rest of the examples that follow:
import boto3
# Create a low-level client representing Amazon SageMaker Runtime
sagemaker_runtime = boto3.client(
"sagemaker-runtime", region_name='aws_region')
# The endpoint name must be unique within
# an AWS Region in your AWS account.
endpoint_name='endpoint-name'

Invoke to Get an Inference Response
The following example uses the invoke_endpoint method to invoke an endpoint with the AWS
SDK for Python (Boto3):
# Gets inference from the model hosted at the specified endpoint:
response = sagemaker_runtime.invoke_endpoint(
EndpointName=endpoint_name,
Body=bytes('{"features": ["This is great!"]}', 'utf-8')
)
# Decodes and prints the response body:
print(response['Body'].read().decode('utf-8'))

This example provide input data in the Body ﬁeld for SageMaker to pass to the model. This data
must be in the same format that was used for training. The example stores the response in the
response variable.

Invoke models

4090


Amazon SageMaker

Developer Guide

The response variable provides access to the HTTP status, the name of the deployed model, and
other ﬁelds. The following snippet prints the HTTPStatusCode:
print(response["HTTPStatusCode"])

Invoke to Stream an Inference Response
If you deployed a model that supports inference streaming, you can invoke the model to receive its
inference payload as a stream of parts. The model delivers these parts incrementally as the model
generates them. When an application receives an inference stream, the application doesn't need to
wait for the model to generate the whole response payload. Instead, the application immediately
starts receiving parts of the response as they become available.
By consuming an inference stream in your application, you can create interactions where your users
perceive the inference to be fast because they get the ﬁrst part immediately. For example, you
could create a chatbot that incrementally shows the text generated by a large language model
(LLM).
To get an inference stream, you can use the invoke_endpoint_with_response_stream
method in the SDK for Python (Boto3). In the response body, the SDK provides an EventStream
object, which gives the inference as a series of PayloadPart objects.
Example Inference Stream
The following example is a stream of PayloadPart objects:
{'PayloadPart': {'Bytes': b'{"outputs": [" a"]}\n'}}
{'PayloadPart': {'Bytes': b'{"outputs": [" challenging"]}\n'}}
{'PayloadPart': {'Bytes': b'{"outputs": [" problem"]}\n'}}
. . .

In each payload part, the Bytes ﬁeld provides a portion of the inference response from the model.
This portion can be any content type that a model generates, such as text, image, or audio data. In
this example, the portions are JSON objects that contain generated text from an LLM.
Usually, the payload part contains a discrete chunk of data from the model. In this example, the
discrete chunks are whole JSON objects. Occasionally, the streaming response splits the chunks
over multiple payload parts, or it combines multiple chunks into one payload part. The following
example shows a chunk of data in JSON format that's split over two payload parts:
Invoke models

4091


When you write application code that processes an inference stream, include logic that handles these occasional splits and combinations of data. As one strategy, you could write code that concatenates the contents of Bytes while your application receives the payload parts. By concatenating the example JSON data here, you would combine the data into a newline-delimeted JSON body. Then, your code could process the stream by parsing the whole JSON object on each line.

The following example shows the newline-delimeted JSON that you would create when you concatenate the example contents of Bytes:

```
{"outputs": [" a"]}
{"outputs": [" challenging"]}
{"outputs": [" problem"]}
...
```

**Example Code to Process an Inference Stream**

The following example Python class, `SmrInferenceStream`, demonstrates how you can process an inference stream that sends text data in JSON format:

```python
import io
import json

# Example class that processes an inference stream:
class SmrInferenceStream:
    def __init__(self, sagemaker_runtime, endpoint_name):
        self.sagemaker_runtime = sagemaker_runtime
        self.endpoint_name = endpoint_name
        # A buffered I/O stream to combine the payload parts:
        self.buff = io.BytesIO()
        self.read_pos = 0

    def stream_inference(self, request_body):
        # Gets a streaming inference response
        # from the specified model endpoint:
        response = self.sagemaker_runtime\
          .invoke_endpoint_with_response_stream(
            endpoint_name=self.endpoint_name,
            data=request_body
        )
        
        # Process the response stream:
        while True:
            chunk = response.read()
            if chunk == b'' or chunk == None:
                break
            self.buff.write(chunk)
```

This example processes the inference stream by doing the following:

- Gets initialized with a SageMaker Runtime client and the name of a model endpoint. Before you can get an inference stream, the model that the endpoint hosts must support inference streaming.
- In the example `stream_inference` method, receives a request body and passes it to the `invoke_endpoint_with_response_stream` method of the SDK.
- Iterates over each event in the `EventStream` object that the SDK returns.
- From each event, gets the contents of the `Bytes` object in the `PayloadPart` object.
- In the example `_write` method, writes to a buffer to concatenate the contents of the `Bytes` objects. The combined contents form a newline-delimited JSON body.
• Uses the example _readlines method to get an iterable series of JSON objects.
• In each JSON object, gets a piece of the inference.
• With the yield expression, returns the pieces incrementally.

The following example creates and uses a SmrInferenceStream object:

```python
request_body = {"inputs": ["Large model inference is"],
                "parameters": {"max_new_tokens": 100,
                               "enable_sampling": true}}
smr_inference_stream = SmrInferenceStream(sagemaker_runtime, endpoint_name)
stream = smr_inference_stream.stream_inference(request_body)
for part in stream:
    print(part, end='')
```

This example passes a request body to the stream_inference method. It iterates over the response to print each piece that the inference stream returns.

The example assumes that the model at the specified endpoint is an LLM that generates text. The output from this example is a body of generated text that prints incrementally:

```
a challenging problem in machine learning. The goal is to . . .
```

**Invoke Your Endpoint by Using the AWS CLI**

You can test your endpoint by running commands with the AWS Command Line Interface (AWS CLI). The AWS CLI supports standard inference requests with the `invoke-endpoint` command, and it supports asynchronous inference requests with the `invoke-endpoint-async` command.

**Note**
The AWS CLI doesn't support streaming inference requests.

The following example uses the `invoke-endpoint` command to send an inference request to a model endpoint:

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name endpoint_name \
```
For the **--endpoint-name** parameter, provide the name you specified for EndpointName when you created your endpoint with CreateEndpoint. For the **--body** parameter, provide input data for SageMaker to pass to the model. The data must be in the same format that was used for training. This example shows how to send binary data to your endpoint.

For more information on when to use `file://` over `fileb://` when passing the contents of a file to a parameter of the AWS CLI, see [Best Practices for Local File Parameters](#).

For more information, and to see additional parameters that you can pass, see `invoke-endpoint` in the [AWS CLI Command Reference](#).

If the `invoke-endpoint` command succeeds it returns a response such as the following:

```
{
  "ContentType": "<content_type>; charset=utf-8",
  "InvokedProductionVariant": "<Variant>"
}
```

If the command doesn't succeed, check whether the input payload is in the correct format.

View the output of the invocation by checking the file output file (`output_file.txt` in this example).

```
more output_file.txt
```

### Manage your endpoints

After deploying your model to an endpoint, you might want to view and manage the endpoint. With SageMaker, you can view the status and details of your endpoint, check metrics and logs to monitor your endpoint's performance, update the models deployed to your endpoint, and more.

The following page describes how to interactively view and make changes to your endpoints using the Amazon SageMaker console or SageMaker Studio.

**Manage endpoints in SageMaker Studio**

In Amazon SageMaker Studio, you can view and manage your SageMaker Hosting endpoints. To learn more about Studio, see [Amazon SageMaker Studio](#).
To find the list of your endpoints in SageMaker Studio do the following:

1. Open the Studio application.
2. In the left navigation pane, choose **Deployments**.
3. From the dropdown menu, choose **Endpoints**.

The **Endpoints** page opens, which lists all of your SageMaker Hosting endpoints. From this page, you can see the endpoints and their **Status**. You can also create a new endpoint, edit an existing endpoint, or delete an endpoint.

To see the details for a specific endpoint, choose an endpoint from the list. On the endpoint's details page, you get an overview like the following screenshot.

![Endpoint summary screenshot](image)

Each endpoint details page contains the following tabs of information:

**Variants (or Models)**

The **Variants** tab (also called the **Models** tab if your endpoint has multiple models deployed) shows you the list of model variants or models currently deployed to your endpoint. The following screenshot shows you what the overview and **Models** section looks like for an endpoint with multiple models deployed.
You can add or edit the settings for each variant or model. You can also select a variant and enable a default auto-scaling policy, which you can edit later in the Auto-scaling tab.

**Settings**

On the Settings tab, you can view the endpoint’s associated AWS IAM role, the AWS KMS key used for encryption (if applicable), the name of your VPC, and the network isolation settings.

**Test inference**

On the Test inference tab, you can send a test inference request to a deployed model. This is useful if you’d like to verify that your endpoint responds to requests as expected.

To test inference, do the following:

1. On the model's Test inference tab, choose one of the following options:
   a. Select Enter the request body if you’d like to test the endpoint and receive a response through the Studio interface.
   b. Select Copy example code (Python) if you’d like to copy an AWS SDK for Python (Boto3) example that you can use to invoke your endpoint from a local environment and receive a response programmatically.
2. For Model, select the model that you want to test on the endpoint.
3. If you chose the Studio interface testing method, then you can also choose your desired Content type for the response from the dropdown.

After configuring your request, then you can either choose Send request (to receive a response through the Studio interface) or Copy to copy the Python example.
Auto-scaling

On the **Auto-scaling** tab, you can view any auto-scaling policies configured for the models hosted on your endpoint. The following screenshot shows you the **Auto-scaling** tab.

You can choose **Edit auto-scaling** to change any of the policies and turn on or turn off the default auto-scaling policy.

To learn more about auto-scaling for real-time endpoints, see [Automatically Scale Amazon SageMaker Models](https://docs.aws.amazon.com/sagemaker/latest/dg/automatically-scale.html). If you’re not sure how to configure an auto-scaling policy for your endpoint,
you can use an Inference Recommender autoscaling recommendations job to get recommendations for an auto-scaling policy.

**Manage endpoints in the SageMaker console**

To view your endpoints in the SageMaker console, do the following:

2. In the left navigation pane, choose **Inference**.
3. From the dropdown list, choose **Endpoints**.
4. On the **Endpoints** page, choose your endpoint.

The endpoint details page should open, showing you a summary of your endpoint and metrics that have been collected for your endpoint.

The following sections describe the tabs on the endpoints details page.

**Monitoring**

After creating a SageMaker Hosting endpoint, you can monitor your endpoint using Amazon CloudWatch, which collects raw data and processes it into readable, near real-time metrics. Using these metrics, you can access historical information and gain a better perspective on how your endpoint is performing. For more information, see the [Amazon CloudWatch User Guide](https://docs.aws.amazon.com/AmazonCloudWatch/latest/monitoring/).  

From the **Monitoring** tab on the endpoint details page, you can view CloudWatch metrics data that has been collected from your endpoint.

The **Monitoring** tab includes the following sections:

- **Operational metrics**: View metrics that track the utilization of your endpoint's resources, such as CPU Utilization and Memory Utilization.
- **Invocation metrics**: View metrics that track the number, health, and status of InvokeEndpoint requests coming to your endpoint, such as Invocation Model Errors and Model Latency.
- **Health metrics**: View metrics that track your endpoint's overall health, such as Invocation Failures and Notification Failures.

For detailed descriptions of each metric, see [Monitor SageMaker with CloudWatch](https://docs.aws.amazon.com/sagemaker/latest/dg/metrics-monitoring.html).
The following screenshot shows the **Operational metrics** section for a serverless endpoint.
You can adjust the **Period** and **Statistic** that you want to track for the metrics in a given section, as well as the length of time for which you want to view metrics data. You can also add and remove metric widgets from the view for each section by choosing **Add widget**. In the **Add widget** dialog box, you can select and deselect the metrics that you want to see.

The metrics that are available may depend on your endpoint type. For example, serverless endpoints have some metrics that aren’t available for real-time endpoints. For more specific metrics information by endpoint type, see the following pages:

- Monitor a serverless endpoint
- Monitor an asynchronous endpoint
- CW Metrics for Multi-Model Endpoint Deployments
- Inference Pipeline Logs and Metrics

**Settings**

You can choose the **Settings** tab to view additional information about your endpoint, such as the data capture settings, the endpoint configuration, and tags.

**Alarms**

From the **Alarms** tab on your endpoint details page, you can view and create simple static threshold metric alarms, where you specify a threshold value for a metric. If the metric breaches the threshold value, the alarm goes into the ALARM state. For more information about CloudWatch alarms, see **Using Amazon CloudWatch alarms**.

In the **Endpoint summary** section, you can view the **Alarms** field, which tells you how many alarms are currently active on your endpoint.

To view which alarms are in the ALARM state, choose the **Alarms** tab. The **Alarms** tab shows you a full list of your endpoint alarms, along with details about their status and conditions. The following screenshot shows a list of alarms in this section that have been configured for an endpoint.
An alarm’s status can be **In alarm**, **OK**, or **Insufficient data** if there isn’t enough metrics data being collected.

To create a new alarm for your endpoint, do the following:

1. In the **Alarms** tab, choose **Create alarm**.
2. The **Create alarm** page opens. For **Alarm name**, enter a name for the alarm.
3. (Optional) Enter a description for the alarm.
4. For **Metric**, choose the CloudWatch metric that you want the alarm to track.
5. For **Variant name**, choose the endpoint model variant that you want to monitor.
6. For **Statistic**, choose one of the available statistics for the metric you selected.
7. For **Period**, choose the time period to use for calculating each statistical value. For example, if you choose the Average statistic and a 5 minute period, each data point monitored by the alarm is the average of the metric’s data points at 5 minute intervals.
8. For **Evaluation periods**, enter the number of data points that you want the alarm to consider when evaluating whether to enter the alarm state or not.
9. For **Condition**, choose the conditional that you want to use for your alarm threshold.
10. For **Threshold value**, enter the desired value for your threshold.
11. (Optional) For **Notification**, you can choose **Add notification** to create or specify an Amazon SNS topic that receives a notification when your alarm state changes.
12. Choose **Create alarm**.
After creating your alarm, you can return to the **Alarms** tab to view its status at any time. From this section, you can also select the alarm and either **Edit** or **Delete** it.

### Hosting options

The following topics describe available SageMaker realtime hosting options along with how to set up, invoke, and delete each hosting option.

**Topics**

- [Host a single model](#)
- [Host multiple models in one container behind one endpoint](#)
- [Host multiple models which use different containers behind one endpoint](#)
- [Host models along with pre-processing logic as serial inference pipeline behind one endpoint](#)
- [Delete Endpoints and Resources](#)

#### Host a single model

You can create, update, and delete real-time inference endpoints that host a single model with Amazon SageMaker Studio, the AWS SDK for Python (Boto3), the SageMaker Python SDK, or the AWS CLI. For procedures and code examples, see [Deploy models for real-time inference](#).

#### Host multiple models in one container behind one endpoint

Multi-model endpoints provide a scalable and cost-effective solution to deploying large numbers of models. They use the same fleet of resources and a shared serving container to host all of your models. This reduces hosting costs by improving endpoint utilization compared with using single-model endpoints. It also reduces deployment overhead because Amazon SageMaker manages loading models in memory and scaling them based on the traffic patterns to your endpoint.

The following diagram shows how multi-model endpoints work compared to single-model endpoints.
Multi-model endpoints are ideal for hosting a large number of models that use the same ML framework on a shared serving container. If you have a mix of frequently and infrequently accessed models, a multi-model endpoint can efficiently serve this traffic with fewer resources and higher cost savings. Your application should be tolerant of occasional cold start-related latency penalties that occur when invoking infrequently used models.

Multi-model endpoints support hosting both CPU and GPU backed models. By using GPU backed models, you can lower your model deployment costs through increased usage of the endpoint and its underlying accelerated compute instances.

Multi-model endpoints also enable time-sharing of memory resources across your models. This works best when the models are fairly similar in size and invocation latency. When this is the case, multi-model endpoints can effectively use instances across all models. If you have models that have significantly higher transactions per second (TPS) or latency requirements, we recommend hosting them on dedicated endpoints.

You can use multi-model endpoints with the following features:
• **AWS PrivateLink** and VPCs

• **Auto scaling**

• **Serial inference pipelines** (but only one multi-model enabled container can be included in an inference pipeline)

• A/B testing

You can't use multi-model-enabled containers with Amazon Elastic Inference.

You can use the AWS SDK for Python (Boto) or the SageMaker console to create a multi-model endpoint. For CPU backed multi-model endpoints, you can create your endpoint with custom-built containers by integrating the [Multi Model Server](https://docs.aws.amazon.com/sagemaker/latest/dg/multi-model-endpoint-definition.html) library.

**Topics**

• **Supported algorithms, frameworks, and instances**

• **Sample notebooks for multi-model endpoints**

• **How multi-model endpoints work**

• **Setting SageMaker multi-model endpoint model caching behavior**

• **Instance recommendations for multi-model endpoint deployments**

• **Create a Multi-Model Endpoint**

• **Invoke a Multi-Model Endpoint**

• **Add or Remove Models**

• **Build Your Own Container for SageMaker Multi-Model Endpoints**

• **Multi-Model Endpoint Security**

• **CloudWatch Metrics for Multi-Model Endpoint Deployments**

• **Set Auto Scaling Policies for Multi-Model Endpoint Deployments**

**Supported algorithms, frameworks, and instances**

For information about the algorithms, frameworks, and instance types that you can use with multi-model endpoints, see the following sections.
Supported algorithms, frameworks, and instances for multi-model endpoints using CPU backed instances

The inference containers for the following algorithms and frameworks support multi-model endpoints:

- XGBoost Algorithm
- K-Nearest Neighbors (k-NN) Algorithm
- Linear Learner Algorithm
- Random Cut Forest (RCF) Algorithm
- Use TensorFlow with Amazon SageMaker
- Use Scikit-learn with Amazon SageMaker
- Use Apache MXNet with Amazon SageMaker
- Use PyTorch with Amazon SageMaker

To use any other framework or algorithm, use the SageMaker inference toolkit to build a container that supports multi-model endpoints. For information, see Build Your Own Container for SageMaker Multi-Model Endpoints.

Multi-model endpoints support all of the CPU instance types.

Supported algorithms, frameworks, and instances for multi-model endpoints using GPU backed instances

Hosting multiple GPU backed models on multi-model endpoints is supported through the SageMaker Triton Inference server. This supports all major inference frameworks such as NVIDIA® TensorRT™, PyTorch, MXNet, Python, ONNX, XGBoost, scikit-learn, RandomForest, OpenVINO, custom C++, and more.

To use any other framework or algorithm, you can use Triton backend for Python or C++ to write your model logic and serve any custom model. After you have the server ready, you can start deploying 100s of Deep Learning models behind one endpoint.

Multi-model endpoints support the following GPU instance types:
<table>
<thead>
<tr>
<th>Instance family</th>
<th>Instance type</th>
<th>vCPUs</th>
<th>GiB of memory per vCPU</th>
<th>GPUs</th>
<th>GPU memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>p2</td>
<td>ml.p2.xlarge</td>
<td>4</td>
<td>15.25</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>p3</td>
<td>ml.p3.2xlarge</td>
<td>8</td>
<td>7.62</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>g5</td>
<td>ml.g5.xlarge</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>24</td>
</tr>
<tr>
<td>g5</td>
<td>ml.g5.2xlarge</td>
<td>8</td>
<td>4</td>
<td>1</td>
<td>24</td>
</tr>
<tr>
<td>g5</td>
<td>ml.g5.4xlarge</td>
<td>16</td>
<td>4</td>
<td>1</td>
<td>24</td>
</tr>
<tr>
<td>g5</td>
<td>ml.g5.8xlarge</td>
<td>32</td>
<td>4</td>
<td>1</td>
<td>24</td>
</tr>
<tr>
<td>g5</td>
<td>ml.g5.16xlarge</td>
<td>64</td>
<td>4</td>
<td>1</td>
<td>24</td>
</tr>
<tr>
<td>g4dn</td>
<td>ml.g4dn.xlarge</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>g4dn</td>
<td>ml.g4dn.2xlarge</td>
<td>8</td>
<td>4</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>g4dn</td>
<td>ml.g4dn.4xlarge</td>
<td>16</td>
<td>4</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>g4dn</td>
<td>ml.g4dn.8xlarge</td>
<td>32</td>
<td>4</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>g4dn</td>
<td>ml.g4dn.16xlarge</td>
<td>64</td>
<td>4</td>
<td>1</td>
<td>16</td>
</tr>
</tbody>
</table>

**Sample notebooks for multi-model endpoints**

To learn more about how to use multi-model endpoints, you can try the following sample notebooks:

- Examples for multi-model endpoints using CPU backed instances:
- **Multi-Model Endpoint XGBoost Sample Nootebok** – This notebook shows how to deploy multiple XGBoost models to an endpoint.

- **Multi-Model Endpoints BYOC Sample Notebook** – This notebook shows how to set up and deploy a customer container that supports multi-model endpoints in SageMaker.

- Example for multi-model endpoints using GPU backed instances:
  - **Run mulitple deep learning models on GPUs with Amazon SageMaker Multi-model endpoints (MME)** – This notebook shows how to use an NVIDIA Triton Inference container to deploy ResNet-50 models to a multi-model endpoint.

For instructions on how to create and access Jupyter notebook instances that you can use to run the previous examples in SageMaker, see [Amazon SageMaker Notebook Instances](#). After you've created a notebook instance and opened it, choose the **SageMaker Examples** tab to see a list of all the SageMaker samples. The multi-model endpoint notebooks are located in the **ADVANCED FUNCTIONALITY** section. To open a notebook, choose its **Use** tab and choose **Create copy**.

For more information about use cases for multi-model endpoints, see the following blogs and resources:

- Video: [Hosting thousands of models on SageMaker](#)
- Video: [SageMaker ML for SaaS](#)
- Blog: [How to scale machine learning inference for multi-tenant SaaS use cases](#)
- Case study: [Veeva Systems](#)

**How multi-model endpoints work**

SageMaker manages the lifecycle of models hosted on multi-model endpoints in the container's memory. Instead of downloading all of the models from an Amazon S3 bucket to the container when you create the endpoint, SageMaker dynamically loads and caches them when you invoke them. When SageMaker receives an invocation request for a particular model, it does the following:

1. Routes the request to an instance behind the endpoint.
2. Downloads the model from the S3 bucket to that instance's storage volume.
3. Loads the model to the container's memory (CPU or GPU, depending on whether you have CPU or GPU backed instances) on that accelerated compute instance. If the model is already loaded in the container's memory, invocation is faster because SageMaker doesn't need to download and load it.
SageMaker continues to route requests for a model to the instance where the model is already loaded. However, if the model receives many invocation requests, and there are additional instances for the multi-model endpoint, SageMaker routes some requests to another instance to accommodate the traffic. If the model isn't already loaded on the second instance, the model is downloaded to that instance's storage volume and loaded into the container's memory.

When an instance's memory utilization is high and SageMaker needs to load another model into memory, it unloads unused models from that instance's container to ensure that there is enough memory to load the model. Models that are unloaded remain on the instance's storage volume and can be loaded into the container's memory later without being downloaded again from the S3 bucket. If the instance's storage volume reaches its capacity, SageMaker deletes any unused models from the storage volume.

To delete a model, stop sending requests and delete it from the S3 bucket. SageMaker provides multi-model endpoint capability in a serving container. Adding models to, and deleting them from, a multi-model endpoint doesn't require updating the endpoint itself. To add a model, you upload it to the S3 bucket and invoke it. You don't need code changes to use it.

**Note**

When you update a multi-model endpoint, initial invocation requests on the endpoint might experience higher latencies as Smart Routing in multi-model endpoints adapt to your traffic pattern. However, once it learns your traffic pattern, you can experience low latencies for most frequently used models. Less frequently used models may incur some cold start latencies since the models are dynamically loaded to an instance.

**Setting SageMaker multi-model endpoint model caching behavior**

By default, multi-model endpoints cache frequently used models in memory (CPU or GPU, depending on whether you have CPU or GPU backed instances) and on disk to provide low latency inference. The cached models are unloaded and/or deleted from disk only when a container runs out of memory or disk space to accommodate a newly targeted model.

You can change the caching behavior of a multi-model endpoint and explicitly enable or disable model caching by setting the parameter `ModelCacheSetting` when you call `create_model`.

We recommend setting the value of the `ModelCacheSetting` parameter to `Disabled` for use cases that do not benefit from model caching. For example, when a large number of models need
to be served from the endpoint but each model is invoked only once (or very infrequently). For such use cases, setting the value of the ModelCacheSetting parameter to Disabled allows higher transactions per second (TPS) for invoke_endpoint requests compared to the default caching mode. Higher TPS in these use cases is because SageMaker does the following after the invoke_endpoint request:

- Asynchronously unloads the model from memory and deletes it from disk immediately after it is invoked.
- Provides higher concurrency for downloading and loading models in the inference container. For both CPU and GPU backed endpoints, the concurrency is a factor of the number of the vCPUs of the container instance.

For guidelines on choosing a SageMaker ML instance type for a multi-model endpoint, see Instance recommendations for multi-model endpoint deployments.

Instance recommendations for multi-model endpoint deployments

There are several items to consider when selecting a SageMaker ML instance type for a multi-model endpoint:

- Provision sufficient Amazon Elastic Block Store (Amazon EBS) capacity for all of the models that need to be served.
- Balance performance (minimize cold starts) and cost (don't over-provision instance capacity). For information about the size of the storage volume that SageMaker attaches for each instance type for an endpoint and for a multi-model endpoint, see Host instance storage volumes.
- For a container configured to run in MultiModel mode, the storage volume provisioned for its instances are larger than the default SingleModel mode. This allows more models to be cached on the instance storage volume than in SingleModel mode.

When choosing a SageMaker ML instance type, consider the following:

- Multi-model endpoints are currently supported for all CPU instances types and on single-GPU instance types.
- For the traffic distribution (access patterns) to the models that you want to host behind the multi-model endpoint, along with the model size (how many models could be loaded in memory on the instance), keep the following information in mind:
• Think of the amount of memory on an instance as the cache space for models to be loaded, and think of the number of vCPUs as the concurrency limit to perform inference on the loaded models (assuming that invoking a model is bound to CPU).

• For CPU backed instances, the number of vCPUs impacts your maximum concurrent invocations per instance (assuming that invoking a model is bound to CPU). A higher amount of vCPUs enables you to invoke more unique models concurrently.

• For GPU backed instances, a higher amount of instance and GPU memory enables you to have more models loaded and ready to serve inference requests.

• For both CPU and GPU backed instances, have some "slack" memory available so that unused models can be unloaded, and especially for multi-model endpoints with multiple instances. If an instance or an Availability Zone fails, the models on those instances will be rerouted to other instances behind the endpoint.

• Determine your tolerance to loading/download times:
  • d instance type families (for example, m5d, c5d, or r5d) and g5s come with an NVMe (non-volatile memory express) SSD, which offers high I/O performance and might reduce the time it takes to download models to the storage volume and for the container to load the model from the storage volume.
  • Because d and g5 instance types come with an NVMe SSD storage, SageMaker does not attach an Amazon EBS storage volume to these ML compute instances that hosts the multi-model endpoint. Auto scaling works best when the models are similarly sized and homogenous, that is when they have similar inference latency and resource requirements.

You can also use the following guidance to help you optimize model loading on your multi-model endpoints:

**Choosing an instance type that can't hold all of the targeted models in memory**

In some cases, you might opt to reduce costs by choosing an instance type that can't hold all of the targeted models in memory at once. SageMaker dynamically unloads models when it runs out of memory to make room for a newly targeted model. For infrequently requested models, you sacrifice dynamic load latency. In cases with more stringent latency needs, you might opt for larger instance types or more instances. Investing time up front for performance testing and analysis helps you to have successful production deployments.

**Evaluating your model cache hits**
Amazon CloudWatch metrics can help you evaluate your models. For more information about metrics you can use with multi-model endpoints, see CloudWatch Metrics for Multi-Model Endpoint Deployments.

You can use the Average statistic of the ModelCacheHit metric to monitor the ratio of requests where the model is already loaded. You can use the SampleCount statistic for the ModelUnloadingTime metric to monitor the number of unload requests sent to the container during a time period. If models are unloaded too frequently (an indicator of thrashing, where models are being unloaded and loaded again because there is insufficient cache space for the working set of models), consider using a larger instance type with more memory or increasing the number of instances behind the multi-model endpoint. For multi-model endpoints with multiple instances, be aware that a model might be loaded on more than 1 instance.

Create a Multi-Model Endpoint

You can use the SageMaker console or the AWS SDK for Python (Boto) to create a multi-model endpoint. To create either a CPU or GPU backed endpoint through the console, see the console procedure in the following sections. If you want to create a multi-model endpoint with the AWS SDK for Python (Boto), use either the CPU or GPU procedure in the following sections. The CPU and GPU workflows are similar but have several differences, such as the container requirements.

Topics

- Create a multi-model endpoint (console)
- Create a multi-model endpoint using CPUs with the AWS SDK for Python (Boto3)
- Create a multi-model endpoint using GPUs with the AWS SDK for Python (Boto3)

Create a multi-model endpoint (console)

You can create both CPU and GPU backed multi-model endpoints through the console. Use the following procedure to create a multi-model endpoint through the SageMaker console.

To create a multi-model endpoint (console)

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Model, and then from the Inference group, choose Create model.
3. For Model name, enter a name.
4. For IAM role, choose or create an IAM role that has the AmazonSageMakerFullAccess IAM policy attached.
5. In the **Container definition** section, for **Provide model artifacts and inference image options**, choose **Use multiple models**.
6. For the **Inference container image**, enter the Amazon ECR path for your desired container image.

   For GPU models, you must use a container backed by the NVIDIA Triton Inference Server. For a list of container images that work with GPU backed endpoints, see the [NVIDIA Triton Inference Containers (SM support only)](https://docs.aws.amazon.com/sagemaker/latest/dg/triton_containers.html). For more information about the NVIDIA Triton Inference Server, see [Use Triton Inference Server with SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/triton_container.html).

7. Choose **Create model**.

8. Deploy your multi-model endpoint as you would a single model endpoint. For instructions, see [Deploy the Model to SageMaker Hosting Services](https://docs.aws.amazon.com/sagemaker/latest/dg/deploy-model.html).

### Create a multi-model endpoint using CPUs with the AWS SDK for Python (Boto3)

Use the following section to create a multi-model endpoint backed by CPU instances. You create a multi-model endpoint using the Amazon SageMaker `create_model`, `create_endpoint_config`, and `create_endpoint` APIs just as you would create a single model endpoint, but with two changes. When defining the model container, you need to pass a new `Mode` parameter value, `MultiModel`. You also need to pass the `ModelDataUrl` field that specifies the prefix in Amazon S3 where the model artifacts are located, instead of the path to a single model artifact, as you would when deploying a single model.

For a sample notebook that uses SageMaker to deploy multiple XGBoost models to an endpoint, see [Multi-Model Endpoint XGBoost Sample Notebook](https://github.com/aws-samples/amazon-sagemaker-endpoint). The following procedure outlines the key steps used in that sample to create a CPU backed multi-model endpoint.

### To deploy the model (AWS SDK for Python (Boto 3))

1. Get a container with an image that supports deploying multi-model endpoints. For a list of built-in algorithms and framework containers that support multi-model endpoints, see [Supported algorithms, frameworks, and instances](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-recognize.html). For this example, we use the **K-Nearest Neighbors (k-NN) Algorithm** built-in algorithm. We call the SageMaker Python SDK utility function `image_uris.retrieve()` to get the address for the K-Nearest Neighbors built-in algorithm image.

```python
import sagemaker
region = sagemaker_session.boto_region_name
```
image = sagemaker.image_uris.retrieve("knn", region=region)
container = {
    'Image': image,
    'ModelDataUrl': 's3://<BUCKET_NAME>/<PATH_TO_ARTIFACTS>',
    'Mode': 'MultiModel'
}

2. Get an AWS SDK for Python (Boto3) SageMaker client and create the model that uses this container.

    import boto3
    sagemaker_client = boto3.client('sagemaker')
    response = sagemaker_client.create_model(
        ModelName = '<MODEL_NAME>',
        ExecutionRoleArn = role,
        Containers = [container])

3. (Optional) If you are using a serial inference pipeline, get the additional container(s) to include in the pipeline, and include it in the Containers argument of CreateModel:

    preprocessor_container = {
        'Image':
            '<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/<PREPROCESSOR_IMAGE>:<TAG>'
    }

    multi_model_container = {
        'Image':
            '<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/<IMAGE>:<TAG>',
        'ModelDataUrl': 's3://<BUCKET_NAME>/<PATH_TO_ARTIFACTS>',
        'Mode': 'MultiModel'
    }

    response = sagemaker_client.create_model(
        ModelName = '<MODEL_NAME>',
        ExecutionRoleArn = role,
        Containers = [preprocessor_container, multi_model_container]
    )

⚠️ Note

You can use only one multi-model-enabled endpoint in a serial inference pipeline.
4. (Optional) If your use case does not benefit from model caching, set the value of the ModelCacheSetting field of the MultiModelConfig parameter to Disabled, and include it in the Container argument of the call to create_model. The value of the ModelCacheSetting field is Enabled by default.

```python
container = {
    'Image': image,
    'ModelDataUrl': 's3://<BUCKET_NAME>/PATH_TO_ARTIFACTS',
    'Mode': 'MultiModel'
    'MultiModelConfig': {
        // Default value is 'Enabled'
        'ModelCacheSetting': 'Disabled'
    }
}

response = sagemaker_client.create_model(
    ModelName = '<MODEL_NAME>',
    ExecutionRoleArn = role,
    Containers = [container]
)
```

5. Configure the multi-model endpoint for the model. We recommend configuring your endpoints with at least two instances. This allows SageMaker to provide a highly available set of predictions across multiple Availability Zones for the models.

```python
response = sagemaker_client.create_endpoint_config(
    EndpointConfigName = '<ENDPOINT_CONFIG_NAME>',
    ProductionVariants=[
        {  
            'InstanceType': 'ml.m4.xlarge',
            'InitialInstanceCount': 2,
            'InitialVariantWeight': 1,
            'ModelName': '<MODEL_NAME>',
            'VariantName': 'AllTraffic'
        }
    ]
)
```
You can use only one multi-model-enabled endpoint in a serial inference pipeline.

6. Create the multi-model endpoint using the `EndpointName` and `EndpointConfigName` parameters.

```python
response = sagemaker_client.create_endpoint(
    EndpointName       = '<ENDPOINT_NAME>',
    EndpointConfigName = '<ENDPOINT_CONFIG_NAME>'
)
```

Create a multi-model endpoint using GPUs with the AWS SDK for Python (Boto3)

Use the following section to create a GPU backed multi-model endpoint. You create a multi-model endpoint using the Amazon SageMaker `create_model`, `create_endpoint_config`, and `create_endpoint` APIs similarly to creating single model endpoints, but there are several changes. When defining the model container, you need to pass a new `Mode` parameter value, `MultiModel`. You also need to pass the `ModelDataUrl` field that specifies the prefix in Amazon S3 where the model artifacts are located, instead of the path to a single model artifact, as you would when deploying a single model. For GPU backed multi-model endpoints, you also must use a container with the NVIDIA Triton Inference Server that is optimized for running on GPU instances. For a list of container images that work with GPU backed endpoints, see the NVIDIA Triton Inference Containers (SM support only).

For an example notebook that demonstrates how to create a multi-model endpoint backed by GPUs, see Run mulitple deep learning models on GPUs with Amazon SageMaker Multi-model endpoints (MME).

The following procedure outlines the key steps to create a GPU backed multi-model endpoint.

**To deploy the model (AWS SDK for Python (Boto 3))**

1. Define the container image. To create a multi-model endpoint with GPU support for ResNet models, define the container to use the NVIDIA Triton Server image. This container supports multi-model endpoints and is optimized for running on GPU instances. We call the SageMaker Python SDK utility function `image_uris.retrieve()` to get the address for the image. For example:
import sagemaker
region = sagemaker_session.boto_region_name

// Find the sagemaker-tritonserver image at
// resnet50/triton_resnet50.ipynb
// Find available tags at https://github.com/aws/deep-learning-containers/blob/
// master/available_images.md#nvidia-triton-inference-containers-sm-support-only

image = "<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/sagemaker-
        tritonserver:<TAG>".format(
    account_id=account_id_map[region], region=region
)

container = {
    'Image':      image,
    'ModelDataUrl': 's3://<BUCKET_NAME>/<PATH_TO_ARTIFACTS>',
    'Mode':        'MultiModel',
    "Environment": {
        "SAGEMAKER_TRITON_DEFAULT_MODEL_NAME": "resnet"},
}

2. Get an AWS SDK for Python (Boto3) SageMaker client and create the model that uses this container.

import boto3
sagemaker_client = boto3.client('sagemaker')
response = sagemaker_client.create_model(
    ModelName     = '<MODEL_NAME>',
    ExecutionRoleArn = role,
    Containers     = [container])

3. (Optional) If you are using a serial inference pipeline, get the additional container(s) to include in the pipeline, and include it in the Containers argument of CreateModel:

preprocessor_container = {
    'Image':
    '<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/<PREPROCESSOR_IMAGE>:<TAG>'
}

multi_model_container = {
    'Image':
    '<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/<IMAGE>:<TAG>'
}
Note

You can use only one multi-model-enabled endpoint in a serial inference pipeline.

4. (Optional) If your use case does not benefit from model caching, set the value of the ModelCacheSetting field of the MultiModelConfig parameter to Disabled, and include it in the Container argument of the call to create_model. The value of the ModelCacheSetting field is Enabled by default.

```python
container = {
    'Image': image,
    'ModelDataUrl': 's3://<BUCKET_NAME>/PATH_TO_ARTIFACTS',
    'Mode': 'MultiModel'
    'MultiModelConfig': {
        // Default value is 'Enabled'
        'ModelCacheSetting': 'Disabled'
    }
}

response = sagemaker_client.create_model(
    ModelName = '<MODEL_NAME>',
    ExecutionRoleArn = role,
    Containers = [preprocessor_container, multi_model_container]
)
```

5. Configure the multi-model endpoint with GPU backed instances for the model. We recommend configuring your endpoints with more than one instance to allow for high availability and higher cache hits.

```python
response = sagemaker_client.create_endpoint_config(
    EndpointConfigName = '<ENDPOINT_CONFIG_NAME>',
)```
6. Create the multi-model endpoint using the EndpointName and EndpointConfigName parameters.

   ```
   response = sagemaker_client.create_endpoint(
       EndpointName       = '<ENDPOINT_NAME>',
       EndpointConfigName = '<ENDPOINT_CONFIG_NAME>'
   )
   ```

Invoke a Multi-Model Endpoint

To invoke a multi-model endpoint, use the `invoke_endpoint` from the SageMaker Runtime just as you would invoke a single model endpoint, with one change. Pass a new TargetModel parameter that specifies which of the models at the endpoint to target. The SageMaker Runtime InvokeEndpoint request supports `X-Amzn-SageMaker-Target-Model` as a new header that takes the relative path of the model specified for invocation. The SageMaker system constructs the absolute path of the model by combining the prefix that is provided as part of the `CreateModel` API call with the relative path of the model.

The following procedures are the same for both CPU and GPU-backed multi-model endpoints.

AWS SDK for Python (Boto 3)

   The following example prediction request uses the [AWS SDK for Python (Boto 3)](https://boto3.amazonaws.com/v1/documentation/api/latest/index.html) in the sample notebook.

   ```
   response = runtime_sagemaker_client.invoke_endpoint(
       EndpointName       = '<ENDPOINT_NAME>',
       ContentType        = 'text/csv',
       TargetModel        = '<MODEL_FILENAME>.tar.gz',
       Body               = body)
   ```
AWS CLI

The following example shows how to make a CSV request with two rows using the AWS Command Line Interface (AWS CLI):

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name "<ENDPOINT_NAME>" \
  --body "1.0,2.0,5.0"$\n
  "2.0,3.0,4.0" \
  --content-type "text/csv" \
  --target-model "<MODEL_NAME>.tar.gz" \
  output_file.txt
```

An `output_file.txt` with information about your inference requests is made if the inference was successful. For more examples on how to make predictions with the AWS CLI, see [Making predictions with the AWS CLI](#) in the SageMaker Python SDK documentation.

The multi-model endpoint dynamically loads target models as needed. You can observe this when running the [MME Sample Notebook](#) as it iterates through random invocations against multiple target models hosted behind a single endpoint. The first request against a given model takes longer because the model has to be downloaded from Amazon Simple Storage Service (Amazon S3) and loaded into memory. This is called a cold start, and it is expected on multi-model endpoints to optimize for better price performance for customers. Subsequent calls finish faster because there's no additional overhead after the model has loaded.

**Note**

For GPU backed instances, the HTTP response code with 507 from the GPU container indicates a lack of memory or other resources. This causes unused models to be unloaded from the container in order to load more frequently used models.

**Retry Requests on ModelNotReadyException Errors**

The first time you call `invoke_endpoint` for a model, the model is downloaded from Amazon Simple Storage Service and loaded into the inference container. This makes the first call take longer to return. Subsequent calls to the same model finish faster, because the model is already loaded.
SageMaker returns a response for a call to `invoke_endpoint` within 60 seconds. Some models are too large to download within 60 seconds. If the model does not finish loading before the 60 second timeout limit, the request to `invoke_endpoint` returns with the error code `ModelNotReadyException`, and the model continues to download and load into the inference container for up to 360 seconds. If you get a `ModelNotReadyException` error code for an `invoke_endpoint` request, retry the request. By default, the AWS SDKs for Python (Boto 3) (using Legacy retry mode) and Java retry `invoke_endpoint` requests that result in `ModelNotReadyException` errors. You can configure the retry strategy to continue retrying the request for up to 360 seconds. If you expect your model to take longer than 60 seconds to download and load into the container, set the SDK socket timeout to 70 seconds. For more information about configuring the retry strategy for the AWS SDK for Python (Boto3), see Configuring a retry mode. The following code shows an example that configures the retry strategy to retry calls to `invoke_endpoint` for up to 180 seconds.

```python
import boto3
from botocore.config import Config

# This example retry strategy sets the retry attempts to 2.
# With this setting, the request can attempt to download and/or load the model
# for up to 180 seconds: 1 original request (60 seconds) + 2 retries (120 seconds)
config = Config(
    read_timeout=70,
    retries={
        'max_attempts': 2  # This value can be adjusted to 5 to go up to the 360s max timeout
    }
)
runtime_sagemaker_client = boto3.client('sagemaker-runtime', config=config)
```

**Add or Remove Models**

You can deploy additional models to a multi-model endpoint and invoke them through that endpoint immediately. When adding a new model, you don't need to update or bring down the endpoint, so you avoid the cost of creating and running a separate endpoint for each new model. The process for adding and removing models is the same for CPU and GPU-backed multi-model endpoints.

SageMaker unloads unused models from the container when the instance is reaching memory capacity and more models need to be downloaded into the container. SageMaker also deletes unused model artifacts from the instance storage volume when the volume is reaching capacity.
and new models need to be downloaded. The first invocation to a newly added model takes longer because the endpoint takes time to download the model from S3 to the container's memory in instance hosting the endpoint.

With the endpoint already running, copy a new set of model artifacts to the Amazon S3 location there you store your models.

```bash
# Add an AdditionalModel to the endpoint and exercise it
aws s3 cp AdditionalModel.tar.gz s3://my-bucket/path/to/artifacts/
```

⚠️ **Important**

To update a model, proceed as you would when adding a new model. Use a new and unique name. Don't overwrite model artifacts in Amazon S3 because the old version of the model might still be loaded in the containers or on the storage volume of the instances on the endpoint. Invocations to the new model could then invoke the old version of the model.

Client applications can request predictions from the additional target model as soon as it is stored in S3.

```python
response = runtime_sagemaker_client.invoke_endpoint(
    EndpointName='<ENDPOINT_NAME>',
    ContentType='text/csv',
    TargetModel='AdditionalModel.tar.gz',
    Body=body)
```

To delete a model from a multi-model endpoint, stop invoking the model from the clients and remove it from the S3 location where model artifacts are stored.

**Build Your Own Container for SageMaker Multi-Model Endpoints**

Refer to the following sections for bringing your own container and dependencies to multi-model endpoints.

**Topics**
- Bring your own dependencies for multi-model endpoints on CPU backed instances
- Bring your own dependencies for multi-model endpoints on GPU backed instances
- Use the SageMaker Inference Toolkit
Bring your own dependencies for multi-model endpoints on CPU backed instances

If none of the pre-built container images serve your needs, you can build your own container for use with CPU backed multi-model endpoints.

Custom Amazon Elastic Container Registry (Amazon ECR) images deployed in Amazon SageMaker are expected to adhere to the basic contract described in Use Your Own Inference Code with Hosting Services that govern how SageMaker interacts with a Docker container that runs your own inference code. For a container to be capable of loading and serving multiple models concurrently, there are additional APIs and behaviors that must be followed. This additional contract includes new APIs to load, list, get, and unload models, and a different API to invoke models. There are also different behaviors for error scenarios that the APIs need to abide by. To indicate that the container complies with the additional requirements, you can add the following command to your Docker file:

```
LABEL com.amazonaws.sagemaker.capabilities.multi-models=true
```

SageMaker also injects an environment variable into the container

```
SAGEMAKER_MULTI_MODEL=true
```

If you are creating a multi-model endpoint for a serial inference pipeline, your Docker file must have the required labels for both multi-models and serial inference pipelines. For more information about serial information pipelines, see Run Real-time Predictions with an Inference Pipeline.

To help you implement these requirements for a custom container, two libraries are available:

- **Multi Model Server** is an open source framework for serving machine learning models that can be installed in containers to provide the front end that fulfills the requirements for the new multi-model endpoint container APIs. It provides the HTTP front end and model management capabilities required by multi-model endpoints to host multiple models within a single container, load models into and unload models out of the container dynamically, and performs inference on a specified loaded model. It also provides a pluggable backend that supports a pluggable custom backend handler where you can implement your own algorithm.

- **SageMaker Inference Toolkit** is a library that bootstraps Multi Model Server with a configuration and settings that make it compatible with SageMaker multi-model endpoints. It also allows
you to tweak important performance parameters, such as the number of workers per model, depending on the needs of your scenario.

**Bring your own dependencies for multi-model endpoints on GPU backed instances**

The bring your own container (BYOC) capability on multi-model endpoints with GPU backed instances is not currently supported by the Multi Model Server and SageMaker Inference Toolkit libraries.

For creating multi-model endpoints with GPU backed instances, you can use the SageMaker supported NVIDIA Triton Inference Server, with the NVIDIA Triton Inference Containers. To bring your own dependencies, you can build your own container with the SageMaker supported NVIDIA Triton Inference Server as the base image to your Docker file:

```bash
FROM 301217895009.dkr.ecr.us-west-2.amazonaws.com/sagemaker-tritonserver:22.07-py3
```

⚠️ **Important**
Containers with the Triton Inference Server are the only supported containers you can use for GPU backed multi-model endpoints.

**Use the SageMaker Inference Toolkit**

**(1)** **Note**
The SageMaker Inference Toolkit is only supported for CPU backed multi-model endpoints.
The SageMaker Inference Toolkit is not currently not supported for GPU backed multi-model endpoints.

Pre-built containers that support multi-model endpoints are listed in the [Supported algorithms, frameworks, and instances](#). If you want to use any other framework or algorithm, you need to build a container. The easiest way to do this is to use the SageMaker Inference Toolkit to extend an existing pre-built container. The SageMaker inference toolkit is an implementation for the multi-model server (MMS) that creates endpoints that can be deployed in SageMaker. For a sample notebook that shows how to set up and deploy a custom container that supports multi-model endpoints in SageMaker, see the Multi-Model Endpoint BYOC Sample Notebook.
Note

The SageMaker inference toolkit supports only Python model handlers. If you want to implement your handler in any other language, you must build your own container that implements the additional multi-model endpoint APIs. For information, see Custom Containers Contract for Multi-Model Endpoints.

To extend a container by using the SageMaker inference toolkit

1. Create a model handler. MMS expects a model handler, which is a Python file that implements functions to pre-process, get predictions from the model, and process the output in a model handler. For an example of a model handler, see model_handler.py from the sample notebook.
2. Import the inference toolkit and use its model_server.start_model_server function to start MMS. The following example is from the dockerd-entrypoint.py file from the sample notebook. Notice that the call to model_server.start_model_server passes the model handler described in the previous step:

```python
def _retry_if_error(exception):
    return isinstance(exception, CalledProcessError or OSError)

@retry(stop_max_delay=1000 * 50,
       retry_on_exception=_retry_if_error)
def _start_mms():
    # by default the number of workers per model is 1, but we can configure it through the
    # environment variable below if desired.
    # os.environ['SAGEMAKER_MODEL_SERVER_WORKERS'] = '2'
    model_server.start_model_server(handler_service='/home/model-server/model_handler.py:handle')

def main():
    if sys.argv[1] == 'serve':
        _start_mms()
```

3. In your Dockerfile, copy the model handler from the first step and specify the Python file from the previous step as the entrypoint in your Dockerfile. The following lines are from the Dockerfile used in the sample notebook:

```bash
# Copy the default custom service file to handle incoming data and inference requests
COPY model_handler.py /home/model-server/model_handler.py

# Define an entrypoint script for the docker image
ENTRYPOINT ["python", "/usr/local/bin/dockerd-entrypoint.py"]
```

4. Build and register your container. The following shell script from the sample notebook builds the container and uploads it to an Amazon Elastic Container Registry repository in your AWS account:

```bash
%%sh

# The name of our algorithm
algorithm_name=demo-sagemaker-multimodel

cd container

account=$(aws sts get-caller-identity --query Account --output text)

# Get the region defined in the current configuration (default to us-west-2 if none defined)
region=$(aws configure get region)
region=${region:-us-west-2}

fullname="${account}.dkr.ecr.${region}.amazonaws.com/${algorithm_name}:latest"

# If the repository doesn't exist in ECR, create it.
aws ecr describe-repositories --repository-names "${algorithm_name}" > /dev/null
```
if [ $? -ne 0 ]
then
    aws ecr create-repository --repository-name "${algorithm_name}" > /dev/null
fi

# Get the login command from ECR and execute it directly
$(aws ecr get-login --region ${region} --no-include-email)

# Build the docker image locally with the image name and then push it to ECR
# with the full name.

docker build -q -t ${algorithm_name} .
docker tag ${algorithm_name} ${fullname}
docker push ${fullname}

You can now use this container to deploy multi-model endpoints in SageMaker.

Topics

- Custom Containers Contract for Multi-Model Endpoints

Custom Containers Contract for Multi-Model Endpoints

To handle multiple models, your container must support a set of APIs that enable Amazon SageMaker to communicate with the container for loading, listing, getting, and unloading models as required. The model\_name is used in the new set of APIs as the key input parameter. The customer container is expected to keep track of the loaded models using model\_name as the mapping key. Also, the model\_name is an opaque identifier and is not necessarily the value of the TargetModel parameter passed into the InvokeEndpoint API. The original TargetModel value in the InvokeEndpoint request is passed to container in the APIs as a X-Amzn-SageMaker-Target-Model header that can be used for logging purposes.

Note

Multi-model endpoints for GPU backed instances are currently supported only with SageMaker's [NVIDIA Triton Inference Server container](https://docs.aws.amazon.com/sagemaker/latest/dg/nvidia-triton-container.html). This container already implements...
the contract defined below. Customers can directly use this container with their multi-model GPU endpoints, without any additional work.

You can configure the following APIs on your containers for CPU backed multi-model endpoints.

**Topics**

- **Load Model API**
- **List Model API**
- **Get Model API**
- **Unload Model API**
- **Invoke Model API**

**Load Model API**

Instructs the container to load a particular model present in the `url` field of the body into the memory of the customer container and to keep track of it with the assigned `model_name`. After a model is loaded, the container should be ready to serve inference requests using this `model_name`.

```plaintext
POST /models HTTP/1.1
Content-Type: application/json
Accept: application/json

{
   "model_name": "{model_name}",
   "url": "/opt/ml/models/{model_name}/model"
}
```

**Note**

If `model_name` is already loaded, this API should return 409. Any time a model cannot be loaded due to lack of memory or to any other resource, this API should return a 507 HTTP status code to SageMaker, which then initiates unloading unused models to reclaim.

**List Model API**

Returns the list of models loaded into the memory of the customer container.
GET /models HTTP/1.1
Accept: application/json

Response =
{
    "models": [
        {
            "modelName": "{model_name}",
            "modelUrl" : "/opt/ml/models/{model_name}/model",
        },
        {
            "modelName": "{model_name}",
            "modelUrl" : "/opt/ml/models/{model_name}/model",
        }....
    ]
}

This API also supports pagination.

GET /models HTTP/1.1
Accept: application/json

Response =
{
    "models": [
        {
            "modelName": "{model_name}",
            "modelUrl" : "/opt/ml/models/{model_name}/model",
        },
        {
            "modelName": "{model_name}",
            "modelUrl" : "/opt/ml/models/{model_name}/model",
        }....
    ]
}

SageMaker can initially call the List Models API without providing a value for `next_page_token`. If a `nextPageToken` field is returned as part of the response, it will be provided as the value for `next_page_token` in a subsequent List Models call. If a `nextPageToken` is not returned, it means that there are no more models to return.
Get Model API

This is a simple read API on the model_name entity.

GET /models/{model_name} HTTP/1.1
Accept: application/json

{
    "modelName" : "{model_name}",
    "modelUrl" : "/opt/ml/models/{model_name}/model",
}

⚠️ Note
If model_name is not loaded, this API should return 404.

Unload Model API

Instructs the SageMaker platform to instruct the customer container to unload a model from memory. This initiates the eviction of a candidate model as determined by the platform when starting the process of loading a new model. The resources provisioned to model_name should be reclaimed by the container when this API returns a response.

DELETE /models/{model_name}

⚠️ Note
If model_name is not loaded, this API should return 404.

Invoke Model API

Makes a prediction request from the particular model_name supplied. The SageMaker Runtime InvokeEndpoint request supports X-Amzn-SageMaker-Target-Model as a new header that takes the relative path of the model specified for invocation. The SageMaker system constructs the absolute path of the model by combining the prefix that is provided as part of the CreateModel API call with the relative path of the model.
POST /models/{model_name}/invoke HTTP/1.1
Content-Type: ContentType
Accept: Accept
X-Amzn-SageMaker-Custom-Attributes: CustomAttributes
X-Amzn-SageMaker-Target-Model: {relativePath}/{artifactName}.tar.gz

⚠️ Note
If model_name is not loaded, this API should return 404.

Additionally, on GPU instances, if InvokeEndpoint fails due to a lack of memory or other resources, this API should return a 507 HTTP status code to SageMaker, which then initiates unloading unused models to reclaim.

**Multi-Model Endpoint Security**

Models and data in a multi-model endpoint are co-located on instance storage volume and in container memory. All instances for Amazon SageMaker endpoints run on a single tenant container that you own. Only your models can run on your multi-model endpoint. It's your responsibility to manage the mapping of requests to models and to provide access for users to the correct target models. SageMaker uses IAM roles to provide IAM identity-based policies that you use to specify allowed or denied actions and resources and the conditions under which actions are allowed or denied.

By default, an IAM principal with InvokeEndpoint permissions on a multi-model endpoint can invoke any model at the address of the S3 prefix defined in the CreateModel operation, provided that the IAM Execution Role defined in operation has permissions to download the model. If you need to restrict InvokeEndpoint access to a limited set of models in S3, you can do one of the following:

- Restrict InvokeEndpoint calls to specific models hosted at the endpoint by using the sagemaker:TargetModel IAM condition key. For example, the following policy allows InvokeEndpoint requests only when the value of the TargetModel field matches one of the specified regular expressions:

```json
{
    "Version": "2012-10-17",
    "Statement": [
```

- Create multi-model endpoints with more restrictive S3 prefixes.

For more information about how SageMaker uses roles to manage access to endpoints and perform operations on your behalf, see [SageMaker Roles](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-create-endpoint.html). Your customers might also have certain data isolation requirements dictated by their own compliance requirements that can be satisfied using IAM identities.

**CloudWatch Metrics for Multi-Model Endpoint Deployments**

Amazon SageMaker provides metrics for endpoints so you can monitor the cache hit rate, the number of models loaded and the model wait times for loading, downloading, and uploading at a multi-model endpoint. Some of the metrics are different for CPU and GPU backed multi-model endpoints, so the following sections describe the Amazon CloudWatch metrics that you can use for each type of multi-model endpoint.


**CloudWatch metrics for CPU backed multi-model endpoints**

You can monitor the following metrics on CPU backed multi-model endpoints.

```json
{
    "Action": [
        "sagemaker:InvokeEndpoint"
    ],
    "Effect": "Allow",
    "Condition": {
        // TargetModel provided must be from this set of values
        "StringLike": {
            "sagemaker:TargetModel": ["company_a/*", "common/*"]
        }
    }
}
```
The AWS/SageMaker namespace includes the following model loading metrics from calls to `InvokeEndpoint`.

Metrics are available at a 1-minute frequency.

For information about how long CloudWatch metrics are retained for, see `GetMetricStatistics` in the *Amazon CloudWatch API Reference*.

### Multi-Model Endpoint Model Loading Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
<th>Units</th>
<th>Valid statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelLoadingWaitTime</td>
<td>The interval of time that an invocation request has waited for the target model to be downloaded, or loaded, or both in order to perform inference.</td>
<td>Microseconds</td>
<td>Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelUnloadingTime</td>
<td>The interval of time that it took to unload the model through the container's <code>UnloadModel</code> API call.</td>
<td>Microseconds</td>
<td>Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelDownloadTime</td>
<td>The interval of time that it took to download the model from Amazon Simple Storage Service (Amazon S3).</td>
<td>Microseconds</td>
<td>Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelLoadingTime</td>
<td>The interval of time that it took to load the model through the container's <code>LoadModel</code> API call.</td>
<td>Microseconds</td>
<td>Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ModelCacheHit</td>
<td>The number of <code>InvokeEndpoint</code> requests sent to the multi-model endpoint for which the model was already loaded. The Average statistic shows the ratio of requests for which the model was already loaded. Units: None Valid statistics: Average, Sum, Sample Count</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Dimensions for Multi-Model Endpoint Model Loading Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EndpointName, VariantName</td>
<td>Filters endpoint invocation metrics for a <code>ProductionVariant</code> of the specified endpoint and variant.</td>
</tr>
</tbody>
</table>

The `/aws/sagemaker/Endpoints` namespaces include the following instance metrics from calls to `InvokeEndpoint`.

Metrics are available at a 1-minute frequency.

For information about how long CloudWatch metrics are retained for, see [GetMetricStatistics](https://docs.aws.amazon.com/AmazonCloudWatch/latest/APIReference/API_GetMetricStatistics.html) in the *Amazon CloudWatch API Reference*.

**Multi-Model Endpoint Model Instance Metrics**

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>LoadedModelCount</td>
<td>The number of models loaded in the containers of the multi-model endpoint. This metric is emitted per instance. The Average statistic with a period of 1 minute tells you the average number of models loaded per instance.</td>
</tr>
</tbody>
</table>
### Metric Description

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Sum statistic</td>
<td>The Sum statistic tells you the total number of models loaded across all instances in the endpoint. The models that this metric tracks are not necessarily unique because a model might be loaded in multiple containers at the endpoint. Units: None Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>CPUUtilization</td>
<td>The sum of each individual CPU core's utilization. The CPU utilization of each core range is 0–100. For example, if there are four CPUs, the range is 0%–400%. For endpoint variants, the value is the sum of the CPU utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
<tr>
<td>MemoryUtilization</td>
<td>The percentage of memory that is used by the containers on an instance. This value range is 0%–100%. For endpoint variants, the value is the sum of the memory utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
<tr>
<td>DiskUtilization</td>
<td>The percentage of disk space used by the containers on an instance. This value range is 0%–100%. For endpoint variants, the value is the sum of the disk space utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
</tbody>
</table>

### CloudWatch metrics for GPU multi-model endpoint deployments

You can monitor the following metrics on GPU backed multi-model endpoints.
The AWS/SageMaker namespace includes the following model loading metrics from calls to `InvokeEndpoint`.

Metrics are available at a 1-minute frequency.

For information about how long CloudWatch metrics are retained for, see `GetMetricStatistics` in the *Amazon CloudWatch API Reference*.

### Multi-Model Endpoint Model Loading Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelLoadingWaitTime</td>
<td>The interval of time that an invocation request has waited for the target model to be downloaded, or loaded, or both in order to perform inference.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelUnloadingTime</td>
<td>The interval of time that it took to unload the model through the container's <code>UnloadModel</code> API call.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelDownloadingTime</td>
<td>The interval of time that it took to download the model from Amazon Simple Storage Service (Amazon S3).</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelLoadingTime</td>
<td>The interval of time that it took to load the model through the container's <code>LoadModel</code> API call.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>
### Metric

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelCacheHit</td>
<td>The number of InvokeEndpoint requests sent to the multi-model endpoint for which the model was already loaded. The Average statistic shows the ratio of requests for which the model was already loaded. Units: None Valid statistics: Average, Sum, Sample Count</td>
</tr>
</tbody>
</table>

### Dimensions for Multi-Model Endpoint Model Loading Metrics

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EndpointName, VariantName</td>
<td>Filters endpoint invocation metrics for a ProductionVariant of the specified endpoint and variant.</td>
</tr>
</tbody>
</table>

The /aws/sagemaker/Endpoints namespaces include the following instance metrics from calls to InvokeEndpoint.

Metrics are available at a 1-minute frequency.

For information about how long CloudWatch metrics are retained for, see GetMetricStatistics in the Amazon CloudWatch API Reference.

### Multi-Model Endpoint Model Instance Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>LoadedModelCount</td>
<td>The number of models loaded in the containers of the multi-model endpoint. This metric is emitted per instance. The Average statistic with a period of 1 minute tells you the average number of models loaded per instance.</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>The Sum statistic</strong></td>
<td>The total number of models loaded across all instances in the endpoint. The models that this metric tracks are not necessarily unique because a model might be loaded in multiple containers at the endpoint. Units: None Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td><strong>CPUUtilization</strong></td>
<td>The sum of each individual CPU core's utilization. The CPU utilization of each core range is 0-100. For example, if there are four CPUs, the CPUUtilization range is 0%–400%. For endpoint variants, the value is the sum of the CPU utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
<tr>
<td><strong>MemoryUtilization</strong></td>
<td>The percentage of memory that is used by the containers on an instance. This value range is 0%-100%. For endpoint variants, the value is the sum of the memory utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
<tr>
<td><strong>GPUUtilization</strong></td>
<td>The percentage of GPU units that are used by the containers on an instance. The value can range between range is 0-100 and is multiplied by the number of GPUs. For example, if there are four GPUs, the GPUUtilization range is 0%–400%. For endpoint variants, the value is the sum of the GPU utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>GPUMemory Utilization</td>
<td>The percentage of GPU memory used by the containers on an instance. The value range is 0-100 and is multiplied by the number of GPUs. For example, if there are four GPUs, the GPUMemoryUtilization range is 0%-400%. For endpoint variants, the value is the sum of the GPU memory utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
<tr>
<td>DiskUtilization</td>
<td>The percentage of disk space used by the containers on an instance. This value range is 0%-100%. For endpoint variants, the value is the sum of the disk space utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
</tbody>
</table>

**Set Auto Scaling Policies for Multi-Model Endpoint Deployments**

SageMaker multi-model endpoints fully support automatic scaling, which manages replicas of models to ensure models scale based on traffic patterns. We recommend that you configure your multi-model endpoint and the size of your instances based on [Instance recommendations for multi-model endpoint deployments](#) and also set up instance based auto scaling for your endpoint. The invocation rates used to trigger an auto-scale event are based on the aggregate set of predictions across the full set of models served by the endpoint. For additional details on setting up endpoint auto scaling, see [Automatically Scale Amazon SageMaker Models](#).

You can set up auto scaling policies with predefined and custom metrics on both CPU and GPU backed multi-model endpoints.

**Note**

SageMaker multi-model endpoint metrics are available at one-minute granularity.
Define a scaling policy

To specify the metrics and target values for a scaling policy, you can configure a target-tracking scaling policy. You can use either a predefined metric or a custom metric.

Scaling policy configuration is represented by a JSON block. You save your scaling policy configuration as a JSON block in a text file. You use that text file when invoking the AWS CLI or the Application Auto Scaling API. For more information about policy configuration syntax, see TargetTrackingScalingPolicyConfiguration in the Application Auto Scaling API Reference.

The following options are available for defining a target-tracking scaling policy configuration.

Use a predefined metric

To quickly define a target-tracking scaling policy for a variant, use the SageMakerVariantInvocationsPerInstance predefined metric. SageMakerVariantInvocationsPerInstance is the average number of times per minute that each instance for a variant is invoked. We strongly recommend using this metric.

To use a predefined metric in a scaling policy, create a target tracking configuration for your policy. In the target tracking configuration, include a PredefinedMetricSpecification for the predefined metric and a TargetValue for the target value of that metric.

The following example is a typical policy configuration for target-tracking scaling for a variant. In this configuration, we use the SageMakerVariantInvocationsPerInstance predefined metric to adjust the number of variant instances so that each instance has an InvocationsPerInstance metric of 70.

```
{"TargetValue": 70.0,
   "PredefinedMetricSpecification":
   {
     "PredefinedMetricType": "InvocationsPerInstance"
   }
}
```

Note

We recommend that you use InvocationsPerInstance while using multi-model endpoints. The TargetValue for this metric depends on your application’s latency.
requirements. We also recommend that you load test your endpoints to set up suitable scaling parameter values. To learn more about load testing and setting up autoscaling for your endpoints, see the blog Configuring autoscaling inference endpoints in Amazon SageMaker.

### Use a custom metric

If you need to define a target-tracking scaling policy that meets your custom requirements, define a custom metric. You can define a custom metric based on any production variant metric that changes in proportion to scaling.

Not all SageMaker metrics work for target tracking. The metric must be a valid utilization metric, and it must describe how busy an instance is. The value of the metric must increase or decrease in inverse proportion to the number of variant instances. That is, the value of the metric should decrease when the number of instances increases.

⚠️ **Important**

Before deploying automatic scaling in production, you must test automatic scaling with your custom metric.

### Example custom metric for a CPU backed multi-model endpoint

The following example is a target-tracking configuration for a scaling policy. In this configuration, for a model named my-model, a custom metric of CPUUtilization adjusts the instance count on the endpoint based on an average CPU utilization of 50% across all instances.

```json
{"TargetValue": 50,
"CustomizedMetricSpecification":
{"MetricName": "CPUUtilization",
"Namespace": "/aws/sagemaker/Endpoints",
"Dimensions": [
{"Name": "EndpointName", "Value": "my-endpoint" },
{"Name": "ModelName","Value": "my-model"}
],
"Statistic": "Average",
"Unit": "Percent"}
}```
Example custom metric for a GPU backed multi-model endpoint

The following example is a target-tracking configuration for a scaling policy. In this configuration, for a model named `my-model`, a custom metric of `GPUUtilization` adjusts the instance count on the endpoint based on an average GPU utilization of 50% across all instances.

```
{"TargetValue": 50,
 "CustomizedMetricSpecification":
  {"MetricName": "GPUUtilization",
   "Namespace": "/aws/sagemaker/Endpoints",
   "Dimensions": [
     {"Name": "EndpointName", "Value": "my-endpoint" },
     {"Name": "ModelName","Value": "my-model"}
   ],
   "Statistic": "Average",
   "Unit": "Percent"
  }
}
```

Add a cooldown period

To add a cooldown period for scaling out your endpoint, specify a value, in seconds, for `ScaleOutCooldown`. Similarly, to add a cooldown period for scaling in your model, add a value, in seconds, for `ScaleInCooldown`. For more information about `ScaleInCooldown` and `ScaleOutCooldown`, see `TargetTrackingScalingPolicyConfiguration` in the Application Auto Scaling API Reference.

The following is an example target-tracking configuration for a scaling policy. In this configuration, the `SageMakerVariantInvocationsPerInstance` predefined metric is used to adjust scaling based on an average of 70 across all instances of that variant. The configuration provides a scale-in cooldown period of 10 minutes and a scale-out cooldown period of 5 minutes.

```
{"TargetValue": 70.0,
 "PredefinedMetricSpecification":
  {"PredefinedMetricType": "SageMakerVariantInvocationsPerInstance" },
  "ScaleInCooldown": 600,
  "ScaleOutCooldown": 300
}
```
Host multiple models which use different containers behind one endpoint

SageMaker multi-container endpoints enable customers to deploy multiple containers, that use different models or frameworks, on a single SageMaker endpoint. The containers can be run in a sequence as an inference pipeline, or each container can be accessed individually by using direct invocation to improve endpoint utilization and optimize costs.

For information about invoking the containers in a multi-container endpoint in sequence, see Host models along with pre-processing logic as serial inference pipeline behind one endpoint.

For information about invoking a specific container in a multi-container endpoint, see Use a multi-container endpoint with direct invocation

Topics

- Create a multi-container endpoint (Boto 3)
- Update a multi-container endpoint
- Delete a multi-container endpoint
- Use a multi-container endpoint with direct invocation

Create a multi-container endpoint (Boto 3)

Create a Multi-container endpoint by calling CreateModel, CreateEndpointConfig, and CreateEndpoint APIs as you would to create any other endpoints. You can run these containers sequentially as an inference pipeline, or run each individual container by using direct invocation.

Multi-container endpoints have the following requirements when you call create_model:

- Use the Containers parameter instead of PrimaryContainer, and include more than one container in the Containers parameter.
- The ContainerHostname parameter is required for each container in a multi-container endpoint with direct invocation.
- Set the Mode parameter of the InferenceExecutionConfig field to Direct for direct invocation of each container, or Serial to use containers as an inference pipeline. The default mode is Serial.

Note

Currently there is a limit of up to 15 containers supported on a multi-container endpoint.
The following example creates a multi-container model for direct invocation.

1. Create container elements and `InferenceExecutionConfig` with direct invocation.

   ```python
   container1 = {
       'Image': '123456789012.dkr.ecr.us-east-1.amazonaws.com/myimage1:mytag',
       'ContainerHostname': 'firstContainer'
   }

   container2 = {
       'Image': '123456789012.dkr.ecr.us-east-1.amazonaws.com/myimage2:mytag',
       'ContainerHostname': 'secondContainer'
   }

   inferenceExecutionConfig = {'Mode': 'Direct'}
   ```

2. Create the model with the container elements and set the `InferenceExecutionConfig` field.

   ```python
   import boto3
   sm_client = boto3.Session().client('sagemaker')

   response = sm_client.create_model(
       ModelName = 'my-direct-mode-model-name',
       InferenceExecutionConfig = inferenceExecutionConfig,
       ExecutionRoleArn = role,
       Containers = [container1, container2]
   )
   ```

To create an endpoint, you would then call `create_endpoint_config` and `create_endpoint` as you would to create any other endpoint.

**Update a multi-container endpoint**

To update a multi-container endpoint, complete the following steps.

1. Call `create_model` to create a new model with a new value for the Mode parameter in the `InferenceExecutionConfig` field.
2. Call `create_endpoint_config` to create a new endpoint config with a different name by using the new model you created in the previous step.

3. Call `update_endpoint` to update the endpoint with the new endpoint config you created in the previous step.

Delete a multi-container endpoint

To delete an endpoint, call `delete_endpoint`, and provide the name of the endpoint you want to delete as the `EndpointName` parameter.

Use a multi-container endpoint with direct invocation

SageMaker multi-container endpoints enable customers to deploy multiple containers to deploy different models on a SageMaker endpoint. You can host up to 15 different inference containers on a single endpoint. By using direct invocation, you can send a request to a specific inference container hosted on a multi-container endpoint.

Topics

- Invoke a multi-container endpoint with direct invocation
- Security with multi-container endpoints with direct invocation
- Metrics for multi-container endpoints with direct invocation
- Autoscale multi-container endpoints
- Troubleshoot multi-container endpoints

Invoke a multi-container endpoint with direct invocation

To invoke a multi-container endpoint with direct invocation, call `invoke_endpoint` as you would invoke any other endpoint, and specify which container you want to invoke by using the `TargetContainerHostname` parameter.

The following example directly invokes the `secondContainer` of a multi-container endpoint to get a prediction.

```python
import boto3
runtime_sm_client = boto3.Session().client('sagemaker-runtime')
response = runtime_sm_client.invoke_endpoint(
    TargetContainerHostname='secondContainer')
```
For each direct invocation request to a multi-container endpoint, only the container with the `TargetContainerHostname` processes the invocation request. You will get validation errors if you do any of the following:

- Specify a `TargetContainerHostname` that does not exist in the endpoint
- Do not specify a value for `TargetContainerHostname` in a request to an endpoint configured for direct invocation
- Specify a value for `TargetContainerHostname` in a request to an endpoint that is not configured for direct invocation.

**Security with multi-container endpoints with direct invocation**

For multi-container endpoints with direct invocation, there are multiple containers hosted in a single instance by sharing memory and a storage volume. It's your responsibility to use secure containers, maintain the correct mapping of requests to target containers, and provide users with the correct access to target containers. SageMaker uses IAM roles to provide IAM identity-based policies that you use to specify whether access to a resource is allowed or denied to that role, and under what conditions. For information about IAM roles, see IAM roles in the AWS Identity and Access Management User Guide. For information about identity-based policies, see Identity-based policies and resource-based policies.

By default, an IAM principal with InvokeEndpoint permissions on a multi-container endpoint with direct invocation can invoke any container inside the endpoint with the endpoint name that you specify when you call `invoke_endpoint`. If you need to restrict `invoke_endpoint` access to a limited set of containers inside a multi-container endpoint, use the `sagemaker:TargetContainerHostname` IAM condition key. The following policies show how to limit calls to specific containers within an endpoint.

The following policy allows `invoke_endpoint` requests only when the value of the `TargetContainerHostname` field matches one of the specified regular expressions.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        
```
The following policy denies `invoke_endpoint` requests when the value of the `TargetContainerHostname` field matches one of the specified regular expressions in the Deny statement.

```
{
   "Version": "2012-10-17",
   "Statement": [
   {
      "Action": [
         "sagemaker:InvokeEndpoint"
      ],
      "Effect": "Allow",
      "Condition": {
         "StringLike": {
            "sagemaker:TargetContainerHostname": ["customIps", "common"]
         }
      }
   },
   {
      "Action": [
         "sagemaker:InvokeEndpoint"
      ],
      "Effect": "Deny",
      "Condition": {
         "StringLike": {
            "sagemaker:TargetContainerHostname": ["special"]
         }
      }
   }
   ]
}
```
For information about SageMaker condition keys, see Condition Keys for SageMaker in the AWS Identity and Access Management User Guide.

## Metrics for multi-container endpoints with direct invocation

In addition to the endpoint metrics that are listed in Monitor Amazon SageMaker with Amazon CloudWatch, SageMaker also provides per-container metrics.

Per-container metrics for multi-container endpoints with direct invocation are located in CloudWatch and categorized into two namespaces: AWS/SageMaker and aws/sagemaker/Endpoints. The AWS/SageMaker namespace includes invocation-related metrics, and the aws/sagemaker/Endpoints namespace includes memory and CPU utilization metrics.

The following table lists the per-container metrics for multi-container endpoints with direct invocation. All the metrics use the [EndpointName, VariantName, ContainerName] dimension, which filters metrics at a specific endpoint, for a specific variant and corresponding to a specific container. These metrics share the same metric names as in those for inference pipelines, but at a per-container level [EndpointName, VariantName, ContainerName].

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Dimension</th>
<th>NameSpace</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invocations</td>
<td>The number of InvokeEndpoint requests sent to a container inside an endpoint. To get the total number of requests sent to that container, use the Sum statistic. Units: None. Valid statistics: Sum, Sample Count.</td>
<td>EndpointName, VariantName, ContainerName</td>
<td>AWS/SageMaker</td>
</tr>
<tr>
<td>Invocation4XX Errors</td>
<td>The number of <code>InvokeEndpoint</code> requests that the model returned a 4xx HTTP response code for on a specific container. For each 4xx response, SageMaker sends a 1. Units: None Valid statistics: Average, Sum</td>
<td>EndpointName, VariantName, ContainerName</td>
<td>AWS/SageMaker</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>Invocation5XX Errors</td>
<td>The number of <code>InvokeEndpoint</code> requests that the model returned a 5xx HTTP response code for on a specific container. For each 5xx response, SageMaker sends a 1. Units: None Valid statistics: Average, Sum</td>
<td>EndpointName, VariantName, ContainerName</td>
<td>AWS/SageMaker</td>
</tr>
<tr>
<td>Container Latency</td>
<td>The time it took for the target container to respond as viewed from SageMaker. Container Latency includes the time it took to send the request, to fetch the response from the model's container, and to complete inference in the container. Units: Microseconds Valid statistics: Average, Sum, Min, Max, Sample Count</td>
<td>EndpointName, VariantName, ContainerName</td>
<td>AWS/SageMaker</td>
</tr>
<tr>
<td>OverheadLatency</td>
<td>The time added to the time taken to respond to a client request by SageMaker for overhead. OverheadLatency is measured from the time that SageMaker receives the request until it returns a response to the client, minus the ModelLatency. Overhead latency can vary depending on request and response payload sizes, request frequency, and authentication or authorization of the request, among other factors. Units: Microseconds Valid statistics: Average, Sum, Min, Max, <code>Sample Count</code></td>
<td>EndpointName, VariantName, ContainerName</td>
<td>AWS/SageMaker</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
<td>Path</td>
<td></td>
</tr>
<tr>
<td>-----------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>CPUUtilization</td>
<td>The percentage of CPU units that are used by each container running on an instance. The value ranges from 0% to 100%, and is multiplied by the number of CPUs. For example, if there are four CPUs, CPUUtilization can range from 0% to 400%. For endpoints with direct invocation, the number of CPUUtilization metrics equals the number of containers in that endpoint. Units: Percent</td>
<td>aws/sagemaker/Endpoints</td>
<td></td>
</tr>
</tbody>
</table>
MemoryUtilization

The percentage of memory that is used by each container running on an instance. This value ranges from 0% to 100%. Similar as CPUUtilization, in endpoints with direct invocation, the number of MemoryUtilization metrics equals the number of containers in that endpoint. Units: Percent

| EndpointName, VariantName, ContainerName | aws/sagemaker/Endpoints |

All the metrics in the previous table are specific to multi-container endpoints with direct invocation. Besides these special per-container metrics, there are also metrics at the variant level with dimension [EndpointName, VariantName] for all the metrics in the table expect ContainerLatency.

**Autoscale multi-container endpoints**

If you want to configure automatic scaling for a multi-container endpoint using the InvocationsPerInstance metric, we recommend that the model in each container exhibits similar CPU utilization and latency on each inference request. This is recommended because if traffic to the multi-container endpoint shifts from a low CPU utilization model to a high CPU utilization model, but the overall call volume remains the same, the endpoint does not scale out and there may not be enough instances to handle all the requests to the high CPU utilization model. For information about automatically scaling endpoints, see [Automatically Scale Amazon SageMaker Models](#).

**Troubleshoot multi-container endpoints**

The following sections can help you troubleshoot errors with multi-container endpoints.
Ping Health Check Errors

With multiple containers, endpoint memory and CPU are under higher pressure during endpoint creation. Specifically, the MemoryUtilization and CPUUtilization metrics are higher than for single-container endpoints, because utilization pressure is proportional to the number of containers. Because of this, we recommend that you choose instance types with enough memory and CPU to ensure that there is enough memory on the instance to have all the models loaded (the same guidance applies to deploying an inference pipeline). Otherwise, your endpoint creation might fail with an error such as **XXX did not pass the ping health check**.

**Missing accept-bind-to-port=true Docker label**

The containers in a multi-container endpoints listen on the port specified in the SAGEMAKER_BIND_TO_PORT environment variable instead of port 8080. When a container runs in a multi-container endpoint, SageMaker automatically provides this environment variable to the container. If this environment variable isn't present, containers default to using port 8080. To indicate that your container complies with this requirement, use the following command to add a label to your Dockerfile:

```
LABEL com.amazonaws.sagemaker.capabilities.accept-bind-to-port=true
```

Otherwise, You will see an error message such as **Your Ecr Image XXX does not contain required com.amazonaws.sagemaker.capabilities.accept-bind-to-port=true Docker label(s)**.

If your container needs to listen on a second port, choose a port in the range specified by the SAGEMAKER_SAFE_PORT_RANGE environment variable. Specify the value as an inclusive range in the format **XXXX-YYYY**, where XXXX and YYYY are multi-digit integers. SageMaker provides this value automatically when you run the container in a multi-container endpoint.

**Host models along with pre-processing logic as serial inference pipeline behind one endpoint**

An **inference pipeline** is a Amazon SageMaker model that is composed of a linear sequence of two to fifteen containers that process requests for inferences on data. You use an inference pipeline to define and deploy any combination of pretrained SageMaker built-in algorithms and your own custom algorithms packaged in Docker containers. You can use an inference pipeline to combine
preprocessing, predictions, and post-processing data science tasks. Inference pipelines are fully managed.

You can add SageMaker Spark ML Serving and scikit-learn containers that reuse the data transformers developed for training models. The entire assembled inference pipeline can be considered as a SageMaker model that you can use to make either real-time predictions or to process batch transforms directly without any external preprocessing.

Within an inference pipeline model, SageMaker handles invocations as a sequence of HTTP requests. The first container in the pipeline handles the initial request, then the intermediate response is sent as a request to the second container, and so on, for each container in the pipeline. SageMaker returns the final response to the client.

When you deploy the pipeline model, SageMaker installs and runs all of the containers on each Amazon Elastic Compute Cloud (Amazon EC2) instance in the endpoint or transform job. Feature processing and inferences run with low latency because the containers are co-located on the same EC2 instances. You define the containers for a pipeline model using the `CreateModel` operation or from the console. Instead of setting one `PrimaryContainer`, you use the `Containers` parameter to set the containers that make up the pipeline. You also specify the order in which the containers are executed.

A pipeline model is immutable, but you can update an inference pipeline by deploying a new one using the `UpdateEndpoint` operation. This modularity supports greater flexibility during experimentation.

For information on how to create an inference pipeline with the SageMaker model registry, see Register and Deploy Models with Model Registry.

There are no additional costs for using this feature. You pay only for the instances running on an endpoint.

**Topics**

- Sample Notebooks for Inference Pipelines
- Feature Processing with Spark ML and Scikit-learn
- Create a Pipeline Model
- Run Real-time Predictions with an Inference Pipeline
- Run Batch Transforms with Inference Pipelines
Sample Notebooks for Inference Pipelines

For an example that shows how to create and deploy inference pipelines, see the Inference Pipeline with Scikit-learn and Linear Learner sample notebook. For instructions on creating and accessing Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances.

To see a list of all the SageMaker samples, after creating and opening a notebook instance, choose the SageMaker Examples tab. There are three inference pipeline notebooks. The first two inference pipeline notebooks just described are located in the advanced_functionality folder and the third notebook is in the sagemaker-python-sdk folder. To open a notebook, choose its Use tab, then choose Create copy.

Feature Processing with Spark ML and Scikit-learn

Before training a model with either Amazon SageMaker built-in algorithms or custom algorithms, you can use Spark and scikit-learn preprocessors to transform your data and engineer features.

Feature Processing with Spark ML

You can run Spark ML jobs with AWS Glue, a serverless ETL (extract, transform, load) service, from your SageMaker notebook. You can also connect to existing EMR clusters to run Spark ML jobs with Amazon EMR. To do this, you need an AWS Identity and Access Management (IAM) role that grants permission for making calls from your SageMaker notebook to AWS Glue.

Note

To see which Python and Spark versions AWS Glue supports, refer to AWS Glue Release Notes.

After engineering features, you package and serialize Spark ML jobs with MLeap into MLeap containers that you can add to an inference pipeline. You don't need to use externally managed Spark clusters. With this approach, you can seamlessly scale from a sample of rows to terabytes of data. The same transformers work for both training and inference, so you don't need to duplicate
preprocessing and feature engineering logic or develop a one-time solution to make the models persist. With inference pipelines, you don’t need to maintain outside infrastructure, and you can make predictions directly from data inputs.

When you run a Spark ML job on AWS Glue, a Spark ML pipeline is serialized into MLeap format. Then, you can use the job with the SparkML Model Serving Container in a SageMaker Inference Pipeline. MLeap is a serialization format and execution engine for machine learning pipelines. It supports Spark, Scikit-learn, and TensorFlow for training pipelines and exporting them to a serialized pipeline called an MLeap Bundle. You can deserialize Bundles back into Spark for batch-mode scoring or into the MLeap runtime to power real-time API services.

For an example that shows how to feature process with Spark ML, see the Train an ML Model using Apache Spark in Amazon EMR and deploy in SageMaker sample notebook.

Feature Processing with Scikit-Learn

You can run and package scikit-learn jobs into containers directly in Amazon SageMaker. For an example of Python code for building a scikit-learn featurizer model that trains on Fisher’s Iris flower data set and predicts the species of Iris based on morphological measurements, see IRIS Training and Prediction with Sagemaker Scikit-learn.

Create a Pipeline Model

To create a pipeline model that can be deployed to an endpoint or used for a batch transform job, use the Amazon SageMaker console or the CreateModel operation.

To create an inference pipeline (console)

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Models, and then choose Create models from the Inference group.
3. On the Create model page, provide a model name, choose an IAM role, and, if you want to use a private VPC, specify VPC values.
4. To add information about the containers in the inference pipeline, choose Add container, then choose Next.

5. Complete the fields for each container in the order that you want to execute them, up to the maximum of fifteen. Complete the Container input options, Location of inference code image, and, optionally, Location of model artifacts, Container host name, and Environmental variables fields.
Container definition 1

Container input options

- Provide model artifacts and inference image.

Provide model artifacts and inference image

Location of inference code image
The registry path where the inference code image is stored in Amazon ECR.

123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage:v1

Location of model artifacts - optional
The URL for the S3 location where model artifacts are stored.

s3://bucket/path-to-your-data/

The path must point to a single gzip compressed tar archive (tar.gz suffix).

Container host name - optional
The DNS host name for the container.

Container definition 2 - optional

Container input options

- Provide model artifacts and inference image.

Provide model artifacts and inference image

Location of inference code image
The registry path where the inference code image is stored in Amazon ECR.

123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage:v1

Location of model artifacts - optional
The URL for the S3 location where model artifacts are stored.

s3://bucket/path-to-your-data/

The path must point to a single gzip compressed tar archive (tar.gz suffix).

Container host name - optional
The DNS host name for the container.
The MyInferencePipelineModel page summarizes the settings for the containers that provide input for the model. If you provided the environment variables in a corresponding container definition, SageMaker shows them in the Environment variables field.
### Model settings

<table>
<thead>
<tr>
<th>Name</th>
<th>ARN</th>
<th>Creation time</th>
<th>IAM role ARN</th>
</tr>
</thead>
</table>

#### Container 1

<table>
<thead>
<tr>
<th>Container Name</th>
<th>Container 1</th>
<th>Model data URL</th>
<th>Scanning status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image</td>
<td>123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage/v1</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

**Environment variables**

<table>
<thead>
<tr>
<th>Key</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>key1</td>
<td>value1</td>
</tr>
<tr>
<td>key2</td>
<td>value2</td>
</tr>
</tbody>
</table>

#### Container 2

<table>
<thead>
<tr>
<th>Container Name</th>
<th>Container 2</th>
<th>Model data URL</th>
<th>Scanning status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image</td>
<td>123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage/v1</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

#### Container 3

<table>
<thead>
<tr>
<th>Container Name</th>
<th>Container 3</th>
<th>Model data URL</th>
<th>Scanning status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image</td>
<td>123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage/v1</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

#### Container 4

<table>
<thead>
<tr>
<th>Container Name</th>
<th>Container 4</th>
<th>Model data URL</th>
<th>Scanning status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image</td>
<td>123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage/v1</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

#### Container 5

<table>
<thead>
<tr>
<th>Container Name</th>
<th>Container 5</th>
<th>Model data URL</th>
<th>Scanning status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image</td>
<td>123456789012.dkr.ecr.us-east-2.amazonaws.com/myimage/v1</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

#### Network

No custom VPC settings applied.

#### Tags

<table>
<thead>
<tr>
<th>Key</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Run Real-time Predictions with an Inference Pipeline

You can use trained models in an inference pipeline to make real-time predictions directly without performing external preprocessing. When you configure the pipeline, you can choose to use the built-in feature transformers already available in Amazon SageMaker. Or, you can implement your own transformation logic using just a few lines of scikit-learn or Spark code.

**MLeap**, a serialization format and execution engine for machine learning pipelines, supports Spark, scikit-learn, and TensorFlow for training pipelines and exporting them to a serialized pipeline called an MLeap Bundle. You can deserialize Bundles back into Spark for batch-mode scoring or into the MLeap runtime to power real-time API services.

The containers in a pipeline listen on the port specified in the `SAGEMAKER_BIND_TO_PORT` environment variable (instead of 8080). When running in an inference pipeline, SageMaker automatically provides this environment variable to containers. If this environment variable isn't present, containers default to using port 8080. To indicate that your container complies with this requirement, use the following command to add a label to your Dockerfile:

```
LABEL com.amazonaws.sagemaker.capabilities.accept-bind-to-port=true
```

If your container needs to listen on a second port, choose a port in the range specified by the `SAGEMAKER_SAFE_PORT_RANGE` environment variable. Specify the value as an inclusive range in the format "**XXXX-YYYY**", where `XXXX` and `YYYY` are multi-digit integers. SageMaker provides this value automatically when you run the container in a multicontainer pipeline.

**Note**

To use custom Docker images in a pipeline that includes SageMaker built-in algorithms, you need an **Amazon Elastic Container Registry (Amazon ECR) policy**. Your Amazon ECR repository must grant SageMaker permission to pull the image. For more information, see **Troubleshoot Amazon ECR Permissions for Inference Pipelines**.

Create and Deploy an Inference Pipeline Endpoint

The following code creates and deploys a real-time inference pipeline model with SparkML and XGBoost models in series using the SageMaker SDK.

```python
from sagemaker.model import Model
from sagemaker.pipeline_model import PipelineModel
```
from sagemaker.sparkml.model import SparkMLModel

sparkml_data = 's3:///{}/{}{}'.format(s3_model_bucket, s3_model_key_prefix, 'model.tar.gz')
sparkml_model = SparkMLModel(model_data=sparkml_data)
xgb_model = Model(model_data=xgb_model.model_data, image=training_image)

model_name = 'serial-inference-' + timestamp_prefix
endpoint_name = 'serial-inference-ep-' + timestamp_prefix
sm_model = PipelineModel(name=model_name, role=role, models=[sparkml_model, xgb_model])
sm_model.deploy(initial_instance_count=1, instance_type='ml.c4.xlarge',
endpoint_name=endpoint_name)

### Request Real-Time Inference from an Inference Pipeline Endpoint

The following example shows how to make real-time predictions by calling an inference endpoint and passing a request payload in JSON format:

```python
import sagemaker
from sagemaker.predictor import json_serializer, json_deserializer, Predictor

payload = {
    "input": [
        {
            "name": "Pclass",
            "type": "float",
            "val": "1.0"
        },
        {
            "name": "Embarked",
            "type": "string",
            "val": "Q"
        },
        {
            "name": "Age",
            "type": "double",
            "val": "48.0"
        },
        {
            "name": "Fare",
            "type": "double",
            "val": "100.67"
        }
    ]
}
```

Hosting options 4164
The response you get from `predictor.predict(payload)` is the model's inference result.

**Realtime inference pipeline example**

You can run this example notebook using the SKLearn predictor that shows how to deploy an endpoint, run an inference request, then deserialize the response. Find this notebook and more examples in the Amazon SageMaker example GitHub repository.

**Run Batch Transforms with Inference Pipelines**

To get inferences on an entire dataset you run a batch transform on a trained model. To run inferences on a full dataset, you can use the same inference pipeline model created and deployed to an endpoint for real-time processing in a batch transform job. To run a batch transform job in a pipeline, you download the input data from Amazon S3 and send it in one or more HTTP requests to the inference pipeline model. For an example that shows how to prepare data for a batch transform, see "Section 2 - Preprocess the raw housing data using Scikit Learn" of the Amazon SageMaker Multi-Model Endpoints using Linear Learner sample notebook. For information about Amazon SageMaker batch transforms, see Use Batch Transform.
Note

To use custom Docker images in a pipeline that includes Amazon SageMaker built-in algorithms, you need an Amazon Elastic Container Registry (ECR) policy. Your Amazon ECR repository must grant SageMaker permission to pull the image. For more information, see Troubleshoot Amazon ECR Permissions for Inference Pipelines.

The following example shows how to run a transform job using the Amazon SageMaker Python SDK. In this example, model_name is the inference pipeline that combines SparkML and XGBoost models (created in previous examples). The Amazon S3 location specified by input_data_path contains the input data, in CSV format, to be downloaded and sent to the Spark ML model. After the transform job has finished, the Amazon S3 location specified by output_data_path contains the output data returned by the XGBoost model in CSV format.

```python
import sagemaker
input_data_path = 's3://{}/{}/*'.format(default_bucket, 'key', 'file_name')
output_data_path = 's3://{}/*'.format(default_bucket, 'key')
transform_job = sagemaker.transformer.Transformer(
    model_name = model_name,
    instance_count = 1,
    instance_type = 'ml.m4.xlarge',
    strategy = 'SingleRecord',
    assemble_with = 'Line',
    output_path = output_data_path,
    base_transform_job_name='inference-pipelines-batch',
    sagemaker_session=sagemaker.Session(),
    accept = CONTENT_TYPE_CSV)
transform_job.transform(data = input_data_path,
    content_type = CONTENT_TYPE_CSV,
    split_type = 'Line')
```

Inference Pipeline Logs and Metrics

Monitoring is important for maintaining the reliability, availability, and performance of Amazon SageMaker resources. To monitor and troubleshoot inference pipeline performance, use Amazon CloudWatch logs and error messages. For information about the monitoring tools that SageMaker provides, see Monitor AWS resources provisioned while using Amazon SageMaker.
Use Metrics to Monitor Multi-container Models

To monitor the multi-container models in Inference Pipelines, use Amazon CloudWatch. CloudWatch collects raw data and processes it into readable, near real-time metrics. SageMaker training jobs and endpoints write CloudWatch metrics and logs in the AWS/SageMaker namespace.

The following tables list the metrics and dimensions for the following:

- Endpoint invocations
- Training jobs, batch transform jobs, and endpoint instances

A *dimension* is a name/value pair that uniquely identifies a metric. You can assign up to 10 dimensions to a metric. For more information on monitoring with CloudWatch, see Monitor Amazon SageMaker with Amazon CloudWatch.

Endpoint Invocation Metrics

The AWS/SageMaker namespace includes the following request metrics from calls to `InvokeEndpoint`.

Metrics are reported at a 1-minute intervals.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invocation4XXErrors</td>
<td>The number of <code>InvokeEndpoint</code> requests that the model returned a 4xx HTTP response code for. For each 4xx response, SageMaker sends a 1.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>Invocation5XXErrors</td>
<td>The number of <code>InvokeEndpoint</code> requests that the model returned a 5xx HTTP response code for. For each 5xx response, SageMaker sends a 1.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Invocations</td>
<td>The number of InvokeEndpoint requests sent to a model endpoint. To get the total number of requests sent to a model endpoint, use the Sum statistic. Units: None Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>InvocationsPerInstance</td>
<td>The number of endpoint invocations sent to a model, normalized by InstanceCount in each ProductionVariant. SageMaker sends 1/numberOfInstances as the value for each request, where numberOfInstances is the number of active instances for the ProductionVariant at the endpoint at the time of the request. Units: None Valid statistics: Sum</td>
</tr>
<tr>
<td>ModelLatency</td>
<td>The time the model or models took to respond. This includes the time it took to send the request, to fetch the response from the model container, and to complete the inference in the container. ModelLatency is the total time taken by all containers in an inference pipeline. Units: Microseconds Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>OverheadLatency</td>
<td>The time added to the time taken to respond to a client request by SageMaker for overhead. OverheadLatency is measured from the time that SageMaker receives the request until it returns a response to the client, minus the ModelLatency. Overhead latency can vary depending on request and response payload sizes, request frequency, and authentication or authorization of the request, among other factors.</td>
</tr>
<tr>
<td>Units: Microseconds</td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ContainerLatency</td>
<td>The time it took for an Inference Pipelines container to respond as viewed from SageMaker. ContainerLatency includes the time it took to send the request, to fetch the response from the model's container, and to complete inference in the container.</td>
</tr>
<tr>
<td>Units: Microseconds</td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>

**Dimensions for Endpoint Invocation Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EndpointName, VariantName, ContainerName</td>
<td>Filters endpoint invocation metrics for a ProductionVariant at the specified endpoint and for the specified variant.</td>
</tr>
</tbody>
</table>

For an inference pipeline endpoint, CloudWatch lists per-container latency metrics in your account as **Endpoint Container Metrics** and **Endpoint Variant Metrics** in the **SageMaker** namespace, as follows. The ContainerLatency metric appears only for inferences pipelines.
For each endpoint and each container, latency metrics display names for the container, endpoint, variant, and metric.

Training Job, Batch Transform Job, and Endpoint Instance Metrics

The namespaces `/aws/sagemaker/TrainingJobs`, `/aws/sagemaker/TransformJobs`, and `/aws/sagemaker/Endpoints` include the following metrics for training jobs and endpoint instances.

Metrics are reported at a 1-minute intervals.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPUUtilization</td>
<td>The percentage of CPU units that are used by the containers running on an instance. The value ranges from 0% to 100%, and is multiplied by the number of CPUs. For example, if there are four CPUs, CPUUtilization can range from 0% to 400%. For training jobs, CPUUtilization is the CPU utilization of the algorithm container running on the instance.</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>CPUUtilization</td>
<td>For batch transform jobs, <strong>CPUUtilization</strong> is the CPU utilization of the transform container running on the instance.</td>
</tr>
<tr>
<td></td>
<td>For multi-container models, <strong>CPUUtilization</strong> is the sum of CPU utilization by all containers running on the instance.</td>
</tr>
<tr>
<td></td>
<td>For endpoint variants, <strong>CPUUtilization</strong> is the sum of CPU utilization by all of the containers running on the instance.</td>
</tr>
<tr>
<td></td>
<td>Units: Percent</td>
</tr>
<tr>
<td>MemoryUtilization</td>
<td>The percentage of memory that is used by the containers running on an instance. This value ranges from 0% to 100%.</td>
</tr>
<tr>
<td></td>
<td>For training jobs, <strong>MemoryUtilization</strong> is the memory used by the algorithm container running on the instance.</td>
</tr>
<tr>
<td></td>
<td>For batch transform jobs, <strong>MemoryUtilization</strong> is the memory used by the transform container running on the instance.</td>
</tr>
<tr>
<td></td>
<td>For multi-container models, <strong>MemoryUtilization</strong> is the sum of memory used by all containers running on the instance.</td>
</tr>
<tr>
<td></td>
<td>For endpoint variants, <strong>MemoryUtilization</strong> is the sum of memory used by all of the containers running on the instance.</td>
</tr>
<tr>
<td></td>
<td>Units: Percent</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>GPUUtilization</td>
<td>The percentage of GPU units that are used by the containers running on an instance. GPUUtilization ranges from 0% to 100% and is multiplied by the number of GPUs. For example, if there are four GPUs, GPUUtilization can range from 0% to 400%. For training jobs, GPUUtilization is the GPU used by the algorithm container running on the instance. For batch transform jobs, GPUUtilization is the GPU used by the transform container running on the instance. For multi-container models, GPUUtilization is the sum of GPU used by all containers running on the instance. For endpoint variants, GPUUtilization is the sum of GPU used by all of the containers running on the instance. Units: Percent</td>
</tr>
<tr>
<td>GPUMemoryUtilization</td>
<td>The percentage of GPU memory used by the containers running on an instance. GPUMemoryUtilization ranges from 0% to 100% and is multiplied by the number of GPUs. For example, if there are four GPUs, GPUMemoryUtilization can range from 0% to 400%. For training jobs, GPUMemoryUtilization is the GPU memory used by the algorithm container running on the instance. For batch transform jobs, GPUMemoryUtilization is the GPU memory used by the transform container running on the instance. For multi-container models, GPUMemoryUtilization is sum of GPU used by all containers running on the instance. For endpoint variants, GPUMemoryUtilization is the sum of the GPU memory used by all of the containers running on the instance. Units: Percent</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>DiskUtilization</td>
<td>The percentage of disk space used by the containers running on an instance. DiskUtilization ranges from 0% to 100%. This metric is not supported for batch transform jobs.</td>
</tr>
</tbody>
</table>

For training jobs, DiskUtilization is the disk space used by the algorithm container running on the instance.

For endpoint variants, DiskUtilization is the sum of the disk space used by all of the provided containers running on the instance.

Units: Percent

Dimensions for Training Job, Batch Transform Job, and Endpoint Instance Metrics

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Host</td>
<td>For training jobs, Host has the format <code>[training-job-name]/algo-[instance-number-in-cluster]</code>. Use this dimension to filter instance metrics for the specified training job and instance. This dimension format is present only in the <code>/aws/sagemaker/TrainingJobs</code> namespace.</td>
</tr>
</tbody>
</table>

For batch transform jobs, Host has the format `[transform-job-name]/[instance-id]`. Use this dimension to filter instance metrics for the specified batch transform job and instance. This dimension format is present only in the `/aws/sagemaker/TransformJobs` namespace.

For endpoints, Host has the format `[endpoint-name]/[production-variant-name]/[instance-id]`. Use this dimension to filter instance metrics for the specified endpoint, variant, and instance. This dimension format is present only in the `/aws/sagemaker/Endpoints` namespace.
To help you debug your training jobs, endpoints, and notebook instance lifecycle configurations, SageMaker also sends anything an algorithm container, a model container, or a notebook instance lifecycle configuration sends to stdout or stderr to Amazon CloudWatch Logs. You can use this information for debugging and to analyze progress.

**Use Logs to Monitor an Inference Pipeline**

The following table lists the log groups and log streams SageMaker sends to Amazon CloudWatch.

A log stream is a sequence of log events that share the same source. Each separate source of logs into CloudWatch makes up a separate log stream. A log group is a group of log streams that share the same retention, monitoring, and access control settings.

**Logs**

<table>
<thead>
<tr>
<th>Log Group Name</th>
<th>Log Stream Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>/aws/sagemaker/</td>
<td>[training-job-name]/algo-[instance-number-in-cluster]-[epoch_timestamp]</td>
</tr>
<tr>
<td>TrainingJobs</td>
<td></td>
</tr>
<tr>
<td>/aws/sagemaker/</td>
<td>[production-variant-name]/[instance-id]</td>
</tr>
<tr>
<td>Endpoints/[EndpointName]</td>
<td></td>
</tr>
<tr>
<td>/aws/sagemaker/</td>
<td>[production-variant-name]/[instance-id]</td>
</tr>
<tr>
<td>TransformJobs</td>
<td>[production-variant-name]/[instance-id]/[container-name provided in the SageMaker model] (For Inference Pipelines) For Inference Pipelines logs, if you don't provide container names, CloudWatch uses <strong>container-1, container-2</strong>, and so on, in the order that containers are provided in the model.</td>
</tr>
<tr>
<td>NotebookInstances</td>
<td>[notebook-instance-name]/[LifecycleConfigHook]</td>
</tr>
<tr>
<td>/aws/sagemaker/</td>
<td>[transform-job-name]/[instance-id]-[epoch_timestamp]</td>
</tr>
<tr>
<td>TransformJobs</td>
<td>[transform-job-name]/[instance-id]-[epoch_timestamp]/data-log</td>
</tr>
</tbody>
</table>

Hosting options
### Log Group Name

<table>
<thead>
<tr>
<th>Log Stream Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>[transform-job-name]/[instance-id]-[epoch_timestamp]/[container-name provided in the SageMaker model] (For Inference Pipelines) For Inference Pipelines logs, if you don't provide container names, CloudWatch uses <strong>container-1, container-2</strong>, and so on, in the order that containers are provided in the model.</td>
</tr>
</tbody>
</table>

### Note

SageMaker creates the /aws/sagemaker/NotebookInstances log group when you create a notebook instance with a lifecycle configuration. For more information, see [Customize a Notebook Instance Using a Lifecycle Configuration Script](#).

For more information about SageMaker logging, see [Log Amazon SageMaker Events with Amazon CloudWatch](#).

### Troubleshoot Inference Pipelines

To troubleshoot inference pipeline issues, use CloudWatch logs and error messages. If you are using custom Docker images in a pipeline that includes Amazon SageMaker built-in algorithms, you might also encounter permissions problems. To grant the required permissions, create an Amazon Elastic Container Registry (Amazon ECR) policy.

### Topics

- [Troubleshoot Amazon ECR Permissions for Inference Pipelines](#)
- [Use CloudWatch Logs to Troubleshoot SageMaker Inference Pipelines](#)
- [Use Error Messages to Troubleshoot Inference Pipelines](#)

### Troubleshoot Amazon ECR Permissions for Inference Pipelines

When you use custom Docker images in a pipeline that includes [SageMaker built-in algorithms](#), you need an [Amazon ECR policy](#). The policy allows your Amazon ECR repository to grant permission for SageMaker to pull the image. The policy must add the following permissions:
Use CloudWatch Logs to Troubleshoot SageMaker Inference Pipelines

SageMaker publishes the container logs for endpoints that deploy an inference pipeline to Amazon CloudWatch at the following path for each container.

/aws/sagemaker/Endpoints/{EndpointName}/{Variant}/{InstanceId}/{ContainerHostname}

For example, logs for this endpoint are published to the following log groups and streams:

**EndpointName:** MyInferencePipelinesEndpoint  
**Variant:** MyInferencePipelinesVariant  
**InstanceId:** i-0179208609ff7e488  
**ContainerHostname:** MyContainerName1 and MyContainerName2

**logGroup:** /aws/sagemaker/Endpoints/MyInferencePipelinesEndpoint  
**logStream:** MyInferencePipelinesVariant/i-0179208609ff7e488/MyContainerName1  
**logStream:** MyInferencePipelinesVariant/i-0179208609ff7e488/MyContainerName2

A log stream is a sequence of log events that share the same source. Each separate source of logs into CloudWatch makes up a separate log stream. A log group is a group of log streams that share the same retention, monitoring, and access control settings.
To see the log groups and streams

2. In the navigation page, choose Logs.
3. In Log Groups, filter on MyInferencePipelinesEndpoint:

   ![Filtering Log Groups](image)

4. To see the log streams, on the CloudWatch Log Groups page, choose MyInferencePipelinesEndpoint, and then Search Log Group.

   ![Searching Log Group](image)

For a list of the logs that SageMaker publishes, see Inference Pipeline Logs and Metrics.

Use Error Messages to Troubleshoot Inference Pipelines

The inference pipeline error messages indicate which containers failed.

If an error occurs while SageMaker is invoking an endpoint, the service returns a ModelError (error code 424), which indicates which container failed. If the request payload (the response from the previous container) exceeds the limit of 5 MB, SageMaker provides a detailed error message, such as:

```
Received response from MyContainerName1 with status code 200. However, the request payload from MyContainerName1 to MyContainerName2 is 6000000 bytes, which has exceeded the maximum limit of 5 MB.
```

If a container fails the ping health check while SageMaker is creating an endpoint, it returns a ClientError and indicates all of the containers that failed the ping check in the last health check.
Delete Endpoints and Resources

Delete endpoints to stop incurring charges.

Delete Endpoint

Delete your endpoint programmatically using AWS SDK for Python (Boto3), with the AWS CLI, or interactively using the SageMaker console.

SageMaker frees up all of the resources that were deployed when the endpoint was created. Deleting an endpoint will not delete the endpoint configuration or the SageMaker model. See [Delete Endpoint Configuration](#) and [Delete Model](#) for information on how to delete your endpoint configuration and SageMaker model.

AWS SDK for Python (Boto3)

Use the `DeleteEndpoint` API to delete your endpoint. Specify the name of your endpoint for the `EndpointName` field.

```python
import boto3

# Specify your AWS Region
aws_region='<aws_region>'

# Specify the name of your endpoint
endpoint_name='<endpoint_name>'

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Delete endpoint
sagemaker_client.delete_endpoint(EndpointName=endpoint_name)
```

AWS CLI

Use the `delete-endpoint` command to delete your endpoint. Specify the name of your endpoint for the `endpoint-name` flag.

```bash
aws sagemaker delete-endpoint --endpoint-name <endpoint-name>
```

SageMaker Console

Delete your endpoint interactively with the SageMaker console.
1. In the SageMaker console at https://console.aws.amazon.com/sagemaker/ navigation menu, choose **Inference**.

2. Choose **Endpoints** from the drop down menu. A list of endpoints created in you AWS account will appear by name, Amazon Resource Name (ARN), creation time, status, and a time stamp of when the endpoint was last updated.

3. Select the endpoint you want to delete.

4. Select the **Actions** dropdown button in the top right corner.

5. Choose **Delete**.

**Delete Endpoint Configuration**

Delete your endpoint configuration programmaticially using AWS SDK for Python (Boto3), with the AWS CLI, or interactively using the SageMaker console. Deleting an endpoint configuration does not delete endpoints created using this configuration. See [Delete Endpoint](#) for information on how to delete your endpoint.

Do not delete an endpoint configuration in use by an endpoint that is live or while the endpoint is being updated or created. You might lose visibility into the instance type the endpoint is using if you delete the endpoint configuration of an endpoint that is active or being created or updated.

**AWS SDK for Python (Boto3)**

Use the [DeleteEndpointConfig](#) API to delete your endpoint. Specify the name of your endpoint configuration for the `EndpointConfigName` field.

```python
import boto3

# Specify your AWS Region
aws_region='<aws_region>'

# Specify the name of your endpoint configuration
endpoint_config_name='<endpoint_name>'

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Delete endpoint configuration
sagemaker_client.delete_endpoint_config(EndpointConfigName=endpoint_config_name)
```
You can optionally use the `DescribeEndpointConfig` API to return information about the name of the your deployed models (production variants) such as the name of your model and the name of the endpoint configuration associated with that deployed model. Provide the name of your endpoint for the `EndpointConfigName` field.

```python
# Specify the name of your endpoint
don_endpoint_name='<endpoint_name>'

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Store DescribeEndpointConfig response into a variable that we can index in the next step.
response =
sagemaker_client.describe_endpoint_config(EndpointConfigName=endpoint_name)

# Delete endpoint
endpoint_config_name = response['ProductionVariants'][0]['EndpointConfigName']

# Delete endpoint configuration
sagemaker_client.delete_endpoint_config(EndpointConfigName=endpoint_config_name)
```

For more information about other response elements returned by `DescribeEndpointConfig`, see `DescribeEndpointConfig` in the SageMaker API Reference guide.

**AWS CLI**

Use the `delete-endpoint-config` command to delete your endpoint configuration. Specify the name of your endpoint configuration for the `endpoint-config-name` flag.

```bash
aws sagemaker delete-endpoint-config \
    --endpoint-config-name <endpoint-config-name>
```

You can optionally use the `describe-endpoint-config` command to return information about the name of the your deployed models (production variants) such as the name of your model and the name of the endpoint configuration associated with that deployed model. Provide the name of your endpoint for the `endpoint-config-name` flag.

```bash
aws sagemaker describe-endpoint-config --endpoint-config-name <endpoint-config-name>
```
This will return a JSON response. You can copy and paste, use a JSON parser, or use a tool built for JSON parsing to obtain the endpoint configuration name associated with that endpoint.

**SageMaker Console**

Delete your endpoint configuration interactively with the SageMaker console.

1. In the SageMaker console at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/) navigation menu, choose **Inference**.
2. Choose **Endpoint configurations** from the dropdown menu. A list of endpoint configurations created in you AWS account will appear by name, Amazon Resource Name (ARN), and creation time.
3. Select the endpoint configuration you want to delete.
4. Select the **Actions** dropdown button in the top right corner.
5. Choose **Delete**.

**Delete Model**

Delete your SageMaker model programmatically using AWS SDK for Python (Boto3), with the AWS CLI, or interactively using the SageMaker console. Deleting a SageMaker model only deletes the model entry that was created in SageMaker. Deleting a model does not delete model artifacts, inference code, or the IAM role that you specified when creating the model.

**AWS SDK for Python (Boto3)**

Use the **DeleteModel** API to delete your SageMaker model. Specify the name of your model for the **ModelName** field.

```python
import boto3

# Specify your AWS Region
aws_region='<aws_region>'

# Specify the name of your endpoint configuration
model_name='<model_name>'

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Delete model
```
You can optionally use the `DescribeEndpointConfig` API to return information about the name of the your deployed models (production variants) such as the name of your model and the name of the endpoint configuration associated with that deployed model. Provide the name of your endpoint for the `EndpointConfigName` field.

```python
# Specify the name of your endpoint
default_endpoint_name='<endpoint_name>'

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Store DescribeEndpointConfig response into a variable that we can index in the next step.
response =
    sagemaker_client.describe_endpoint_config(EndpointConfigName=default_endpoint_name)

# Delete endpoint
model_name = response['ProductionVariants'][0]['ModelName']
sagemaker_client.delete_model(ModelName=model_name)
```

For more information about other response elements returned by `DescribeEndpointConfig`, see `DescribeEndpointConfig` in the SageMaker API Reference guide.

**AWS CLI**

Use the `delete-model` command to delete your SageMaker model. Specify the name of your model for the `model-name` flag.

```bash
aws sagemaker delete-model \
    --model-name <model-name>
```

You can optionally use the `describe-endpoint-config` command to return information about the name of the your deployed models (production variants) such as the name of your model and the name of the endpoint configuration associated with that deployed model. Provide the name of your endpoint for the `endpoint-config-name` flag.

```bash
aws sagemaker describe-endpoint-config --endpoint-config-name <endpoint-config-name>
```
This will return a JSON response. You can copy and paste, use a JSON parser, or use a tool built for JSON parsing to obtain the name of the model associated with that endpoint.

### SageMaker Console

Delete your SageMaker model interactively with the SageMaker console.

1. In the SageMaker console at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/) navigation menu, choose **Inference**.

2. Choose **Models** from the dropdown menu. A list of models created in your AWS account will appear by name, Amazon Resource Name (ARN), and creation time.

3. Select the model you want to delete.

4. Select the **Actions** dropdown button in the top right corner.

5. Choose **Delete**.

### Automatically Scale Amazon SageMaker Models

Amazon SageMaker supports automatic scaling (auto scaling) for your hosted models. *Auto scaling* dynamically adjusts the number of instances provisioned for a model in response to changes in your workload. When the workload increases, auto scaling brings more instances online. When the workload decreases, auto scaling removes unnecessary instances so that you don't pay for provisioned instances that you aren't using.

**Topics**

- [Auto scaling overview](#)
- [Configure model auto scaling with the console](#)
- [Register a model](#)
- [Define a scaling policy](#)
- [Apply a scaling policy](#)
- [Edit a scaling policy](#)
- [Delete a scaling policy](#)
- [Check the status of a scaling activity by describing scaling activities](#)
- [Load testing your auto scaling configuration](#)
- [Use AWS CloudFormation to create a scaling policy](#)
- [Update or delete endpoints that use auto scaling](#)
Auto scaling overview

The following overview provides details on the prerequisites and components used for auto scaling.

Topics

• Prerequisites
• Scaling policy overview
• Scale based on a schedule
• Minimum and maximum scaling limits
• Cooldown period
• Permissions
• Service-linked role
• Related resources

Prerequisites

Before you can use auto scaling, you must have already created an Amazon SageMaker model endpoint. You can have multiple model versions for the same endpoint. Each model is referred to as a production (model) variant. For more information about deploying a model endpoint, see Deploy the Model to SageMaker Hosting Services.

To activate auto scaling for a model, you can use the SageMaker console, the AWS Command Line Interface (AWS CLI), or an AWS SDK through the Application Auto Scaling API.

• If this is your first time configuring scaling for a model, we recommend you Configure model auto scaling with the console.

• When using the AWS CLI or the Application Auto Scaling API, the flow is to register the model as a scalable target, define the scaling policy, and then apply it. On the SageMaker console, under Inference in the navigation pane, choose Endpoints. Find your model's endpoint name and then choose it to find the variant name. You must specify both the endpoint name and the variant name to activate auto scaling for a model.

Scaling policy overview

To use auto scaling, you define a scaling policy that adds and removes the number of instances for your production variant in response to actual workloads.
To automatically scale as workload changes occur, you have two options: target tracking and step scaling policies.

We recommend using target tracking scaling policies. With target tracking, you choose an Amazon CloudWatch metric and target value. Auto scaling creates and manages the CloudWatch alarms for the scaling policy and calculates the scaling adjustment based on the metric and the target value. The policy adds and removes the number of instances as required to keep the metric at, or close to, the specified target value. For example, a scaling policy that uses the predefined `InvocationsPerInstance` metric with a target value of 70 can keep `InvocationsPerInstance` at, or close to 70. For more information, see Target tracking scaling policies in the Application Auto Scaling User Guide.

You can use step scaling when you require an advanced configuration, such as specifying how many instances to deploy under what conditions. Otherwise, using target tracking scaling is preferred as it will be fully automated. Note that step scaling can be managed only from the AWS CLI or the Application Auto Scaling API. For an overview of step scaling policies and how they work, see Step scaling policies in the Application Auto Scaling User Guide.

To create a target tracking scaling policy, you specify the following:

- **Metric** — The CloudWatch metric to track, such as average number of invocations per instance.
- **Target value** — The target value for the metric, such as 70 invocations per instance per minute.

You can create target tracking scaling policies with either predefined metrics or custom metrics. A predefined metric is defined in an enumeration so that you can specify it by name in code or use it in the SageMaker console. Alternatively, you can use either the AWS CLI or the Application Auto Scaling API to apply a target tracking scaling policy based on a predefined or custom metric.

Note that scaling activities are performed with cooldown periods between them to prevent rapid fluctuations in capacity. You can optionally configure the cooldown periods for your scaling policy.

**Scale based on a schedule**

You can also create scheduled actions to perform scaling activities at specific times. You can create scheduled actions that scale one time only or that scale on a recurring schedule. After a scheduled action runs, your scaling policy can continue to make decisions about whether to scale dynamically as workload changes occur. Scheduled scaling can be managed only from the AWS CLI or the Application Auto Scaling API. For more information, see Scheduled scaling in the Application Auto Scaling User Guide.
Minimum and maximum scaling limits

When configuring auto scaling, you must specify your scaling limits before creating a scaling policy. You set limits separately for the minimum and maximum values.

The minimum value must be at least 1, and equal to or less than the value specified for the maximum value.

The maximum value must be equal to or greater than the value specified for the minimum value. SageMaker auto scaling does not enforce a limit for this value.

To determine the scaling limits that you need for typical traffic, test your auto scaling configuration with the expected rate of traffic to your model.

If a variant's traffic becomes zero, SageMaker automatically scales in to the minimum number of instances specified. In this case, SageMaker emits metrics with a value of zero.

There are three options for specifying the minimum and maximum capacity:

1. Use the console to update the **Minimum instance count** and **Maximum instance count** settings.
2. Use the AWS CLI and include the `--min-capacity` and `--max-capacity` options when running the `register-scalable-target` command.
3. Call the `RegisterScalableTarget` API and specify the `MinCapacity` and `MaxCapacity` parameters.

Tip

You can manually scale out by increasing the minimum value, or manually scale in by decreasing the maximum value.

Cooldown period

A *cooldown period* is used to protect against over-scaling when your model is scaling in (reducing capacity) or scaling out (increasing capacity). It does this by slowing down subsequent scaling activities until the period expires. Specifically, it blocks the deletion of instances for scale-in requests, and limits the creation of instances for scale-out requests. For more information, see [Define cooldown periods](https://docs.aws.amazon.com/autoscaling/application/userguide/define-cooldown-periods.html) in the *Application Auto Scaling User Guide*.

You configure the cooldown period in your scaling policy.
If you don't specify a scale-in or a scale-out cooldown period, your scaling policy uses the default, which is 300 seconds for each.

If instances are being added or removed too quickly when you test your scaling configuration, consider increasing this value. You might see this behavior if the traffic to your model has a lot of spikes, or if you have multiple scaling policies defined for a variant.

If instances are not being added quickly enough to address increased traffic, consider decreasing this value.

Permissions

Auto scaling is made possible by a combination of the Amazon SageMaker, Amazon CloudWatch, and Application Auto Scaling APIs. For information about the minimum required permissions, see Application Auto Scaling identity-based policy examples in the Application Auto Scaling User Guide.

The SagemakerFullAccessPolicy IAM policy has all the IAM permissions required to perform auto scaling. For more information about SageMaker IAM permissions, see SageMaker Roles.

If you manage your own permission policy, you must include the following permissions:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:DescribeEndpoint",
                "sagemaker:DescribeEndpointConfig",
                "sagemaker:UpdateEndpointWeightsAndCapacities"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "application-autoscaling:*"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": "iam:CreateServiceLinkedRole",
            "Resource": "*"
        }
    ]
}
```
Auto scaling uses the `AWSServiceRoleForApplicationAutoScaling_SageMakerEndpoint` service-linked role. This service-linked role grants Application Auto Scaling permission to describe the alarms for your policies, to monitor current capacity levels, and to scale the target resource. This role is created for you automatically. For automatic role creation to succeed, you must have permission for the `iam:CreateServiceLinkedRole` action. For more information, see Service-linked roles in the Application Auto Scaling User Guide.

Related resources

For more information about configuring auto scaling, see the following resources:

- [application-autoscaling](https://docs.aws.amazon.com/cli/latest/reference/application-autoscaling/) section of the AWS CLI Command Reference
- [Application Auto Scaling API Reference](https://docs.aws.amazon.com/autoscaling/application/latest/api/index.html)
- [Application Auto Scaling User Guide](https://docs.aws.amazon.com/autoscaling/application/latest/userguide/)

Note

SageMaker recently introduced new inference capabilities built on real-time inference endpoints. You create a SageMaker endpoint with an endpoint configuration that defines
the instance type and initial instance count for the endpoint. Then, create an inference component, which is a SageMaker hosting object that you can use to deploy a model to an endpoint. For information about scaling inference components, see SageMaker adds new inference capabilities to help reduce foundation model deployment costs and latency and Reduce model deployment costs by 50% on average using the latest features of SageMaker on the AWS Blog.

Configure model auto scaling with the console

To configure auto scaling for a model (console)

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the navigation pane, choose Inference, and then choose Endpoints.
3. Choose your endpoint, and then for Endpoint runtime settings, choose the variant.
4. Choose Configure auto scaling.
5. On the Configure variant automatic scaling page, for Variant automatic scaling, do the following:
   a. For Minimum instance count, type the minimum number of instances that you want the scaling policy to maintain. At least 1 instance is required.
   b. For Maximum instance count, type the maximum number of instances that you want the scaling policy to maintain.
6. For Built-in scaling policy, do the following:
   a. For the Target metric, SageMakerVariantInvocationsPerInstance is automatically selected for the metric and cannot be changed.
   b. For the Target value, type the average number of invocations per instance per minute for the model. To determine this value, follow the guidelines in Load testing.
   c. (Optional) For Scale-in cool down (seconds) and Scale-out cool down (seconds), enter the amount of time, in seconds, for each cool down period.
   d. (Optional) Select Disable scale in if you don’t want auto scaling to terminate instances when traffic decreases.
7. Choose Save.
This procedure registers a model as a scalable target with Application Auto Scaling. When you register a model, Application Auto Scaling performs validation checks to ensure the following:

- The model exists
- The permissions are sufficient
- You aren’t registering a variant with an instance that is a burstable performance instance such as T2

**Note**

SageMaker doesn't support auto scaling for burstable instances such as T2, because they already allow for increased capacity under increased workloads. For information about burstable performance instances, see [Amazon EC2 instance types](https://aws.amazon.com/ec2/instance-types/).

## Register a model

Before you add a scaling policy to your model, you first must register your model for auto scaling and define the scaling limits for the model.

The following procedures cover how to register a model (production variant) for auto scaling using the AWS Command Line Interface (AWS CLI) or Application Auto Scaling API.

### Topics

- [Register a model (AWS CLI)](#)
- [Register a model (Application Auto Scaling API)](#)

### Register a model (AWS CLI)

To register your production variant, use the `register-scalable-target` command with the following parameters:

- `--service-namespace`—Set this value to `sagemaker`.
- `--resource-id`—The resource identifier for the model (specifically, the production variant). For this parameter, the resource type is `endpoint` and the unique identifier is the name of the production variant. For example, `endpoint/my-endpoint/variant/my-variant`.
- `--scalable-dimension`—Set this value to `sagemaker:variant:DesiredInstanceCount`.

Automatically scale models
- **--min-capacity**—The minimum number of instances. This value must be set to at least 1 and must be equal to or less than the value specified for **max-capacity**.

- **--max-capacity**—The maximum number of instances. This value must be set to at least 1 and must be equal to or greater than the value specified for **min-capacity**.

**Example**

The following example shows how to register a variant named *my-variant*, running on the *my-endpoint* endpoint, that can be dynamically scaled to have one to eight instances.

```bash
aws application-autoscaling register-scalable-target \
  --service-namespace sagemaker \
  --resource-id endpoint/my-endpoint/variant/my-variant \
  --scalable-dimension sagemaker:variant:DesiredInstanceCount \
  --min-capacity 1 \
  --max-capacity 8
```

**Register a model (Application Auto Scaling API)**

To register your model with Application Auto Scaling, use the [RegisterScalableTarget](https://docs.aws.amazon.com/sagemaker/latest/dg/API_RegisterScalableTarget.html) Application Auto Scaling API action with the following parameters:

- **ServiceNamespace**—Set this value to sagemaker.

- **ResourceId**—The resource identifier for the production variant. For this parameter, the resource type is endpoint and the unique identifier is the name of the variant. For example `endpoint/my-endpoint/variant/my-variant`.

- **ScalableDimension**—Set this value to `sagemaker:variant:DesiredInstanceCount`.

- **MinCapacity**—The minimum number of instances. This value must be set to at least 1 and must be equal to or less than the value specified for **MaxCapacity**.

- **MaxCapacity**—The maximum number of instances. This value must be set to at least 1 and must be equal to or greater than the value specified for **MinCapacity**.

**Example**

The following example shows how to register a variant named *my-variant*, running on the *my-endpoint* endpoint, that can be dynamically scaled to use one to eight instances.
Define a scaling policy

Before you add a scaling policy to your model, save your policy configuration as a JSON block in a text file. You use that text file when invoking the AWS Command Line Interface (AWS CLI) or the Application Auto Scaling API. You can optimize scaling by choosing an appropriate CloudWatch metric. However, before using a custom metric in production, you must test auto scaling with your custom metric.

This section shows you example policy configurations for target tracking scaling policies.

Topics

- Specify a predefined metric (CloudWatch metric: InvocationsPerInstance)
- Define a custom metric (CloudWatch metric: CPUUtilization)
- Define a custom metric (CloudWatch metric: ExplanationsPerInstance)
- Specify cooldown periods

Specify a predefined metric (CloudWatch metric: InvocationsPerInstance)

Example

The following is an example target tracking policy configuration for a variant that keeps the average invocations per instance at 70. Save this configuration in a file named config.json.
Define a custom metric (CloudWatch metric: CPUUtilization)

To create a target tracking scaling policy with a custom metric, specify the metric's name, namespace, unit, statistic, and zero or more dimensions. A dimension consists of a dimension name and a dimension value. You can use any production variant metric that changes in proportion to capacity.

Example

The following example configuration shows a target tracking scaling policy with a custom metric. The policy scales the variant based on an average CPU utilization of 50 percent across all instances. Save this configuration in a file named `config.json`.

```json
{
    "TargetValue": 50.0,
    "CustomizedMetricSpecification":
    {
        "MetricName": "CPUUtilization",
        "Namespace": "/aws/sagemaker/Endpoints",
        "Dimensions": [
            {"Name": "EndpointName", "Value": "my-endpoint" },
            {"Name": "VariantName","Value": "my-variant"}
        ],
        "Statistic": "Average",
        "Unit": "Percent"
    }
}
```

For more information, see `CustomizedMetricSpecification` in the Application Auto Scaling API Reference.
Define a custom metric (CloudWatch metric: ExplanationsPerInstance)

When the endpoint has online explainability activated, it emits an ExplanationsPerInstance metric that outputs the average number of records explained per minute, per instance, for a variant. The resource utilization of explaining records can be more different than that of predicting records. We strongly recommend using this metric for target tracking scaling of endpoints with online explainability activated.

You can create multiple target tracking policies for a scalable target. Consider adding the InvocationsPerInstance policy from the Specify a predefined metric (CloudWatch metric: InvocationsPerInstance) section (in addition to the ExplanationsPerInstance policy). If most invocations don’t return an explanation because of the threshold value set in the EnableExplanations parameter, then the endpoint can choose the InvocationsPerInstance policy. If there is a large number of explanations, the endpoint can use the ExplanationsPerInstance policy.

Example

The following example configuration shows a target tracking scaling policy with a custom metric. The policy scale adjusts the number of variant instances so that each instance has an ExplanationsPerInstance metric of 20. Save this configuration in a file named config.json.

```json
{
   "TargetValue": 20.0,
   "CustomizedMetricSpecification": {
      "MetricName": "ExplanationsPerInstance",
      "Namespace": "AWS/SageMaker",
      "Dimensions": [
         {"Name": "EndpointName", "Value": "my-endpoint" },
         {"Name": "VariantName","Value": "my-variant"}
      ],
      "Statistic": "Sum"
   }
}
```

For more information, see CustomizedMetricSpecification in the Application Auto Scaling API Reference.
Specify cooldown periods

You can optionally define cooldown periods in your target tracking scaling policy by specifying the `ScaleOutCooldown` and `ScaleInCooldown` parameters.

Example

The following is an example target tracking policy configuration for a variant that keeps the average invocations per instance at 70. The policy configuration provides a scale-in cooldown period of 10 minutes (600 seconds) and a scale-out cooldown period of 5 minutes (300 seconds). Save this configuration in a file named `config.json`.

```
{
    "TargetValue": 70.0,
    "PredefinedMetricSpecification": {
        "PredefinedMetricType": "SageMakerVariantInvocationsPerInstance"
    },
    "ScaleInCooldown": 600,
    "ScaleOutCooldown": 300
}
```

For more information, see [TargetTrackingScalingPolicyConfiguration](https://docs.aws.amazon.com/autoscaling/application/APIReference/API_TargetTrackingScalingPolicyConfiguration.html) in the Application Auto Scaling API Reference.

Apply a scaling policy

After you register your model and define a scaling policy, apply the scaling policy to the registered model. This section shows how to apply a scaling policy using the AWS Command Line Interface (AWS CLI) or the Application Auto Scaling API.

Topics

- [Apply a target tracking scaling policy (AWS CLI)](https://docs.aws.amazon.com/autoscaling/application/APIReference/API_PutScalingPolicy.html)
- [Apply a scaling policy (Application Auto Scaling API)](https://docs.aws.amazon.com/autoscaling/application/APIReference/API_PutScalingPolicy.html)

Apply a target tracking scaling policy (AWS CLI)

To apply a scaling policy to your model, use the `put-scaling-policy` AWS CLI command with the following parameters:
• **--policy-name**—The name of the scaling policy.
• **--policy-type**—Set this value to TargetTrackingScaling.
• **--resource-id**—The resource identifier for the variant. For this parameter, the resource type is endpoint and the unique identifier is the name of the variant. For example, endpoint/my-endpoint/variant/my-variant.
• **--service-namespace**—Set this value to sagemaker.
• **--scalable-dimension**—Set this value to sagemaker:variant:DesiredInstanceCount.
• **--target-tracking-scaling-policy-configuration**—The target-tracking scaling policy configuration to use for the model.

**Example**

The following example applies a target tracking scaling policy named *my-scaling-policy* to a variant named *my-variant*, running on the *my-endpoint* endpoint. For the **--target-tracking-scaling-policy-configuration** option, specify the config.json file that you created previously.

```bash
aws application-autoscaling put-scaling-policy \
--policy-name my-scaling-policy \
--policy-type TargetTrackingScaling \
--resource-id endpoint/my-endpoint/variant/my-variant \
--service-namespace sagemaker \
--scalable-dimension sagemaker:variant:DesiredInstanceCount \
--target-tracking-scaling-policy-configuration file://config.json
```

**Apply a scaling policy (Application Auto Scaling API)**

To apply a scaling policy to a variant with the Application Auto Scaling API, use the `PutScalingPolicy` Application Auto Scaling API action with the following parameters:

• **PolicyName**—The name of the scaling policy.
• **ServiceNamespace**—Set this value to sagemaker.
• **ResourceID**—The resource identifier for the variant. For this parameter, the resource type is endpoint and the unique identifier is the name of the variant. For example, endpoint/my-endpoint/variant/my-variant.
• **ScalableDimension**—Set this value to sagemaker:variant:DesiredInstanceCount.
• **PolicyType**—Set this value to `TargetTrackingScaling`.

• **TargetTrackingScalingPolicyConfiguration**—The target-tracking scaling policy configuration to use for the variant.

**Example**

The following example applies a target tracking scaling policy named `my-scaling-policy` to a variant named `my-variant`, running on the `my-endpoint` endpoint. The policy configuration keeps the average invocations per instance at 70.

```
POST / HTTP/1.1
Host: application-autoscaling.us-east-2.amazonaws.com
Accept-Encoding: identity
X-Amz-Target: AnyScaleFrontendService.
X-Amz-Date: 20230506T182145Z
User-Agent: aws-cli/2.0.0 Python/3.7.5 Windows/10 botocore/2.0.0dev4
Content-Type: application/x-amz-json-1.1
Authorization: AUTHPARAMS

{
    "PolicyName": "my-scaling-policy",
    "ServiceNamespace": "sagemaker",
    "ResourceId": "endpoint/my-endpoint/variant/my-variant",
    "ScalableDimension": "sagemaker:variant:DesiredInstanceCount",
    "PolicyType": "TargetTrackingScaling",
    "TargetTrackingScalingPolicyConfiguration": {
        "TargetValue": 70.0,
        "PredefinedMetricSpecification": {
            "PredefinedMetricType": "SageMakerVariantInvocationsPerInstance"
        }
    }
}
```

**Edit a scaling policy**

After creating a scaling policy, you can edit any of its settings except the name.

**Topics**

- **Edit a scaling policy (AWS CLI or Application Auto Scaling API)**
- **Temporarily turn off scaling policies**

**Edit a scaling policy (console)**

To edit a target tracking scaling policy with the AWS Management Console, use the same procedure that you used to [Configure model auto scaling with the console](#).

**Edit a scaling policy (AWS CLI or Application Auto Scaling API)**

You can use the AWS CLI or the Application Auto Scaling API to edit a scaling policy in the same way that you create a new scaling policy. For more information, see [Apply a scaling policy](#).

**Temporarily turn off scaling policies**

After you configure auto scaling, you have the following options if you need to investigate an issue without interference from scaling policies (dynamic scaling):

- Temporarily suspend and then resume scaling activities by calling the `register-scalable-target` CLI command or `RegisterScalableTarget` API action, specifying a Boolean value for both `DynamicScalingInSuspended` and `DynamicScalingOutSuspended`.

**Example**

The following example shows how to suspend scaling policies for a variant named `my-variant`, running on the `my-endpoint` endpoint.

```
aws application-autoscaling register-scalable-target \
  --service-namespace sagemaker \
  --resource-id endpoint/my-endpoint/variant/my-variant \
  --scalable-dimension sagemaker:variant:DesiredInstanceCount \
  --suspended-state '{"DynamicScalingInSuspended":true,"DynamicScalingOutSuspended":true}'
```

- Prevent specific target tracking scaling policies from scaling in your variant by disabling the policy's scale-in portion. This method prevents the scaling policy from deleting instances, while still allowing it to create them as needed.

  Temporarily disable and then enable scale-in activities by editing the policy using the `put-scaling-policy` CLI command or the `PutScalingPolicy` API action, specifying a Boolean value for `DisableScaleIn`.

Automatically scale models
Example

The following is an example of a target tracking configuration for a scaling policy that will scale out but not scale in.

```json
{
    "TargetValue": 70.0,
    "PredefinedMetricSpecification": {
        "PredefinedMetricType": "SageMakerVariantInvocationsPerInstance"
    },
    "DisableScaleIn": true
}
```

Delete a scaling policy

If you no longer need a scaling policy, you can delete it at any time.

Topics

- delete all scaling policies and deregister the model (console)
- Delete a scaling policy (AWS CLI or Application Auto Scaling API)

Delete all scaling policies and deregister the model (console)

To delete all scaling policies and deregister the variant as a scalable target

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. On the navigation pane, choose Endpoints.
3. Choose your endpoint, and then for Endpoint runtime settings, choose the variant.
4. Choose Configure auto scaling.
5. Choose Deregister auto scaling.

Delete a scaling policy (AWS CLI or Application Auto Scaling API)

You can use the AWS CLI or the Application Auto Scaling API to delete a scaling policy from a variant.
Delete a scaling policy (AWS CLI)

To delete a scaling policy from a variant, use the `delete-scaling-policy` command with the following parameters:

- `-policy-name`—The name of the scaling policy.
- `-resource-id`—The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example, `endpoint/my-endpoint/variant/my-variant`.
- `-service-namespace`—Set this value to `sagemaker`.
- `-scalable-dimension`—Set this value to `sagemaker:variant:DesiredInstanceCount`.

Example

The following example deletes a target tracking scaling policy named `my-scaling-policy` from a variant named `my-variant`, running on the `my-endpoint` endpoint.

```
aws application-autoscaling delete-scaling-policy \
    --policy-name my-scaling-policy \
    --resource-id endpoint/my-endpoint/variant/my-variant \
    --service-namespace sagemaker \
    --scalable-dimension sagemaker:variant:DesiredInstanceCount
```

Delete a scaling policy (Application Auto Scaling API)

To delete a scaling policy from your variant, use the `DeleteScalingPolicy` Application Auto Scaling API action with the following parameters:

- `PolicyName`—The name of the scaling policy.
- `ServiceNamespace`—Set this value to `sagemaker`.
- `ResourceID`—The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example, `endpoint/my-endpoint/variant/my-variant`.
- `ScalableDimension`—Set this value to `sagemaker:variant:DesiredInstanceCount`. 
Example

The following example deletes a target tracking scaling policy named *my-scaling-policy* from a variant named *my-variant*, running on the *my-endpoint* endpoint.

```json
POST / HTTP/1.1
Host: application-autoscaling.us-east-2.amazonaws.com
Accept-Encoding: identity
X-Amz-Target: AnyScaleFrontendService.DeleteScalingPolicy
X-Amz-Date: 20230506T182145Z
User-Agent: aws-cli/2.0.0 Python/3.7.5 Windows/10 botocore/2.0.0dev4
Content-Type: application/x-amz-json-1.1
Authorization: AUTHPARAMS

{
    "PolicyName": "my-scaling-policy",
    "ServiceNamespace": "sagemaker",
    "ResourceId": "endpoint/my-endpoint/variant/my-variant",
    "ScalableDimension": "sagemaker:variant:DesiredInstanceCount"
}
```

Check the status of a scaling activity by describing scaling activities

You can check the status of a scaling activity for your auto scaled endpoint by describing scaling activities. Application Auto Scaling provides descriptive information about the scaling activities in the specified namespace from the previous six weeks. For more information, see Scaling activities for Application Auto Scaling in the Application Auto Scaling User Guide.

To check the status of a scaling activity, use the `describe-scaling-activities` command. You can't check the status of a scaling activity using the console.

Topics

- Describe scaling activities (AWS CLI)
- Identify blocked scaling activities from instance quotas (AWS CLI)

Describe scaling activities (AWS CLI)

To describe scaling activities for all SageMaker resources that registered with Application Auto Scaling, use the `describe-scaling-activities` command, specifying `sagemaker` for the `--service-namespace` option.
aws application-autoscaling describe-scaling-activities \
--service-namespace sagemaker

To describe scaling activities for a specific resource, include the --resource-id option.

aws application-autoscaling describe-scaling-activities \
--service-namespace sagemaker \
--resource-id endpoint/my-endpoint/variant/my-variant

The following example shows the output produced when you run this command.

```json
{
    "ActivityId": "activity-id",
    "ServiceNamespace": "sagemaker",
    "ResourceId": "endpoint/my-endpoint/variant/my-variant",
    "ScalableDimension": "sagemaker:variant:DesiredInstanceCount",
    "Description": "string",
    "Cause": "string",
    "StartTime": timestamp,
    "EndTime": timestamp,
    "StatusCode": "string",
    "StatusMessage": "string"
}
```

**Identify blocked scaling activities from instance quotas (AWS CLI)**

When you scale out (add more instances), you might reach your account-level instance quota. You can use the `describe-scaling-activities` command to check whether you have reached your instance quota. When you exceed your quota, auto scaling is blocked.

To check if you have reached your instance quota, use the `describe-scaling-activities` command and specify the resource ID for the --resource-id option.

aws application-autoscaling describe-scaling-activities \
--service-namespace sagemaker \
--resource-id endpoint/my-endpoint/variant/my-variant

Within the return syntax, check the **StatusCode** and **StatusMessage** keys and their associated values. StatusCode returns Failed. Within StatusMessage there is a message indicating that
the account-level service quota was reached. The following is an example of what that message might look like:

```
{
    "ActivityId": "activity-id",
    "ServiceNamespace": "sagemaker",
    "ResourceId": "endpoint/my-endpoint/variant/my-variant",
    "ScalableDimension": "sagemaker:variant:DesiredInstanceCount",
    "Description": "string",
    "Cause": "minimum capacity was set to 110",
    "StartTime": timestamp,
    "EndTime": timestamp,
    "StatusCode": "Failed",
    "StatusMessage": "Failed to set desired instance count to 110. Reason: The account-level service limit 'ml.xx.xxxxxx for endpoint usage' is 1000 Instances, with current utilization of 997 Instances and a request delta of 20 Instances. Please contact AWS support to request an increase for this limit. (Service: AmazonSageMaker; Status Code: 400; Error Code: ResourceLimitExceeded; Request ID: request-id)."
}
```

**Load testing your auto scaling configuration**

Perform load tests to choose a scaling configuration that works the way you want.

The following guidelines for load testing assume you are using a scaling policy that uses the predefined target metric SageMakerVariantInvocationsPerInstance.

**Topics**

- [Determine the performance characteristics](#)
- [Calculate the target load](#)

**Determine the performance characteristics**

Perform load testing to find the peak InvocationsPerInstance that your model's production variant can handle, and the latency of requests, as concurrency increases.

This value depends on the instance type chosen, payloads that clients of your model typically send, and the performance of any external dependencies your model has.
To find the peak requests-per-second (RPS) your model's production variant can handle and latency of requests

1. Set up an endpoint with your model using a single instance. For information about how to set up an endpoint, see Deploy the Model to SageMaker Hosting Services.

2. Use a load testing tool to generate an increasing number of parallel requests, and monitor the RPS and model latency in the output of the load testing tool.

Note

You can also monitor requests-per-minute instead of RPS. In that case don't multiply by 60 in the equation to calculate SageMakerVariantInvocationsPerInstance shown below.

When the model latency increases or the proportion of successful transactions decreases, this is the peak RPS that your model can handle.

Calculate the target load

After you find the performance characteristics of the variant, you can determine the maximum RPS we should allow to be sent to an instance. The threshold used for scaling must be less than this maximum value. Use the following equation in combination with load testing to determine the correct value for the SageMakerVariantInvocationsPerInstance target metric in your scaling configuration.

\[ \text{SageMakerVariantInvocationsPerInstance} = (\text{MAX}_\text{RPS} \times \text{SAFETY}_\text{FACTOR}) \times 60 \]

Where MAX_RPS is the maximum RPS that you determined previously, and SAFETY_FACTOR is the safety factor that you chose to ensure that your clients don't exceed the maximum RPS. Multiply by 60 to convert from RPS to invocations-per-minute to match the per-minute CloudWatch metric that SageMaker uses to implement auto scaling (you don't need to do this if you measured requests-per-minute instead of requests-per-second).
Note

SageMaker recommends that you start testing with a SAFETY_FACTOR of 0.5. Test your scaling configuration to ensure it operates in the way you expect with your model for both increasing and decreasing customer traffic on your endpoint.

Use AWS CloudFormation to create a scaling policy

The following example shows how to configure model auto scaling on an endpoint using AWS CloudFormation.

```yaml
Endpoint:
  Type: "AWS::SageMaker::Endpoint"
  Properties:
    EndpointName: yourEndpointName
    EndpointConfigName: yourEndpointConfigName

ScalingTarget:
  Type: "AWS::ApplicationAutoScaling::ScalableTarget"
  Properties:
    MaxCapacity: 10
    MinCapacity: 2
    ResourceId: endpoint/my-endpoint/variant/my-variant
    RoleARN: arn
    ScalableDimension: sagemaker:variant:DesiredInstanceCount
    ServiceNamespace: sagemaker

ScalingPolicy:
  Type: "AWS::ApplicationAutoScaling::ScalingPolicy"
  Properties:
    PolicyName: my-scaling-policy
    PolicyType: TargetTrackingScaling
    ScalingTargetId:
      Ref: ScalingTarget
    TargetTrackingScalingPolicyConfiguration:
      TargetValue: 70.0
      ScaleInCooldown: 600
      ScaleOutCooldown: 30
    PredefinedMetricSpecification:
      PredefinedMetricType: SageMakerVariantInvocationsPerInstance
```

Automatically scale models
For more information, see Create Application Auto Scaling resources with AWS CloudFormation in the Application Auto Scaling User Guide.

Update or delete endpoints that use auto scaling

Topics

- Update endpoints that use auto scaling
- Delete endpoints configured for auto scaling

Update endpoints that use auto scaling

When you update an endpoint, Application Auto Scaling checks to see whether any of the models on that endpoint are targets for auto scaling. If the update would change the instance type for any model that is a target for auto scaling, the update fails.

In the AWS Management Console, you see a warning that you must deregister the model from auto scaling before you can update it. If you are trying to update the endpoint by calling the UpdateEndpoint API, the call fails. Before you update the endpoint, delete any scaling policies configured for it and deregister the variant as a scalable target by calling the DeregisterScalableTarget Application Auto Scaling API action. After you update the endpoint, you can register the updated variant as a scalable target and attach a scaling policy.

There is one exception. If you change the model for a variant that is configured for auto scaling, Amazon SageMaker auto scaling allows the update. This is because changing the model doesn’t typically affect performance enough to change scaling behavior. If you do update a model for a variant configured for auto scaling, ensure that the change to the model doesn’t significantly affect performance and scaling behavior.

When you update SageMaker endpoints that have auto scaling applied, complete the following steps:

To update an endpoint that has auto scaling applied

1. Deregister the endpoint as a scalable target by calling DeregisterScalableTarget.
2. Because auto scaling is blocked while the update operation is in progress (or if you turned off auto scaling in the previous step), you might want to take the additional precaution of increasing the number of instances for your endpoint during the update. To do this, update the instance counts for the production variants hosted at the endpoint by calling UpdateEndpointWeightsAndCapacities.
3. Call **DescribeEndpoint** repeatedly until the value of the EndpointStatus field of the response is InService.

4. Call **DescribeEndpointConfig** to get the values of the current endpoint config.

5. Create a new endpoint config by calling **CreateEndpointConfig**. For the production variants where you want to keep the existing instance count or weight, use the same variant name from the response from the call to **DescribeEndpointConfig** in the previous step. For all other values, use the values that you got as the response when you called **DescribeEndpointConfig** in the previous step.

6. Update the endpoint by calling **UpdateEndpoint**. Specify the endpoint config you created in the previous step as the EndpointConfig field. If you want to retain the variant properties like instance count or weight, set the value of the RetainAllVariantProperties parameter to True. This specifies that production variants with the same name will are updated with the most recent DesiredInstanceCount from the response from the call to DescribeEndpoint, regardless of the values of the InitialInstanceCount field in the new EndpointConfig.

7. (Optional) Re-activate auto scaling by calling **RegisterScalableTarget** and **PutScalingPolicy**.

**Note**

Steps 1 and 7 are required only if you are updating an endpoint with the following changes:

- Changing the instance type for a production variant that has auto scaling configured
- Removing a production variant that has auto scaling configured.

**Delete endpoints configured for auto scaling**

If you delete an endpoint, Application Auto Scaling checks to see whether any of the models on that endpoint are targets for auto scaling. If any are and you have permission to deregister the model, Application Auto Scaling deregisters those models as scalable targets without notifying you. If you use a custom permission policy that doesn't provide permission for the **DeregisterScalableTarget** action, you must request access to this action before deleting the endpoint.
Note

As an IAM user, you might not have sufficient permission to delete an endpoint if another user configured auto scaling for a variant on that endpoint.

Host instance storage volumes

When you create an endpoint, Amazon SageMaker attaches an Amazon Elastic Block Store (Amazon EBS) storage volume to Amazon EC2 instances that hosts the endpoint. The size of the storage volume is scalable, and storage options are divided into two categories: SSD-backed storage and HDD-backed storage.

For more information about Amazon EBS storages and features, see the following pages.

- Amazon EBS Features
- Amazon EBS User Guide

For a full list of the host instance storage volumes, see Host Instance Storage Volumes Table

Note

Amazon SageMaker attaches an Amazon Elastic Block Store (Amazon EBS) storage volume to Amazon EC2 instances only when you create Asynchronous inference or Real-time inference endpoint types. For more information on customizing Amazon EBS storage volume, see SageMaker endpoint parameters for large model inference.

Safely validate models in production

With SageMaker, you can test multiple models or model versions behind the same endpoint using variants. A variant consists of an ML instance and the serving components specified in a SageMaker model. You can have multiple variants behind an endpoint. Each variant can have a different instance type or a SageMaker model that can be autoscaled independently of the others. The models within the variants can be trained using different datasets, different algorithms, different ML frameworks, or any combination of all of these. All the variants behind an endpoint share the same inference code. SageMaker supports two types of variants, production variants and shadow variants.
If you have multiple production variants behind an endpoint, then you can allocate a portion of your inference requests to each variant. Each request is routed to only one of the production variants. The production variant to which the request was routed provides the response to the caller. You can compare how the production variants perform relative to each other.

You can also have a shadow variant corresponding to a production variant behind an endpoint. A portion of the inference requests that goes to the production variant is replicated to the shadow variant. The responses of the shadow variant are logged for comparison and not returned to the caller. This lets you test the performance of the shadow variant without exposing the caller to the response produced by the shadow variant.

Topics

- Production variants
- Shadow variants

Production variants

In production ML workflows, data scientists and engineers frequently try to improve performance using various methods, such as Perform Automatic Model Tuning with SageMaker, training on additional or more-recent data, improving feature selection, using better updated instances and serving containers. You can use production variants to compare your models, instances and containers, and choose the best performing candidate to respond to inference requests.

With SageMaker multi-variant endpoints you can distribute endpoint invocation requests across multiple production variants by providing the traffic distribution for each variant, or you can invoke a specific variant directly for each request. In this topic, we look at both methods for testing ML models.

Topics

- Test models by specifying traffic distribution
- Test models by invoking specific variants
- Model A/B test example

Test models by specifying traffic distribution

To test multiple models by distributing traffic between them, specify the percentage of the traffic that gets routed to each model by specifying the weight for each production variant in the
endpoint configuration. For information, see `CreateEndpointConfig`. The following diagram shows how this works in more detail.

**Production Variants**

![Diagram showing production variants for a SageMaker endpoint]

- **Variant Name**: ProductionVariant1  
  **Variant Weight**: 0.7  
  **Model Name**: ExampleModel1  
  **Instance Type**: ml.m5.xlarge  
  **Instance Count**: 1

- **Variant Name**: ProductionVariant2  
  **Variant Weight**: 0.2  
  **Model Name**: ExampleModel2  
  **Instance Type**: ml.m5.xlarge  
  **Instance Count**: 2

- **Variant Name**: ProductionVariant3  
  **Variant Weight**: 0.1  
  **Model Name**: ExampleModel3  
  **Instance Type**: ml.m5.xlarge  
  **Instance Count**: 1

---

**Test models by invoking specific variants**

To test multiple models by invoking specific models for each request, specify the specific version of the model you want to invoke by providing a value for the `TargetVariant` parameter when you call `InvokeEndpoint`. SageMaker ensures that the request is processed by the production variant you specify. If you have already provided traffic distribution and specify a value for the `TargetVariant` parameter, the targeted routing overrides the random traffic distribution. The following diagram shows how this works in more detail.
Model A/B test example

Performing A/B testing between a new model and an old model with production traffic can be an effective final step in the validation process for a new model. In A/B testing, you test different variants of your models and compare how each variant performs. If the newer version of the model delivers better performance than the previously existing version, replace the old version of the model with the new version in production.

The following example shows how to perform A/B model testing. For a sample notebook that implements this example, see "A/B Testing ML models in production."

Step 1: Create and deploy models

First, we define where our models are located in Amazon S3. These locations are used when we deploy our models in subsequent steps:

```python
model_url = f"s3://{path_to_model_1}"
model_url2 = f"s3://{path_to_model_2}"```

Safely validate models in production
Next, we create the model objects with the image and model data. These model objects are used to deploy production variants on an endpoint. The models are developed by training ML models on different data sets, different algorithms or ML frameworks, and different hyperparameters:

```python
from sagemaker.amazon.amazon_estimator import get_image_uri

model_name = f"DEMO-xgb-churn-pred-{datetime.now():%Y-%m-%d-%H-%M-%S}" 
model_name2 = f"DEMO-xgb-churn-pred2-{datetime.now():%Y-%m-%d-%H-%M-%S}" 
image_uri = get_image_uri(boto3.Session().region_name, 'xgboost', '0.90-1') 
image_uri2 = get_image_uri(boto3.Session().region_name, 'xgboost', '0.90-2')

sm_session.create_model(  
    name=model_name,  
    role=role,  
    container_defs={  
        'Image': image_uri,  
        'ModelDataUrl': model_url  
    }  
)

sm_session.create_model(  
    name=model_name2,  
    role=role,  
    container_defs={  
        'Image': image_uri2,  
        'ModelDataUrl': model_url2  
    }  
)
```

We now create two production variants, each with its own different model and resource requirements (instance type and counts). This enables you to also test models on different instance types.

We set an initial_weight of 1 for both variants. This means that 50% of requests go to Variant1, and the remaining 50% of requests to Variant2. The sum of weights across both variants is 2 and each variant has weight assignment of 1. This means that each variant receives 1/2, or 50%, of the total traffic.

```python
from sagemaker.session import production_variant
```
variant1 = production_variant(
    model_name=model_name,
    instance_type="ml.m5.xlarge",
    initial_instance_count=1,
    variant_name='Variant1',
    initial_weight=1,
)

variant2 = production_variant(
    model_name=model_name2,
    instance_type="ml.m5.xlarge",
    initial_instance_count=1,
    variant_name='Variant2',
    initial_weight=1,
)

Finally we’re ready to deploy these production variants on a SageMaker endpoint.

```python
endpoint_name = f"DEMO-xgb-churn-pred-{datetime.now():%Y-%m-%d-%H-%M-%S}"
print(f"EndpointName={endpoint_name}"

sm_session.endpoint_from_production_variants(
    name=endpoint_name,
    production_variants=[variant1, variant2]
)
```

**Step 2: Invoke the deployed models**

Now we send requests to this endpoint to get inferences in real time. We use both traffic distribution and direct targeting.

First, we use traffic distribution that we configured in the previous step. Each inference response contains the name of the production variant that processes the request, so we can see that traffic to the two production variants is roughly equal.

```python
# get a subset of test data for a quick test
!tail -120 test_data/test-dataset-input-cols.csv > test_data/
test_sample_tail_input_cols.csv
print(f"Sending test traffic to the endpoint {endpoint_name}. \nPlease wait...")
```
with open('test_data/test_sample_tail_input_cols.csv', 'r') as f:
    for row in f:
        print('.', end='", flush=True)
        payload = row.rstrip('"
')
        sm_runtime.invoke_endpoint(
            EndpointName=endpoint_name,
            ContentType="text/csv",
            Body=payload
        )
        time.sleep(0.5)

print('Done!')

SageMaker emits metrics such as Latency and Invocations for each variant in Amazon CloudWatch. For a complete list of metrics that SageMaker emits, see Monitor Amazon SageMaker with Amazon CloudWatch. Let's query CloudWatch to get the number of invocations per variant, to show how invocations are split across variants by default:

<table>
<thead>
<tr>
<th>Variant 1</th>
<th>Variant 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>44.0</td>
<td>60.0</td>
</tr>
<tr>
<td>7.0</td>
<td>9.0</td>
</tr>
</tbody>
</table>

Now let's invoke a specific version of the model by specifying Variant1 as the TargetVariant in the call to invoke_endpoint.

print(f'Sending test traffic to the endpoint {endpoint_name}. Please wait...')
with open('test_data/test_sample_tail_input_cols.csv', 'r') as f:
    for row in f:
        print('.', end='", flush=True)
payload = row.rstrip('
')
sm_runtime.invoke_endpoint(
    EndpointName=endpoint_name,
    ContentType="text/csv",
    Body=payload,
    TargetVariant="Variant1"
)
time.sleep(0.5)

To confirm that all new invocations were processed by Variant1, we can query CloudWatch to get the number of invocations per variant. We see that for the most recent invocations (latest timestamp), all requests were processed by Variant1, as we had specified. There were no invocations made for Variant2.

<table>
<thead>
<tr>
<th>Timestamp</th>
<th>Variant1</th>
<th>Variant2</th>
</tr>
</thead>
<tbody>
<tr>
<td>2020-06-05 15:34:00+00:00</td>
<td>44.0</td>
<td>60.0</td>
</tr>
<tr>
<td>2020-06-05 15:35:00+00:00</td>
<td>7.0</td>
<td>9.0</td>
</tr>
<tr>
<td>2020-06-05 15:36:00+00:00</td>
<td>99.0</td>
<td>NaN</td>
</tr>
</tbody>
</table>

Step 3: Evaluate model performance

To see which model version performs better, let's evaluate the accuracy, precision, recall, F1 score, and Receiver operating characteristic/Area under the curve for each variant. First, let's look at these metrics for Variant1:
Now let's look at the metrics for Variant2:

- **Accuracy**: 0.958333333333334
- **Precision**: 0.8947368421052632
- **Recall**: 0.85
- **F1 Score**: 0.8717948717948718
- **AUC**: 0.92

For most of our defined metrics, Variant2 is performing better, so this is the one that we want to use in production.

**Step 4: Increase traffic to the best model**

Now that we have determined that Variant2 performs better than Variant1, we shift more traffic to it. We can continue to use TargetVariant to invoke a specific model variant, but a simpler approach is to update the weights assigned to each variant by calling `UpdateEndpointWeightsAndCapacities`. This changes the traffic distribution to your production.
variants without requiring updates to your endpoint. Recall from the setup section that we set variant weights to split traffic 50/50. The CloudWatch metrics for the total invocations for each variant below show us the invocation patterns for each variant:

![Graph showing invocation patterns for Variant 1 and Variant 2](image)

Now we shift 75% of the traffic to Variant 2 by assigning new weights to each variant using `UpdateEndpointWeightsAndCapacities`. SageMaker now sends 75% of the inference requests to Variant 2 and remaining 25% of requests to Variant 1.

```python
sm.update_endpoint_weights_and_capacities(
    EndpointName=endpoint_name,
    DesiredWeightsAndCapacities=[
        {
            "DesiredWeight": 25,
            "VariantName": variant1["VariantName"]
        },
        {
            "DesiredWeight": 75,
            "VariantName": variant2["VariantName"]
        }
    ]
)
```

The CloudWatch metrics for total invocations for each variant shows us higher invocations for Variant 2 than for Variant 1:
We can continue to monitor our metrics, and when we're satisfied with a variant's performance, we can route 100% of the traffic to that variant. We use `UpdateEndpointWeightsAndCapacities` to update the traffic assignments for the variants. The weight for `Variant1` is set to 0 and the weight for `Variant2` is set to 1. SageMaker now sends 100% of all inference requests to `Variant2`.

```python
sm.update_endpoint_weights_and_capacities(
    EndpointName=endpoint_name,
    DesiredWeightsAndCapacities=[
        {
            "DesiredWeight": 0,
            "VariantName": variant1["VariantName"]
        },
        {
            "DesiredWeight": 1,
            "VariantName": variant2["VariantName"]
        }
    ]
)
```

The CloudWatch metrics for the total invocations for each variant show that all inference requests are being processed by `Variant2` and there are no inference requests processed by `Variant1`. 
You can now safely update your endpoint and delete Variant1 from your endpoint. You can also continue testing new models in production by adding new variants to your endpoint and following steps 2 - 4.

**Shadow variants**

You can use SageMaker Model Shadow Deployments to create long running shadow variants to validate any new candidate component of your model serving stack before promoting it to production. The following diagram shows how shadow variants work in more detail.
Deploy shadow variants

The following code example shows how you can programmatically deploy shadow variants. Replace the *user placeholder text* in the example with your own information.

1. Create two SageMaker models: one for your production variant, and one for your shadow variant.

```python
import boto3
from sagemaker import get_execution_role, Session

aws_region = "aws-region"

boto_session = boto3.Session(region_name=aws_region)
sagemaker_client = boto_session.client("sagemaker")

role = get_execution_role()
```
bucket = Session(boto_session).default_bucket()

model_name1 = "name-of-your-first-model"
model_name2 = "name-of-your-second-model"

sagemaker_client.create_model(
    ModelName = model_name1,
    ExecutionRoleArn = role,
    Containers=[
        {
            "Image": "ecr-image-uri-for-first-model",
            "ModelDataUrl": "s3-location-of-trained-first-model"
        }
    ]
)

sagemaker_client.create_model(
    ModelName = model_name2,
    ExecutionRoleArn = role,
    Containers=[
        {
            "Image": "ecr-image-uri-for-second-model",
            "ModelDataUrl": "s3-location-of-trained-second-model"
        }
    ]
)

2. Create an endpoint configuration. Specify both your production and shadow variants in the configuration.

endpoint_config_name = name-of-your-endpoint-config

create_endpoint_config_response = sagemaker_client.create_endpoint_config(
    EndpointConfigName=endpoint_config_name,
    ProductionVariants=[
        {
            "VariantName": "name-of-your-production-variant",
            "ModelName": model_name1,
            "InstanceType": "ml.m5.xlarge",
            "InitialInstanceCount": 1,
            "InitialVariantWeight": 1,
        }
    ]
)
ShadowProductionVariants=[
    {
        "VariantName": name-of-your-shadow-variant,
        "ModelName": model_name2,
        "InstanceType": "ml.m5.xlarge",
        "InitialInstanceCount": 1,
        "InitialVariantWeight": 1,
    }
]
)

3. Create an endpoint.

create_endpoint_response = sm.create_endpoint(
    EndpointName=name-of-your-endpoint,
    EndpointConfigName=endpoint_config_name,
)

Online Explainability with SageMaker Clarify

This guide shows how to configure online explainability with SageMaker Clarify. With SageMaker real-time inference endpoints, you can analyze explainability in real time, continuously. The online explainability function fits into the Deploy to production part of the Amazon SageMaker Machine Learning workflow.

How Clarify Online Explainability Works

The following graphic depicts SageMaker architecture for hosting an endpoint that serves explainability requests. It depicts interactions between an endpoint, the model container, and the SageMaker Clarify explainer.
Here's how Clarify online explainability works. The application sends a REST-style `InvokeEndpoint` request to the SageMaker Runtime Service. The service routes this request to a SageMaker endpoint to obtain predictions and explanations. Then, the service receives the response from the endpoint. Lastly, the service sends the response back to the application.

To increase the endpoint availability, SageMaker automatically attempts to distribute endpoint instances in multiple Availability Zones, according to the instance count in the endpoint configuration. On an endpoint instance, upon a new explainability request, the SageMaker Clarify explainer calls the model container for predictions. Then it computes and returns the feature attributions.

Here are the four steps to create an endpoint that uses SageMaker Clarify online explainability:

1. Check if your pre-trained SageMaker model is compatible with online explainability by following the **pre-check** steps.
2. **Create an endpoint configuration** with the SageMaker Clarify explainer configuration using the `CreateEndpointConfig` API.
3. **Create an endpoint** and provide the endpoint configuration to SageMaker using the `CreateEndpoint` API. The service launches the ML compute instance and deploys the model as specified in the configuration.
4. **Invoke the endpoint**: After the endpoint is in service, call the SageMaker Runtime API `InvokeEndpoint` to send requests to the endpoint. The endpoint then returns explanations and predictions.

### Pre-check the model container

This section shows how to pre-check the model container inputs and outputs for compatibility before configuring an endpoint. The SageMaker Clarify explainer is **model agnostic**, but it has requirements for model container input and output.

> **Note**
> You can increase efficiency by configuring your container to support batch requests, which support two or more records in a single request. For example, a single record is a single line of CSV data, or a single line of JSON Lines data. SageMaker Clarify will attempt to send a mini-batch of records to the model container first before falling back to single record requests.

### Model container input

#### CSV

The model container supports input in CSV with MIME type: `text/csv`. The following table shows example inputs that SageMaker Clarify supports.

<table>
<thead>
<tr>
<th>Model container input (string representation)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>'1,2,3,4'</td>
<td>Single record that uses four numerical features.</td>
</tr>
<tr>
<td>'1,2,3,4\n5,6,7,8'</td>
<td>Two records, separated by line break '\n'.</td>
</tr>
<tr>
<td>&quot;&quot;This is a good product&quot;,5'</td>
<td>Single record that contains a text feature and a numerical feature.</td>
</tr>
<tr>
<td>&quot;&quot;This is a good product&quot;,5\n&quot;Bad shopping experience&quot;,1'</td>
<td>Two records.</td>
</tr>
</tbody>
</table>
JSON Lines

SageMaker also supports input in **JSON Lines dense format** with MIME type: application/jsonlines, as shown in the following table.

<table>
<thead>
<tr>
<th>Model container input</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>'{&quot;data&quot;:{&quot;features&quot;:[1,2,3,4]}}'</td>
<td>Single record; a list of features can be extracted by JMESPath expression data.features .</td>
</tr>
</tbody>
</table>
| '{"data":{"features":[1,2,3,4]}}
{"data":{"features":[5,6,7,8]}}' | Two records.                                                             |
| '{"features":["This is a good product",5]}'              | Single record; a list of features can be extracted by JMESPath expression features. |
| '{"features":["This is a good product",5]}
{"features":["Bad shopping experience",1]}' | Two records.                                                             |

**Model container output**

Your model container output should also be in either CSV, or JSON Lines dense format. Additionally the model container should include the probabilities of the input records, which SageMaker Clarify uses to compute feature attributions.

The following data examples are for model container outputs in **CSV format**.

**Probability only**

For regression and binary classification problems, the model container outputs a single probability value (score) of the predicted label. These probabilities can be extracted using column index 0. For multi-class problems, the model container outputs a list of probabilities (scores). For multi-class problems, if no index is provided, all values are extracted.
### Predicted label and probabilities

The model container outputs the predicted label followed by its probability in **CSV** format. The probabilities can be extracted using index 1.

<table>
<thead>
<tr>
<th>Model container input</th>
<th>Model container output</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'1,0.6'</td>
</tr>
<tr>
<td>Two records</td>
<td>'1,0.6\n0.3'</td>
</tr>
</tbody>
</table>

### Predicted labels header and probabilities

A multi-class model container trained by Autopilot can be configured to output the **string representation** of the list of predicted labels and probabilities in **CSV** format. In the following example, the probabilities can be extracted by index 1. The label headers can be extracted by index 1, and the label headers can be extracted using index 0.

<table>
<thead>
<tr>
<th>Model container input</th>
<th>Model container output</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>&quot;['cat','dog','fish'],[0.1,0.6,0.3]&quot;</td>
</tr>
</tbody>
</table>
The following data examples are for model container outputs in JSON Lines format.

Probability only

In this example, the model container outputs the probability that can be extracted by JMESPath expression score in JSON Lines format.

Predicted label and probabilities

In this example, a multi-class model container outputs a list of label headers along with a list of probabilities in JSON Lines format. The probabilities can be extracted by JMESPath expression probability, and the label headers can be extracted by JMESPath expression predicted labels.
Predicted labels header and probabilities

In this example, a multi-class model container outputs a list of label headers and probabilities in **JSON Lines** format. The probabilities can be extracted by JMESPath expression `probability`, and the label headers can be extracted by JMESPath expression `predicted_labels`.

<table>
<thead>
<tr>
<th>Model container input</th>
<th>Model container output</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;predicted_labels&quot;: [&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;], &quot;probabilities&quot;: [0.1, 0.6, 0.3]}'</td>
</tr>
<tr>
<td>Two records</td>
<td>'{&quot;predicted_labels&quot;: [&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;], &quot;probabilities&quot;: [0.1, 0.6, 0.3]},&quot;predicted_labels&quot;: [&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;], &quot;probabilities&quot;: [0.2, 0.5, 0.3]}'</td>
</tr>
</tbody>
</table>

**Model container validation**

We recommend that you deploy your model to a SageMaker real-time inference endpoint, and send requests to the endpoint. Manually examine the requests (model container inputs) and responses (model container outputs) to make sure that both are compliant with the requirements in the **Model Container Input** section and **Model Container Output** section. If your model container supports batch requests, you can start with a single record request, and then try two or more records.

The following commands show how to request a response using the AWS CLI. The AWS CLI is pre-installed in SageMaker Studio Classic, and SageMaker Notebook instances. If you need to install the AWS CLI, follow this [installation guide](#).

```
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name $ENDPOINT_NAME \
  --content-type $CONTENT_TYPE \
  --accept $ACCEPT_TYPE \
  --body $REQUEST_DATA \
  $CLI_BINARY_FORMAT  \
  /dev/stderr 1>/dev/null
```

The parameters are defined, as follows:
• $ENDPOINT NAME: The name of the endpoint.
• $CONTENT_TYPE: The MIME type of the request (model container input).
• $ACCEPT_TYPE: The MIME type of the response (model container output).
• $REQUEST_DATA: The requested payload string.
• $CLI_BINARY_FORMAT: The format of the command line interface (CLI) parameter. For AWS CLI v1, this parameter should remain blank. For v2, this parameter should be set to --cli-binary-format raw-in-base64-out.

Note

AWS CLI v2 passes binary parameters as base64-encoded strings default.

The following examples use AWS CLI v1:

Request and response in CSV format

- The request consists of a single record and the response is its probability value.

```
aws sagemaker-runtime invoke-endpoint \
   --endpoint-name test-endpoint-sagemaker-xgboost-model \
   --content-type text/csv \
   --accept text/csv \
   --body '1,2,3,4' \
   /dev/stderr 1>/dev/null
```

Output:

```
0.6
```

- The request consists of two records, and the response includes their probabilities, and the model separates the probabilities by a comma. The $'content' expression in the --body tells the command to interpret \n in the content as a line break.

```
aws sagemaker-runtime invoke-endpoint \
   --endpoint-name test-endpoint-sagemaker-xgboost-model \
   --content-type text/csv \
   --accept text/csv \
   --body $'1,2,3,4\n5,6,7,8' \
```

Clarify online explainability
The request consists of two records, the response includes their probabilities, and the model separates the probabilities with a line break.

```
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-1 \
  --content-type text/csv \
  --accept text/csv \
  --body $'1,2,3,4
5,6,7,8' \
/dev/stderr 1>/dev/null
```

Output:

```
0.6
0.3
```

The request consists of two records, the response includes their probabilities, and the model separates the probabilities with a line break.

The request consists of a single record, and the response is probability values (multi-class model, three classes).

```
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-1 \
  --content-type text/csv \
  --accept text/csv \
  --body '1,2,3,4' \
/dev/stderr 1>/dev/null
```

Output:

```
0.1
0.6
0.3
```

The request consists of two records, and the response includes their probability values (multi-class model, three classes).
The request consists of two records, and the response includes predicted label and probability.

```bash
cat $'1,2,3,4
5,6,7,8' 
/dev/stderr 1>/dev/null
```

Output:

0.1,0.6,0.3
0.2,0.5,0.3

The request consists of two records and the response includes label headers and probabilities.

```bash
cat $'1,2,3,4
5,6,7,8' 
/dev/stderr 1>/dev/null
```

Output:

1,0.6
0,0.3

The request consists of two records and the response includes label headers and probabilities.

```bash
cat $'1,2,3,4
5,6,7,8' 
/dev/stderr 1>/dev/null
```

Output:

"['cat','dog','fish']","[0.1,0.6,0.3]"
"['cat','dog','fish']","[0.2,0.5,0.3]"
Request and response in JSON Lines format

- The request consists of a single record and the response is its probability value.

```bash
aws sagemaker-runtime invoke-endpoint
   --endpoint-name test-endpoint-jsonlines
   --content-type application/jsonlines
   --accept application/jsonlines
   --body '{"features":["This is a good product",5]}'
/dev/stderr 1>/dev/null
```

Output:

```json
{"score":0.6}
```

- The request contains two records, and the response includes predicted label and probability.

```bash
aws sagemaker-runtime invoke-endpoint
   --endpoint-name test-endpoint-jsonlines-2
   --content-type application/jsonlines
   --accept application/jsonlines
   --body $'{"features":[1,2,3,4]}\n{"features":[5,6,7,8]}'}
/dev/stderr 1>/dev/null
```

Output:

```json
{"predicted_label":1,"probability":0.6}
{"predicted_label":0,"probability":0.3}
```

- The request contains two records and the response includes label headers and probabilities.

```bash
aws sagemaker-runtime invoke-endpoint
   --endpoint-name test-endpoint-jsonlines-3
   --content-type application/jsonlines
   --accept application/jsonlines
   --body $'{"data":{"features":[1,2,3,4]}}\n{"data":{"features":[5,6,7,8]}}'}
/dev/stderr 1>/dev/null
```

Output:
Request and response in different formats

- The request is in CSV format and the response is in JSON Lines format:

```bash
aws sagemaker-runtime invoke-endpoint \
--endpoint-name test-endpoint-csv-in-jsonlines-out \
--content-type text/csv \
--accept application/jsonlines \
--body $'1,2,3,4\n5,6,7,8' \
/dev/stderr 1>/dev/null
```

Output:

```json
{"probability":0.6}
{"probability":0.3}
```

- The request is in JSON Lines format and the response is in CSV format:

```bash
aws sagemaker-runtime invoke-endpoint \
--endpoint-name test-endpoint-jsonlines-in-csv-out \
--content-type application/jsonlines \
--accept text/csv \
--body $'{"features":[1,2,3,4]}\n{"features":[5,6,7,8]}' \
/dev/stderr 1>/dev/null
```

Output:

```
0.6
0.3
```

After the validations are complete, delete the testing endpoint.
Configure and create an endpoint

Create a new endpoint configuration to fit your model, and use this configuration to create the endpoint. You can use the model container validated in the pre-check step to create an endpoint and enable the SageMaker Clarify online explainability feature.

Use the sagemaker_client object to create an endpoint using the CreateEndpointConfig API. Set the member ClarifyExplainerConfig inside the ExplainerConfig parameter as follows:

```python
sagemaker_client.create_endpoint_config(
    EndpointConfigName='name-of-your-endpoint-config',
    ExplainerConfig={
        'ClarifyExplainerConfig': {
            'EnableExplanations': 'true',
            'InferenceConfig': {
                ...
            },
            'ShapConfig': {
                ...
            }
        },
        ProductionVariants=[{
            'VariantName': 'AllTraffic',
            'ModelName': 'name-of-your-model',
            'InitialInstanceCount': 1,
            'InstanceType': 'ml.m5.xlarge',
        }]
    }
)

sagemaker_client.create_endpoint(
    EndpointName='name-of-your-endpoint',
    EndpointConfigName='name-of-your-endpoint-config'
)
```

The first call to the sagemaker_client object creates a new endpoint configuration with the explainability feature enabled. The second call uses the endpoint configuration to launch the endpoint.
Note
You can also host multiple models in one container behind a SageMaker real-time inference multi-model endpoint and configure online explainability with SageMaker Clarify.

The EnableExplanations expression

The EnableExplanations parameter is a JMESPath Boolean expression string. It is evaluated for each record in the explainability request. If this parameter is evaluated to be true, then the record will be explained. If this parameter is evaluated to be false, then explanations are not be generated.

SageMaker Clarify deserializes the model container output for each record into a JSON compatible data structure, and then uses the EnableExplanations parameter to evaluate the data.

Notes

There are two options for records depending on the format of the model container output.

- If the model container output is in CSV format, then a record is loaded as a JSON array.
- If the model container output is in JSON Lines format, then a record is loaded as a JSON object.

The EnableExplanations parameter is a JMESPath expression that can be passed either during the InvokeEndpoint or CreateEndpointConfig operations. If the JMESPath expression that you supplied is not valid, the endpoint creation will fail. If the expression is valid, but the expression evaluation result is unexpected, then the endpoint will be created successfully, but an error will be generated when the endpoint is invoked. Test your EnableExplanations expression by using the InvokeEndpoint API, and then apply it to the endpoint configuration.

The following are some examples of valid EnableExplanations expression. In the examples, a JMESPath expression encloses a literal using backtick characters. For example, `true` means true.
<table>
<thead>
<tr>
<th>Expression (string representation)</th>
<th>Model container output (string representation)</th>
<th>Evaluation result (Boolean)</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>'true'</code></td>
<td>(N/A)</td>
<td>True</td>
<td>Activate online explainability unconditionally.</td>
</tr>
<tr>
<td><code>'false'</code></td>
<td>(N/A)</td>
<td>False</td>
<td>Deactivate online explainability unconditionally.</td>
</tr>
<tr>
<td><code>[1] &gt; </code>0.5`'</td>
<td>'1,0.6'</td>
<td>True</td>
<td>For each record, the model container outputs its predicted label and probability. Explains a record if its probability (at index 1) is greater than 0.5.</td>
</tr>
<tr>
<td><code>'probability &gt; </code>0.5`'</td>
<td>'{&quot;predicted_label&quot;:1,&quot;probability&quot;:0.6}'</td>
<td>True</td>
<td>For each record, the model container outputs JSON data. Explain a record if its probability is greater than 0.5.</td>
</tr>
<tr>
<td>`!contains(probabilities[:1], max(probabilities))'</td>
<td>'{&quot;probabilities&quot;: [0.4, 0.1, 0.4], &quot;labels&quot;: [&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;]}'</td>
<td>False</td>
<td>For a multi-class model: Explains a record if its predicted label (the class that has the max probability value) is the last class. Literally, the expression means that the max probability</td>
</tr>
</tbody>
</table>
Expression (string representation)  | Model container output (string representation) | Evaluation result (Boolean) | Meaning
--- | --- | --- | ---

### Synthetic dataset

SageMaker Clarify uses the Kernel SHAP algorithm. Given a record (also called a sample or an instance) and the SHAP configuration, the explainer first generates a synthetic dataset. SageMaker Clarify then queries the model container for the predictions of the dataset, and then computes and returns the feature attributions. The size of the synthetic dataset affects the runtime for the Clarify explainer. Larger synthetic datasets take more time to obtain model predictions than smaller ones.

The synthetic dataset size is determined by the following formula:

\[
\text{Synthetic dataset size} = \text{SHAP baseline size} \times n\_samples
\]

The SHAP baseline size is the number of records in the SHAP baseline data. This information is taken from the `ShapBaselineConfig`.

The size of \( n\_samples \) is set by the parameter `NumberOfSamples` in the explainer configuration and the number of features. If the number of features is \( n\_features \), then \( n\_samples \) is the following:

\[
n\_samples = \text{MIN}(\text{NumberOfSamples}, 2^{n\_features} - 2)
\]

The following shows \( n\_samples \) if `NumberOfSamples` is not provided.

\[
n\_samples = \text{MIN}(2\times n\_features + 2^{11}, 2^{n\_features} - 2)
\]

For example, a tabular record with 10 features has a SHAP baseline size of 1. If `NumberOfSamples` is not provided, the synthetic dataset contains 1022 records. If the record has 20 features, the synthetic dataset contains 2088 records.
For NLP problems, n_features is equal to the number of non-text features plus the number of text units.

**Note**

The InvokeEndpoint API has a request timeout limit. If the synthetic dataset is too large, the explainer may not be able to complete the computation within this limit. If necessary, use the previous information to understand and reduce the SHAP baseline size and NumberOfSamples. If your model container is set up to handle batch requests, then you can also adjust the value of MaxRecordCount.

**Invoke the endpoint**

After the endpoint is running, use the SageMaker Runtime **InvokeEndpoint** API in the SageMaker Runtime service to send requests to, or invoke the endpoint. In response, the requests are handled as explainability requests by the SageMaker Clarify explainer.

**Note**

To invoke an endpoint, choose one of the following options:

- For instructions to use Boto3 or the AWS CLI to invoke an endpoint, see [Invoke models for real-time inference](#).
- To use the SageMaker SDK for Python to invoke an endpoint, see the **Predictor** API.

**Request**

The InvokeEndpoint API has an optional parameter EnableExplanations, which is mapped to the HTTP header X-Amzn-SageMaker-Enable-Explanations. If this parameter is provided, it overrides the EnableExplanations parameter of the ClarifyExplainerConfig.

**Note**

The ContentType and Accept parameters of the InvokeEndpoint API are required. Supported formats include MIME type text/csv and application/jsonlines.
Use the `sagemaker_runtime_client` to send a request to the endpoint, as follows:

```python
response = sagemaker_runtime_client.invoke_endpoint(
    EndpointName='name-of-your-endpoint',
    EnableExplanations='true',
    ContentType='text/csv',
    Accept='text/csv',
    Body='1,2,3,4',  # single record (of four numerical features)
)
```

For multi-model endpoints, pass an additional `TargetModel` parameter in the previous example request to specifies which model to target at the endpoint. The multi-model endpoint dynamically loads target models as needed. For more information about multi-model endpoints, see [Host multiple models in one container behind one endpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/multi-model-endpoint.html). See the [SageMaker Clarify Online Explainability on Multi-Model Endpoint Sample Notebook](https://github.com/aws-samples/sage-maker-clarify-explainability-multi-model-endpoint) for an example of how to set up and invoke multiple target models from a single endpoint.

**Response**

If the endpoint is created with `ExplainerConfig`, then a new response schema is used. This new schema is different from, and is not compatible with, an endpoint that lacks the `ExplainerConfig` parameter provided.

The MIME type of the response is `application/json`, and the response payload can be decoded from UTF-8 bytes to a JSON object. The following shows the members of this JSON object are as follows:

- **version**: The version of the response schema in string format. For example, `1.0`.
- **predictions**: The predictions that the request makes have the following:
  - **content_type**: The MIME type of the predictions, referring to the `ContentType` of the model container response.
  - **data**: The predictions data string delivered as the payload of the model container response for the request.
  - **label_headers**: The label headers from the `LabelHeaders` parameter. This is provided in either the explainer configuration or the model container output.
  - **explanations**: The explanations provided in the request payload. If no records are explained, then this member returns the empty object `{}`.
kernel_shap: A key that refers to an array of Kernel SHAP explanations for each record in the request. If a record is not explained, the corresponding explanation is null.

The kernel_shap element has the following members:

- feature_header: The header name of the features provided by the FeatureHeaders parameter in the explainer configuration ExplainerConfig.
- feature_type: The feature type inferred by explainer or provided in the FeatureTypes parameter in the ExplainerConfig. This element is only available for NLP explainability problems.
- attributions: An array of attribution objects. Text features can have multiple attribution objects, each for a unit. The attribution object has the following members:
  - attribution: A list of probability values, given for each class.
  - description: The description of the text units, available only for NLP explainability problems.
  - partial_text: The portion of the text explained by the explainer.
  - start_idx: A zero-based index to identify the array location of the beginning of the partial text fragment.

Code examples: SDK for Python

This section provides sample code to create and invoke an endpoint that uses SageMaker Clarify online explainability. These code examples use the AWS SDK for Python.

Tabular data

The following example uses tabular data and a SageMaker model called model_name. In this example, the model container accepts data in CSV format, and each record has four numerical features. In this minimal configuration, for demonstration purposes only, the SHAP baseline data is set to zero. Refer to SHAP Baselines for Explainability to learn how to choose more appropriate values for ShapBaseline.

Configure the endpoint, as follows:

```python
endpoint_config_name = 'tabular_explainer_endpoint_config'
response = sagemaker_client.create_endpoint_config(
```
Use the endpoint configuration to create an endpoint, as follows:

```python
endpoint_name = 'tabular_explainer_endpoint'
response = sagemaker_client.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name,
)
```

Use the `DescribeEndpoint` API to inspect the progress of creating an endpoint, as follows:

```python
response = sagemaker_client.describe_endpoint(
    EndpointName=endpoint_name,
)
response['EndpointStatus']
```

After the endpoint status is "InService", invoke the endpoint with a test record, as follows:

```python
response = sagemaker_runtime_client.invoke_endpoint(
    EndpointName=endpoint_name,
    ContentType='text/csv',
    Accept='text/csv',
    Body='1,2,3,4',
)
```
Note

In the previous code example, for multi-model endpoints, pass an additional TargetModel parameter in the request to specify which model to target at the endpoint.

Assume that the response has a status code of 200 (no error), and load the response body, as follows:

```python
import codecs
import json
json.load(codecs.getreader('utf-8')(response['Body']))
```

The default action for the endpoint is to explain the record. The following shows example output in the returned JSON object.

```json
{
  "version": "1.0",
  "predictions": {
    "content_type": "text/csv; charset=utf-8",
    "data": "0.0006380207487381"
  },
  "explanations": {
    "kernel_shap": [
      {
        "attribution": [-0.00433456]
      },
      {
        "attribution": [-0.005369821]
      },
      {
        "attribution": [-0.005639821]
      }
    ]
  }
}
```
Use the EnableExplanations parameter to enable on-demand explanations, as follows:

```python
response = sagemaker_runtime_client.invoke_endpoint(
    EndpointName=endpoint_name,
    ContentType='text/csv',
    Accept='text/csv',
    Body='1,2,3,4',
    EnableExplanations='[0]>0.8',
)
```

**Note**

In the previous code example, for multi-model endpoints, pass an additional `TargetModel` parameter in the request to specify which model to target at the endpoint.

In this example, the prediction value is less than the threshold value of 0.8, so the record is not explained:

```json
{
    "version": "1.0",
    "predictions": {
        "content_type": "text/csv; charset=utf-8",
        "data": "0.6380207487381995"
    }
}
```
Use visualization tools to help interpret the returned explanations. The following image shows how SHAP plots can be used to understand how each feature contributes to the prediction. The base value on the diagram, also called the expected value, is the mean predictions of the training dataset. Features that push the expected value higher are red, and features that push the expected value lower are blue. See [SHAP additive force layout](#) for additional information.

See the [full example notebook for tabular data](#).

**Text data**

This section provides a code example to create and invoke an online explainability endpoint for text data. The code example uses SDK for Python.

The following example uses text data and a SageMaker model called `model_name`. In this example, the model container accepts data in CSV format, and each record is a single string.

```python
import sagemaker

endpoint_config_name = 'text_explainer_endpoint_config'
response = sagemaker_client.create_endpoint_config(  
    EndpointConfigName=endpoint_config_name,  
    ProductionVariants=[{  
        'VariantName': 'AllTraffic',  
        'ModelName': model_name,  
        'InitialInstanceCount': 1,  
        'InstanceType': 'ml.m5.xlarge',  
    }],  
    ExplainerConfig={  
        'ClarifyExplainerConfig': {  
            'InferenceConfig': {  
                'FeatureTypes': ['text'],  
                'MaxRecordCount': 100,  
            },  
        },
    },
)```
ShapBaseline: A special token reserved for natural language processing (NLP) processing.

FeatureTypes: Identifies the feature as text. If this parameter is not provided, the explainer will attempt to infer the feature type.

TextConfig: Specifies the unit of granularity and language for the analysis of text features. In this example, the language is English, and granularity token means a word in English text.

NumberOfSamples: A limit to set the upper bounds of the size of the synthetic dataset.

MaxRecordCount: The maximum number of records in a request that the model container can handle. This parameter is set to stabilize performance.

Use the endpoint configuration to create the endpoint, as follows:

```python
epsilon endpoint_name = 'text_explainer_endpoint'
response = sagemaker_client.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name,
)
```

After the status of the endpoint becomes InService, invoke the endpoint. The following code sample uses a test record as follows:

```python
response = sagemaker_runtime_client.invoke_endpoint(
    EndpointName=endpoint_name,
    ContentType='text/csv',
    Accept='text/csv',
    Body='"This is a good product"',
)"
If the request completes successfully, the response body will return a valid JSON object that's similar to the following:

```json
{
  "version": "1.0",
  "predictions": {
    "content_type": "text/csv",
    "data": "0.9766594\n"
  },
  "explanations": {
    "kernel_shap": [
      {
        "attributions": [
          {
            "attribution": [-0.007270948666666712],
            "description": {
              "partial_text": "This",
              "start_idx": 0
            }
          },
          {
            "attribution": [-0.018199033666666628],
            "description": {
              "partial_text": "is",
              "start_idx": 5
            }
          },
          {
            "attribution": [0.01970993241666666],
            "description": {
              "partial_text": "a",
              "start_idx": 8
            }
          }
        ]
      }
    ]
  }
}
```
Use visualization tools to help interpret the returned text attributions. The following image shows how the captum visualization utility can be used to understand how each word contributes to the prediction. The higher the color saturation, the higher the importance given to the word. In this example, a highly saturated bright red color indicates a strong negative contribution. A highly saturated green color indicates a strong positive contribution. The color white indicates that the word has a neutral contribution. See the captum library for additional information on parsing and rendering the attributions.

**Legend:** ■ Negative □ Neutral ■ Positive

<table>
<thead>
<tr>
<th>True Label</th>
<th>Predicted Label</th>
<th>Attribution Label</th>
<th>Attribution Score</th>
<th>Word Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1 (0.57)</td>
<td>True</td>
<td>1.47</td>
<td>This is a <strong>good</strong> product</td>
</tr>
</tbody>
</table>

See the full example notebook for text data.
Troubleshooting guide

If you encounter errors using SageMaker Clarify online explainability, consult the topics in this section.

InvokeEndpoint API fails with the error "ReadTimeoutError:Read timeout on endpoint..."

This error means that the request could not be completed within the 60-second time limit set by the request timeout.

To reduce the request latency, try the following:

- Tune the model's performance during inference. For example, SageMaker Neo can optimize models for inference.
- Allow the model container to handle batch requests.
- Use a larger MaxRecordCount to reduce the number of calls from the explainer to the model container. This will reduce network latency and overhead.
- Use an instance type that has more resources allocated to it. Alternately, assign more instances to the endpoint to help balance the load.
- Reduce the number of records inside a single InvokeEndpoint request.
- Reduce the number of records in the baseline data.
- Use a smaller NumberOfSamples value to reduce the size of the synthetic dataset. For more information about how the number of samples affects your synthetic dataset, see Synthetic dataset.

Serverless Inference

Amazon SageMaker Serverless Inference is a purpose-built inference option that enables you to deploy and scale ML models without configuring or managing any of the underlying infrastructure. On-demand Serverless Inference is ideal for workloads which have idle periods between traffic spurts and can tolerate cold starts. Serverless endpoints automatically launch compute resources and scale them in and out depending on traffic, eliminating the need to choose instance types or manage scaling policies. This takes away the undifferentiated heavy lifting of selecting and managing servers. Serverless Inference integrates with AWS Lambda to offer you high availability, built-in fault tolerance and automatic scaling. With a pay-per-use model, Serverless Inference is a cost-effective option if you have an infrequent or unpredictable traffic pattern. During times
when there are no requests, Serverless Inference scales your endpoint down to 0, helping you to minimize your costs. For more information about pricing for on-demand Serverless Inference, see Amazon SageMaker Pricing.

Optionally, you can also use Provisioned Concurrency with Serverless Inference. Serverless Inference with provisioned concurrency is a cost-effective option when you have predictable bursts in your traffic. Provisioned Concurrency allows you to deploy models on serverless endpoints with predictable performance, and high scalability by keeping your endpoints warm. SageMaker ensures that for the number of Provisioned Concurrency that you allocate, the compute resources are initialized and ready to respond within milliseconds. For Serverless Inference with Provisioned Concurrency, you pay for the compute capacity used to process inference requests, billed by the millisecond, and the amount of data processed. You also pay for Provisioned Concurrency usage, based on the memory configured, duration provisioned, and the amount of concurrency enabled. For more information about pricing for Serverless Inference with Provisioned Concurrency, see Amazon SageMaker Pricing.

You can integrate Serverless Inference with your MLOps Pipelines to streamline your ML workflow, and you can use a serverless endpoint to host a model registered with Model Registry.

Serverless Inference is generally available in 21 AWS Regions: US East (N. Virginia), US East (Ohio), US West (N. California), US West (Oregon), Africa (Cape Town), Asia Pacific (Hong Kong), Asia Pacific (Mumbai), Asia Pacific (Tokyo), Asia Pacific (Seoul), Asia Pacific (Osaka), Asia Pacific (Singapore), Asia Pacific (Sydney), Canada (Central), Europe (Frankfurt), Europe (Ireland), Europe (London), Europe (Paris), Europe (Stockholm), Europe (Milan), Middle East (Bahrain), South America (São Paulo). For more information about Amazon SageMaker regional availability, see the AWS Regional Services List.

How it works

The following diagram shows the workflow of on-demand Serverless Inference and the benefits of using a serverless endpoint.
When you create an on-demand serverless endpoint, SageMaker provisions and manages the compute resources for you. Then, you can make inference requests to the endpoint and receive model predictions in response. SageMaker scales the compute resources up and down as needed to handle your request traffic, and you only pay for what you use.

For Provisioned Concurrency, Serverless Inference also integrates with Application Auto Scaling, so that you can manage Provisioned Concurrency based on a target metric or on a schedule. For more information, see [Automatically scale Provisioned Concurrency for a serverless endpoint](#).

The following sections provide additional details about Serverless Inference and how it works.

**Topics**

- [Container support](#)
- [Memory size](#)
- [Concurrent invocations](#)
- [Minimizing cold starts](#)
- [Feature exclusions](#)

**Container support**

For your endpoint container, you can choose either a SageMaker-provided container or bring your own. SageMaker provides containers for its built-in algorithms and prebuilt Docker images for some of the most common machine learning frameworks, such as Apache MXNet, TensorFlow, PyTorch, and Chainer. For a list of available SageMaker images, see [Available Deep Learning Containers Images](#). If you are bringing your own container, you must modify it to work with
SageMaker. For more information about bringing your own container, see Adapting Your Own Inference Container.

The maximum size of the container image you can use is 10 GB. For serverless endpoints, we recommend creating only one worker in the container and only loading one copy of the model. Note that this is unlike real-time endpoints, where some SageMaker containers may create a worker for each vCPU to process inference requests and load the model in each worker.

If you already have a container for a real-time endpoint, you can use the same container for your serverless endpoint, though some capabilities are excluded. To learn more about the container capabilities that are not supported in Serverless Inference, see Feature exclusions. If you choose to use the same container, SageMaker escrows (retains) a copy of your container image until you delete all endpoints that use the image. SageMaker encrypts the copied image at rest with a SageMaker-owned AWS KMS key.

**Memory size**

Your serverless endpoint has a minimum RAM size of 1024 MB (1 GB), and the maximum RAM size you can choose is 6144 MB (6 GB). The memory sizes you can choose are 1024 MB, 2048 MB, 3072 MB, 4096 MB, 5120 MB, or 6144 MB. Serverless Inference auto-assigns compute resources proportional to the memory you select. If you choose a larger memory size, your container has access to more vCPUs. Choose your endpoint's memory size according to your model size. Generally, the memory size should be at least as large as your model size. You may need to benchmark in order to choose the right memory selection for your model based on your latency SLAs. For a step by step guide to benchmark, see Introducing the Amazon SageMaker Serverless Inference Benchmarking Toolkit. The memory size increments have different pricing; see the Amazon SageMaker pricing page for more information.

Regardless of the memory size you choose, your serverless endpoint has 5 GB of ephemeral disk storage available. For help with container permissions issues when working with storage, see Troubleshooting.

**Concurrent invocations**

On-demand Serverless Inference manages predefined scaling policies and quotas for the capacity of your endpoint. Serverless endpoints have a quota for how many concurrent invocations can be processed at the same time. If the endpoint is invoked before it finishes processing the first request, then it handles the second request concurrently.
The total concurrency that you can share between all serverless endpoints in your account depends on your region:

- For the US East (Ohio), US East (N. Virginia), US West (Oregon), Asia Pacific (Singapore), Asia Pacific (Sydney), Asia Pacific (Tokyo), Europe (Frankfurt), and Europe (Ireland) Regions, the total concurrency you can share between all serverless endpoints per Region in your account is 1000.

- For the US West (N. California), Africa (Cape Town), Asia Pacific (Hong Kong), Asia Pacific (Mumbai), Asia Pacific (Osaka), Asia Pacific (Seoul), Canada (Central), Europe (London), Europe (Milan), Europe (Paris), Europe (Stockholm), Middle East (Bahrain), and South America (São Paulo) Regions, the total concurrency per Region in your account is 500.

You can set the maximum concurrency for a single endpoint up to 200, and the total number of serverless endpoints you can host in a Region is 50. The maximum concurrency for an individual endpoint prevents that endpoint from taking up all of the invocations allowed for your account, and any endpoint invocations beyond the maximum are throttled.

**Note**

Provisioned Concurrency that you assign to a serverless endpoint should always be less than or equal to the maximum concurrency that you assigned to that endpoint.

To learn how to set the maximum concurrency for your endpoint, see [Create an endpoint configuration](#). For more information about quotas and limits, see [Amazon SageMaker endpoints and quotas](#) in the *AWS General Reference*. To request a service limit increase, contact [AWS Support](#). For instructions on how to request a service limit increase, see [Supported Regions and Quotas](#).

**Minimizing cold starts**

If your on-demand Serverless Inference endpoint does not receive traffic for a while and then your endpoint suddenly receives new requests, it can take some time for your endpoint to spin up the compute resources to process the requests. This is called a *cold start*. Since serverless endpoints provision compute resources on demand, your endpoint may experience cold starts. A cold start can also occur if your concurrent requests exceed the current concurrent request usage. The cold start time depends on your model size, how long it takes to download your model, and the start-up time of your container.
To monitor how long your cold start time is, you can use the Amazon CloudWatch metric `OverheadLatency` to monitor your serverless endpoint. This metric tracks the time it takes to launch new compute resources for your endpoint. To learn more about using CloudWatch metrics with serverless endpoints, see Monitor a serverless endpoint.

You can minimize cold starts by using Provisioned Concurrency. SageMaker keeps the endpoint warm and ready to respond in milliseconds, for the number of Provisioned Concurrency that you allocated.

**Feature exclusions**

Some of the features currently available for SageMaker Real-time Inference are not supported for Serverless Inference, including GPUs, AWS marketplace model packages, private Docker registries, Multi-Model Endpoints, VPC configuration, network isolation, data capture, multiple production variants, Model Monitor, and inference pipelines.

You cannot convert your instance-based, real-time endpoint to a serverless endpoint. If you try to update your real-time endpoint to serverless, you receive a `ValidationError` message. You can convert a serverless endpoint to real-time, but once you make the update, you cannot roll it back to serverless.

**Getting started**

You can create, update, describe, and delete a serverless endpoint using the SageMaker console, the AWS SDKs, the Amazon SageMaker Python SDK, and the AWS CLI. You can invoke your endpoint using the AWS SDKs, the Amazon SageMaker Python SDK, and the AWS CLI. For serverless endpoints with Provisioned Concurrency, you can use Application Auto Scaling to auto scale Provisioned Concurrency based on a target metric or a schedule. For more information about how to set up and use a serverless endpoint, read the guide Create, invoke, update, and delete a serverless endpoint. For more information on auto scaling serverless endpoints with Provisioned Concurrency, see Automatically scale Provisioned Concurrency for a serverless endpoint.

⚠️ **Note**

Application Auto Scaling for Serverless Inference with Provisioned Concurrency is currently not supported on AWS CloudFormation.
Example notebooks and blogs

For Jupyter notebook examples that show end-to-end serverless endpoint workflows, see the Serverless Inference example notebooks.

Create, invoke, update, and delete a serverless endpoint

Unlike other SageMaker real-time endpoints, Serverless Inference manages compute resources for you, reducing complexity so you can focus on your ML model instead of on managing infrastructure. The following guide highlights the key capabilities of serverless endpoints: how to create, invoke, update, describe, or delete an endpoint. You can use the SageMaker console, the AWS SDKs, the Amazon SageMaker Python SDK, or the AWS CLI to manage your serverless endpoints.

Topics

- Prerequisites
- Create a serverless endpoint
- Invoke a serverless endpoint
- Update a serverless endpoint
- Describe a serverless endpoint
- Delete a serverless endpoint

Prerequisites

Before you can create a serverless endpoint, complete the following prerequisites.

1. **Set up an AWS account.** You first need an AWS account and an AWS Identity and Access Management administrator user. For instructions on how to set up an AWS account, see How do I create and activate a new AWS account?. For instructions on how to secure your account with an IAM administrator user, see Creating your first IAM admin user and user group in the IAM User Guide.

2. **Create an Amazon S3 bucket.** You use an Amazon S3 bucket to store your model artifacts. To learn how to create a bucket, see Create your first S3 bucket in the Amazon S3 User Guide.

3. **Upload your model artifacts to your S3 bucket.** For instructions on how to upload your model to your bucket, see Upload an object to your bucket in the Amazon S3 User Guide.
4. **Create an IAM role for Amazon SageMaker.** Amazon SageMaker needs access to the S3 bucket that stores your model. Create an IAM role with a policy that gives SageMaker read access to your bucket. The following procedure shows how to create a role in the console, but you can also use the CreateRole API from the IAM User Guide. For information on giving your role more granular permissions based on your use case, see SageMaker Roles.

   a. Sign in to the IAM console.
   b. In the navigation tab, choose Roles.
   c. Choose Create Role.
   d. For Select type of trusted entity, choose AWS service and then choose SageMaker.
   e. Choose Next: Permissions and then choose Next: Tags.
   f. (Optional) Add tags as key-value pairs if you want to have metadata for the role.
   g. Choose Next: Review.
   h. For Role name, enter a name for the new role that is unique within your AWS account. You cannot edit the role name after creating the role.
   i. (Optional) For Role description, enter a description for the new role.
   j. Choose Create role.

5. **Attach S3 bucket permissions to your SageMaker role.** After creating an IAM role, attach a policy that gives SageMaker permission to access the S3 bucket containing your model artifacts.

   a. In the IAM console navigation tab, choose Roles.
   b. From the list of roles, search for the role you created in the previous step by name.
   c. Choose your role, and then choose Attach policies.
   d. For Attach permissions, choose Create policy.
   e. In the Create policy view, select the JSON tab.
   f. Add the following policy statement into the JSON editor. Make sure to replace `<your-bucket-name>` with the name of the S3 bucket that stores your model artifacts. If you want to restrict the access to a specific folder or file in your bucket, you can also specify the Amazon S3 folder path, for example, `<your-bucket-name>/<model-folder>`.

   ```json
   {
     "Version": "2012-10-17",
     "Statement": [
   ```
g. Choose **Next: Tags**.

h. (Optional) Add tags in key-value pairs to the policy.

i. Choose **Next: Review**.

j. For **Name**, enter a name for the new policy.

k. (Optional) Add a **Description** for the policy.

l. Choose **Create policy**.

m. After creating the policy, return to **Roles** in the [IAM console](https://console.aws.amazon.com/iam) and select your SageMaker role.

n. Choose **Attach policies**.

o. For **Attach permissions**, search for the policy you created by name. Select it and choose **Attach policy**.

6. **Select a prebuilt Docker container image or bring your own.** The container you choose serves inference on your endpoint. SageMaker provides containers for built-in algorithms and prebuilt Docker images for some of the most common machine learning frameworks, such as Apache MXNet, TensorFlow, PyTorch, and Chainer. For a full list of the available SageMaker images, see [Available Deep Learning Containers Images](https://docs.aws.amazon.com/sagemaker/latest/dg/available-docker-images.html).

   If none of the existing SageMaker containers meet your needs, you may need to create your own Docker container. For information about how to create your Docker image and make it compatible with SageMaker, see [Use your own inference code](https://docs.aws.amazon.com/sagemaker/latest/dg/serverless-infrastructure.html). To use your container with a serverless endpoint, the container image must reside in an Amazon ECR repository within the same AWS account that creates the endpoint.

7. **(Optional) Register your model with Model Registry.** [SageMaker Model Registry](https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry.html) helps you catalog and manage versions of your models for use in ML pipelines. For more information about registering a version of your model, see [Create a Model Group](https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry-create-model-group.html) and [Register a Model Version](https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry-register-model-version.html). For an example of a Model Registry and Serverless Inference workflow, see the following [example notebook](https://github.com/awslabs/aws-serverless-inference-with-sagemaker/blob/master/example.ipynb).
8. **(Optional) Bring an AWS KMS key.** When setting up a serverless endpoint, you have the option to specify a KMS key that SageMaker uses to encrypt your Amazon ECR image. Note that the key policy for the KMS key must grant access to the IAM role you specify when setting up your endpoint. To learn more about KMS keys, see the [AWS Key Management Service Developer Guide](https://docs.aws.amazon.com/kms/latest/userguide/)

**Create a serverless endpoint**

To create a serverless endpoint, you can use the Amazon SageMaker console, the APIs, or the AWS CLI. You can create a serverless endpoint using a similar process as a real-time endpoint.

**Topics**

- Create a model
- Create an endpoint configuration
- Create an endpoint

**Create a model**

To create your model, you must provide the location of your model artifacts and container image. You can also use a model version from SageMaker Model Registry. The examples in the following sections show you how to create a model using the `CreateModel` API, Model Registry, and the Amazon SageMaker console.

**To create a model (using Model Registry)**

Model Registry is a feature of SageMaker that helps you catalog and manage versions of your model for use in ML pipelines. To use Model Registry with Serverless Inference, you must first register a model version in a Model Registry model group. To learn how to register a model in Model Registry, follow the procedures in [Create a Model Group](https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry-create-model-group.html) and [Register a Model Version](https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry-register-model-version.html).

The following example requires you to have the ARN of a registered model version and uses the [AWS SDK for Python (Boto3)](https://boto3.amazonaws.com/v1/documentation/api/latest/index.html) to call the `CreateModel` API. For Serverless Inference, Model Registry is currently only supported by the AWS SDK for Python (Boto3). For the example, specify the following values:

- For `model_name`, enter a name for the model.
- For `sagemaker_role`, you can use the default SageMaker-created role or a customized SageMaker IAM role from Step 4 of the [Prerequisites](https://docs.aws.amazon.com/sagemaker/latest/dg/ccc-environment-configuration.html) section.
• For ModelPackageName, specify the ARN for your model version, which must be registered to a model group in Model Registry.

```python
#Setup
import boto3
import sagemaker
region = boto3.Session().region_name
client = boto3.client("sagemaker", region_name=region)

#Role to give SageMaker permission to access AWS services.
sagemaker_role = sagemaker.get_execution_role()

#Specify a name for the model
model_name = "<name-for-model>"

#Specify a Model Registry model version
container_list = [
    {
        "ModelPackageName": <model-version-arn>
    }
]

#Create the model
response = client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    container_list
)
```

**To create a model (using API)**

The following example uses the [AWS SDK for Python (Boto3)](https://boto3.amazonaws.com/v1/documentation/api/latest/index.html) to call the [CreateModel](https://boto3.amazonaws.com/v1/documentation/api/latest/reference/services/sagemaker.html#SageMaker.Client.create_model) API. Specify the following values:

- For `sagemaker_role`, you can use the default SageMaker-created role or a customized SageMaker IAM role from Step 4 of the [Prerequisites](#) section.
- For `model_url`, specify the Amazon S3 URI to your model.
- For `container`, retrieve the container you want to use by its Amazon ECR path. This example uses a SageMaker-provided XGBoost container. If you have not selected a SageMaker container or brought your own, see Step 6 of the [Prerequisites](#) section for more information.
• For model_name, enter a name for the model.

```python
#Setup
import boto3
import sagemaker
region = boto3.Session().region_name
client = boto3.client("sagemaker", region_name=region)

#Role to give SageMaker permission to access AWS services.
sagemaker_role = sagemaker.get_execution_role()

#Get model from S3
model_url = "s3://DOC-EXAMPLE-BUCKET/models/model.tar.gz"

#Get container image (prebuilt example)
from sagemaker import image_uris
container = image_uris.retrieve("xgboost", region, "0.90-1")

#Create model
model_name = "<name-for-model>"

response = client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    Containers = [
        {
            "Image": container,
            "Mode": "SingleModel",
            "ModelDataUrl": model_url,
        }
    ]
)
```

To create a model (using the console)

1. Sign in to the Amazon SageMaker console.
2. In the navigation tab, choose Inference.
3. Next, choose Models.
4. Choose Create model.
5. For Model name, enter a name for the model that is unique to your account and AWS Region.
6. For IAM role, either select an IAM role you have already created (see Prerequisites) or allow SageMaker to create one for you.
7. In Container definition 1, for Container input options, select Provide model artifacts and input location.

8. For Provide model artifacts and inference image options, select Use a single model.

9. For Location of inference code image, enter an Amazon ECR path to a container. The image must either be a SageMaker-provided first party image (e.g. TensorFlow, XGBoost) or an image that resides in an Amazon ECR repository within the same account in which you are creating the endpoint. If you do not have a container, go back to Step 6 of the Prerequisites section for more information.

10. For Location of model artifacts, enter the Amazon S3 URI to your ML model. For example, 
    s3://DOC-EXAMPLE-BUCKET/models/model.tar.gz.

11. (Optional) For Tags, add key-value pairs to create metadata for your model.

12. Choose Create model.

Create an endpoint configuration

After you create a model, create an endpoint configuration. You can then deploy your model using the specifications in your endpoint configuration. In the configuration, you specify whether you want a real-time or serverless endpoint. To create a serverless endpoint configuration, you can use the Amazon SageMaker console, the CreateEndpointConfig API, or the AWS CLI. The API and console approaches are outlined in the following sections.

To create an endpoint configuration (using API)

The following example uses the AWS SDK for Python (Boto3) to call the CreateEndpointConfig API. Specify the following values:

- For EndpointConfigName, choose a name for the endpoint configuration. The name should be unique within your account in a Region.
- (Optional) For KmsKeyId, use the key ID, key ARN, alias name, or alias ARN for an AWS KMS key that you want to use. SageMaker uses this key to encrypt your Amazon ECR image.
- For ModelName, use the name of the model you want to deploy. It should be the same model that you used in the Create a model step.
- For ServerlessConfig:
  - Set MemorySizeInMB to 2048. For this example, we set the memory size to 2048 MB, but you can choose any of the following values for your memory size: 1024 MB, 2048 MB, 3072 MB, 4096 MB, 5120 MB, or 6144 MB.
• Set MaxConcurrency to 20. For this example, we set the maximum concurrency to 20. The maximum number of concurrent invocations you can set for a serverless endpoint is 200, and the minimum value you can choose is 1.

• (Optional) To use Provisioned Concurrency, set ProvisionedConcurrency to 10. For this example, we set the Provisioned Concurrency to 10. The ProvisionedConcurrency number for a serverless endpoint must be lower than or equal to the MaxConcurrency number. You can leave it empty if you want to use on-demand Serverless Inference endpoint. You can dynamically scale Provision Concurrency. For more information, see Automatically scale Provisioned Concurrency for a serverless endpoint.

```python
response = client.create_endpoint_config(
    EndpointConfigName="<your-endpoint-configuration>",
    KmsKeyId="arn:aws:kms:us-east-1:123456789012:key/143ef68f-76fd-45e3-abba-ed28fc8d3d5e",
    ProductionVariants=[
        {
            "ModelName": "<your-model-name>",
            "VariantName": "AllTraffic",
            "ServerlessConfig": {
                "MemorySizeInMB": 2048,
                "MaxConcurrency": 20,
                "ProvisionedConcurrency": 10,
            }
        }
    ]
)
```

To create an endpoint configuration (using the console)

1. Sign in to the Amazon SageMaker console.
2. In the navigation tab, choose Inference.
3. Next, choose Endpoint configurations.
4. Choose Create endpoint configuration.
5. For Endpoint configuration name, enter a name that is unique within your account in a Region.
6. For Type of endpoint, select Serverless.
Create endpoint configuration

To deploy models to Amazon SageMaker, first create an endpoint configuration. In the configuration, specify which models to
deploy, and the relative traffic weighting and hardware requirements for each. See Deploying a Model on Amazon SageMaker
Hosting Services. Learn more about the API

**Endpoint configuration**

Endpoint configuration name

Maximum of 63 alphanumeric characters. Can include hyphens (-), but not spaces. Must be unique within
your account in an AWS Region.

Type of endpoint
- Provisioned
- Serverless

Encryption key - optional
Encrypt your data. Choose an existing KMS key or enter a key's ARN.

Default SageMaker Role

**Variants**

- **Provisioned Concurrency**
  Serverless endpoints now supports provisioned concurrency. After selecting a production variant click edit in the actions column
  below to set the provisioned concurrency for your production variant. Learn more

**Production**

<table>
<thead>
<tr>
<th>Model name</th>
<th>Training job</th>
<th>Variant name</th>
<th>Memory Size</th>
<th>Max Concurrency</th>
<th>Provisioned Concurrency</th>
<th>Actions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

There are currently no resources

Create production variant

**Tags - optional**

<table>
<thead>
<tr>
<th>Key</th>
<th>Value</th>
<th>Remove</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Add tag
7. For **Production variants**, choose **Add model**.

8. Under **Add model**, select the model you want to use from the list of models and then choose **Save**.

9. After adding your model, under **Actions**, choose **Edit**.

10. For **Memory size**, choose the memory size you want in GB.

```
Edit Production Variant
```

- **Model name**: test-gb-gamma-model
- **Variant name**: variant-name-1
- **Memory Size**: 1 GB
- **Max Concurrency**: 20

*Provisioned concurrency setting - optional*

Provisioned concurrency enables you to deploy models on serverless endpoints with predictable performance and high scalability. For the set number of concurrent invocations, SageMaker will keep underlying compute warm and ready to respond instantaneously without cold starts.

Numeric values only. Provisioned concurrency must be <= the Max Concurrency set for the production variant.

11. For **Max Concurrency**, enter your desired maximum concurrent invocations for the endpoint. The maximum value you can enter is 200 and the minimum is 1.
12. (Optional) To use Provisioned Concurrency, enter the desired number of concurrent invocations in the **Provisioned Concurrency setting** field. The number of provisioned concurrent invocations must be less than or equal to the number of maximum concurrent invocations.

13. Choose **Save**.

14. (Optional) For **Tags**, enter key-value pairs if you want to create metadata for your endpoint configuration.

15. Choose **Create endpoint configuration**.

**Create an endpoint**

To create a serverless endpoint, you can use the [Amazon SageMaker console](https://console.aws.amazon.com), the `CreateEndpoint` API, or the AWS CLI. The API and console approaches are outlined in the following sections. Once you create your endpoint, it can take a few minutes for the endpoint to become available.

**To create an endpoint (using API)**

The following example uses the [AWS SDK for Python (Boto3)](https://boto3.amazonaws.com/v1/documentation/api/latest/) to call the `CreateEndpoint` API. Specify the following values:

- For **EndpointName**, enter a name for the endpoint that is unique within a Region in your account.
- For **EndpointConfigName**, use the name of the endpoint configuration that you created in the previous section.

```python
response = client.create_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-endpoint-config>"
)
```

**To create an endpoint (using the console)**

1. Sign in to the [Amazon SageMaker console](https://console.aws.amazon.com).
2. In the navigation tab, choose **Inference**.
3. Next, choose **Endpoints**.
4. Choose **Create endpoint**.
5. For **Endpoint name**, enter a name than is unique within a Region in your account.
6. For **Attach endpoint configuration**, select **Use an existing endpoint configuration**.

7. For **Endpoint configuration**, select the name of the endpoint configuration you created in the previous section and then choose **Select endpoint configuration**.

8. (Optional) For **Tags**, enter key-value pairs if you want to create metadata for your endpoint.

9. Choose **Create endpoint**.
Create and configure endpoint

To deploy models to Amazon SageMaker, first create an endpoint. Provide an endpoint configuration to specify which models to deploy and the hardware requirements for each. See Deploying a Model on Amazon SageMaker Hosting Services. Learn more about the API.

**Endpoint**

**Endpoint name**

Your application uses this name to access this endpoint.

Maximum of 63 alphanumeric characters. Can include hyphens (-), but not spaces. Must be unique within your account in an AWS Region.

---

**Attach endpoint configuration**

- Use an existing endpoint configuration
  
  Use an existing endpoint configuration or clone an endpoint configuration

- Create a new endpoint configuration
  
  Add models and configure the instance and initial weight for each model.

---

**Endpoint configuration**

<table>
<thead>
<tr>
<th>Endpoint configuration name</th>
<th>Encryption key</th>
</tr>
</thead>
<tbody>
<tr>
<td>new-ex-342</td>
<td>-</td>
</tr>
</tbody>
</table>

**Variants**

**Production**

<table>
<thead>
<tr>
<th>Model name</th>
<th>Training job</th>
<th>Variant name</th>
<th>Memory Size</th>
<th>Max Concurrency</th>
<th>Provisioned Concurrency</th>
</tr>
</thead>
<tbody>
<tr>
<td>my-model</td>
<td></td>
<td>var-name-23</td>
<td>1 GB</td>
<td>20</td>
<td>10</td>
</tr>
</tbody>
</table>

---

**Tags - optional**

<table>
<thead>
<tr>
<th>Key</th>
<th>Value</th>
<th>Remove</th>
</tr>
</thead>
</table>

Create, invoke, update, and delete a serverless endpoint

**Add tag**
Invoke a serverless endpoint

In order to perform inference using a serverless endpoint, you must send an HTTP request to the endpoint. You can use the `InvokeEndpoint` API or the AWS CLI, which make a POST request to invoke your endpoint. The maximum request and response payload size for serverless invocations is 4 MB. For serverless endpoints:

- The model must download and the server must respond successfully to `/ping` within 3 minutes.
- The timeout for the container to respond to inference requests to `/invocations` is 1 minute.

To invoke an endpoint

The following example uses the AWS SDK for Python (Boto3) to call the `InvokeEndpoint` API. Note that unlike the other API calls in this guide, for InvokeEndpoint, you must use SageMaker Runtime Runtime as the client. Specify the following values:

- For `endpoint_name`, use the name of the in-service serverless endpoint you want to invoke.
- For `content_type`, specify the MIME type of your input data in the request body (for example, `application/json`).
- For `payload`, use your request payload for inference. Your payload should be in bytes or a file-like object.

```python
runtime = boto3.client("sagemaker-runtime")
endpoint_name = "<your-endpoint-name>"
content_type = "<request-mime-type>"
payload = <your-request-body>

response = runtime.invoke_endpoint(
    EndpointName=endpoint_name,
    ContentType=content_type,
    Body=payload
)
```

Update a serverless endpoint

Before updating your endpoint, create a new endpoint configuration or use an existing endpoint configuration. The endpoint configuration is where you specify the changes for your update. Then,
you can update your endpoint with the SageMaker console, the UpdateEndpoint API, or the AWS CLI. The process for updating a serverless endpoint is the same as the process for updating a real-time endpoint. Note that when updating your endpoint, you can experience cold starts when making requests to the endpoint because SageMaker must re-initialize your container and model.

You may want to update an on-demand serverless endpoint to a serverless endpoint with provisioned concurrency or adjust the Provisioned Concurrency value for an existing serverless endpoint with provisioned concurrency. For both cases, you will have to create a new serverless endpoint configuration with the desired value for Provisioned Concurrency, and apply UpdateEndpoint to the existing serverless endpoint. For more information on creating a new serverless endpoint configuration with Provisioned Concurrency, see Create an endpoint configuration.

If you want to remove Provisioned Concurrency from a serverless endpoint, you will have to create a new endpoint configuration without specifying any value for Provisioned Concurrency, and then apply UpdateEndpoint to the endpoint.

⚠️ Note

Updating a real-time inference endpoint to either an on-demand serverless endpoint or a serverless endpoint with Provisioned Concurrency is currently not supported.

Update the endpoint

After creating a new serverless endpoint configuration you can use the AWS SDK for Python (Boto3) or the SageMaker console to update an existing serverless endpoint. Examples of how to update your endpoint using the AWS SDK for Python (Boto3) and the SageMaker console are outlined in the following sections.

To update the endpoint (using Boto3)

The following example uses the AWS SDK for Python (Boto3) to call the update_endpoint method. Specify at least the following parameters when calling the method:

- For EndpointName, use the name of the endpoint you’re updating.
- For EndpointConfigName, use the name of the endpoint configuration that you want to use for the update.
response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<new-endpoint-config>",
)

To update the endpoint (using the console)

1. Sign in to the Amazon SageMaker console.
2. In the navigation tab, choose Inference.
3. Next, choose Endpoints.
4. From the list of endpoints, select the endpoint you want to update.
5. Choose Change in Endpoint configuration settings section.
6. For Change the Endpoint configuration, choose Use an existing endpoint configuration.
7. From the list of endpoint configurations, select the one you want to use for your update.
8. Choose Select endpoint configuration.
9. Choose Update endpoint.

Describe a serverless endpoint

You might want to retrieve information about your endpoint, including details such as the endpoint’s ARN, current status, deployment configuration, and failure reasons. You can find information about your endpoint using the SageMaker console, the DescribeEndpoint API, or the AWS CLI.

To describe an endpoint (using API)

The following example uses the AWS SDK for Python (Boto3) to call the DescribeEndpoint API. For EndpointName, use the name of the endpoint you want to check.

response = client.describe_endpoint(
    EndpointName="<your-endpoint-name>",
)

To describe an endpoint (using the console)

1. Sign in to the Amazon SageMaker console.
2. In the navigation tab, choose **Inference**.
3. Next, choose **Endpoints**.
4. From the list of endpoints, choose the endpoint you want to check.

The endpoint page contains the information about your endpoint.

**Delete a serverless endpoint**

You can delete your serverless endpoint using the SageMaker console, the DeleteEndpoint API, or the AWS CLI. The following examples show you how to delete your endpoint through the API and the SageMaker console.

**To delete an endpoint (using API)**

The following example uses the AWS SDK for Python (Boto3) to call the DeleteEndpoint API. For EndpointName, use the name of the serverless endpoint you want to delete.

```python
response = client.delete_endpoint(
    EndpointName="<your-endpoint-name>",
)
```

**To delete an endpoint (using the console)**

1. Sign in to the Amazon SageMaker console.
2. In the navigation tab, choose **Inference**.
3. Next, choose **Endpoints**.
4. From the list of endpoints, select the endpoint you want to delete.
5. Choose the **Actions** drop-down list, and then choose **Delete**.
6. When prompted again, choose **Delete**.

Your endpoint should now begin the deletion process.

**Monitor a serverless endpoint**

To monitor your serverless endpoint, you can use Amazon CloudWatch alarms. CloudWatch is a service that collects metrics in real time from your AWS applications and resources. An alarm watches metrics as they are collected and gives you the ability to pre-specify a threshold and the
actions to take if that threshold is breached. For example, your CloudWatch alarm can send you a notification if your endpoint breaches an error threshold. By setting up CloudWatch alarms, you gain visibility into the performance and functionality of your endpoint. For more information about CloudWatch alarms, see Using Amazon CloudWatch alarms in the Amazon CloudWatch User Guide.

Monitoring with CloudWatch

The metrics below are an exhaustive list of metrics for serverless endpoints. Any metric not listed below is not published for serverless endpoints. For information about the following metrics, see Monitor Amazon SageMaker with Amazon CloudWatch.

Common endpoint metrics

These CloudWatch metrics are the same as the metrics published for real-time endpoints.

The OverheadLatency metric tracks all additional latency that SageMaker added which includes the cold start time for launching new compute resources for your serverless endpoint. Compared to on-demand serverless endpoints, the OverheadLatency for serverless endpoints with provision concurrency is generally significantly less.

Serverless endpoints can also use the Invocations4XXErrors, Invocations5XXErrors, Invocations, ModelLatency, ModelSetupTime and MemoryUtilization metrics. To learn more about these metrics, see SageMaker Endpoint Invocation Metrics.

Common serverless endpoint metrics

These CloudWatch metrics are published for both on-demand serverless endpoints and serverless endpoint with Provisioned Concurrency.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Unit/Stats</th>
</tr>
</thead>
</table>
| ServerlessConcurrencyExecutionsUtilization | The number of concurrent executions divided by the maximum concurrency. | Units: None
Valid statistics: Average, Max, Min |

Serverless endpoint with Provisioned Concurrency metrics

These CloudWatch metrics are published for serverless endpoints with Provisioned Concurrency.
<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Unit/Stats</th>
</tr>
</thead>
<tbody>
<tr>
<td>ServerlessProvisionedConcurrencyExecutions</td>
<td>The number of concurrent executions handled by the endpoint.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
</tr>
<tr>
<td>ServerlessProvisionedConcurrencyUtilization</td>
<td>The number of concurrent executions divided by the allocated Provisioned Concurrency.</td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
</tr>
<tr>
<td>ServerlessProvisionedConcurrencyInvocations</td>
<td>The number of InvokeEndpoint requests handled by Provisioned Concurrency.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
</tr>
<tr>
<td>ServerlessProvisionedConcurrencySpilloverInvocations</td>
<td>The number of InvokeEndpoint requests not handled by Provisioned Concurrency, that is handled by on-demand Serverless Inference.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
</tr>
</tbody>
</table>

**Logs**

If you want to monitor the logs from your endpoint for debugging or progress analysis, you can use Amazon CloudWatch Logs. The SageMaker-provided log group that you can use for serverless endpoints is /aws/sagemaker/Endpoints/[EndpointName]. For more information about using CloudWatch Logs in SageMaker, see [Log Amazon SageMaker Events with Amazon CloudWatch](#). To learn more about CloudWatch Logs, see [What is Amazon CloudWatch Logs?](#) in the [Amazon CloudWatch Logs User Guide](#).

**Automatically scale Provisioned Concurrency for a serverless endpoint**

Amazon SageMaker automatically scales in or out on-demand serverless endpoints. For serverless endpoints with Provisioned Concurrency you can use Application Auto Scaling to scale up or down the Provisioned Concurrency based on your traffic profile, thus optimizing costs.
The following are the prerequisites to autoscale Provisioned Concurrency on serverless endpoints:

- **Register a model**
- **Define a scaling policy**
- **Apply a scaling policy**

Before you can use autoscaling, you must have already deployed a model to a serverless endpoint with Provisioned Concurrency. Deployed models are referred to as *production variants*. See [Create an endpoint configuration](#) and [Create an endpoint](#) for more information about deploying a model to a serverless endpoint with Provisioned Concurrency. To specify the metrics and target values for a scaling policy, you must configure a scaling policy. For more information on how to define a scaling policy, see [Define a scaling policy](#). After registering your model and defining a scaling policy, apply the scaling policy to the registered model. For information on how to apply the scaling policy, see [Apply a scaling policy](#).

For details on other prerequisites and components used with autoscaling, see the [Auto scaling overview](#) section in the SageMaker autoscaling documentation.

### Register a model

To add autoscaling to a serverless endpoint with Provisioned Concurrency, you first must register your model (production variant) using AWS CLI or Application Auto Scaling API.

**Register a model (AWS CLI)**

To register your model, use the `register-scalable-target` AWS CLI command with the following parameters:

- **--service-namespace** – Set this value to sagemaker.
- **--resource-id** – The resource identifier for the model (specifically the production variant). For this parameter, the resource type is endpoint and the unique identifier is the name of the production variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- **--scalable-dimension** – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.
- **--min-capacity** – The minimum number of Provisioned Concurrency for the model. Set `--min-capacity` to at least 1. It must be equal to or less than the value specified for `--max-capacity`.

Automatically scale Provisioned Concurrency for a serverless endpoint
• **--max-capacity** – The maximum number of Provisioned Concurrency that should be enabled through Application Auto Scaling. Set **--max-capacity** to a minimum of 1. It must be greater than or equal to the value specified for **--min-capacity**.

The following example shows how to register a model named MyVariant that is dynamically scaled to have 1 to 10 Provisioned Concurrency value:

```bash
aws application-autoscaling register-scalable-target \
  --service-namespace sagemaker \
  --scalable-dimension sagemaker:variant:DesiredProvisionedConcurrency \
  --resource-id endpoint/MyEndpoint/variant/MyVariant \
  --min-capacity 1 \
  --max-capacity 10
```

**Register a model (Application Auto Scaling API)**

To register your model, use the `RegisterScalableTarget` Application Auto Scaling API action with the following parameters:

- **ServiceNamespace** – Set this value to `sagemaker`.
- **ResourceId** – The resource identifier for the model (specifically the production variant). For this parameter, the resource type is `endpoint` and the unique identifier is the name of the production variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- **ScalableDimension** – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.
- **MinCapacity** – The minimum number of Provisioned Concurrency for the model. Set **MinCapacity** to at least 1. It must be equal to or less than the value specified for **MaxCapacity**.
- **MaxCapacity** – The maximum number of Provisioned Concurrency that should be enabled through Application Auto Scaling. Set **MaxCapacity** to a minimum of 1. It must be greater than or equal to the value specified for **MinCapacity**.

The following example shows how to register a model named MyVariant that is dynamically scaled to have 1 to 10 Provisioned Concurrency value:

```plaintext
POST / HTTP/1.1

Automatically scale Provisioned Concurrency for a serverless endpoint

4274
Define a scaling policy

To specify the metrics and target values for a scaling policy, you can configure a target-tracking scaling policy. Define the scaling policy as a JSON block in a text file. You can then use that text file when invoking the AWS CLI or the Application Auto Scaling API. To quickly define a target-tracking scaling policy for a serverless endpoint, use the SageMakerVariantProvisionedConcurrencyUtilization predefined metric.

```
{
    "TargetValue": 0.5,
    "PredefinedMetricSpecification": [
        "PredefinedMetricType": "SageMakerVariantProvisionedConcurrencyUtilization"
    ],
    "ScaleOutCooldown": 1,
    "ScaleInCooldown": 1
}
```

Apply a scaling policy

After registering your model, you can apply a scaling policy to your serverless endpoint with Provisioned Concurrency. See Apply a target-tracking scaling policy to apply a target-tracking scaling policy that you have defined. If the traffic flow to your serverless endpoint has a predictable...
routine then instead of applying a target-tracking scaling policy you might want to schedule scaling actions at specific times. For more information on scheduling scaling actions, see Scheduled scaling.

**Apply a target-tracking scaling policy**

You can use the AWS Management Console, AWS CLI or the Application Auto Scaling API to apply a target-tracking scaling policy to your serverless endpoint with Provisioned Concurrency.

**Apply a target-tracking scaling policy (AWS CLI)**

To apply a scaling policy to your model, use the `put-scaling-policy` AWS CLI command with the following parameters:

- `--policy-name` – The name of the scaling policy.
- `--policy-type` – Set this value to `TargetTrackingScaling`.
- `--resource-id` – The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- `--service-namespace` – Set this value to `sagemaker`.
- `--scalable-dimension` – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.
- `--target-tracking-scaling-policy-configuration` – The target-tracking scaling policy configuration to use for the model.

The following example shows how to apply a target-tracking scaling policy named `MyScalingPolicy` to a model named `MyVariant`. The policy configuration is saved in a file named `scaling-policy.json`.

```
aws application-autoscaling put-scaling-policy \
  --policy-name MyScalingPolicy \
  --policy-type TargetTrackingScaling \
  --service-namespace sagemaker \
  --scalable-dimension sagemaker:variant:DesiredProvisionedConcurrency \
  --resource-id endpoint/MyEndpoint/variant/MyVariant \
  --target-tracking-scaling-policy-configuration file://[file-location]/scaling-policy.json
```
Apply a target-tracking scaling policy (Application Auto Scaling API)

To apply a scaling policy to your model, use the PutScalingPolicy Application Auto Scaling API action with the following parameters:

- **PolicyName** – The name of the scaling policy.
- **PolicyType** – Set this value to TargetTrackingScaling.
- **ResourceId** – The resource identifier for the variant. For this parameter, the resource type is endpoint and the unique identifier is the name of the variant. For example endpoint/MyEndpoint/variant/MyVariant.
- **ServiceNamespace** – Set this value to sagemaker.
- **ScalableDimension** – Set this value to sagemaker:variant:DesiredProvisionedConcurrency.
- **TargetTrackingScalingPolicyConfiguration** – The target-tracking scaling policy configuration to use for the model.

The following example shows how to apply a target-tracking scaling policy named MyScalingPolicy to a model named MyVariant. The policy configuration is saved in a file named scaling-policy.json.

```json
{
    "PolicyName": "MyScalingPolicy",
    "ServiceNamespace": "sagemaker",
    "ResourceId": "endpoint/MyEndpoint/variant/MyVariant",
    "ScalableDimension": "sagemaker:variant:DesiredProvisionedConcurrency",
    "PolicyType": "TargetTrackingScaling",
    "TargetTrackingScalingPolicyConfiguration": {
        "TargetValue": 0.5,
        "PredefinedMetricSpecification":
    }
}
```

Automatically scale Provisioned Concurrency for a serverless endpoint
Apply a target-tracking scaling policy (AWS Management Console)

To apply a target-tracking scaling policy with the AWS Management Console:

1. Sign in to the [Amazon SageMaker console](https://sagemaker.console.aws.amazon.com/sagemaker/)
2. In the navigation panel, choose **Inference**.
3. Choose **Endpoints** to view a list of all of your endpoints.
4. Choose the endpoint to which you want to apply the scaling policy. A page with the settings of the endpoint will appear, with the models (production variant) listed under **Endpoint runtime settings section**.
5. Select the production variant to which you want to apply the scaling policy, and choose **Configure auto scaling**. The **Configure variant automatic scaling** dialog box appears.
Automatically scale Provisioned Concurrency for a serverless endpoint

### Configure variant automatic scaling

**Variant automatic scaling** [Learn more](#)

<table>
<thead>
<tr>
<th>Variant name</th>
<th>Current max concurrency</th>
<th>Current provisioned concurrency</th>
</tr>
</thead>
<tbody>
<tr>
<td>variant-name-1</td>
<td>20</td>
<td>11</td>
</tr>
</tbody>
</table>

**Minimum provisioned concurrency**
**Maximum provisioned concurrency**

**IAM role**
Amazon SageMaker uses the following service-linked role for automatic scaling. [Learn more](#)

AWS_ServiceRoleForApplicationAutoScaling_SageMakerEndpoint

### Built-in scaling policy

**Built-in scaling policy** [Learn more](#)

**Policy name**
SageMakerServerlessEndpoint ProvisionedConcurrency Scaling Policy

**Target metric**
SageMakerVariant ProvisionedConcurrentUtilization

**Target value**

**Scale in cool down (seconds) - optional**
300

**Scale out cool down (seconds) - optional**
300

**Disable scale in**
Select if you don't want automatic scaling to delete instances when traffic decreases. [Learn more](#)

### Custom scaling policy

**Custom scaling policy** [Learn more](#)

There are no custom scaling policies for this variant.
6. Enter the minimum and maximum Provisioned Concurrency values in the **Minimum provisioned concurrency** and **Maximum provisioned concurrency** fields, respectively, in the **Variant automatic scaling** section. Minimum Provisioned Concurrency must be less than or equal to maximum Provisioned Concurrency.

7. Enter the target value in the **Target value** field for the target metric, **SageMakerVariantProvisionedConcurrencyUtilization**.

8. (Optional) Enter scale in cool down and scale out cool down values (in seconds) in **Scale in cool down** and **Scale out cool down** fields respectively.

9. (Optional) Select **Disable scale in** if you don’t want auto scaling to delete instance when traffic decreases.

10. Select **Save**.

**Scheduled scaling**

If the traffic to your serverless endpoint with Provisioned Concurrency follows a routine pattern you might want to schedule scaling actions at specific times, to scale in or scale out Provisioned Concurrency. You can use the AWS CLI or the Application Auto Scaling to schedule scaling actions.

**Scheduled scaling (AWS CLI)**

To apply a scaling policy to your model, use the `put-scheduled-action` AWS CLI; command with the following parameters:

- `--schedule-action-name` – The name of the scaling action.
- `--schedule` – A cron expression that specifies the start and end times of the scaling action with a recurring schedule.
- `--resource-id` – The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- `--service-namespace` – Set this value to `sagemaker`.
- `--scalable-dimension` – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.
- `--scalable-target-action` – The target of the scaling action.

The following example shows how to add a scaling action named `MyScalingAction` to a model named `MyVariant` on a recurring schedule. On the specified schedule (every day at 12:15 PM
UTC), if the current Provisioned Concurrency is below the value specified for MinCapacity. Application Auto Scaling scales out the Provisioned Concurrency to the value specified by MinCapacity.

```bash
aws application-autoscaling put-scheduled-action
  --scheduled-action-name 'MyScalingAction' 
  --schedule 'cron(15 12 * * ? *)' 
  --service-namespace sagemaker 
  --resource-id endpoint/MyEndpoint/variant/MyVariant 
  --scalable-dimension sagemaker:variant:DesiredProvisionedConcurrency 
  --scalable-target-action 'MinCapacity=10'
```

**Scheduled scaling (Application Auto Scaling API)**

To apply a scaling policy to your model, use the `PutScheduledAction` Application Auto Scaling API action with the following parameters:

- **ScheduleActionName** – The name of the scaling action.
- **Schedule** – A cron expression that specifies the start and end times of the scaling action with a recurring schedule.
- **ResourceId** – The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- **ServiceNamespace** – Set this value to `sagemaker`.
- **ScalableDimension** – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.
- **ScalableTargetAction** – The target of the scaling action.

The following example shows how to add a scaling action named `MyScalingAction` to a model named `MyVariant` on a recurring schedule. On the specified schedule (every day at 12:15 PM UTC), if the current Provisioned Concurrency is below the value specified for MinCapacity. Application Auto Scaling scales out the Provisioned Concurrency to the value specified by MinCapacity.

```plaintext
POST / HTTP/1.1
Host: autoscaling.us-east-2.amazonaws.com
```
Delete a scaling policy

You can delete a scaling policy with the AWS Management Console, the AWS CLI, or the Application Auto Scaling API. For more information on deleting a scaling policy with the AWS Management Console, see Delete a scaling policy in the SageMaker autoscaling documentation.

Delete a scaling policy (AWS CLI)

To apply a scaling policy to your model, use the delete-scaling-policy AWS CLI; command with the following parameters:

- --policy-name – The name of the scaling policy.
- --resource-id – The resource identifier for the variant. For this parameter, the resource type is endpoint and the unique identifier is the name of the variant. For example endpoint/MyEndpoint/variant/MyVariant.
- --service-namespace – Set this value to sagemaker.
- --scalable-dimension – Set this value to sagemaker:variant:DesiredProvisionedConcurrency.

The following example deletes scaling policy named MyScalingPolicy from a model named MyVariant.
aws application-autoscaling delete-scaling-policy \
  --policy-name MyScalingPolicy \
  --service-namespace sagemaker \
  --scalable-dimension sagemaker:variant:DesiredProvisionedConcurrency \
  --resource-id endpoint/MyEndpoint/variant/MyVariant

Delete a scaling policy (Application Auto Scaling API)

To delete a scaling policy to your model, use the DeleteScalingPolicy Application Auto Scaling API action with the following parameters:

- **PolicyName** – The name of the scaling policy.
- **ResourceId** – The resource identifier for the variant. For this parameter, the resource type is *endpoint* and the unique identifier is the name of the variant. For example *endpoint/MyEndpoint/variant/MyVariant*.
- **ServiceNamespace** – Set this value to *sagemaker*.
- **ScalableDimension** – Set this value to *sagemaker:variant:DesiredProvisionedConcurrency*.

The following example uses the Application Auto Scaling API to delete a scaling policy named *MyScalingPolicy* from a model named *MyVariant*.

```
POST / HTTP/1.1
Host: autoscaling.us-east-2.amazonaws.com
Accept-Encoding: identity
X-Amz-Target: AnyScaleFrontendService.DeleteScalingPolicy
X-Amz-Date: 20160506T182145Z
User-Agent: aws-cli/1.10.23 Python/2.7.11 Darwin/15.4.0 botocore/1.4.8
Content-Type: application/x-amz-json-1.1
Authorization: AUTHPARAMS

{
  "PolicyName": "MyScalingPolicy",
  "ServiceNamespace": "sagemaker",
  "ResourceId": "endpoint/MyEndpoint/variant/MyVariant",
  "ScalableDimension": "sagemaker:variant:DesiredProvisionedConcurrency",
}
```
Deregister a model

You can deregister a model with the AWS Management Console, the AWS CLI, or the Application Auto Scaling API.

Deregister a model (AWS CLI)

To deregister a model from Application Auto Scaling, use the `deregister-scalable-target` AWS CLI command with the following parameters:

- `--resource-id` – The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- `--service-namespace` – Set this value to `sagemaker`.
- `--scalable-dimension` – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.

The following example deregisters a model named `MyVariant` from Application Auto Scaling.

```bash
aws application-autoscaling deregister-scalable-target \
   --service-namespace sagemaker \
   --scalable-dimension sagemaker:variant:DesiredProvisionedConcurrency \
   --resource-id endpoint/MyEndpoint/variant/MyVariant
```

Deregister a model (Application Auto Scaling API)

To deregister a model from Application Auto Scaling use the `DeregisterScalableTarget` Application Auto Scaling API action with the following parameters:

- `ResourceId` – The resource identifier for the variant. For this parameter, the resource type is `endpoint` and the unique identifier is the name of the variant. For example `endpoint/MyEndpoint/variant/MyVariant`.
- `ServiceNamespace` – Set this value to `sagemaker`.
- `ScalableDimension` – Set this value to `sagemaker:variant:DesiredProvisionedConcurrency`.
The following example uses the Application Auto Scaling API to deregister a model named MyVariant from Application Auto Scaling.

```plaintext
POST / HTTP/1.1
Host: autoscaling.us-east-2.amazonaws.com
Accept-Encoding: identity
X-Amz-Target: AnyScaleFrontendService.DeregisterScalableTarget
X-Amz-Date: 20160506T182145Z
User-Agent: aws-cli/1.10.23 Python/2.7.11 Darwin/15.4.0 botocore/1.4.8
Content-Type: application/x-amz-json-1.1
Authorization: AUTHPARAMS

{
  "ServiceNamespace": "sagemaker",
  "ResourceId": "endpoint/MyEndpoint/variant/MyVariant",
  "ScalableDimension": "sagemaker:variant:DesiredProvisionedConcurrency",
}
```

**Deregister a model (AWS Management Console)**

To deregister a model (production variant) with the AWS Management Console:

1. Open the [Amazon SageMaker console](https://console.aws.amazon.com/sagemaker/).
2. In the navigational panel, choose **Inference**.
3. Choose **Endpoints** to view a list of your endpoints.
4. Choose the serverless endpoint hosting the production variant. A page with the settings of the endpoint will appear, with the production variants listed under **Endpoint runtime settings** section.
5. Select the production variant that you want to deregister, and choose **Configure auto scaling**. The **Configure variant automatic scaling** dialog box appears.
6. Choose **Deregister auto scaling**.

**Troubleshooting**

If you are having trouble with Serverless Inference, refer to the following troubleshooting tips.
Container issues

If the container you use for a serverless endpoint is the same one you used on an instance-based endpoint, your container may not have permissions to write files. This can happen for the following reasons:

- Your serverless endpoint fails to create or update due to a ping health check failure.
- The Amazon CloudWatch logs for the endpoint show that the container is failing to write to some file or directory due to a permissions error.

To fix this issue, you can try to add read, write, and execute permissions for other on the file or directory and then rebuild the container. You can perform the following steps to complete this process:

1. In the Dockerfile you used to build your container, add the following command: `RUN chmod o+rwx <file or directory name>`
2. Rebuild the container.
3. Upload the new container image to Amazon ECR.
4. Try to create or update the serverless endpoint again.

Asynchronous inference

Amazon SageMaker Asynchronous Inference is a capability in SageMaker that queues incoming requests and processes them asynchronously. This option is ideal for requests with large payload sizes (up to 1GB), long processing times (up to one hour), and near real-time latency requirements. Asynchronous Inference enables you to save on costs by autoscaling the instance count to zero when there are no requests to process, so you only pay when your endpoint is processing requests.

How It Works

Creating an asynchronous inference endpoint is similar to creating real-time inference endpoints. You can use your existing SageMaker models and only need to specify the AsyncInferenceConfig object while creating your endpoint configuration with the EndpointConfig field in the CreateEndpointConfig API. The following diagram shows the architecture and workflow of Asynchronous Inference.
To invoke the endpoint, you need to place the request payload in Amazon S3 and provide a pointer to this payload as a part of the `InvokeEndpointAsync` request. Upon invocation, SageMaker queues the request for processing and returns an identifier and output location as a response. Upon processing, SageMaker places the result in the Amazon S3 location. You can optionally choose to receive success or error notifications with Amazon SNS. For more information about how to set up asynchronous notifications, see [Check prediction results](#).

**Note**

The presence of an asynchronous inference configuration (`AsyncInferenceConfig`) object in the endpoint configuration implies that the endpoint can only receive asynchronous invocations.

**How Do I Get Started?**

If you are a first-time user of Amazon SageMaker Asynchronous Inference, we recommend that you do the following:

- Read [Create, invoke, and update an Asynchronous Endpoint](#) for information on how to create, invoke, update, and delete an asynchronous endpoint.
Explore the Asynchronous Inference example notebook in the aws/amazon-sagemaker-examples GitHub repository.

Note that if your endpoint uses any of the features listed in this Exclusions page, you cannot use Asynchronous Inference.

Create, invoke, and update an Asynchronous Endpoint

This guide demonstrates the prerequisites you must satisfy to create an asynchronous endpoint, along with how to create, invoke, and delete your asynchronous endpoints. You can create, update, delete, and invoke asynchronous endpoints with the AWS SDKs and the Amazon SageMaker Python SDK.

Topics

- Prerequisites
- Create an Asynchronous Inference Endpoint
- Invoke an Asynchronous Endpoint
- Update an Asynchronous Endpoint
- Delete an Asynchronous Endpoint

Prerequisites

To use asynchronous endpoints, first make sure you have met these prerequisites.

1. Create an IAM role for Amazon SageMaker.

   Asynchronous Inference needs access to your Amazon S3 bucket URI. To facilitate this, create an IAM role that can run SageMaker and has permission to access Amazon S3 and Amazon SNS. Using this role, SageMaker can run under your account and access your Amazon S3 bucket and Amazon SNS topics.

   You can create an IAM role by using the IAM console, AWS SDK for Python (Boto3), or AWS CLI. The following is an example of how to create an IAM role and attach the necessary policies with the IAM console.

   a. Sign in to the AWS Management Console and open the IAM console at https://console.aws.amazon.com/iam/.
b. In the navigation pane of the IAM console, choose Roles, and then choose Create role.

c. For Select type of trusted entity, choose AWS service.

d. Choose the service that you want to allow to assume this role. In this case, choose SageMaker. Then choose Next: Permissions.

   • This automatically creates an IAM policy that grants access to related services such as Amazon S3, Amazon ECR, and CloudWatch Logs.

e. Choose Next: Tags.

f. (Optional) Add metadata to the role by attaching tags as key–value pairs. For more information about using tags in IAM, see Tagging IAM resources.

g. Choose Next: Review.

h. Type in a Role name.

i. If possible, type a role name or role name suffix. Role names must be unique within your AWS account. They are not distinguished by case. For example, you cannot create roles named both PRODROLE and prodrole. Because other AWS resources might reference the role, you cannot edit the name of the role after it has been created.

j. (Optional) For Role description, type a description for the new role.

k. Review the role and then choose Create role.

   Note the SageMaker role ARN. To find the role ARN using the console, do the following:

   i. Go to the IAM console: https://console.aws.amazon.com/iam/

   ii. Select Roles.

   iii. Search for the role you just created by typing in the name of the role in the search field.

   iv. Select the role.

   v. The role ARN is at the top of the Summary page.

2. Add Amazon SageMaker, Amazon S3 and Amazon SNS Permissions to your IAM Role.

   Once the role is created, grant SageMaker, Amazon S3, and optionally Amazon SNS permissions to your IAM role.

   Choose Roles in the IAM console. Search for the role you created by typing in your role name in the Search field.
a. Choose your role.

b. Next, choose **Attach Policies**.

c. Amazon SageMaker Asynchronous Inference needs permission to perform the following actions: "sagemaker:CreateModel", "sagemaker:CreateEndpointConfig", "sagemaker:CreateEndpoint", and "sagemaker:InvokeEndpointAsync".

These actions are included in the AmazonSageMakerFullAccess policy. Add this policy to your IAM role. Search for AmazonSageMakerFullAccess in the **Search** field. Select AmazonSageMakerFullAccess.

d. Choose **Attach policy**.

e. Next, choose **Attach Policies** to add Amazon S3 permissions.

f. Select **Create policy**.

g. Select the **JSON** tab.

h. Add the following policy statement:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Action": [
            "s3:GetObject",
            "s3:PutObject",
            "s3:AbortMultipartUpload",
            "s3:ListBucket"
         ],
         "Effect": "Allow",
         "Resource": "arn:aws:s3:::bucket_name/*"
      }
   ]
}
```

i. Choose **Next: Tags**.

j. Type in a **Policy name**.

k. Choose **Create policy**.

l. Repeat the same steps you completed to add Amazon S3 permissions in order to add Amazon SNS permissions. For the policy statement, attach the following:
3. **Upload your inference data (e.g., machine learning model, sample data) to Amazon S3.**

4. **Select a prebuilt Docker inference image or create your own Inference Docker Image.**

   SageMaker provides containers for its built-in algorithms and prebuilt Docker images for some of the most common machine learning frameworks, such as Apache MXNet, TensorFlow, PyTorch, and Chainer. For a full list of the available SageMaker images, see [Available Deep Learning Containers Images](#). If you choose to use a SageMaker provided container, you can increase the endpoint timeout and payload sizes from the default by setting the environment variables in the container. To learn how to set the different environment variables for each framework, see the Create a Model step of creating an asynchronous endpoint.

   If none of the existing SageMaker containers meet your needs and you don't have an existing container of your own, you may need to create a new Docker container. See [Use your own inference code](#) for information on how to create your Docker image.

5. **Create an Amazon SNS topic (optional)**

   Create an Amazon Simple Notification Service (Amazon SNS) topic that sends notifications about requests that have completed processing. Amazon SNS is a notification service for messaging-oriented applications, with multiple subscribers requesting and receiving "push" notifications of time-critical messages via a choice of transport protocols, including HTTP, Amazon SQS, and email. You can specify Amazon SNS topics when you create an EndpointConfig object when you specify AsyncInferenceConfig using the EndpointConfig API.

   Follow the steps to create and subscribe to an Amazon SNS topic.
a. Using Amazon SNS console, create a topic. For instructions, see [Creating an Amazon SNS topic](#) in the Amazon Simple Notification Service Developer Guide.

b. Subscribe to the topic. For instructions, see [Subscribing to an Amazon SNS topic](#) in the Amazon Simple Notification Service Developer Guide.

c. When you receive email requesting that you confirm your subscription to the topic, confirm the subscription.

d. Note the topic Amazon Resource Name (ARN). The Amazon SNS topic you created is another resource in your AWS account, and it has a unique ARN. The ARN is in the following format:

   ```
   arn:aws:sns:aws-region:account-id:topic-name
   ```

   For more information about Amazon SNS, see the [Amazon SNS Developer Guide](#).

**Create an Asynchronous Inference Endpoint**

Create an asynchronous endpoint the same way you would create an endpoint using SageMaker hosting services:

- Create a model in SageMaker with `CreateModel`.
- Create an endpoint configuration with `CreateEndpointConfig`.
- Create an HTTPS endpoint with `CreateEndpoint`.

To create an endpoint, you first create a model with `CreateModel`, where you point to the model artifact and a Docker registry path (Image). You then create a configuration using `CreateEndpointConfig` where you specify one or more models that were created using the CreateModel API to deploy and the resources that you want SageMaker to provision. Create your endpoint with `CreateEndpoint` using the endpoint configuration specified in the request. You can update an asynchronous endpoint with the `UpdateEndpoint` API. Send and receive inference requests from the model hosted at the endpoint with `InvokeEndpointAsync`. You can delete your endpoints with the `DeleteEndpoint` API.

For a full list of the available SageMaker Images, see [Available Deep Learning Containers Images](#). See [Use your own inference code](#) for information on how to create your Docker image.
Create a Model

The following example shows how to create a model using the AWS SDK for Python (Boto3). The first few lines define:

- **sagemaker_client**: A low-level SageMaker client object that makes it easy to send and receive requests to AWS services.
- **sagemaker_role**: A string variable with the SageMaker IAM role Amazon Resource Name (ARN).
- **aws_region**: A string variable with the name of your AWS region.

```python
import boto3

# Specify your AWS Region
aws_region = '<aws_region>'

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# Role to give SageMaker permission to access AWS services.
sagemaker_role = 'arn:aws:iam::<account>:role/*'
```

Next, specify the location of the pre-trained model stored in Amazon S3. In this example, we use a pre-trained XGBoost model named demo-xgboost-model.tar.gz. The full Amazon S3 URI is stored in a string variable `model_url`:

```python
# Create a variable w/ the model S3 URI
s3_bucket = '<your-bucket-name>'  # Provide the name of your S3 bucket
bucket_prefix = 'saved_models'
model_s3_key = f'{bucket_prefix}/demo-xgboost-model.tar.gz'

# Specify S3 bucket w/ model
model_url = f's3://{s3_bucket}/{model_s3_key}'
```

Specify a primary container. For the primary container, you specify the Docker image that contains inference code, artifacts (from prior training), and a custom environment map that the inference code uses when you deploy the model for predictions.

In this example, we specify an XGBoost built-in algorithm container image:

```python
from sagemaker import image_uris
```
Create a model in Amazon SageMaker with `create_model`. Specify the following:

- **ModelName**: A name for your model (in this example it is stored as a string variable called `model_name`).
- **ExecutionRoleArn**: The Amazon Resource Name (ARN) of the IAM role that Amazon SageMaker can assume to access model artifacts and Docker images for deployment on ML compute instances or for batch transform jobs.
- **PrimaryContainer**: The location of the primary Docker image containing inference code, associated artifacts, and custom environment maps that the inference code uses when the model is deployed for predictions.

```python
model_name = '<The_name_of_the_model>'

# Create model
create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    PrimaryContainer = {
        'Image': container,
        'ModelDataUrl': model_url,
    }
)
```

See [CreateModel](#) description in the SageMaker API Reference Guide for a full list of API parameters.

If you're using a SageMaker provided container, you can increase the model server timeout and payload sizes from the default values to the framework-supported maximums by setting environment variables in this step. You might not be able to leverage the maximum timeout and payload sizes that Asynchronous Inference supports if you don't explicitly set these variables. The following example shows how you can set the environment variables for a PyTorch Inference container based on TorchServe.

```python
model_name = '<The_name_of_the_model>'
```
# Create model
create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    PrimaryContainer = {
        'Image': container,
        'ModelDataUrl': model_url,
        'Environment': {
            'TS_MAX_REQUEST_SIZE': '100000000',
            'TS_MAX_RESPONSE_SIZE': '100000000',
            'TS_DEFAULT_RESPONSE_TIMEOUT': '1000'
        }
    }
)

After you finish creating your endpoint, you should test that you've set the environment variables correctly by printing them out from your `inference.py` script. The following table lists the environment variables for several frameworks that you can set to change the default values.

<table>
<thead>
<tr>
<th>Framework</th>
<th>Environment variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>PyTorch 1.8 (based on TorchServe)</td>
<td>'TS_MAX_REQUEST_SIZE': '100000000'</td>
</tr>
<tr>
<td></td>
<td>'TS_MAX_RESPONSE_SIZE': '100000000'</td>
</tr>
<tr>
<td></td>
<td>'TS_DEFAULT_RESPONSE_TIMEOUT': '1000'</td>
</tr>
<tr>
<td>PyTorch 1.4 (based on MMS)</td>
<td>'MMS_MAX_REQUEST_SIZE': '10000000000'</td>
</tr>
<tr>
<td></td>
<td>'MMS_MAX_RESPONSE_SIZE': '10000000000'</td>
</tr>
<tr>
<td></td>
<td>'MMS_DEFAULT_RESPONSE_TIMEOUT': '900'</td>
</tr>
<tr>
<td>HuggingFace Inference Container (based on MMS)</td>
<td>'MMS_MAX_REQUEST_SIZE': '20000000000'</td>
</tr>
<tr>
<td></td>
<td>'MMS_MAX_RESPONSE_SIZE': '20000000000'</td>
</tr>
<tr>
<td></td>
<td>'MMS_DEFAULT_RESPONSE_TIMEOUT': '900'</td>
</tr>
</tbody>
</table>
Create an Endpoint Configuration

Once you have a model, create an endpoint configuration with `CreateEndpointConfig`. Amazon SageMaker hosting services uses this configuration to deploy models. In the configuration, you identify one or more models, created using with `CreateModel`, to deploy the resources that you want Amazon SageMaker to provision. Specify the AsyncInferenceConfig object and provide an output Amazon S3 location for `OutputConfig`. You can optionally specify Amazon SNS topics on which to send notifications about prediction results. For more information about Amazon SNS topics, see Configuring Amazon SNS.

The following example shows how to create an endpoint configuration using AWS SDK for Python (Boto3):

```python
import datetime
from time import gmtime, strftime

# Create an endpoint config name. Here we create one based on the date
# so it we can search endpoints based on creation time.
endpoint_config_name = f"XGBoostEndpointConfig-{strftime('%Y-%m-%d-%H-%M-%S',
    gmtime())}"

# The name of the model that you want to host. This is the name that you specified when
# creating the model.
model_name='<The_name_of_your_model>'

create_endpoint_config_response = sagemaker_client.create_endpoint_config(
    EndpointConfigName=endpoint_config_name, # You will specify this name in a
    CreateEndpoint request.
    # List of ProductionVariant objects, one for each model that you want to host at
    # this endpoint.
    ProductionVariants=[
        {
            "VariantName": "variant1", # The name of the production variant.
            "ModelName": model_name,
            "InstanceType": "ml.m5.xlarge", # Specify the compute instance type.
            "InitialInstanceCount": 1 # Number of instances to launch initially.
        }
    ],
    AsyncInferenceConfig={
        "OutputConfig": {
            # Location to upload response outputs when no location is provided in the
            request.
        }
    }
)
```
"S3OutputPath": f"s3://{s3_bucket}/{bucket_prefix}/output"
# (Optional) specify Amazon SNS topics
"NotificationConfig": {
}
},
"ClientConfig": {
   # (Optional) Specify the max number of inflight invocations per instance
   # If no value is provided, Amazon SageMaker will choose an optimal value
   for you
   "MaxConcurrentInvocationsPerInstance": 4
}
)

print(f"Created EndpointConfig: {create_endpoint_config_response['EndpointConfigArn']}")

In the aforementioned example, you specify the following keys for OutputConfig for the AsyncInferenceConfig field:

- **S3OutputPath**: Location to upload response outputs when no location is provided in the request.
- **NotificationConfig**: (Optional) SNS topics that post notifications to you when an inference request is successful (SuccessTopic) or if it fails (ErrorTopic).

You can also specify the following optional argument for ClientConfig in the AsyncInferenceConfig field:

- **MaxConcurrentInvocationsPerInstance**: (Optional) The maximum number of concurrent requests sent by the SageMaker client to the model container.

**Create Endpoint**

Once you have your model and endpoint configuration, use the CreateEndpoint API to create your endpoint. The endpoint name must be unique within an AWS Region in your AWS account.

The following creates an endpoint using the endpoint configuration specified in the request. Amazon SageMaker uses the endpoint to provision resources and deploy models.
# The name of the endpoint. The name must be unique within an AWS Region in your AWS account.
endpoint_name = 'endpoint-name'

# The name of the endpoint configuration associated with this endpoint.
endpoint_config_name='endpoint-config-name'

create_endpoint_response = sagemaker_client.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name)

When you call the CreateEndpoint API, Amazon SageMaker Asynchronous Inference sends a test notification to check that you have configured an Amazon SNS topic. Amazon SageMaker Asynchronous Inference also sends test notifications after calls to UpdateEndpoint and UpdateEndpointWeightsAndCapacities. This lets SageMaker check that you have the required permissions. The notification can simply be ignored. The test notification has the following form:

```json
{
    "eventVersion":"1.0",
    "eventSource":"aws:sagemaker",
    "eventName":"TestNotification"
}
```

**Invoke an Asynchronous Endpoint**

Get inferences from the model hosted at your asynchronous endpoint with InvokeEndpointAsync.

**Note**

If you have not done so already, upload your inference data (e.g., machine learning model, sample data) to Amazon S3.

Specify the following fields in your request:

- For InputLocation, specify the location of your inference data.
- For EndpointName, specify the name of your endpoint.
• (Optional) For InvocationTimeoutSeconds, you can set the max timeout for the requests. You can set this value to a maximum of 3600 seconds (one hour) on a per-request basis. If you don't specify this field in your request, by default the request times out at 15 minutes.

```python
# Create a low-level client representing Amazon SageMaker Runtime
sagemaker_runtime = boto3.client("sagemaker-runtime", region_name=<aws_region>)

# Specify the location of the input. Here, a single SVM sample
input_location = "s3://bucket-name/test_point_0.libsvm"

# The name of the endpoint. The name must be unique within an AWS Region in your AWS account.
endpoint_name='<endpoint-name>'

# After you deploy a model into production using SageMaker hosting services, your client applications use this API to get inferences from the model hosted at the specified endpoint.
response = sagemaker_runtime.invoke_endpoint_async(
    EndpointName=endpoint_name,
    InputLocation=input_location,
    InvocationTimeoutSeconds=3600)
```

You receive a response as a JSON string with your request ID and the name of the Amazon S3 bucket that will have the response to the API call after it is processed.

**Update an Asynchronous Endpoint**

Update an asynchronous endpoint with the UpdateEndpoint API. When you update an endpoint, SageMaker first provisions and switches to the new endpoint configuration you specify before it deletes the resources that were provisioned in the previous endpoint configuration. Do not delete an EndpointConfig with an endpoint that is live or while the UpdateEndpoint or CreateEndpoint operations are being performed on the endpoint.

```python
# The name of the endpoint. The name must be unique within an AWS Region in your AWS account.
endpoint_name='<endpoint-name>'

# The name of the endpoint configuration associated with this endpoint.
endpoint_config_name='<endpoint-config-name>'

sagemaker_client.update_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name,
    # Additional parameters as needed
)`
When Amazon SageMaker receives the request, it sets the endpoint status to **Updating**. After updating the asynchronous endpoint, it sets the status to **InService**. To check the status of an endpoint, use the **DescribeEndpoint** API. For a full list of parameters you can specify when updating an endpoint, see the **UpdateEndpoint** API.

### Delete an Asynchronous Endpoint

Delete an asynchronous endpoint in a similar manner to how you would delete a SageMaker hosted endpoint with the **DeleteEndpoint** API. Specify the name of the asynchronous endpoint you want to delete. When you delete an endpoint, SageMaker frees up all of the resources that were deployed when the endpoint was created. Deleting a model does not delete model artifacts, inference code, or the IAM role that you specified when creating the model.

Delete your SageMaker model with the **DeleteModel** API or with the SageMaker console.

#### Boto3

```python
import boto3

# Create a low-level SageMaker service client.
sagemaker_client = boto3.client('sagemaker', region_name=<aws_region>)
sagemaker_client.delete_endpoint(EndpointName='<endpoint-name>')
```

#### SageMaker console

2. Expand the **Inference** dropdown list.
3. Select **Endpoints**.
4. Search for endpoint in the **Search endpoints** search bar.
5. Select your endpoint.
6. Choose **Delete**.

In addition to deleting the asynchronous endpoint, you might want to clear up other resources that were used to create the endpoint, such as the Amazon ECR repository (if you created a custom inference image), the SageMaker model, and the asynchronous endpoint configuration itself.
Monitor asynchronous endpoint

You can monitor SageMaker using Amazon CloudWatch, which collects raw data and processes it into readable, near real-time metrics. With Amazon CloudWatch, you can access historical information and gain a better perspective on how your web application or service is performing. For more information about Amazon CloudWatch, see What is Amazon CloudWatch?

Monitoring with CloudWatch

The metrics below are an exhaustive list of metrics for asynchronous endpoints and are in the the AWS/SageMaker namespace. Any metric not listed below is not published if the endpoint is enabled for asynchronous inference. Such metrics include (but are not limited to):

- OverheadLatency
- Invocations
- InvocationsPerInstance

Common Endpoint Metrics

These metrics are the same as the metrics published for real-time endpoints today. For more information about other metrics in Amazon CloudWatch, see Monitor SageMaker with Amazon CloudWatch.

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Unit/Stats</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invocation4XXErrors</td>
<td>The number of requests where the model returned a 4xx HTTP response code. For each 4xx response, 1 is sent; otherwise, 0 is sent.</td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
<td></td>
</tr>
<tr>
<td>Invocation5XXErrors</td>
<td>The number of InvokeEndpoint requests where the model returned a 5xx HTTP response code. For each 5xx response, 1 is sent; otherwise, 0 is sent.</td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
<td></td>
</tr>
</tbody>
</table>
## Metric Name

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Unit/Stats</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelLatency</td>
<td>The interval of time taken by a model to respond as viewed from SageMaker. This interval includes the local communication times taken to send the request and to fetch the response from the container of a model and the time taken to complete the inference in the container.</td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
<td></td>
</tr>
</tbody>
</table>

### Asynchronous Inference Endpoint Metrics

These metrics are published for endpoints enabled for asynchronous inference. The following metrics are published with the EndpointName dimension:

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Unit/Stats</th>
</tr>
</thead>
<tbody>
<tr>
<td>ApproximateBacklog Size</td>
<td>The number of items in the queue for an endpoint that are currently being processed or yet to be processed.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
<td></td>
</tr>
<tr>
<td>ApproximateBacklog SizePerInstance</td>
<td>Number of items in the queue divided by the number of instances behind an endpoint. This metric is primarily used for setting up application autoscaling for an async-enabled endpoint.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
<td></td>
</tr>
<tr>
<td>ApproximateAgeOfOldestRequest</td>
<td>Age of the oldest request in the queue.</td>
<td>Units: Seconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Max, Min</td>
<td></td>
</tr>
</tbody>
</table>
### Metric Name

- **HasBacklogWithoutCapacity**

  The value of this metric is 1 when there are requests in the queue but zero instances behind the endpoint. The value is 0 at all other times. You can use this metric for autoscaling your endpoint up from zero instances upon receiving a new request in the queue.

  **Unit/Stats**
  - Units: Count
  - Valid statistics: Average

### The following metrics are published with the EndpointName and VariantName dimensions:

<table>
<thead>
<tr>
<th>Metric Name</th>
<th>Description</th>
<th>Unit/Stats</th>
</tr>
</thead>
<tbody>
<tr>
<td>RequestDownloadFailures</td>
<td>When an inference failure occurs due to an issue downloading the request from Amazon S3.</td>
<td>Units: Count, Valid statistics: Sum</td>
</tr>
<tr>
<td>ResponseUploadFailures</td>
<td>When an inference failure occurs due to an issue uploading the response to Amazon S3.</td>
<td>Units: Count, Valid statistics: Sum</td>
</tr>
<tr>
<td>NotificationFailures</td>
<td>When an issue occurs publishing notifications.</td>
<td>Units: Count, Valid statistics: Sum</td>
</tr>
<tr>
<td>RequestDownloadLatency</td>
<td>Total time to download the request payload.</td>
<td>Units: Microseconds, Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ResponseUploadLatency</td>
<td>Total time to upload the response payload.</td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td>Metric Name</td>
<td>Description</td>
<td>Unit/Stats</td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>------------------------------------------------</td>
</tr>
<tr>
<td>ExpiredRequests</td>
<td>Number of requests in the queue that fail due to reaching their specified request TTL.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>InvocationFailures</td>
<td>If an invocation fails for any reason.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>InvocationsProcessed</td>
<td>Number of async invocations processed by the endpoint.</td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>TimeInBacklog</td>
<td>Total time the request was queued before being processed. This does not include the actual processing time (i.e. downloading time, uploading time, model latency).</td>
<td>Units: Milliseconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>TotalProcessingTime</td>
<td>Time the inference request was received by SageMaker to the time the request finished processing. This includes time in backlog and time to upload and send response notifications, if any.</td>
<td>Units: Milliseconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>

Amazon SageMaker Asynchronous Inference also includes host-level metrics. For information on host-level metrics, see [SageMaker Jobs and Endpoint Metrics](#).
Logs

In addition to the Model container logs that are published to Amazon CloudWatch in your account, you also get a new platform log for tracing and debugging inference requests.

The new logs are published under the Endpoint Log Group:

/aws/sagemaker/Endpoints/[EndpointName]

The log stream name consists of:

[production-variant-name]/[instance-id]/data-log.

Log lines contain the request's inference ID so that errors can be easily mapped to a particular request.

Check prediction results

There are several ways you can check predictions results from your asynchronous endpoint. Some options are:

1. Amazon SNS topics.
2. Check for outputs in your Amazon S3 bucket.

Amazon SNS Topics

Amazon SNS is a notification service for messaging-oriented applications, with multiple subscribers requesting and receiving "push" notifications of time-critical messages via a choice of transport protocols, including HTTP, Amazon SQS, and email. Amazon SageMaker Asynchronous Inference posts notifications when you create an endpoint with CreateEndpointConfig and specify an Amazon SNS topic.

Note

In order to receive Amazon SNS notifications, your IAM role must have sns:Publish permissions. See the Prerequisites for information on requirements you must satisfy to use Asynchronous Inference.
To use Amazon SNS to check prediction results from your asynchronous endpoint, you first need to create a topic, subscribe to the topic, confirm your subscription to the topic, and note the Amazon Resource Name (ARN) of that topic. For detailed information on how to create, subscribe, and find the Amazon ARN of an Amazon SNS topic, see Configuring Amazon SNS.

Provide the Amazon SNS topic ARN(s) in the AsyncInferenceConfig field when you create an endpoint configuration with CreateEndpointConfig. You can specify both an Amazon SNS ErrorTopic and a SuccessTopic.

```python
import boto3

sagemaker_client = boto3.client('sagemaker', region_name=<aws_region>)

sagemaker_client.create_endpoint_config(
    EndpointConfigName=<endpoint_config_name>, # You specify this name in a CreateEndpoint request.
    ProductionVariants=[
        {
            "VariantName": "variant1", # The name of the production variant.
            "ModelName": "model_name",
            "InstanceType": "ml.m5.xlarge", # Specify the compute instance type.
            "InitialInstanceCount": 1 # Number of instances to launch initially.
        }
    ],
    AsyncInferenceConfig={
        "OutputConfig": {
            "S3OutputPath": "s3://<bucket>/<output_directory>"
        },
        "NotificationConfig": {
        }
    }
)
```

After creating your endpoint and invoking it, you receive a notification from your Amazon SNS topic. For example, if you subscribed to receive email notifications from your topic, you receive an
email notification every time you invoke your endpoint. The following example shows the JSON content of a successful invocation email notification.

```json
{
    "awsRegion": "us-east-1",
    "eventTime": "2022-01-25T22:46:00.608Z",
    "receivedTime": "2022-01-25T22:46:00.455Z",
    "invocationStatus": "Completed",
    "requestParameters": {
        "contentType": "text/csv",
        "endpointName": "<example-endpoint>",
        "inputLocation": "s3://<bucket>/<input-directory>/input-data.csv"
    },
    "responseParameters": {
        "contentType": "text/csv; charset=utf-8",
        "outputLocation": "s3://<bucket>/<output_directory>/prediction.out"
    },
    "inferenceId": "11111111-2222-3333-4444-555555555555",
    "eventVersion": "1.0",
    "eventSource": "aws:sagemaker",
    "eventName": "InferenceResult"
}
```

### Check Your S3 Bucket

When you invoke an endpoint with `InvokeEndpointAsync`, it returns a response object. You can use the response object to get the Amazon S3 URI where your output is stored. With the output location, you can use a SageMaker Python SDK SageMaker session class to programmatically check for an output.

The following stores the output dictionary of `InvokeEndpointAsync` as a variable named `response`. With the response variable, you then get the Amazon S3 output URI and store it as a string variable called `output_location`.

```python
import uuid
import boto3

sagemaker_runtime = boto3.client("sagemaker-runtime", region_name=<aws_region>)

# Specify the S3 URI of the input. Here, a single SVM sample
input_location = "s3://bucket-name/test_point_0.libsvm"
```
response = sagemaker_runtime.invoke_endpoint_async(
    EndpointName='<endpoint-name>',
    InputLocation=input_location,
    InferenceId=str(uuid.uuid4()),
    ContentType="text/libsvm" #Specify the content type of your data
)

output_location = response['OutputLocation']
print(f"OutputLocation: {output_location}")

For information about supported content types, see Common Data Formats for Inference.

With the Amazon S3 output location, you can then use a SageMaker Python SDK SageMaker Session Class to read in Amazon S3 files. The following code example shows how to create a function (get_output) that repeatedly attempts to read a file from the Amazon S3 output location:

```python
import sagemaker
import urllib, time
from botocore.exceptions import ClientError

sagemaker_session = sagemaker.session.Session()

def get_output(output_location):
    output_url = urllib.parse.urlparse(output_location)
    bucket = output_url.netloc
    key = output_url.path[1:]
    while True:
        try:
            return sagemaker_session.read_s3_file(
                bucket=bucket,
                key_prefix=key)
        except ClientError as e:
            if e.response['Error']['Code'] == 'NoSuchKey':
                print("waiting for output...")
                time.sleep(2)
            else:
                continue
    raise

output = get_output(output_location)
print(f"Output: {output}")
```

Check prediction results 4308
Autoscale an asynchronous endpoint

Amazon SageMaker supports automatic scaling (autoscaling) your asynchronous endpoint. Autoscaling dynamically adjusts the number of instances provisioned for a model in response to changes in your workload. Unlike other hosted models Amazon SageMaker supports, with Asynchronous Inference you can also scale down your asynchronous endpoints instances to zero. Requests that are received when there are zero instances are queued for processing once the endpoint scales up.

To autoscale your asynchronous endpoint you must at a minimum:

- Register a deployed model (production variant).
- Define a scaling policy.
- Apply the autoscaling policy.

Before you can use autoscaling, you must have already deployed a model to a SageMaker endpoint. Deployed models are referred to as a production variant. See Deploy the Model to SageMaker Hosting Services for more information about deploying a model to an endpoint. To specify the metrics and target values for a scaling policy, you configure a scaling policy. For information on how to define a scaling policy, see Define a scaling policy. After registering your model and defining a scaling policy, apply the scaling policy to the registered model. For information on how to apply the scaling policy, see Apply a scaling policy.

For more information on how to define an optional additional scaling policy that scales up your endpoint upon receiving a request after your endpoint has been scaled down to zero, see Optional: Define a scaling policy that scales up from zero for new requests. If you don't specify this optional policy, then your endpoint only initiates scaling up from zero after the number of backlog requests exceeds the target tracking value.

For details on other prerequisites and components used with autoscaling, see the Prerequisites section in the SageMaker autoscaling documentation.

⚠️ Note

If you attach multiple scaling policies to the same autoscaling group, you might have scaling conflicts. When a conflict occurs, Amazon EC2 Auto Scaling chooses the policy that provisions the largest capacity for both scale out and scale in. For more information
Define a scaling policy

To specify the metrics and target values for a scaling policy, you configure a target-tracking scaling policy. Define the scaling policy as a JSON block in a text file. You use that text file when invoking the AWS CLI or the Application Auto Scaling API. For more information about policy configuration syntax, see `TargetTrackingScalingPolicyConfiguration` in the Application Auto Scaling API Reference.

For asynchronous endpoints SageMaker strongly recommends that you create a policy configuration for target-tracking scaling for a variant. In this configuration example, we use a custom metric, `CustomizedMetricSpecification`, called `ApproximateBacklogSizePerInstance`.

```
TargetTrackingScalingPolicyConfiguration={
    'TargetValue': 5.0, # The target value for the metric. Here the metric is: ApproximateBacklogSizePerInstance
    'CustomizedMetricSpecification': {
        'MetricName': 'ApproximateBacklogSizePerInstance',
        'Namespace': 'AWS/SageMaker',
        'Dimensions': [
            {'Name': 'EndpointName', 'Value': <endpoint_name> }
        ],
        'Statistic': 'Average',
    }
}
```

Define a scaling policy that scales to zero

The following shows you how to both define and register your endpoint variant with application autoscaling using the AWS SDK for Python (Boto3). After defining a low-level client object representing application autoscaling with Boto3, we use the `RegisterScalableTarget` method to register the production variant. We set `MinCapacity` to 0 because Asynchronous Inference enables you to autoscale to 0 when there are no requests to process.

```
# Common class representing application autoscaling for SageMaker
```
client = boto3.client('application-autoscaling')

# This is the format in which application autoscaling references the endpoint
resource_id='endpoint/' + '<endpoint_name>' + '/variant/' + '<variant1>'

# Define and register your endpoint variant
response = client.register_scalable_target(
    ServiceNamespace='sagemaker',
    ResourceId=resource_id,
    ScalableDimension='sagemaker:variant:DesiredInstanceCount', # The number of EC2
    instances for your Amazon SageMaker model endpoint variant.
    MinCapacity=0,
    MaxCapacity=5
)

For detailed description about the Application Autoscaling API, see the Application Scaling Boto3
documentation.

Optional: Define a scaling policy that scales up from zero for new requests

You might have a use case where you have sporadic requests or periods with low numbers of
requests. If your endpoint has been scaled down to zero instances during these periods, then
your endpoint won’t scale up again until the number of requests in the queue exceeds the target
specified in your scaling policy. This can result in long waiting times for requests in the queue. The
following section shows you how to create an additional scaling policy that scales your endpoint
up from zero instances after receiving any new request in the queue. Your endpoint will be able to
respond to new requests more quickly instead of waiting for the queue size to exceed the target.

To create a scaling policy for your endpoint that scales up from zero instances, do the following:

1. Create a scaling policy that defines the desired behavior, which is to scale up your endpoint
   when it’s at zero instances but has requests in the queue. The following shows you how to
define a scaling policy called HasBacklogWithoutCapacity-ScalingPolicy using the
AWS SDK for Python (Boto3). When the queue is greater than zero and the current instance
count for your endpoint is also zero, the policy scales your endpoint up. In all other cases, the
policy does not affect scaling for your endpoint.

    response = client.put_scaling_policy(
        PolicyName="HasBacklogWithoutCapacity-ScalingPolicy",
        ServiceNamespace="sagemaker", # The namespace of the service that provides the
        resource.
ResourceId=resource_id, # Endpoint name
ScalableDimension="sagemaker:variant:DesiredInstanceCount", # SageMaker supports only Instance Count
PolicyType="StepScaling", # 'StepScaling' or 'TargetTrackingScaling'
StepScalingPolicyConfiguration={
  "AdjustmentType": "ChangeInCapacity", # Specifies whether the ScalingAdjustment value in the StepAdjustment property is an absolute number or a percentage of the current capacity.
  "MetricAggregationType": "Average", # The aggregation type for the CloudWatch metrics.
  "Cooldown": 300, # The amount of time, in seconds, to wait for a previous scaling activity to take effect.
  "StepAdjustments": # A set of adjustments that enable you to scale based on the size of the alarm breach.
  [
    {
      "MetricIntervalLowerBound": 0,
      "Scaling Adjustment": 1
    }
  ],
}
)

2. Create a CloudWatch alarm with the custom metric HasBacklogWithoutCapacity. When triggered, the alarm initiates the previously defined scaling policy. For more information about the HasBacklogWithoutCapacity metric, see Asynchronous Inference Endpoint Metrics.

```python
response = cw_client.put_metric_alarm(
    AlarmName=step_scaling_policy_alarm_name,
    MetricName='HasBacklogWithoutCapacity',
    Namespace='AWS/SageMaker',
    Statistic='Average',
    EvaluationPeriods=2,
    DatapointsToAlarm=2,
    Threshold=1,
    ComparisonOperator='GreaterThanOrEqualToThreshold',
    TreatMissingData='missing',
    Dimensions=[
        { 'Name':'EndpointName', 'Value':endpoint_name },
    ],
    Period=60,
    AlarmActions=[step_scaling_policy_arn]
)
```
You should now have a scaling policy and CloudWatch alarm that scale up your endpoint from zero instances whenever your queue has pending requests.

## Troubleshooting

The following FAQs can help you troubleshoot issues with your Amazon SageMaker Asynchronous Inference endpoints.

**Q: I have autoscaling enabled. How can I find the instance count behind the endpoint at any given point?**

You can use the following methods to find the instance count behind your endpoint:

- You can use the SageMaker [DescribeEndpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/API_DescribeEndpoint.html) API to describe the number of instances behind the endpoint at any given point in time.
- You can get the instance count by viewing your Amazon CloudWatch metrics. View the **metrics for your endpoint instances**, such as CPUUtilization or MemoryUtilization and check the sample count statistic for a 1 minute period. The count should be equal to the number of active instances. The following screenshot shows the CPUUtilization metric graphed in the CloudWatch console, where the **Stat**ic is set to **Sample count**, the **Period** is set to 1 minute, and the resulting count is 5.

![CloudWatch Instance Count](image)

**Q: What are the common tunable environment variables for SageMaker containers?**

The following tables outline the common tunable environment variables for SageMaker containers by framework type.

### TensorFlow
<table>
<thead>
<tr>
<th>Environment variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAGEMAKER_TFS_INSTANCE_COUNT</td>
<td>For TensorFlow-based models, the <code>tensorflow_model_server</code> binary is the operational piece that is responsible for loading a model in memory, running inputs against a model graph, and deriving outputs. Typically, a single instance of this binary is launched to serve models in an endpoint. This binary is internally multi-threaded and spawns multiple threads to respond to an inference request. In certain instances, if you observe that the CPU is respectably utilized (over 30% utilized) but the memory is underutilized (less than 10% utilization), increasing this parameter might help. Increasing the number of <code>tensorflow_model_servers</code> available to serve typically increases the throughput of an endpoint.</td>
</tr>
<tr>
<td>SAGEMAKER_TFS_FRACTIONAL_GPU_MEM_MARGIN</td>
<td>This parameter governs the fraction of the available GPU memory to initialize CUDA/cuDNN and other GPU libraries. 0.2 means 20% of the available GPU memory is reserved for initializing CUDA/cuDNN and other GPU libraries, and 80% of the available GPU memory is allocated equally across the TF processes. GPU memory is pre-allocated unless the <code>allow_growth</code> option is enabled.</td>
</tr>
<tr>
<td>SAGEMAKER_TFS_INTER_OP_PARALLELISM</td>
<td>This ties back to the <code>inter_op_parallelism_threads</code> variable. This variable determines the number of threads used by independent non-blocking operations. 0 means that the system picks an appropriate number.</td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>SAGEMAKER_TFS_INTRA_OP_PARALLELISM</td>
<td>This ties back to the <code>intra_op_parallelism_threads</code> variable. This determines the number of threads that can be used for certain operations like matrix multiplication and reductions for speedups. A value of 0 means that the system picks an appropriate number.</td>
</tr>
<tr>
<td>SAGEMAKER_GUNICORN_WORKERS</td>
<td>This governs the number of worker processes that Gunicorn is requested to spawn for handling requests. This value is used in combination with other parameters to derive a set that maximizes inference throughput. In addition to this, the <code>SAGEMAKER_GUNICORN_WORKER_CLASS</code> governs the type of workers spawned, typically async or gevent.</td>
</tr>
<tr>
<td>SAGEMAKER_GUNICORN_WORKER_CLASS</td>
<td>This governs the number of worker processes that Gunicorn is requested to spawn for handling requests. This value is used in combination with other parameters to derive a set that maximizes inference throughput. In addition to this, the <code>SAGEMAKER_GUNICORN_WORKER_CLASS</code> governs the type of workers spawned, typically async or gevent.</td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>OMP_NUM_THREADS</td>
<td>Python internally uses OpenMP for implementing multithreading within processes. Typically, threads equivalent to the number of CPU cores are spawned. But when implemented on top of Simultaneous Multi Threading (SMT), such Intel’s HyperThreading, a certain process might oversubscribe a particular core by spawning twice as many threads as the number of actual CPU cores. In certain cases, a Python binary might end up spawning up to four times as many threads as available processor cores. Therefore, an ideal setting for this parameter, if you have oversubscribed available cores using worker threads, is 1, or half the number of CPU cores on a CPU with SMT turned on.</td>
</tr>
<tr>
<td>TF_DISABLE_MKL</td>
<td>In some cases, turning off MKL can speed up inference if TF_DISABLE_MKL and TF_DISABLE_POOL_ALLOCATOR are set to 1.</td>
</tr>
<tr>
<td>TF_DISABLE_POOL_ALLOCATOR</td>
<td></td>
</tr>
</tbody>
</table>

**PyTorch**

<table>
<thead>
<tr>
<th>Environment variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAGEMAKER_TS_MAX_BATCH_DELAY</td>
<td>This is the maximum batch delay time TorchServe waits to receive.</td>
</tr>
<tr>
<td>SAGEMAKER_TS_BATCH_SIZE</td>
<td>If TorchServe doesn’t receive the number of requests specified in batch_size before the timer runs out, it sends the requests that were received to the model handler.</td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>SAGEMAKER_TS_MIN_WORKERS</td>
<td>The minimum number of workers to which TorchServe is allowed to scale down.</td>
</tr>
<tr>
<td>SAGEMAKER_TS_MAX_WORKERS</td>
<td>The maximum number of workers to which TorchServe is allowed to scale up.</td>
</tr>
<tr>
<td>SAGEMAKER_TS_RESPONSE_TIMEOUT</td>
<td>The time delay, after which inference times out in absence of a response.</td>
</tr>
<tr>
<td>SAGEMAKER_TS_MAX_REQUEST_SIZE</td>
<td>The maximum payload size for TorchServe.</td>
</tr>
<tr>
<td>SAGEMAKER_TS_MAX_RESPONSE_SIZE</td>
<td>The maximum response size for TorchServe.</td>
</tr>
</tbody>
</table>

**Multi Model Server (MMS)**

<table>
<thead>
<tr>
<th>Environment variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>job_queue_size</td>
<td>This parameter is useful to tune when you have a scenario where the type of the inference request payload is large, and due to the size of payload being larger, you may have higher heap memory consumption of the JVM in which this queue is being maintained. Ideally you might want to keep the heap memory requirements of JVM lower and allow Python workers to allot more memory for actual model serving. JVM is only for receiving the HTTP requests, queuing them, and dispatching them to the Python-based workers for inference. If you increase the <code>job_queue_size</code>, you might end up increasing the heap memory consumption of the JVM and ultimately taking away memory from the host that could have been used by</td>
</tr>
</tbody>
</table>
### Environment variable: `default_workers_per_model`

This parameter is for the backend model serving and might be valuable to tune since this is the critical component of the overall model serving, based on which the Python processes spawn threads for each Model. If this component is slower (or not tuned properly), the front-end tuning might not be effective.

---

**Q: How do I make sure my container supports Asynchronous Inference?**

You can use the same container for Asynchronous Inference that you do for Real-Time Inference or Batch Transform. You should confirm that the timeouts and payload size limits on your container are set to handle larger payloads and longer timeouts.

**Q: What are the limits specific to Asynchronous Inference, and can they be adjusted?**

Refer to the following limits for Asynchronous Inference:

- **Payload size limit:** 1 GB
- **Timeout limit:** A request can take up to 60 minutes.
- **Queue message TimeToLive (TTL):** 6 hours
- **Number of messages that can be put inside Amazon SQS:** Unlimited. However, there is a quota of 120,000 for the number of in-flight messages for a standard queue, and 20,000 for a FIFO queue.

**Q: What metrics are best to define for autoscaling on Asynchronous Inference? Can I have multiple scaling policies?**

In general, with Asynchronous Inference, you can scale out based on invocations or instances. For invocation metrics, it's a good idea to look at your `ApproximateBacklogSize`, which is a metric that defines the number of items in your queue that have yet to be processed. You can
utilize this metric or your InvocationsPerInstance metric to understand what TPS you may be getting throttled at. At the instance level, check your instance type and its CPU/GPU utilization to define when to scale out. If a singular instance is above 60-70% capacity, this is often a good sign that you are saturating your hardware.

We don't recommend having multiple scaling policies, as these can conflict and lead to confusion at the hardware level, causing delays when scaling out.

**Q: Why is my asynchronous endpoint terminating an instance as Unhealthy and the update requests from autoscaling are failing?**

Check if your container is able to handle ping and invoke requests concurrently. SageMaker invoke requests take approximately 3 minutes, and in this duration, usually multiple ping requests end up failing due to the timeout causing SageMaker to detect your container as Unhealthy.

**Q: Can MaxConcurrentInvocationsPerInstance work for my BYOC model container with the nginx/gunicorn/flask settings?**

Yes. MaxConcurrentInvocationsPerInstance is a feature of asynchronous endpoints. This does not depend on the custom container implementation. MaxConcurrentInvocationsPerInstance controls the rate at which invoke requests are sent to the customer container. If this value is set as 1, then only 1 request is sent to the container at a time, no matter how many workers are on the customer container.

**Q: How can I debug model server errors (500) on my asynchronous endpoint?**

The error means that the customer container returned an error. SageMaker does not control the behavior of customer containers. SageMaker simply returns the response from the ModelContainer and does not retry. If you want, you can configure the invocation to retry on failure. We suggest that you turn on container logging and check your container logs to find the root cause of the 500 error from your model. Check the corresponding CPUUtilization and MemoryUtilization metrics at the point of failure as well. You can also configure the S3FailurePath to the model response in Amazon SNS as part of the Async Error Notifications to investiage failures.

**Q: How can I know if MaxConcurrentInvocationsPerInstance=1 takes effect? Are there any metrics that I can check?**

You can check the metric InvocationsProcessed, which should align with the number of invocations that you expect to be processed in a minute based on single concurrency.
Q: How can I track the success and failures of my invocation requests? What are the best practices?

The best practice is to enable Amazon SNS, which is a notification service for messaging-oriented applications, with multiple subscribers requesting and receiving "push" notifications of time-critical messages from a choice of transport protocols, including HTTP, Amazon SQS, and email. Asynchronous Inference posts notifications when you create an endpoint with `CreateEndpointConfig` and specify an Amazon SNS topic.

To use Amazon SNS to check prediction results from your asynchronous endpoint, you first need to create a topic, subscribe to the topic, confirm your subscription to the topic, and note the Amazon Resource Name (ARN) of that topic. For detailed information on how to create, subscribe, and find the Amazon ARN of an Amazon SNS topic, see Configuring Amazon SNS in the Amazon SNS Developer Guide. For more information about how to use Amazon SNS with Asynchronous Inference, see Check prediction results.

Q: Can I define a scaling policy that scales up from zero instances upon receiving a new request?

Yes. Asynchronous Inference provides a mechanism to scale down to zero instances when there are no requests. If your endpoint has been scaled down to zero instances during these periods, then your endpoint won't scale up again until the number of requests in the queue exceeds the target specified in your scaling policy. This can result in long waiting times for requests in the queue. In such cases, if you want to scale up from zero instances for new requests less than the queue target specified, you can use an additional scaling policy called `HasBacklogWithoutCapacity`. For more information about how to define this scaling policy, see Autoscale an asynchronous endpoint.

Q: I'm getting an error that the instance type is not supported for Asynchronous Inference. What are the instance types Asynchronous Inference supports?

For an exhaustive list of instances supported by Asynchronous Inference per region, see SageMaker pricing. Check if the required instance is available in your region before proceeding.

**Use Batch Transform**

Use batch transform when you need to do the following:

- Preprocess datasets to remove noise or bias that interferes with training or inference from your dataset.
• Get inferences from large datasets.
• Run inference when you don't need a persistent endpoint.
• Associate input records with inferences to assist the interpretation of results.

To filter input data before performing inferences or to associate input records with inferences about those records, see Associate Prediction Results with Input Records. For example, you can filter input data to provide context for creating and interpreting reports about the output data.

Topics
• Use Batch Transform to Get Inferences from Large Datasets
• Speed up a Batch Transform Job
• Use Batch Transform to Test Production Variants
• Batch Transform Sample Notebooks
• Associate Prediction Results with Input Records
• Storage in Batch Transform
• Troubleshooting

Use Batch Transform to Get Inferences from Large Datasets

Batch transform automatically manages the processing of large datasets within the limits of specified parameters. For example, suppose that you have a dataset file, input1.csv, stored in an S3 bucket. The content of the input file might look like the following example.

Record1-Attribute1, Record1-Attribute2, Record1-Attribute3, ..., Record1-AttributeM
Record2-Attribute1, Record2-Attribute2, Record2-Attribute3, ..., Record2-AttributeM
Record3-Attribute1, Record3-Attribute2, Record3-Attribute3, ..., Record3-AttributeM
...
RecordN-Attribute1, RecordN-Attribute2, RecordN-Attribute3, ..., RecordN-AttributeM

When a batch transform job starts, SageMaker initializes compute instances and distributes the inference or preprocessing workload between them. Batch Transform partitions the Amazon S3 objects in the input by key and maps Amazon S3 objects to instances. When you have multiple files, one instance might process input1.csv, and another instance might process the file named input2.csv. If you have one input file but initialize multiple compute instances, only one instance processes the input file and the rest of the instances are idle.
You can also split input files into mini-batches. For example, you might create a mini-batch from input1.csv by including only two of the records.

Record3-Attribute1, Record3-Attribute2, Record3-Attribute3, ..., Record3-AttributeM
Record4-Attribute1, Record4-Attribute2, Record4-Attribute3, ..., Record4-AttributeM

**Note**

SageMaker processes each input file separately. It doesn't combine mini-batches from different input files to comply with the MaxPayloadInMB limit.

To split input files into mini-batches when you create a batch transform job, set the **SplitType** parameter value to **Line**. If **SplitType** is set to **None** or if an input file can't be split into mini-batches, SageMaker uses the entire input file in a single request. Note that Batch Transform doesn't support CSV-formatted input that contains embedded newline characters. You can control the size of the mini-batches by using the **BatchStrategy** and **MaxPayloadInMB** parameters. MaxPayloadInMB must not be greater than 100 MB. If you specify the optional **MaxConcurrentTransforms** parameter, then the value of (MaxConcurrentTransforms * MaxPayloadInMB) must also not exceed 100 MB.

If the batch transform job successfully processes all of the records in an input file, it creates an output file with the same name and the **.out** file extension. For multiple input files, such as input1.csv and input2.csv, the output files are named input1.csv.out and input2.csv.out. The batch transform job stores the output files in the specified location in Amazon S3, such as s3://awsexamplebucket/output/.

The predictions in an output file are listed in the same order as the corresponding records in the input file. The output file input1.csv.out, based on the input file shown earlier, would look like the following.

Inference1-Attribute1, Inference1-Attribute2, Inference1-Attribute3, ..., Inference1-AttributeM
Inference2-Attribute1, Inference2-Attribute2, Inference2-Attribute3, ..., Inference2-AttributeM
Inference3-Attribute1, Inference3-Attribute2, Inference3-Attribute3, ..., Inference3-AttributeM
If you set `SplitType` to `Line`, you can set the `AssembleWith` parameter to `Line` to concatenate the output records with a line delimiter. This does not change the number of output files. The number of output files is equal to the number of input files, and using `AssembleWith` does not merge files. If you don't specify the `AssembleWith` parameter, by default the output records are concatenated in a binary format.

When the input data is very large and is transmitted using HTTP chunked encoding, to stream the data to the algorithm, set `MaxPayloadInMB` to 0. Amazon SageMaker built-in algorithms don't support this feature.

For information about using the API to create a batch transform job, see the `CreateTransformJob` API. For more information about the correlation between batch transform input and output objects, see `OutputDataConfig`. For an example of how to use batch transform, see (Optional) Make Prediction with Batch Transform.

**Speed up a Batch Transform Job**

If you are using the `CreateTransformJob` API, you can reduce the time it takes to complete batch transform jobs by using optimal values for parameters such as `MaxPayloadInMB`, `MaxConcurrentTransforms`, or `BatchStrategy`. The ideal value for `MaxConcurrentTransforms` is equal to the number of compute workers in the batch transform job. If you are using the SageMaker console, you can specify these optimal parameter values in the `Additional configuration` section of the `Batch transform job configuration` page. SageMaker automatically finds the optimal parameter settings for built-in algorithms. For custom algorithms, provide these values through an `execution-parameters` endpoint.

**Use Batch Transform to Test Production Variants**

To test different models or various hyperparameter settings, create a separate transform job for each new model variant and use a validation dataset. For each transform job, specify a unique model name and location in Amazon S3 for the output file. To analyze the results, use Inference Pipeline Logs and Metrics.
Batch Transform Sample Notebooks

For a sample notebook that uses batch transform with a principal component analysis (PCA) model to reduce data in a user-item review matrix, followed by the application of a density-based spatial clustering of applications with noise (DBSCAN) algorithm to cluster movies, see Batch Transform with PCA and DBSCAN Movie Clusters. For instructions on creating and accessing Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After creating and opening a notebook instance, choose the SageMaker Examples tab to see a list of all the SageMaker examples. The topic modeling example notebooks that use the NTM algorithms are located in the Advanced functionality section. To open a notebook, choose its Use tab, then choose Create copy.

Associate Prediction Results with Input Records

When making predictions on a large dataset, you can exclude attributes that aren't needed for prediction. After the predictions have been made, you can associate some of the excluded attributes with those predictions or with other input data in your report. By using batch transform to perform these data processing steps, you can often eliminate additional preprocessing or postprocessing. You can use input files in JSON and CSV format only.

Topics

- Workflow for Associating Inferences with Input Records
- Use Data Processing in Batch Transform Jobs
- Supported JSONPath Operators
- Batch Transform Examples

Workflow for Associating Inferences with Input Records

The following diagram shows the workflow for associating inferences with input records.
To associate inferences with input data, there are three main steps:

1. Filter the input data that is not needed for inference before passing the input data to the batch transform job. Use the `InputFilter` parameter to determine which attributes to use as input for the model.

2. Associate the input data with the inference results. Use the `JoinSource` parameter to combine the input data with the inference.

3. Filter the joined data to retain the inputs that are needed to provide context for interpreting the predictions in the reports. Use `OutputFilter` to store the specified portion of the joined dataset in the output file.

**Use Data Processing in Batch Transform Jobs**

When creating a batch transform job with `CreateTransformJob` to process data:

1. Specify the portion of the input to pass to the model with the `InputFilter` parameter in the `DataProcessing` data structure.

2. Join the raw input data with the transformed data with the `JoinSource` parameter.
3. Specify which portion of the joined input and transformed data from the batch transform job to include in the output file with the `OutputFilter` parameter.

4. Choose either JSON- or CSV-formatted files for input:
   - For JSON- or JSON Lines-formatted input files, SageMaker either adds the `SageMakerOutput` attribute to the input file or creates a new JSON output file with the `SageMakerInput` and `SageMakerOutput` attributes. For more information, see [DataProcessing](#).
   - For CSV-formatted input files, the joined input data is followed by the transformed data and the output is a CSV file.

If you use an algorithm with the `DataProcessing` structure, it must support your chosen format for both input and output files. For example, with the `TransformOutput` field of the `CreateTransformJob` API, you must set both the `ContentType` and `Accept` parameters to one of the following values: `text/csv`, `application/json`, or `application/jsonlines`. The syntax for specifying columns in a CSV file and specifying attributes in a JSON file are different. Using the wrong syntax causes an error. For more information, see [Batch Transform Examples](#). For more information about input and output file formats for built-in algorithms, see [Use Amazon SageMaker Built-in Algorithms or Pre-trained Models](#).

The record delimiters for the input and output must also be consistent with your chosen file input. The `SplitType` parameter indicates how to split the records in the input dataset. The `AssembleWith` parameter indicates how to reassemble the records for the output. If you set input and output formats to `text/csv`, you must also set the `SplitType` and `AssembleWith` parameters to `line`. If you set the input and output formats to `application/jsonlines`, you can set both `SplitType` and `AssembleWith` to `line`.

For CSV files, you cannot use embedded newline characters. For JSON files, the attribute name `SageMakerOutput` is reserved for output. The JSON input file can't have an attribute with this name. If it does, the data in the input file might be overwritten.

### Supported JSONPath Operators

To filter and join the input data and inference, use a JSONPath subexpression. SageMaker supports only a subset of the defined JSONPath operators. The following table lists the supported JSONPath operators. For CSV data, each row is taken as a JSON array, so only index based JSONPaths can be applied, e.g. `$[0]$, `$[1:]$`. CSV data should also follow [RFC format](#).
<table>
<thead>
<tr>
<th>JSONPath Operator</th>
<th>Description</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>$</td>
<td>The root element to a query. This operator is required at the beginning of all path expressions.</td>
<td>$</td>
</tr>
<tr>
<td>.&lt;name&gt;</td>
<td>A dot-notated child element.</td>
<td>$.id</td>
</tr>
<tr>
<td>*</td>
<td>A wildcard. Use in place of an attribute name or numeric value.</td>
<td>$.id.*</td>
</tr>
<tr>
<td>['&lt;name&gt;' (,',&lt;name&gt;')]</td>
<td>A bracket-notated element or multiple child elements.</td>
<td>$['id','SageMakerOutput']</td>
</tr>
<tr>
<td>[&lt;number&gt; (,'&lt;number&gt;')]]</td>
<td>An index or array of indexes. Negative index values are also supported. A -1 index refers to the last element in an array.</td>
<td>$[1],[1,3,5]</td>
</tr>
<tr>
<td>[&lt;start&gt;:&lt;end&gt;]</td>
<td>An array slice operator. The array slice() method extracts a section of an array and returns a new array. If you omit &lt;start&gt;, SageMaker uses the first element of the array. If you omit &lt;end&gt;, SageMaker uses the last element of the array.</td>
<td>$[2:5],[::5],[2:]</td>
</tr>
</tbody>
</table>

When using the bracket-notation to specify multiple child elements of a given field, additional nesting of children within brackets is not supported. For example, $.field1.['child1','child2'] is supported while $.field1.['child1','child2.grandchild'] is not.

For more information about JSONPath operators, see [JsonPath](https://github.com/json-path/json-path) on GitHub.

**Batch Transform Examples**

The following examples show some common ways to join input data with prediction results.

**Topics**
• **Example: Output Only Inferences**

• **Example: Output Inferences Joined with Input Data**

• **Example: Output Inferences Joined with Input Data and Exclude the ID Column from the Input (CSV)**

• **Example: Output Inferences Joined with an ID Column and Exclude the ID Column from the Input (CSV)**

**Example: Output Only Inferences**

By default, the `DataProcessing` parameter doesn't join inference results with input. It outputs only the inference results.

If you want to explicitly specify to not join results with input, use the Amazon SageMaker Python SDK and specify the following settings in a transformer call.

```python
sm_transformer = sagemaker.transformer.Transformer(...) sm_transformer.transform(..., input_filter="$", join_source= "None", output_filter="$")
```

To output inferences using the AWS SDK for Python, add the following code to your `CreateTransformJob` request. The following code mimics the default behavior.

```json
{
    "DataProcessing": {
        "InputFilter": "$",
        "JoinSource": "None",
        "OutputFilter": "$"
    }
}
```

**Example: Output Inferences Joined with Input Data**

If you're using the Amazon SageMaker Python SDK to combine the input data with the inferences in the output file, specify the `assemble_with` and `accept` parameters when initializing the transformer object. When you use the transform call, specify `Input` for the `join_source` parameter, and specify the `split_type` and `content_type` parameters as well. The `split_type` parameter must have the same value as `assemble_with`, and the `content_type` parameter must have the same value as `accept`. For more information about the parameters and their accepted values, see the Transformer page in the Amazon SageMaker Python SDK.
sm_transformer = sagemaker.transformer.Transformer(..., assemble_with="Line", accept="text/csv")
sm_transformer.transform(..., join_source="Input", split_type="Line", content_type="text/csv")

If you're using the AWS SDK for Python (Boto 3), join all input data with the inference by adding the following code to your `CreateTransformJob` request. The values for `Accept` and `ContentType` must match, and the values for `AssembleWith` and `SplitType` must also match.

```json
{
    "DataProcessing": {
        "JoinSource": "Input"
    },
    "TransformOutput": {
        "Accept": "text/csv",
        "AssembleWith": "Line"
    },
    "TransformInput": {
        "ContentType": "text/csv",
        "SplitType": "Line"
    }
}
```

For JSON or JSON Lines input files, the results are in the `SageMakerOutput` key in the input JSON file. For example, if the input is a JSON file that contains the key-value pair `{"key":1}`, the data transform result might be `{"label":1}`.

SageMaker stores both in the input file in the `SageMakerInput` key.

```json
{
    "key":1,
    "SageMakerOutput":{"label":1}
}
```

Note

The joined result for JSON must be a key-value pair object. If the input isn't a key-value pair object, SageMaker creates a new JSON file. In the new JSON file, the input data is stored in the `SageMakerInput` key and the results are stored as the `SageMakerOutput` value.
For a CSV file, for example, if the record is [1, 2, 3], and the label result is [1], then the output file would contain [1, 2, 3, 1].

**Example: Output Inferences Joined with Input Data and Exclude the ID Column from the Input (CSV)**

If you are using the Amazon SageMaker Python SDK to join your input data with the inference output while excluding an ID column from the transformer input, specify the same parameters from the preceding example as well as a JSONPath subexpression for the `input_filter` in your transformer call. For example, if your input data includes five columns and the first one is the ID column, use the following transform request to select all columns except the ID column as features. The transformer still outputs all of the input columns joined with the inferences. For more information about the parameters and their accepted values, see the Transformer page in the Amazon SageMaker Python SDK.

```python
sm_transformer = sagemaker.transformer.Transformer(..., assemble_with="Line", accept="text/csv")
sm_transformer.transform(..., split_type="Line", content_type="text/csv",
    input_filter="$[1:]", join_source="Input")
```

If you are using the AWS SDK for Python (Boto 3), add the following code to your `CreateTransformJob` request.

```json
{
    "DataProcessing": {
        "InputFilter": "$[1:]",
        "JoinSource": "Input"
    },
    "TransformOutput": {
        "Accept": "text/csv",
        "AssembleWith": "Line"
    },
    "TransformInput": {
        "ContentType": "text/csv",
        "SplitType": "Line"
    }
}
```

To specify columns in SageMaker, use the index of the array elements. The first column is index 0, the second column is index 1, and the sixth column is index 5.
To exclude the first column from the input, set `InputFilter` to "$[1:]". The colon (:) tells SageMaker to include all of the elements between two values, inclusive. For example, $[1:4]$ specifies the second through fifth columns.

If you omit the number after the colon, for example, [5:], the subset includes all columns from the 6th column through the last column. If you omit the number before the colon, for example, [:5], the subset includes all columns from the first column (index 0) through the sixth column.

**Example: Output Inferences Joined with an ID Column and Exclude the ID Column from the Input (CSV)**

If you are using the Amazon SageMaker Python SDK, you can specify the output to join only specific input columns (such as the ID column) with the inferences by specifying the `output_filter` in the transformer call. The `output_filter` uses a JSONPath subexpression to specify which columns to return as output after joining the input data with the inference results. The following request shows how you can make predictions while excluding an ID column and then join the ID column with the inferences. Note that in the following example, the last column (-1) of the output contains the inferences. If you are using JSON files, SageMaker stores the inference results in the attribute `SageMakerOutput`. For more information about the parameters and their accepted values, see the Transformer page in the Amazon SageMaker Python SDK.

```python
sm_transformer = sagemaker.transformer.Transformer(…, assemble_with="Line", accept="text/csv")
sm_transformer.transform(…, split_type="Line", content_type="text/csv",
input_filter="$[1:]", join_source="Input", output_filter="$[0,-1]"
)
```

If you are using the AWS SDK for Python (Boto 3), join only the ID column with the inferences by adding the following code to your `CreateTransformJob` request.

```json
{
    "DataProcessing": {
        "InputFilter": "$[1:]",
        "JoinSource": "Input",
        "OutputFilter": "$[0,-1]"
    },
    "TransformOutput": {
        "Accept": "text/csv",
        "AssembleWith": "Line"
    },
    "TransformInput": {
```
Warning

If you are using a JSON-formatted input file, the file can't contain the attribute name SageMake\textbackslash rOutput. This attribute name is reserved for the inferences in the output file. If your JSON-formatted input file contains an attribute with this name, values in the input file might be overwritten with the inference.

Storage in Batch Transform

When you run a batch transform job, Amazon SageMaker attaches an Amazon Elastic Block Store storage volume to Amazon EC2 instances that process your job. The volume stores your model, and the size of the storage volume is fixed at 30 GB. You have the option to encrypt your model at rest in the storage volume.

Note

If you have a large model, you may encounter an InternalServerError.

For more information about Amazon EBS storage and features, see the following pages:

- Amazon EBS in the Amazon EC2 User Guide for Linux Instances
- Amazon EBS volumes in the Amazon EC2 User Guide for Linux Instances

Note

G4dn instances come with their own local SSD storage. To learn more about G4dn instances, see the Amazon EC2 G4 Instances page.
Troubleshooting

If you are having errors in Amazon SageMaker Batch Transform, refer to the following troubleshooting tips.

Max timeout errors

If you are getting max timeout errors when running batch transform jobs, try the following:

• Begin with the single-record BatchStrategy, a batch size of the default (6 MB) or smaller which you specify in the MaxPayloadInMB parameter, and a small sample dataset. Tune the maximum timeout parameter InvocationsTimeoutInSeconds (which has a maximum of 1 hour) until you receive a successful invocation response.

• After you receive a successful invocation response, increase the MaxPayloadInMB (which has a maximum of 100 MB) and the InvocationsTimeoutInSeconds parameters together to find the maximum batch size that can support your desired model timeout. You can use either the single-record or multi-record BatchStrategy in this step.

⚠️ Note

Exceeding the MaxPayloadInMB limit causes an error. This might happen with a large dataset if it can't be split, the SplitType parameter is set to none, or individual records within the dataset exceed the limit.

• (Optional) Tune the MaxConcurrentTransforms parameter, which specifies the maximum number of parallel requests that can be sent to each instance in a batch transform job. However, the value of MaxConcurrentTransforms * MaxPayloadInMB must not exceed 100 MB.

Incomplete output

SageMaker uses the Amazon S3 Multipart Upload API to upload results from a batch transform job to Amazon S3. If an error occurs, the uploaded results are removed from Amazon S3. In some cases, such as when a network outage occurs, an incomplete multipart upload might remain in Amazon S3. An incomplete upload might also occur if you have multiple input files but some of the files can't be processed by SageMaker Batch Transform. The input files that couldn't be processed won't have corresponding output files in Amazon S3.
To avoid incurring storage charges, we recommend that you add the S3 bucket policy to the S3 bucket lifecycle rules. This policy deletes incomplete multipart uploads that might be stored in the S3 bucket. For more information, see Object Lifecycle Management.

**Job shows as failed**

If a batch transform job fails to process an input file because of a problem with the dataset, SageMaker marks the job as failed. If an input file contains a bad record, the transform job doesn’t create an output file for that input file because doing so prevents it from maintaining the same order in the transformed data as in the input file. When your dataset has multiple input files, a transform job continues to process input files even if it fails to process one. The processed files still generate useable results.

If you are using your own algorithms, you can use placeholder text, such as ERROR, when the algorithm finds a bad record in an input file. For example, if the last record in a dataset is bad, the algorithm places the placeholder text for that record in the output file.

**Model parallelism and large model inference**

State-of-the-art deep learning models for applications such as natural language processing (NLP) are large, typically with tens or hundreds of billions of parameters. Larger models are often more accurate, which makes them attractive to machine learning practitioners. However, these models are often too large to fit on a single accelerator or GPU device, making it difficult to achieve low-latency inference. You can avoid this memory bottleneck by using model parallelism techniques to partition a model across multiple accelerators or GPUs.

Amazon SageMaker includes specialized deep learning containers (DLCs), libraries, and tooling for model parallelism and large model inference (LMI). In the following sections, you can find resources to get started with LMI on SageMaker.

**Topics**

- Deep learning containers for large model inference
- SageMaker endpoint parameters for large model inference
- Large model inference tutorials
- Configurations and settings
- Choosing instance types for large model inference
Deep learning containers for large model inference

SageMaker maintains deep learning containers (DLCs) with popular open source libraries for hosting large models such as GPT, T5, OPT, BLOOM, and Stable Diffusion on AWS infrastructure. With these DLCs you can use third party libraries such as DeepSpeed, Accelerate, TensorRT-LLM, and FasterTransformer to partition model parameters using model parallelism techniques to leverage the memory of multiple GPUs for inference. The following table lists the DLCs available with SageMaker for large model inference (LMI). We recommend that you start with these DLCs for LMI on SageMaker. These include components, libraries, and drivers that have been optimized and tested for use on SageMaker.

<table>
<thead>
<tr>
<th>DLC URI</th>
<th>Libraries</th>
<th>Tuning parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.25.0-tensorrtlm0.5.0-cu122</td>
<td>PyTorch 2.1.0</td>
<td>TensorRT-LLM</td>
</tr>
<tr>
<td></td>
<td>DJL Serving 0.25.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TensorRT toolkit 0.5.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Accelerate 0.23.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Transformers 4.34.0</td>
<td></td>
</tr>
<tr>
<td>763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.25.0-deepspeed0.11.0-cu118</td>
<td>PyTorch 2.0.1</td>
<td>DeepSpeed</td>
</tr>
<tr>
<td></td>
<td>DJL Serving 0.25.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>DeepSpeed 0.11.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Accelerate 0.23.0</td>
<td></td>
</tr>
<tr>
<td>DLC URI</td>
<td>Libraries</td>
<td>Tuning parameters</td>
</tr>
<tr>
<td>------------------------------------------------------------------------</td>
<td>----------------------------------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>763104351884.dkr.e region.amazonaws.com/djl-inference:0.25.0-neuronx-sdk2.15.0</td>
<td>Hugging Face Transformers 4.34.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PyTorch 1.13.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>DJL Serving 0.25.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TransformersNeuronX 0.8.268</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AWS Neuron SDK 2.15.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>NeuronX Distributed 0.5.0</td>
<td></td>
</tr>
<tr>
<td>763104351884.dkr.e region.amazonaws.com/djl-inference:0.23.0-fastertransformer5.3.0-cu118</td>
<td>PyTorch 2.0.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>DJL Serving 0.23.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>FasterTransformer 5.3.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Accelerate 0.20.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Transformers 4.30.1</td>
<td></td>
</tr>
<tr>
<td>763104351884.dkr.e region.amazonaws.com/djl-inference:0.23.0-deepspeed0.9.5-cu118</td>
<td>PyTorch 2.0.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>DJL Serving 0.23.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>DeepSpeed 0.9.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Accelerate 0.20.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hugging Face Transformers 4.30.2</td>
<td></td>
</tr>
</tbody>
</table>
In addition to PyTorch, LMI DLCs include libraries to facilitate large model inference. SageMaker supports the following categories of libraries.

- **Model zoo** – Model zoos provide simple API access to pre-trained models. SageMaker provides the following model zoos:
  - [Hugging Face Transformers](https://huggingface.co/) is a popular library for pre-trained deep learning models that use a transformer architecture such as GPT, OPT, and BLOOM.
  - [Hugging Face Diffusers](https://huggingface.co/) is a library with pre-trained deep learning models that use a diffusion technique such as Stable Diffusion.

- **Model parallelism and inference optimization libraries** – These libraries handle model parallel inference by partitioning a model artifact so that its comprising parameters can be spread across multiple GPUs. SageMaker supports the following model parallelism and inference optimization libraries:
  - [TensorRT-LLM](https://github.com/nvidia/tensorrt llama) is the latest open-source library from NVIDIA which is available as part of LMI DLCs release (0.25.0). This library enables state-of-the-art optimizations, like SmoothQuant, FP8 and continuous batching for large language models when using NVIDIA GPUs.

  TensorRT-LLM supports deployments ranging from single GPU to multi-GPU configurations, with additional performance gains possible through techniques like tensor parallelism. Leveraging TensorRT-LLM through the SageMaker LMI DLC enables you to optimize the performance of your large language models and provide your users with a more responsive experience.

  We optimized the TensorRT-LLM library to speed up inference, and we created a toolkit that supports just-in-time model conversion. You can use this toolkit to provide a Hugging Face
model ID and deploy the model end to end. The toolkit also supports continuous batching with streaming. You can compile the Llama-2 7B and 13B models in around 1-2 minutes, and you can compile the 70B model in around 7 minutes.

To avoid the overhead of compiling when you set up a SageMaker endpoint and scale your instances, you can use Ahead of Time (AOT) compilation. For more information about preparing the model, see TensorRT-LLM ahead-of-time compilation of models tutorial.

We also accept any TensorRT LLM model built for Triton Server that can be used with LMI DLC. For examples about deploying Llama-2 70B using the TRT-LLM 0.25.0 LMI container, see the notebooks.

- **DeepSpeed Inference** is an open-source inference optimization library. It includes model partitioning schemes for model parallelism with supported models, including many transformer models. It also has optimized kernels for popular models such as OPT, GPT, and BLOOM that can significantly improve inference latency. The version of DeepSpeed in the LMI DLCs is optimized and tested to work on SageMaker. It includes several enhancements, including support for BF16 precision models.

- **Hugging Face Accelerate** in an open-source model parallel inference library. It supports model parallelism for most models in the Hugging Face Transformers library.

- **FasterTransformer** is an open source library from Nvidia that provides an accelerated engine for efficiently running transformer-based neural network inference. It has been designed to handle large models that require multiple GPUs and nodes in a distributed manner. The library includes an optimized version of the transformer block, which comprises both the encoder and decoder parts, enabling you to run the inference of full encoder-decoder architectures like T5, as well as encoder-only models like BERT and decoder-only models like GPT.

- **Model server** – Model servers handle an inference request end to end. They accept requests, invoke pre-processing and post-processing scripts, and respond to users. Model servers that are compatible with model parallelism, also organize workers and threads across multiple devices. SageMaker supports the following model servers:

  - **DJL-Serving** is an open-source, high-performance model server powered by DJL. It takes multiple deep learning models or workflows, and makes them available through an HTTP endpoint. Versions 0.19 and above are supported by SageMaker and work with Amazon EC2 instances with multiple GPUs to facilitate LMI with model parallelism.
**Supported instance types**

AWS LMI DLCs support the p4d, p3, g5, and g4dn instance types.

**SageMaker endpoint parameters for large model inference**

You can customize the following parameters to facilitate low-latency large model inference (LMI) with SageMaker:

- **Maximum Amazon EBS volume size on the instance (VolumeSizeInGB)** – If the size of the model is larger than 30 GB and you are using an instance without a local disk, you should increase this parameter to be slightly larger than the size of your model.

- **Health check timeout quota (ContainerStartupHealthCheckTimeoutInSeconds)** – If your container is correctly set up and the CloudWatch logs indicate a health check timeout, you should increase this quota so the container has enough time to respond to health checks.

- **Model download timeout quota (ModelDataDownloadTimeoutInSeconds)** – If the size of your model is larger than 40 GB, then you should increase this quota to provide sufficient time to download the model from Amazon S3 to the instance.

For more information on low latency inference with large models, see [Deploy large models on Amazon SageMaker using DJL Serving and DeepSpeed model parallel inference](#). The following code snippet demonstrates how to programatically configure the aforementioned parameters. Replace the *italicized placeholder text* in the example with your own information.

```python
import boto3

aws_region = "aws-region"
sagemaker_client = boto3.client('sagemaker', region_name=aws_region)

# The name of the endpoint. The name must be unique within an AWS Region in your AWS account.
endpoint_name = "endpoint-name"

# Create an endpoint config name.
endpoint_config_name = "endpoint-config-name"

# The name of the model that you want to host.
model_name = "the-name-of-your-model"

instance_type = "instance-type"
```
sagemaker_client.create_endpoint_config(
    EndpointConfigName = endpoint_config_name
    ProductionVariants=[
        {
            "VariantName": "variant1", # The name of the production variant.
            "ModelName": model_name,
            "InstanceType": instance_type, # Specify the compute instance type.
            "InitialInstanceCount": 1, # Number of instances to launch initially.
            "VolumeSizeInGB": 256, # Specify the size of the Amazon EBS volume.
            "ModelDataDownloadTimeoutInSeconds": 1800, # Specify the model download timeout in seconds.
            "ContainerStartupHealthCheckTimeoutInSeconds": 1800, # Specify the health checkup timeout in seconds
        },
    ],
)

sagemaker_client.create_endpoint(EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name)

For more information about the keys for ProductionVariants, see ProductionVariant.

**Large model inference tutorials**

The following tutorials demonstrate the use of different model parallelization and inference optimization libraries available inside a large model inference (LMI) DLC. You can use these to get started with LMI on SageMaker.

**Topics**
- Large model inference with DeepSpeed and DJL Serving
- Large model inference with FasterTransformer and DJL Serving
- Large model inference with TorchServe
- Additional resources to get started

**Large model inference with DeepSpeed and DJL Serving**

This tutorial demonstrates how to deploy large models with DJL Serving using DeepSpeed and Hugging Face Accelerate model parallelization frameworks. This example uses a GPT-J model with
6 billion parameters and an ml.g5 instance. You can modify this to work with other models and instance types. Replace the italicized placeholder text in the examples with your own information.

Topics

- Large model inference container image with DJL Serving
- Preparing your model artifacts
- Deploy the model using the SageMaker SDK

Large model inference container image with DJL Serving

Large model inference (LMI) DLCs are available as Docker images on the Amazon Elastic Container Registry (Amazon ECR). These containers include the necessary components, libraries, and drivers to host large models on SageMaker or using Amazon EC2 infrastructure. You can find a list of LMI DLCs in the Large model inference section of the DLCs list. This tutorial uses the following container:

<table>
<thead>
<tr>
<th>Framework</th>
<th>Job type</th>
<th>CPU/GPU</th>
<th>Python version options</th>
<th>Example URL</th>
</tr>
</thead>
<tbody>
<tr>
<td>DJL Serving 0.25.0 with DeepSpeed 0.11.0, Hugging</td>
<td>Inference</td>
<td>GPU</td>
<td>3.9 (py39)</td>
<td>763104351884.dkr.ecr.us-east-1.amazonaws.com/djl-inference:0.25.0-deepspeed0.11.0-cu118</td>
</tr>
<tr>
<td>Face Transformers 4.34.0, and Hugging Face Accelerate 0.23.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

LMI DLCs are regularly updated and tested on SageMaker. You can directly reference these Docker image URLs when you host on SageMaker, selecting the container that suits your need the most. Each LMI DLC image includes DeepSpeed, Accelerate, Transformers and DJL Serving.
Preparing your model artifacts

LMI DLCs use DJL Serving to serve your model for inference. You have to configure DJL Serving and package your model in a format supported by DJL Serving and DeepSpeed for LMI.

1. Create a `serving.properties` configuration file to indicate to DJL Serving which model parallelization and inference optimization libraries you would like to use. You can find the configuration options for both DeepSpeed and Hugging Face Accelerate in Configurations and settings. Modify the following example `serving.properties` file below to suit your needs.

```properties
engine=DeepSpeed
#engine=Python # for Hugging Face Accelerate, vLLM rolling batch
#engine=MPI # lmi-dist rolling batch option
option.entryPoint=djl_python.deepspeed
#option.entryPoint=djl_python.huggingface # for Hugging Face Accelerate
option.tensor_parallel_degree=2
option.model_id=EleutherAI/gpt-j-6B
option.max_rolling_batch_size=64
option.rolling_batch=deepspeed
option.task=text-generation
option.dtype=fp16
```

For a complete list of configuration options, see Configurations and settings. The following list describes some of the options used in the example `serving.properties` file.

- **option.entryPoint** – This option is used to specify which handler offered by DJL Serving you would like to use. The possible values are huggingface, deepspeed, and stable-diffusion. Here we use deepspeed.

- **option.model_id** – Only provide this option if you also provide `option.entryPoint`. The value of this option is either the Hugging Face ID of a model or the S3 URL of the model artifacts. DJL Serving uses this value to download the model from Hugging Face or the S3 URL.

- **option.task** – Model architectures generally require you specify a task as one of the parameters in `transformers.pipeline`.

- **option.rolling_batch** – Select a rolling batch strategy. The value `auto` will make the handler choose the strategy based on the provided configuration. If you specify `option.rolling_batch=djl_python.huggingface` and `auto`, one of the following options will be chosen automatically based on the model: `lmi-dist`, or `scheduler`. If engine is DeepSpeed, then deepspeed rolling batch is chosen.
For more information, see the DeepSpeed, and Hugging Face handler implementations.

2. Package your model artifacts into a compressed .tar file. DJL Serving and DeepSpeed expect the model artifacts to be packaged and formatted in a specific way. This example uses the following directory structure for this purpose.

```
- deepspeed-gptj  # root directory
  - serving.properties
  - model.py # your custom handler file, if you choose not to use the handlers provided with DJL Serving
  - model binary files # if you do not want to use option.model_id
```

The root directory contains the serving.properties file, along with any other necessary files. This tutorial only needs the serving.properties file because it uses DJL Serving's handler and the model_id option to download the model directly from Hugging Face. The following code snippet packages the model artifacts into a compressed .tar file.

```
mkdir deepspeed-gptj
 cp serving.properties deepspeed-gptj/
tar -czvf deepspeed-gptj.tar.gz deepspeed-gptj
aws s3 sync deepspeed-gptj.tar.gz s3://djl-sm-test/tutorial/
```

Deploy the model using the SageMaker SDK

This example uses SageMaker to handle the end to end model deployment process to an inference endpoint.

1. Create a SageMaker session.

```
import boto3
import sagemaker

aws_region = "aws-region"
sagemaker_session =
    sagemaker.Session(boto_session=boto3.Session(region_name=aws_region))
```
2. Create a model in SageMaker.

```python
from sagemaker.model import Model
from sagemaker import image_uris, get_execution_role

role = get_execution_role()

def create_model(model_name, model_s3_url):
    # Get the DJL DeepSpeed image uri
    image_uri = image_uris.retrieve(
        framework="djl-deepspeed",
        region=sagemaker_session.boto_session.region_name,
        version="0.25.0"
    )
    model = Model(
        image_uri=image_uri,
        model_data=model_s3_url,
        role=role,
        name=model_name,
        sagemaker_session=sagemaker_session,
    )
    return model
```

The g5.12xlarge instance has 4 Nvidia A10G GPUs. The `serving.properties` in [Preparing your model artifacts](#) specified a tensor parallel degree of 2. DJL Serving automatically loads 2 copies of the model with 2 tensor parallel partitions to make use of all the 4 GPUs on the instance. This doubles the throughput.

3. Deploy the model to a g5.12xlarge instance.

```python
from sagemaker import serializers, deserializers

def deploy_model(model, _endpoint_name):
    model.deploy(
        initial_instance_count=1,
        instance_type="ml.g5.12xlarge",
        endpoint_name=_endpoint_name
    )
    predictor = sagemaker.Predictor(
        endpoint_name=_endpoint_name,
        sagemaker_session=sagemaker_session,
        serializer=serializers.JSONSerializer(),
```
Large model inference with FasterTransformer and DJL Serving

This tutorial demonstrates how to deploy a T5 model with large model inference (LMI) deep learning containers (DLCs), DJL Serving, and the FasterTransformer model parallelization framework. Here we use a flan-t5-xl model with 3 billion parameters and an ml.g5 instance. You can modify this to work with other variants of T5 models and instance types. Replace the italicized placeholder text in the examples with your own information.
Large model inference container image with FasterTransformer backend for DJL Serving

LMI DLCs are available as Docker images on the Amazon Elastic Container Registry (Amazon ECR). These containers include the necessary components, libraries, and drivers to host large models on SageMaker or using Amazon EC2 infrastructure. You can find a list of LMI DLCs in the Large model inference section of the DLCs list. This tutorial uses the following container:

<table>
<thead>
<tr>
<th>Framework</th>
<th>Job type</th>
<th>CPU/GPU</th>
<th>Python version options</th>
<th>Example URL</th>
</tr>
</thead>
<tbody>
<tr>
<td>DJL Serving 0.23.0 with FasterTransformer 5.3, Hugging Face Transformers 4.30.1.</td>
<td>Inference</td>
<td>GPU</td>
<td>3.9 (py39)</td>
<td>763104351884.dkr.ecr.us-west-2.amazonaws.com/djl-inference:0.23.0-fastertransformer5.3.0-cu118</td>
</tr>
</tbody>
</table>

LMI DLCs are regularly updated and tested on SageMaker. You can directly reference these Docker image URLs when you host on SageMaker, selecting the container that suits your need the most. Each LMI DLC image includes DeepSpeed, Accelerate, Transformers and DJL Serving.

Preparing your model artifacts

LMI DLCs use DJL Serving to serve your model for inference. You must configure DJL Serving and package your model in a format supported by DJL Serving and FasterTransformer.

1. Create a `serving.properties` configuration file to indicate to DJL Serving which model parallelization and inference optimization libraries you would like to use.

   ```
   engine=FasterTransformer
   option.tensor_parallel_degree=2
   option.model_id=google/flan-t5-xl
   option.dtype=fp32
   ```

   For a complete list of configuration options, see Configurations and settings. The following list describes some of the options used in the aforementioned example.
• **option.tensor_parallel_degree** – This option specifies number of tensor parallel partitions performed on the model.

• **option.model_id** – The value of this option is either the Hugging Face ID of a model or the S3 URL of the model artifacts. DJL Serving uses this value to download the model from Hugging Face or the S3 URL.

• **option.dtype** – This option specifies data type that you want to run inference with.

2. Create a `model.py` handler script with a `handle` method that will be invoked by DJL Serving when inference requests are received. This tutorial uses the following `model.py`:

```python
import fastertransformer as ft
from djl_python import Input, Output
from transformers import AutoTokenizer
import logging
import os

model = None

def load_model(properties):
    model = ft.init_inference(model_name, tensor_parallel_degree,
                              pipeline_parallel_degree, dtype)
    return model

def handle(inputs: Input):
    global model

    if not model:
        model = load_model(inputs.get_properties())

    if inputs.is_empty():
        # Model server makes an empty call to warmup the model on startup
        return None
```

LMI tutorials
data = inputs.get_as_json()
input_text = data["text"]
result = model.pipeline_generate(input_text)
return Output().add(result)

3. Package your model artifacts into a compressed tar file. DJL Serving expects the model artifacts to be packaged and formatted in a specific way. This example uses the following directory structure for this purpose.

```
- deepspeed-gptj # root directory
  - serving.properties
  - model.py # your custom handler file, if you choose not to use the handlers provided with DJL Serving
```

The root directory contains the `serving.properties` and `model.py` files. The following code snippet packages the model artifacts into a compressed .tar file.

```
mkdir fastertransformer-t5
cp serving.properties fastertransformer-t5/
cp model.py fastertransformer-t5/
tar -czvf fastertransformer-t5.tar.gz fastertransformer-t5/
aws s3 cp fastertransformer-t5.tar.gz s3://djl-sm-test/tutorial/
```

**Deploy the model using the SageMaker SDK**

This example uses SageMaker to handle the end to end model deployment process to an inference endpoint.

1. Create a SageMaker session.

```
import boto3
import sagemaker

aws_region = "aws-region"
sagemaker_session = sagemaker.Session(boto_session=boto3.Session(region_name=aws_region))
```

2. Create a model in SageMaker.
from sagemaker.model import Model
from sagemaker import image_uris, get_execution_role

role = get_execution_role()

def create_model(model_name, model_s3_url):
    image_uri = "763104351884.dkr.ecr.us-west-2.amazonaws.com/djl-inference:0.23.0-fastertransformer5.3.0-cu118"
    model = Model(
        image_uri=image_uri,
        model_data=model_s3_url,
        role=role,
        name=model_name,
        sagemaker_session=sagemaker_session,
    )
    return model

The g5.12xlarge instance has 4 Nvidia A10G GPUs. The serving.properties in Preparing your model artifacts specified a tensor parallel degree of 2. DJL Serving automatically loads 2 copies of the model with 2 tensor parallel partitions to make use of all the 4 GPUs on the instance. This doubles the throughput.

3. Deploy the model to a g5.12xlarge instance.

from sagemaker import serializers, deserializers

def deploy_model(model, _endpoint_name):
    model.deploy(
        initial_instance_count=1,
        instance_type="ml.g5.12xlarge",
        endpoint_name=_endpoint_name
    )
    predictor = sagemaker.Predictor(
        endpoint_name=_endpoint_name,
        sagemaker_session=sagemaker_session,
        serializer=serializers.JSONSerializer(),
        deserializer=deserializers.JSONDeserializer()
    )
    return predictor

4. Make a prediction
_model_name = "ft-flan-t5-xl"
_model_s3_url = "s3://djl-sm-test/tutorial/fastertransformer-t5.tar.gz"
_endpoint_name="ft-flan-t5-xl"

model = create_model(_model_name, _model_s3_url)
predictor = deploy_model(model, _endpoint_name)
print(predictor.predict(
    {
        "text" : ["translate English to German: The house is wonderful."],
    }
))

5. Spin down the resources

predictor.delete_endpoint(delete_endpoint_config=True)

Large model inference with TorchServe

This tutorial demonstrates how to deploy large models and serve inference in Amazon SageMaker with TorchServe on GPUs. This example deploys the OPT-30b model to an ml.g5 instance. You can modify this to work with other models and instance types. Replace the italicized placeholder text in the examples with your own information.

TorchServe is a powerful open platform for large distributed model inference. By supporting popular libraries like PyTorch, native PiPPy, DeepSpeed, and HuggingFace Accelerate, it offers uniform handler APIs that remain consistent across distributed large model and non-distributed model inference scenarios. For more information, see TorchServe’s large model inference documentation.

Deep learning containers with TorchServe

To deploy a large model with TorchServe on SageMaker, you can use one of the SageMaker deep learning containers (DLCs). By default, TorchServe is installed in all AWS PyTorch DLCs. During model loading, TorchServe can install specialized libraries tailored for large models such as PiPPy, DeepSpeed, and Accelerate.

The following table lists all of the SageMaker DLCs with TorchServe.
<table>
<thead>
<tr>
<th>DLC category</th>
<th>Framework</th>
<th>Hardware</th>
<th>Example URL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SageMaker Framework Containers</strong></td>
<td>PyTorch 2.0.0+</td>
<td>CPU, GPU</td>
<td>763104351884.dkr.ecr.us-east-1.amazonaws.com/pytorch-inference:2.0.1-gpu-py310-cu118-ubuntu20.04-sagemaker</td>
</tr>
<tr>
<td><strong>SageMaker Framework Graviton Containers</strong></td>
<td>PyTorch 2.0.0+</td>
<td>CPU</td>
<td>763104351884.dkr.ecr.us-east-1.amazonaws.com/pytorch-inference-graviton:2.0.1-cpu-py310-ubuntu20.04-sagemaker</td>
</tr>
<tr>
<td><strong>StabilityAI Inference Containers</strong></td>
<td>PyTorch 2.0.0+</td>
<td>GPU</td>
<td>763104351884.dkr.ecr.us-east-1.amazonaws.com/stabilityai-pytorch-inference:2.0.1-sgm0.1.0-gpu-py310-cu118-ubuntu20.04-sagemaker</td>
</tr>
</tbody>
</table>
Getting started

Before deploying your model, complete the prerequisites. You can also configure your model parameters and customize the handler code.

Prerequisites

To get started, ensure that you have the following prerequisites:

1. Ensure you have access to an AWS account. Set up your environment so that the AWS CLI can access your account through either an AWS IAM user or an IAM role. We recommend using an IAM role. For the purposes of testing in your personal account, you can attach the following managed permissions policies to the IAM role:

   - AmazonEC2ContainerRegistryFullAccess
   - AmazonEC2FullAccess
   - AWSServiceRoleForAmazonEKSNodegroup
   - AmazonSageMakerFullAccess
   - AmazonS3FullAccess

   For more information about attaching IAM policies to a role, see Adding and removing IAM identity permissions in the AWS IAM User Guide.

2. Locally configure your dependencies, as shown in the following examples.

   a. Install version 2 of the AWS CLI:

   ```bash
   # Install the latest AWS CLI v2 if it is not installed
   !curl "https://awscli.amazonaws.com/awscli-exe-linux-x86_64.zip" -o "awscliv2.zip" !unzip awscliv2.zip
   #Follow the instructions to install v2 on the terminal
   !cat aws/README.md
   
   # If already installed, update your client
   #%pip install sagemaker pip --upgrade --quiet
   !pip install -U sagemaker
   !pip install -U boto
   !pip install -U botocore
   ```
Configure model settings and parameters

TorchServe uses `torchrun` to set up the distributed environment for model parallel processing. TorchServe has the capability to support multiple workers for a large model. By default, TorchServe uses a round-robin algorithm to assign GPUs to a worker on a host. In the case of large model inference, the number of GPUs assigned to each worker is automatically calculated based on the number of GPUs specified in the `model_config.yaml` file. The environment variable `CUDA_VISIBLE_DEVICES`, which specifies the GPU device IDs that are visible at a given time, is set based on this number.

For example, suppose there are 8 GPUs on a node and one worker needs 4 GPUs on a node (`nproc_per_node=4`). In this case, TorchServe assigns four GPUs to the first worker (`CUDA_VISIBLE_DEVICES="0,1,2,3"`) and four GPUs to the second worker (`CUDA_VISIBLE_DEVICES="4,5,6,7"`).

In addition to this default behavior, TorchServe provides the flexibility for users to specify GPUs for a worker. For instance, if you set the variable `deviceIds: [2,3,4,5]` in the `model config YAML file`, and set `nproc_per_node=2`, then TorchServe assigns `CUDA_VISIBLE_DEVICES="2,3"` to the first worker and `CUDA_VISIBLE_DEVICES="4,5"` to the second worker.

In the following `model_config.yaml` example, we configure both front-end and back-end parameters for the `OPT-30b` model. The configured front-end parameters are `parallelType`, `deviceType`, `deviceIds` and `torchrun`. For more detailed information about the front-end parameters you can configure, see the PyTorch GitHub documentation. The back-end configuration is based on a YAML map that allows for free-style customization. For the back-end parameters, we define the DeepSpeed configuration and additional parameters used by custom handler code.

```yaml
# TorchServe front-end parameters
minWorkers: 1
maxWorkers: 1
maxBatchDelay: 100
responseTimeout: 1200
parallelType: "tp"
deviceType: "gpu"

# example of user specified GPU deviceIds
deviceIds: [0,1,2,3] # sets CUDA_VISIBLE_DEVICES
```
torchrun:
    nproc-per-node: 4

# TorchServe back-end parameters
deespeed:
    config: ds-config.json
    checkpoint: checkpoints.json

handler: # parameters for custom handler code
    model_name: "facebook/opt-30b"
    model_path: "model/models--facebook--opt-30b/snapshots/
ceea0a90ac0f6fae7c2c34bcb40477438c152546"
    max_length: 50
    max_new_tokens: 10
    manual_seed: 40

Customize handlers

TorchServe offers base handlers and handler utilities for large model inference built with popular libraries. The following example demonstrates how the custom handler class TransformersSeqClassifierHandler extends BaseDeepSpeedHandler and uses the handler utilities. For a full code example, see the custom_handler.py code on the PyTorch GitHub documentation.

class TransformersSeqClassifierHandler(BaseDeepSpeedHandler, ABC):
    
    """Transformers handler class for sequence, token classification and question answering."
    """

    def __init__(self):
        super(TransformersSeqClassifierHandler, self).__init__()
        self.max_length = None
        self.max_new_tokens = None
        self.tokenizer = None
        self.initialized = False

    def initialize(self, ctx: Context):
        """In this initialize function, the HF large model is loaded and partitioned using DeepSpeed.
        Args:
            ctx (context): It is a JSON Object containing information pertaining to the model artifacts parameters."""
super().initialize(ctx)
model_dir = ctx.system_properties.get("model_dir")
self.max_length = int(ctx.model_yaml_config["handler"]['max_length'])
self.max_new_tokens = int(ctx.model_yaml_config["handler"]['max_new_tokens'])
model_name = ctx.model_yaml_config["handler"]['model_name']
model_path = ctx.model_yaml_config["handler"]['model_path']
seed = int(ctx.model_yaml_config["handler"]['manual_seed'])
torch.manual_seed(seed)

logger.info("Model %s loading tokenizer", ctx.model_name)
self.tokenizer = AutoTokenizer.from_pretrained(model_name)
self.tokenizer.pad_token = self.tokenizer.eos_token
config = AutoConfig.from_pretrained(model_name)
with torch.device("meta"):
    self.model = AutoModelForCausalLM.from_config(config, torch_dtype=torch.float16)
self.model = self.model.eval()

ds_engine = get_ds_engine(self.model, ctx)
self.model = ds_engine.module
logger.info("Model %s loaded successfully", ctx.model_name)
self.initialized = True

def preprocess(self, requests):
    """
    Basic text preprocessing, based on the user's choice of application mode.
    Args:
        requests (list): A list of dictionaries with a "data" or "body" field, each
                       containing the input text to be processed.
    Returns:
        tuple: A tuple with two tensors: the batch of input ids and the batch of
               attention masks.
    """
    def inference(self, input_batch):
        """
        Predicts the class (or classes) of the received text using the serialized
        transformers checkpoint.
        Args:
    """
input_batch (tuple): A tuple with two tensors: the batch of input ids and the batch of attention masks, as returned by the preprocess function.

Returns:
list: A list of strings with the predicted values for each input text in the batch.

"""

def postprocess(self, inference_output):
""""Post Process Function converts the predicted response into Torchserve readable format.

Args:
inference_output (list): It contains the predicted response of the input text.

Returns:
(list): Returns a list of the Predictions and Explanations.

"""

Prepare your model artifacts

Before deploying your model on SageMaker, you must package your model artifacts. For large models, we recommend that you use the PyTorch torch-model-archiver tool with the argument --archive-format no-archive, which skips compressing model artifacts. The following example saves all of the model artifacts to a new folder named opt/.

```
torch-model-archiver --model-name opt --version 1.0 --handler custom_handler.py --extra-files ds-config.json -r requirements.txt --config-file opt/model-config.yaml --archive-format no-archive
```

Once the opt/ folder is created, download the OPT-30b model to the folder using the PyTorch Download_model tool.

```
cd opt
python path_to/Download_model.py --model_path model --model_name facebook/opt-30b --revision main
```

Lastly, upload the model artifacts to an Amazon S3 bucket.

```
aws s3 cp opt {your_s3_bucket}/opt --recursive
```
You should now have model artifacts stored in Amazon S3 that are ready to deploy to a SageMaker endpoint.

**Deploy the model using the SageMaker Python SDK**

After preparing your model artifacts, you can deploy your model to a SageMaker Hosting endpoint. This section describes how to deploy a single large model to an endpoint and make streaming response predictions. For more information about streaming responses from endpoints, see [Invoke real-time endpoints](#).

To deploy your model, complete the following steps:

1. Create a SageMaker session, as shown in the following example.

   ```python
   import boto3
   import sagemaker
   from sagemaker import Model, image_uris, serializers, deserializers

   boto3_session=boto3.session.Session(region_name="us-west-2")
   smr = boto3.client('sagemaker-runtime-demo')
   sm = boto3.client('sagemaker')
   role = sagemaker.get_execution_role()  # execution role for the endpoint
   sess = sagemaker.session.Session(boto3_session, sagemaker_client=sm,
                                    sagemaker_runtime_client=smr)  # SageMaker session for interacting with different AWS APIs
   region = sess._region_name    # region name of the current SageMaker Studio Classic environment
   account = sess.account_id()   # account_id of the current SageMaker Studio Classic environment

   # Configuration:
   bucket_name = sess.default_bucket()
   prefix = "torchserve"
   output_path = f"s3://{bucket_name}/{prefix}"
   print(f'\naccount={account}, region={region}, role={role},
output_path={output_path}')
   ```

2. Create an uncompressed model in SageMaker, as shown in the following example.

   ```python
   from datetime import datetime
   instance_type = "ml.g5.24xlarge"
   endpoint_name = sagemaker.utils.name_from_base("ts-opt-30b")
   ```
s3_uri = {your_s3_bucket}/opt

model = Model(
    name="torchserve-opt-30b" + datetime.now().strftime("%Y-%m-%d-%H-%M-%S"),
    # Enable SageMaker uncompressed model artifacts
    model_data={
        "S3DataSource": {
            "S3Uri": s3_uri,
            "S3DataType": "S3Prefix",
            "CompressionType": "None",
        }
    },
    image_uri=container,
    role=role,
    sagemaker_session=sess,
    env={"TS_INSTALL_PY_DEP_PER_MODEL": "true"},
)

print(model)

3. Deploy the model to an Amazon EC2 instance, as shown in the following example.

```python
model.deploy(
    initial_instance_count=1,
    instance_type=instance_type,
    endpoint_name=endpoint_name,
    volume_size=512, # increase the size to store large model
    model_data_download_timeout=3600, # increase the timeout to download large model
    container_startup_health_check_timeout=600, # increase the timeout to load large model
)
```

4. Initialize a class to process the streaming response, as shown in the following example.

```python
import io

class Parser:
    """
    A helper class for parsing the byte stream input.
    
    The output of the model will be in the following format:
    """
    b'"outputs": [" a"]'
```
While usually each PayloadPart event from the event stream will contain a byte array with a full json, this is not guaranteed and some of the json objects may be split across PayloadPart events. For example:
```
{'PayloadPart': {'Bytes': b'"outputs": [" challenging"]\n'}}
```
```
{'PayloadPart': {'Bytes': b'"outputs": [" problem"]\n'}}
```

This class accounts for this by concatenating bytes written via the 'write' function and then exposing a method which will return lines (ending with a '\n' character) within the buffer via the 'scan_lines' function. It maintains the position of the last read position to ensure that previous bytes are not exposed again.

```python
def __init__(self):
    self.buff = io.BytesIO()
    self.read_pos = 0

def write(self, content):
    self.buff.seek(0, io.SEEK_END)
    self.buff.write(content)
    data = self.buff.getvalue()

def scan_lines(self):
    self.buff.seek(self.read_pos)
    for line in self.buff.readlines():
        if line[-1] != b'\n':
            self.read_pos += len(line)
            yield line[:-1]

def reset(self):
    self.read_pos = 0
```

5. Test a streaming response prediction, as shown in the following example.
import json

body = "Today the weather is really nice and I am planning on".encode('utf-8')
resp = smr.invoke_endpoint_with_response_stream(EndpointName=endpoint_name,
    Body=body, ContentType="application/json")
event_stream = resp['Body']
parser = Parser()
for event in event_stream:
    parser.write(event['PayloadPart']['Bytes'])
for line in parser.scan_lines():
    print(line.decode("utf-8"), end=' ')

You have now deployed your model to a SageMaker endpoint and should be able to invoke it for responses. For more information about SageMaker real-time endpoints, see Host a single model.

Additional resources to get started

For more information about using SageMaker LMI DLCs, see the following resources:

- Blog: Improve throughput performance of Llama 2 models using Amazon SageMaker
- Blog: Improve performance of Falcon models with Amazon SageMaker
- Blog: Build an image-to-text generative AI application using multimodality models on Amazon SageMaker
- Blog: Deploy large language models on AWS Inferentia2 using large model inference containers
- Blog: Deploy BLOOM-176B and OPT-30B on Amazon SageMaker with large model inference Deep Learning Containers and DeepSpeed
- Blog: Deploy large models on Amazon SageMaker using DJL Serving and DeepSpeed model parallel inference
- Example notebooks: AWS SageMaker examples GitHub repository

Configurations and settings

The following sections describe the configuration options you can use inside the serving.properties file.
## DJL Serving general settings

The following table identifies the general DJL Serving configuration options that you can use in `serving.properties`, irrespective of whatever handler you choose.

### Common parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>engine</td>
<td>Yes</td>
<td>The runtime engine of the code. MPI is an engine that allows the model server to start distributed processes to load the model. This is used in some of the frameworks that are supported for LMI. As of 0.25.0, the TRTLLM, LMI-Dist and DeepSpeed frameworks use MPI engine. vLLM, TransformersNeuron X, Optimum Neuron, HuggingFace Accelerate uses Python engine.</td>
<td>Python, MPI, DeepSpeed (deprecated)</td>
</tr>
<tr>
<td>option.model_dir</td>
<td>No</td>
<td>The directory path to load the model. The default is set to the current path with model files. On SageMaker, this is set to the location where SageMaker is installed.</td>
<td>Default: <code>/opt/djl/ml</code></td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>-----------------------</td>
<td>----------</td>
<td>------------------------------------------------------------------</td>
<td>---------------</td>
</tr>
</tbody>
</table>
| option.model_id       | No       | The value of this option will be the Hugging Face ID of a model or the S3 URL of the model artifacts. DJL Serving will use the ID to download the model from Hugging Face or the S3 URL. DJL Serving uses s5cmd to download the model from the bucket, which is generally faster. | google/flan-t5-xl, s3://<my-bucket>/google/flan-t5-xl  
Default: None |
<p>| option.dtype          | No       | The datatype to which you plan to cast the model. The default is fp16. You can also set to bf16 if you are using G5, P4D and newer GPU machines. | fp16, fp32, bf16, int8 (only used in LMI-Dist) |</p>
<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
</table>
| option.tensor_parallel_degree | No       | The number of GPUs (model slicing number) to shard the model. If you are using LLMs, you should set this value to achieve the best performance. If you don't know what the value should be, start trying from "max" (split the model to maximum number of GPUs on the machine). | Default for DeepSpeed, Transformer-Neuron X : 1  
Default for HuggingFace Accelerate: -1  
Default for TrtLLM Container: max |
<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
</table>
| `option.rolling_batch` | No       | Also commonly known as continuous batching. Enables iteration-level batching using one of the supported strategies. This allows concurrent requests that arrived at different times to merge as a batch to run inference with the model server. | DeepSpeed container: auto, scheduler, lmi-dist, vllm, deepspeed  
Neuron container: auto  
TRTLLM container: trtllm |

Disabled for DeepSpeed container by default given there are many choices of backends.  
For the TensorRT container, rolling batch is enabled by default.  
For TransformersNeuronX, this is disabled by default.
<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
</table>
| option.max_rolling_batch_size | No       | The maximum concurrent requests or batches that the model can take. The model server will feed to the python processes `<max_rolling_batch_size>` to prevent GPU OOM. Customer can still send more requests to the model server. The requests more than `<max_rolling_batch_size>` will be queued and fed to Python until they are finished. Note: This is a model-specific configuration. If you set `<max_rolling_batch_size>` on Model A, and there are N copies of Model A inside the container. Then the model server can deal up to `<max_rolling_batch_size> x N` requests. | Default: 32 (for all engines except DeepSpeed)  
Default: 4 (for DeepSpeed) |
## Advanced parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.trust_remote_code</td>
<td>No</td>
<td>Set to <code>true</code> to use a Hugging Face Hub model with custom code. We set a default value to <code>false</code> to prevent malicious code execution from Hugging Face Hub.</td>
<td>Default: <code>false</code></td>
</tr>
<tr>
<td>option.revision</td>
<td>No</td>
<td>Use a particular version or commit hash of a Hugging Face Hub model</td>
<td>Default: None</td>
</tr>
<tr>
<td>option.entryPoint</td>
<td></td>
<td>Defines which built-in model loading handler to use. You can also use a custom model handler (model.py) in the model directory. Specify as <code>djl_python.deepspeed</code>, <code>djl_python.hugging_face</code>, <code>djl_python.transformers_neuron</code>, <code>djl_python.tensorrt_llm</code>, <code>djl_python.stable_diffusion</code> or the path to your customized handler. Different DLCs we offer choose the entry point for you. For example, if you use TensorRTL LM DLC, it uses</td>
<td>djl_python.deepspeed, djl_python.hugging_face, djl_python.transformers_neuron, djl_python.tensorrt_llm, djl_python.stable_diffusion</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>---------------------</td>
<td>----------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>option.parallel_loading</td>
<td>No</td>
<td>It loads the workers in parallel and reduces the model loading time if your model can fit in CPU memory with multiple processes. Note: If you are loading N copies of model at the same time, then the peak CPU memory can go up to N x model_size and cause CPU OOM.</td>
<td>Default: false</td>
</tr>
<tr>
<td>option.model_loading_timeout</td>
<td>No</td>
<td>Sets the limit on the time model can load before model server timeout. Default time is 30 min (1800s). If you are using on SageMaker and you do expect the model takes longer to load, you should also set container _startup_health_check_timeout=&lt;model_loading_timeout&gt; on SageMaker</td>
<td>Default: 1800</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>-------------------------</td>
<td>----------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>job_queue_size</td>
<td>No</td>
<td>Specifies the job queue size at the model level. The job queue is typically used to deal with concurrent requests that go beyond what the backend model server can take. The model server will queue requests up to &lt;job_queue_size&gt;.</td>
<td>Default: 1000</td>
</tr>
<tr>
<td>option.output_formatter</td>
<td>No</td>
<td>Use only if option.rolling_batch is enabled. Defines which output format the model server will be sent as the result. If no value being set, we are sending back JSON in tokens.</td>
<td>json, jsonlines</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default: json</td>
<td></td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>------</td>
<td>----------</td>
<td>-------------</td>
<td>--------------</td>
</tr>
<tr>
<td><code>option.enable_streaming</code> (Deprecated since 0.25.0)</td>
<td>No</td>
<td>From 0.25.0, we started to deprecate this parameter. Enables response streaming for static batching. Use <code>huggingface</code> to enable HuggingFace-like streaming output. RollingBatch already does token-streaming by default. Setting this value for RollingBatch has no effect.</td>
<td><code>false, true, huggingface</code></td>
</tr>
</tbody>
</table>

### Advanced parameters: dynamic batching

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>batch_size</code></td>
<td>No</td>
<td>Dynamic-request-level batching. This is commonly used in non-text inference to wait for requests to come to build a batch. It helps with handling concurrent requests more efficiently. Dynamic batching cannot be used with rolling batching given they</td>
<td>Default: 1</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>----------------------</td>
<td>----------</td>
<td>------------------------------------------------------------------------------</td>
<td>--------------------------------</td>
</tr>
<tr>
<td>are different batching algorithms defined by the model server.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>option.max_batch_delay</td>
<td>No</td>
<td>The maximum delay for batch aggregation in milliseconds.</td>
<td>Default: 100</td>
</tr>
<tr>
<td>option.max_idle_time</td>
<td>No</td>
<td>The maximum idle time in seconds before the worker thread is scaled down.</td>
<td>Default: 60</td>
</tr>
</tbody>
</table>

**Hugging Face handler settings**

If you use the Hugging Face handler provided by DJL Serving, you can configure the options in the following table in serving.properties.

**Common parameters**

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.device_map</td>
<td>No</td>
<td>Enables fitting the model across multiple GPUs.</td>
<td>auto, balanced, balanced_low_0, sequential</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>------</td>
<td>----------</td>
<td>-------------</td>
<td>---------------</td>
</tr>
<tr>
<td>option.low_cpu_mem_usage</td>
<td>No</td>
<td>Reduces CPU memory usage when loading models. We recommend that you set this to TRUE.</td>
<td>TRUE</td>
</tr>
<tr>
<td>option.quantize</td>
<td>No</td>
<td>Quantize the model with the supported quantization methods.</td>
<td>bitsandbytes4, bitsandbytes8</td>
</tr>
<tr>
<td>option.task</td>
<td>No</td>
<td>The task used in Hugging Face for different pipelines.</td>
<td>text-generation</td>
</tr>
</tbody>
</table>

### Advanced parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.decoding_strategy</td>
<td>No</td>
<td>Specifies the decoding method for sample, greedy and contrastive search.</td>
<td>sample, greedy, contrastive</td>
</tr>
<tr>
<td>option.disable_flash_attn</td>
<td>No</td>
<td>Used to toggle between using huggingface flash_attention or not. Note</td>
<td>Default: true</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>------</td>
<td>----------</td>
<td>-------------</td>
<td>---------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>that huggingface flash attention can be affected by padding.</td>
<td></td>
</tr>
<tr>
<td>option.load_in_4bit</td>
<td>No</td>
<td>Uses bitsandbytes quantization. Supported only on certain models.</td>
<td>Deprecated. Use option.quantize=bitsandbytes4 instead. Default: false</td>
</tr>
<tr>
<td>option.load_in_8bit</td>
<td>No</td>
<td>Uses bitsandbytes quantization. Supported only on certain models.</td>
<td>Deprecated. Use option.quantize=bitsandbytes8 instead. Default: false</td>
</tr>
<tr>
<td>option.max_sparsity</td>
<td>No</td>
<td>Used in the max-sparsity thresholding mechanism. Limits the max_sparsity in the token sequence caused by padding.</td>
<td>0.01, 0.5 Default: 0.33</td>
</tr>
<tr>
<td>option.max_splits</td>
<td>No</td>
<td>Used in the max-sparsity thresholding mechanism. Limits the max number of batch splits, where each split has its own inference call.</td>
<td>1, 5 Default: 3</td>
</tr>
</tbody>
</table>
# LMI-Dist handler settings

If you use LMI-Dist for the rolling batch option with DJL Serving, you can configure the options in the following table in `serving.properties`.

## Common parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.quantize</td>
<td>No</td>
<td>Use <code>&lt;option.quantize&gt;</code> quantize technology with the model. <code>gptq quantize</code> requires loading a <code>gptq</code> model.</td>
<td>bitsandbytes8, gptq</td>
</tr>
</tbody>
</table>

- `bitsandbytes` is deprecated. Use `bitsandbytes8` instead. Both the options are the same.

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.paged_attention</td>
<td>No</td>
<td>Use PagedAttention or not. Default is always use. Disable this if you plan to run on G4 or older GPU architecture</td>
<td>Default: true</td>
</tr>
</tbody>
</table>

## Advanced parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.max_rolling_prefill_tokens</td>
<td>No</td>
<td>Limits the number of tokens for caching. This needs to be tuned based on batch size and input sequence length to</td>
<td>Default: None</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>------</td>
<td>----------</td>
<td>-------------</td>
<td>---------------</td>
</tr>
<tr>
<td>(Deprecated since 0.25.0)</td>
<td></td>
<td>avoid GPU OOM. Currently we are calculating the best value for you. From 0.25.0, this is no longer required. If you still plan to calculate this value manually, use this formula: [(\text{max_input_tokens} + \text{max_output_tokens}) \times \text{batch_size} = \text{&lt;max_rolling_batch_prefill_tokens&gt;}.] For example, 512 input, 512 output and batch size 16, would set 16384. Setting this value does not mean you have to send input 512 and output 512. You can still do 1024 input, 512 output with batch size 10 for your request [(16384 / (1024+512)) = 10.6]</td>
<td></td>
</tr>
</tbody>
</table>

**vLLM handler settings**

If you use vLLM for the rolling batch option with DJL Serving, you can configure the options in the following table in `serving.properties`. 
<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.quantize</td>
<td>No</td>
<td>Quantize the model with the supported quantization methods</td>
<td>awq</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default: None</td>
<td></td>
</tr>
<tr>
<td>option.max_rolling</td>
<td>No</td>
<td>Limits the number of tokens for caching. This needs to be tuned based on batch size and input sequence length to avoid GPU OOM. If you don't set, vLLM will try to find a good number to fit in. If you still plan to calculate this value manually, use this formula: (max_input_tokens + max_output_tokens) * batch_size = &lt;max_rolling_batch_prefill_tokens&gt;. For example, 512 input, 512 output and batch size 16, would set 16384. Setting this value does not mean you have to send input 512 and output 512. You can still do 1024 input, 512 output.</td>
<td>Default: None</td>
</tr>
</tbody>
</table>
### DeepSpeed handler settings

If you use the DeepSpeed handler provided by DJL Serving, you can configure the options in the following table in `serving.properties`.

#### Common parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.max_tokens</td>
<td>No</td>
<td>Total number of tokens (input and output) with which DeepSpeed can work. The number of output tokens is the difference between the total number of tokens and the number of input tokens. By default we set the value to 1024. If you are looking for long sequence generation, you may want to set this to higher value, such as 2048 or 4096.</td>
<td>1024</td>
</tr>
<tr>
<td>option.quantize</td>
<td>No</td>
<td>Specify this option to quantize your model using the supported quantization methods.</td>
<td>dynamic_int8, smoothquant</td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------</td>
<td>-------------------------------------------------------------------------------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>option.methods</td>
<td></td>
<td>ion methods in DeepSpeed. SmoothQuant is our special offering to provide quantization with better quality.</td>
<td></td>
</tr>
<tr>
<td>option.task</td>
<td>No</td>
<td>The task used in Hugging Face for different pipelines. The default is <code>text-generation</code>.</td>
<td><code>text-generation</code></td>
</tr>
</tbody>
</table>

Advanced parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.checkpoint</td>
<td>No</td>
<td>Path to DeepSpeed compatible checkpoint file.</td>
<td><code>ds_inference_checkpoint.json</code></td>
</tr>
<tr>
<td>option.enable_cuda_graph</td>
<td>No</td>
<td>Activates capturing the CUDA graph of the forward pass to accelerate.</td>
<td><code>TRUE</code></td>
</tr>
<tr>
<td>option.low_cpu_mem_usage</td>
<td>No</td>
<td>Reduce CPU memory usage when loading models. We recommend that you set this to <code>TRUE</code>.</td>
<td><code>TRUE</code></td>
</tr>
<tr>
<td>option.return_tuple</td>
<td>No</td>
<td>Whether transformer layers need to return a tuple or a tensor.</td>
<td><code>FALSE</code></td>
</tr>
<tr>
<td>Item</td>
<td>Required</td>
<td>Description</td>
<td>Example value</td>
</tr>
<tr>
<td>-------------------------------------</td>
<td>----------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td>option.smoothquant_alpha</td>
<td>No</td>
<td>If smoothquant is provided in option.quantize, you can provide this alpha value. If not provided, DeepSpeed will choose one for you</td>
<td>Any float value between 0 and 1</td>
</tr>
<tr>
<td>option.training_mp_size</td>
<td>No</td>
<td>If the model was trained with DeepSpeed, this indicates the tensor parallelism degree with which the model was trained. Can be different than the tensor parallel degree desired for inference.</td>
<td>2</td>
</tr>
<tr>
<td>option.triangular_masking</td>
<td>No</td>
<td>Whether to use triangular masking for the attention mask. This is application or model specific.</td>
<td>FALSE</td>
</tr>
</tbody>
</table>

**FasterTransformer handler settings**

If you use the FasterTransformer handler provided by DJL Serving, you can configure the options in the following table in serving.properties.

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.model_dir</td>
<td>No</td>
<td>The directory path to load the model. The default is set to Default: /opt/djl/ml</td>
<td>Default: /opt/djl/ml</td>
</tr>
</tbody>
</table>
### Amazon SageMaker Developer Guide

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>the current path with model files.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>option.model_id</td>
<td>No</td>
<td>This overrides option.model_dir if set. Used to download the model from Hugging Face or an S3 bucket</td>
<td>google/flan-t5-xl, s3://my-bucket/google/flan-t5-xl</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default: None</td>
<td></td>
</tr>
</tbody>
</table>

### Neuronx LMI handler settings

If you use the Neuronx LMI handler provided by DJL Serving, you can configure the options in the following table in `serving.properties`.

#### Common parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.load_in_8bit</td>
<td>No</td>
<td>Specify this option to quantize your model using the supported quantization methods in TransformerNeuronX.</td>
<td>True, False</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default: False</td>
<td></td>
</tr>
<tr>
<td>option.n_positions</td>
<td>No</td>
<td>Maximum input plus output token length per request. You may want to set this value larger if you plan to do inference with long input or output tokens.</td>
<td>Default: 128</td>
</tr>
</tbody>
</table>
## Advanced parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.context_length_estimate</td>
<td>No</td>
<td>Estimated context input length for Llama models. Customer can specify different bucket sizes to increase the KV cache reusability. This will help to improve latency.</td>
<td>Default: None</td>
</tr>
<tr>
<td>option.load_split_model</td>
<td>No</td>
<td>Toggle to <code>True</code> when using model artifacts that have already been split for neuron compilation or loading.</td>
<td>Default: False</td>
</tr>
<tr>
<td>option.low_cpu_mem_usage</td>
<td>No</td>
<td>Reduce CPU memory usage when loading models.</td>
<td>Default: false</td>
</tr>
<tr>
<td>option.neuron_optimize_level</td>
<td>No</td>
<td>Neuron runtime compiler optimization level, determines the type of optimizations applied during compilation. The higher the optimization level, the more time spent on compilation. But in exchange, you get better latency and throughput.</td>
<td>1, 2, 3</td>
</tr>
</tbody>
</table>

**Default:** 2
### Common parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.unroll</td>
<td>No</td>
<td>Unroll the model graph for compilation. With unroll=No, the compiler will have more opportunities optimize across the layers.</td>
<td>Default: None</td>
</tr>
<tr>
<td>option.max_input_len</td>
<td>No</td>
<td>The maximum input token size you expect the model to have per request. This is a compilation parameter that sets to the model to just-in-time compilation. If you set this value too low, the model will be unable to consume the long input.</td>
<td>The Default values for Llama is 512</td>
</tr>
<tr>
<td>option.max_output_len</td>
<td>No</td>
<td>The maximum output token size you expect the model to have per request. This</td>
<td>The Default values for Llama is 512</td>
</tr>
</tbody>
</table>

### TensorRT-LLM handler settings

If you use the TensorRT-LLM handler provided by DJL Serving, you can configure the options in the following table in `serving.properties`.

---

Configurations and settings
### Item

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>is a compilation parameter that sets the model to just-in-time compilation. If you set this value too low, the model will be unable to produce tokens beyond the value you set.</td>
<td></td>
</tr>
</tbody>
</table>

| Option.use_custom_all_reduce | No | Use a custom all reduce kernel for GPUs that have NVLink enabled. This can help to speed up model inference speed with better communication. Turn this on by setting true on P4D, P4De, P5 and other GPUs that are NVLink connected. | Default: false |

---

### Advanced parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.tokens_per_block</td>
<td>No</td>
<td>Tokens per block to be used in paged attention algorithms.</td>
<td>Default: 64</td>
</tr>
</tbody>
</table>

<p>| Option.batch_scheduler_policy | No | The scheduler policy of the Tensorrt-LLM batch manager. | max_utilization, guarantee d_no_evict |</p>
<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.kv_cache_free_gpu_mem_fraction</td>
<td>No</td>
<td>Fraction of free GPU memory allocated for kv cache. The larger value you set, the more memory the model will try to take over on the GPU. The more memory preserved, the larger KV Cache size we can use and that means longer input+output sequence or larger batch size.</td>
<td>Float number between 0 and 1. Default is 0.95</td>
</tr>
</tbody>
</table>
| option.max_num_sequences                            | No       | Maximum number of input requests processed in the batch. We will apply max_rolling_batch_size as the value for it if you don't set this. Generally you don't have to touch it unless you really want the model to be compiled to a batch size that not the same as model server set. | Integer greater than 0  
Default value is the batch size set while building the TensorRT engine |
<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.enable_trt_overlap</td>
<td>No</td>
<td>Parameter to overlap the execution of batches of requests. It may have a</td>
<td>True, False.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>negative impact on performance when the number of requests is too small.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>During our experiment, we saw more negative impact to turn this on than off.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default: False</td>
<td></td>
</tr>
</tbody>
</table>

**Advanced parameters: quantization**

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>option.quantize</td>
<td>No</td>
<td>Currently only supports smoothquant for Llama models with just-in-time</td>
<td>smoothquant</td>
</tr>
<tr>
<td></td>
<td></td>
<td>compilation mode.</td>
<td></td>
</tr>
<tr>
<td>option.smoothquant_alpha</td>
<td>No</td>
<td>The smoothquant alpha parameter.</td>
<td>Default: 0.8</td>
</tr>
<tr>
<td>option.smoothquant_per_token</td>
<td>No</td>
<td>This is only applied when option.quantize is set to smoothquant. This</td>
<td>True, False.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>enables choosing, at run time, a custom</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Default: False</td>
<td></td>
</tr>
</tbody>
</table>
### Choosing instance types for large model inference

When deploying deep learning models, we typically balance the cost of hosting these models against the performance in terms of latency, throughput, and accuracy. A core input to this equation is the SageMaker instance type. SageMaker offers many instance types with different GPU devices. For a given model, there are likely multiple suitable instances for hosting the model for inference. Benchmarking can help you to decide with which instance to proceed.

<table>
<thead>
<tr>
<th>Item</th>
<th>Required</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>smoothquant scaling factor for each token</td>
<td>No</td>
<td>smoothquant scaling factor for each token. This is usually a little slower, but more accurate</td>
<td></td>
</tr>
<tr>
<td>option.smoothquant_per_channel</td>
<td>No</td>
<td>This is only applied when option.quantize is set to smoothquant. This enables choosing, at run time, a custom smoothquant scaling factor for each token. This is usually a little slower, but more accurate</td>
<td>True, False. Default: False.</td>
</tr>
<tr>
<td>option.multi_query_mode</td>
<td>No</td>
<td>This is only needed when option.quantize is set to smoothquant. This should be set for models that support multi-query-attention, such as llama-70b.</td>
<td>True, False. Default: False.</td>
</tr>
</tbody>
</table>

---

Choose a suitable instance type for hosting your large model for inference.
The following sections provide some guidelines that you can follow to determine which instance type to choose for hosting large models. To use these guidelines, you should know the following characteristics of your use case:

- Model architecture or type, such as OPT, GPTJ, BLOOM, or Bert
- Data type precision, such as fp32, fp16, bf16, or int8
- Model size in MB or GB
- Input and output token sizes

**Determining possible instance types**

When choosing an instance type, consider the size of the model as well as available GPU devices. The large model inference (LMI) containers are currently only supported on instances with NVIDIA GPUs, and are not supported on Graviton instances. For the most up-to-date information on available GPU instances, see Recommended GPU Instances.

When deploying large models, the ideal situation is to fit the model on a single GPU. This is the best option with respect to performance as it eliminates the overhead of communication between GPU devices. For some models it is simply impossible to fit them on a single GPU due to model size. For other models, they may fit on a single GPU, but it may be more cost effective to partition the model across multiple cheaper GPUs.

The following sections show how you can develop a metric to filter the list of available GPUs to ones that are likely to work for your use case.

**Determining possible instance types based on data type**

The GPUs available on SageMaker differ in their native support of data types. If you are planning to deploy your model with bf16, choose an instance with GPU devices that support compute capability 7.5+. If you are planning to deploy your model with int8, we highly recommend that you select an instance with GPU devices that support compute capability 7.5+ as these GPUs contain int8 tensor cores. However, you can still deploy with int8 using either HuggingFace Accelerate (using bitsandbytes quantization) or DeepSpeed (using ZeroQuant quantization) on lower compute capabilities. You should expect lower performance if you go this route compared to using GPUs with native support for int8 math.

To verify the compute capability of the GPU, see Your GPU Compute Capability on NVIDIA's website.
Estimating a lower bound for the memory required to host a model

To further filter the list of GPUs, determine how large your model is for the data type precision with which you wish to host the model. If the size of the model can fit on a single GPU, and low latency is the highest priority, we recommend choosing an instance with a GPU that has sufficient memory to host this model. You can estimate a lower bound for the memory required to host your model based on the number of parameters in your model and the data type. This lower bound memory estimate (LBME) represents the bare minimum memory in bytes required to load the model parameters into the GPU memory. Calculate the LBME using the following equations:

- int8 – LBME = number of parameters
- fp16 and bf16 – LBME = 2 × number of parameters
- fp32 – LBME = 4 × number of parameters

Determining possible instances based on number of partitions

Determining the level of partitioning to use with your model comes down to the following factors:

- Size of the model
- Cost you are willing to pay for an instance
- Availability of a given instance
- Your latency requirements

For example, the EleutherAI/GPT-NEOX-20b model takes about 45 GB to host in fp16. You can deploy this model using a p4de.24xlarge instance without any sharing since the GPU memory available per device is 80 GB. This is the only GPU currently on AWS that would support fitting this model fully on the device. With 8GPUs on that instance, you can host 8 copies of the model. Alternatively you can also deploy this model with 2-way partitioning on a g5.12xlarge With 4 GPUs, you can host 2 copies of the model. Using 4 g5.12xlarge instances to host 8 copies of this model compared to 1 p4de.24xlarge instance is close to half the cost (though the remaining GPU memory on the p4de.24xlarge supports larger batch sizes). While performance is likely lower on the g5.12xlarge, it might make more sense from a cost perspective.

Some instances may not be readily available in certain AWS Regions. You can check the availability of an instance with a Region/Availability Zone using the AWS CLI:
Fewer partitions generally results in lower overall latency. The best performance will come from single GPU run, but we highly recommend that you experiment with partitioning to understand the latency impact for your specific model.

You can narrow down the list of possible instances by considering the LBME you calculated in the previous step, and your desired number of partitions. If possible, consider multiple partition values to keep many instance types in consideration. For example, if the LBME is 30 GB, some estimates for different levels of partitioning are:

- Single GPU deployment on p4d.24xlarge (40 GB per GPU) or p4de.24xlarge (80 GB per GPU)
- 2 GPUs deployment on p3.8xlarge (16 GB per GPU) or g5.12xlarge (24 GB per GPU)

Choosing an engine

LMI deep learning containers (DLCs) provide support for DeepSpeed, FasterTransformer and HuggingFace Accelerate backends. All of these frameworks can be used to deploy and host large models partitioned across multiple GPUs. You can typically expect higher performance (lower latency or higher throughput) with DeepSpeed or FasterTransformer, but these engines do not offer optimized inference for all model architectures. All frameworks also implement model parallelism differently. DeepSpeed and FasterTransformer uses tensor parallelism, which is typically more performant at the cost of higher GPU memory usage. HuggingFace Accelerate uses pipeline parallelism which uses less memory but is also less performant.

We recommend that you use DeepSpeed or FasterTransformer when possible because the fused CUDA kernels significantly improve performance compared to using HuggingFace Accelerate. DeepSpeed currently offers fused kernel implementations for the following model architectures:

- Bert
- DistilBert
- GPT Neo, GPT Neo X, GPT2
- OPT
- BLOOM
FasterTransformer offers fused kernel implementations for the following model architectures:

- GPT2
- OPT
- BLOOM
- T5

For other model architectures there aren't any significant difference. We recommend that you experiment with both DeepSpeed and HuggingFace Accelerate to determine which engine works best for your model. To see which models FasterTransformer supports, see the FasterTransformer support matrix.

To choose an instance type, adjust your LBME for the specific engines. Estimating the expected GPU usage also differs between DeepSpeed, FasterTransformer and HuggingFace Accelerate due to differences in how they run model parallelism.

DeepSpeed and FasterTransformer uses tensor parallelism. Some modules, such as embeddings, do not support tensor parallelism. These modules are replicated across all GPUs. HuggingFace Accelerate uses naive pipeline partitioning, which does not result in replicated modules. Thus with DeepSpeed or FasterTransformer you can typically expect memory requirements to increase with the tensor parallelism degree.

The following sections show how you can adjust your LBME to get a loaded model memory estimate (LMME), which is an estimate of the memory required per GPU to load your model.

**HuggingFace Accelerate**

With HuggingFace Accelerate, the pipeline partitioning does not significantly increase the memory required to load the model. No sub-modules need to be duplicated across GPUs.

- For single GPU deployment, $\text{LMME} = 1.15 \times \text{LBME}$.
- For multiple GPUs deployment, $\text{LMME} = (1.15 + 0.035 \times \text{number of partitions}) \times \text{LBME}$. Here the number of partitions is assumed to be a multiple of 2.
DeepSpeed or FasterTransformer

With DeepSpeed or FasterTransformer, the tensor parallel partitioning results in some sub-modules of the model being duplicated since not every module supports tensor parallelism. The level of module replication strongly depends on the model architecture.

- For single GPU deployment, \( \text{LMME} = 1.20 \times \text{LBME} \).
- For multiple GPUs deployment, \( \text{LMME} = \text{multiplier} \times \text{LBME} \), where the multiplier strongly depends on the model architecture, and the data type. For some popular architectures, we present the recommended multipliers in the following table:

<table>
<thead>
<tr>
<th>Model architecture</th>
<th>fp32</th>
<th>fp16</th>
<th>int8</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPT variants</td>
<td>1.45</td>
<td>1.55</td>
<td>1.55</td>
</tr>
<tr>
<td>Bloom variants</td>
<td>1.5</td>
<td>1.55</td>
<td>1.55</td>
</tr>
<tr>
<td>OPT variants</td>
<td>1.55</td>
<td>1.75</td>
<td>1.75</td>
</tr>
<tr>
<td>Other models</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Adjusting loaded model memory estimate for sequence length and batch size

During inference more memory is required for the input tensors, intermediate tensors, and output tensors, than what your LMME suggests. You also need to account for additional memory required as a result of partitioning the model.

The runtime model memory estimate (RMME), defined in the following equation, accounts for the additional memory requirement based on the sequence length (total number of input and output tokens):

\[
\text{RMME per GPU} = \text{multiplier} \times \text{LMME} / \text{number of partitions}
\]

The recommended multipliers listed below are based on a sequence length of 1024. If you plan to use shorter or longer sequences you can further adjust these multipliers by multiplying them by \( \text{sequence length} / 1024 \).

- GPT variants – \( 1.1 + 0.05 \times (\text{number of partitions} - 1) \)
• BLOOM variants \(-1.15 + 0.05 \times (\text{number of partitions} - 1)\)
• OPT variants \(-1.20 + 0.1 \times (\text{number of partitions} - 1)\)

If you plan on accommodating batch sizes larger than 1, multiply the aforementioned multiplier by your expected batch size, to get the final multiplier for calculating RMME per GPU.

**Finalizing possible instance types**

With the RMME per GPU you can filter the list of available GPUs to ones that are likely to work for your use case. RMME per GPU is an intentionally cautious metric to account for the performance variance between different model architectures with different data types and engine framework versions. If you find that any instances are barely excluded by your estimate, we recommend trying them out.

**Deploying uncompressed models**

When deploying ML models, one option is to archive and compress the model artifacts into a 
.tar.gz format. Although this method works well for small models, compressing a large model artifact with hundreds of billions of parameters and then decompressing it on an endpoint can take a significant amount of time. For large model inference, we recommend that you deploy uncompressed ML model. This guide shows how you can deploy uncompressed ML model.

To deploy uncompressed ML models, upload all model artifacts to Amazon S3 and organize them under a common Amazon S3 prefix. A Amazon S3 prefix is a string of characters at the beginning of an Amazon S3 object key name, separated from the rest of the name by a delimiter. For more information on Amazon S3 prefix, see Organizing objects using prefixes.

For deploying with SageMaker, you must use slash (/) as the delimiter. You have to ensure that only artifacts associated with your ML model are organized with the prefix. For ML models with a single uncompressed artifact, the prefix will be identical to the key name. You can check which objects are associated with your prefix with the AWS CLI:

```bash
aws s3 ls --recursive s3://bucket/prefix
```

After uploading the model artifacts to Amazon S3 and organizing them under a common prefix, you can specify their location as part of the ModelDataSource field when you invoke the CreateModel request. SageMaker will automatically download the uncompressed model artifacts.
to /opt/ml/model for inference. For more information about the rules that SageMaker uses when downloading the artifacts, see S3ModelDataSource.

The following code snippet shows how you can invoke the CreateModel API when deploying an uncompressed model. Replace the italicized user text with your own information.

```python
model_name = "model-name"
sagemaker_role = "arn:aws:iam::123456789012:role/SageMakerExecutionRole"
container = "123456789012.dkr.ecr.us-west-2.amazonaws.com/inference-image:latest"

create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    PrimaryContainer = {
        "Image": container,
        "ModelDataSource": {
            "S3DataSource": {
                "S3Uri": "s3://my-bucket/prefix/to/model/data/",
                "S3DataType": "S3Prefix",
                "CompressionType": "None",
            },
        },
    },
)
```

The aforementioned example assumes that your model artifacts are organized under a common prefix. If instead your model artifact is a single uncompressed Amazon S3 object, then change "S3Uri" to point to the Amazon S3 object, and change "S3DataType" to "S3Object".

Note
---


Large model inference FAQs
---

Refer to the following FAQ items for answers to commonly asked questions about large model inference (LMI) with SageMaker.
Q: When should I use LMI DLCs?

A: Large model inference deep learning containers (LMI DLCs) include tested versions of popular model parallelization and optimization libraries for hosting models with tens or hundreds of billions of parameters. You should use an LMI DLC if your deep learning model requires multiple accelerators for hosting or if you want to accelerate inference with a popular, supported model such as GPT, Bloom, and OPT.

Q: Who can I contact if something doesn't work?

A: If, after reviewing the documentation, you continue to have problems hosting your large model, contact AWS Support.

Q: Can I use large model inference deep learning containers (LMI DLCs) outside of SageMaker?

A: AWS LMI DLCs have been tested on and designed for SageMaker, but can be used on Amazon EC2 with supported instance types.

Q: What model formats can I use with LMI DLCs?

AWS LMI DLCs include a convenience function to load a Hugging Face model format. You can bring models in another format, such as Megatron checkpoint, but you need to write logic to load that format or convert to a Hugging Face format.

Q: How can I switch between model parallelization libraries such as Hugging Face Accelerate and DeepSpeed?

If you're using the DJL handler (like in the tutorial example), then switch the default engine and handler in the serving.properties file. For example, modify the engine and option.entryPoint settings to switch from DeepSpeed to Hugging Face. In general, you can switch the engine and use the same tensor parallel degree.

Large model inference troubleshooting

If you run into any problem or error, you can try to use the following list to troubleshoot. If the problem persists, contact us at LMI-DLC-feedback@amazon.com.

Topics
- Downloading a model to my instance takes a long time
- Inference latency or throughput performance is poor
- SageMaker endpoint failed to start with a timeout or out of memory error
Downloading a model to my instance takes a long time

Consider uploading your model artifacts to Amazon S3 and mention the S3 object in `option.model_id` in your `serving.properties` configuration file. This downloading method uses `s5cmd` and speeds up data downloading. Make sure that the container has permission to access the specified S3 bucket.

Inference latency or throughput performance is poor

Even with model parallelism, large models can still have poor performance such as high latency or low throughput. First, if your model is supported, consider using the DeepSpeed engine that includes optimized kernels for inference. This can improve performance by up to 3 times. Next, consider using a more powerful instance type, such as a `p4d.24xlarge` instance. Finally, consider using a smaller model. To achieve the best performance and lowest cost, try to find the smallest model that meets the accuracy bar for your use case, as more parameters can result in higher latency, lower throughput, or higher cost. Consider fine-tuning a smaller model to improve accuracy.

SageMaker endpoint failed to start with a timeout or out of memory error

Large models can require more time to load to memory. You must adjust default endpoint configurations to handle this incremental time. If an endpoint fails a health check or is out of memory, check that you have configured your endpoint appropriately. For example, if Amazon CloudWatch Logs indicate a health check timeout, you should increase the `ContainerStartupHealthCheckTimeoutInSeconds` parameter of `ProductionVariants` during the `create_endpoint_config` step of hosting on SageMaker.

For more information on these parameters, see [SageMaker endpoint parameters for large model inference](#).

Release notes for large model inference deep learning containers

You can find a list of deep learning containers (DLCs) supported by SageMaker on the [AWS DLC GitHub repository](#). Below are the associated release notes.

Topics

- [LMI DLC Release Notes: November 27, 2023 (1)](#)
- [LMI DLC Release Notes: November 27, 2023 (2)](#)
LMI DLC Release Notes: November 27, 2023 (1)

Container URI

- 763104351884.dkr.ecr.us-east-1.amazonaws.com/djl-inference:0.25.0-deepspeed0.11.0-cu11

Highlights

- Added iteration-level batching support to DeepSpeed backend
- Upgraded to vllm 2.0 to lmi-dist backend
- Added dynamic warmup feature to lmi-dist backend

Version updates

- DJL Serving 0.25.0
- DeepSpeed 0.11.0
- HuggingFace Accelerate 0.23.0
- HuggingFace Transformers 4.34.0

Supported model architectures

- Models supported by DeepSpeed 0.11.0 using the DeepSpeed engine
- Models supported by HuggingFace Accelerate using the Accelerate engine
- Iteration-level batching support for GPT, LLaMA, LLaMA 2, Falcon, and T5 model architectures using the MPI engine
• Paged Attention and Flash Attention support for GPT, LLaMA, LLaMA 2, Falcon and T5 model architectures using the MPI engine

Known issues
• While using lmi-dist on p5 instance on Amazon SageMaker, disable aiccl by using the environment variable `ENV USE_AICCL_BACKEND=false` to avoid model server crashing on model load.

LMI DLC Release Notes: November 27, 2023 (2)

Container URI
• 763104351884.dkr.ecr.us-east-1.amazonaws.com/djl-inference:0.25.0-tensorrtllm0.5.0-cu122

Highlights
• Supports iteration-level/inflight batching of requests
• Supports response streaming
• Added support for on-the-fly TensorRT-LLM model compilation so that users can provide just Hugging Face model id for Llama and Falcon models
• Accepts user created model repositories with compiled TensorRT-LLM artifacts for faster loading

Version updates
• DJL Serving 0.25.0
• TensorRT-LLM 0.5.0
• PyTorch 2.0.1
• Python 3.10
• CUDA 12.2

Supported model architectures
• LLAMA and Falcon models with support for on-the-fly model compilation
• We have not tested other model types yet. If you want to use other than Llama and Falcon models, they should be compiled ahead of time and tested before production.

Known issues

• Using the repetition_penalty sampling parameter randomly crashes the program. It is recommended to not use repetition_penalty in this version.

LMI DLC Release Notes: November 27, 2023 (3)

Container URI

• 763104351884.dkr.ecr.us-east-1.amazonaws.com/djl-inference:0.25.0-neuronx-sdk2.15.0

Highlights

• Precompiled model support

Version updates

• DJL Serving 0.25.0
• NeuronSDK 2.15.0
• HuggingFace Accelerate 0.23.0
• HuggingFace Transformers 4.34.0

Supported model architectures

• GPT-2
• GPT-J
• GPT-NeoX
• OPT
• Bloom
• Llama
• Llama-2 (7b and 13b)
LMI DLC Release Notes: August 4, 2023 (1)

Container URI

- 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.23.0-fastertransformer5.3.0-cu118

Highlights

- Added support for serving GPT-NeoX models.
- New containers support response streaming.
- FasterTransformer handler updated to support PEFT and LORA.
- DJLServing provides integration to the Triton inference backend through the java-cpp API.

Version updates

- DJL Serving 0.23.0
- PyTorch 2.0.1
- FasterTransformer 5.3.0
- Hugging Face Accelerate 0.20.3
- Hugging Face Transformers 4.30.1
- Hugging Face Diffusers 0.12.0
- Python 3.9
- CUDA 11.8

Supported model architectures

- GPT
- GPT-J
- GPT-NeoX
- T5
- OPT
- Bloom
LMI DLC Release Notes: August 4, 2023 (2)

Container URI

- 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.23.0-deepspeed0.9.5-cu118

Highlights

- Upgrades the DeepSpeed DLC container to DeepSpeed 0.9.5.
- Improvements made to enable 2-3x memory savings over oss DeepSpeed.
- Added PyTorch-2.0.1, PEFT, and LORA support to DeepSpeed default handlers. This new release of DeepSpeed adds support for LLaMA v1 models.
- Added iteration-level batching support to the Hugging Face backend for GPT, LLaMA, Falcon, and T5 model architectures. This improves model serving throughput.

Version updates

- DJL Serving 0.23.0
- DeepSpeed 0.9.5
- Hugging Face Accelerate 0.20.3
- Hugging Face Transformers 4.30.2
- Python 3.9
- CUDA 11.8

Supported model architectures

- Models supported by DeepSpeed 0.9.5 using the DeepSpeed engine
- Models supported by HuggingFace Accelerate using the Accelerate engine
- Iteration-level batching support for GPT, LLaMA, LLaMA 2, Falcon, and T5 model architectures using the MPI engine
- Paged Attention and Flash Attention support for GPT, LLaMA, LLaMA 2, Falcon and T5 model architectures using the MPI engine
LMI DLC Release Notes: August 4, 2023 (3)

Container URI

- 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.23.0-neuronx-sdk2.12.0

Highlights

- [Experimental] Added support for GPT-NeoX models.
- [Experimental] Added support for BLOOM models.
- [Prototype] Added support for LLaMA models.
- Added support for more flexible tensor-parallel configurations to GPT2, OPT, and BLOOM.
- Added multi-query/multi-group attention support for GPT2.

Version updates

- DJL Serving 0.23.0
- Torch Neuronx 1.13.1.1.9.0
- Transformers NeuronX 0.4.149
- Hugging Face Accelerate 0.20.3
- Hugging Face Transformers 4.30.1
- Hugging Face Diffusers 0.14.0

Supported model architectures

- OPT
- GPT2
- GPT-J
- GPT-NeoX
- LLaMA
- LLaMA-2 7B and 13B
- Bloom
LMI DLC Release Notes: March 7, 2023

Container URI

- 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.21.0-fastertransformer5.3.0-cu117

Highlights

- The FasterTransformer (FT) model parallel library is now available in LMI DLCs, adding support for popular models such as flan-t5-xxl and flan-ul2. FT is an open-source library from Nvidia that provides an accelerated engine for efficiently running transformer-based neural network inference. It is designed to handle large models that require multiple GPUs and nodes in a distributed manner. The library includes an optimized version of the transformer block, which comprises both the encoder and decoder parts, enabling you to run the inference of full encoder-decoder architectures like T5, as well as encoder-only models like BERT and decoder-only models like GPT.

Version updates

- DJL Serving 0.21.0
- FasterTransformer 5.3.0
- Hugging Face Accelerate 0.16.0
- Hugging Face Transformers 4.26.0
- Hugging Face Diffusers 0.12.0
- Python 3.9
- CUDA 11.6

Known issues

- In this first release, you may see long model loading times when using the FT library. Make sure to configure health check and timeout values (see SageMaker endpoint parameters for large model inference) to allow sufficient time to load the model. In our tests, 1 hour was sufficient. Future releases will address this long model loading time with FT.
LMI DLC Release Notes: February 24, 2023

Container URI

- 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.21.0-deepspeed0.8.0-cu117

Highlights

- SageMaker Python SDK integration is new in this release, enabling no-code deployment of large language models

  Ahead-of-time partitioning support with DeepSpeed reduces endpoint start up time by separating model partitioning from runtime

Version updates

- DJL Serving 0.21.0
- DeepSpeed 0.8.0
- Hugging Face Accelerate 0.16.0
- Hugging Face Transformers 4.26.0
- Hugging Face Diffusers 0.12.0
- Python 3.9
- CUDA 11.6

LMI DLC Release Notes: December 16, 2022

Container URI

- 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.20.0-deepspeed0.7.5-cu116

Highlights
• Added BF16 precision support for models supported by DeepSpeed inference (Bloom, GPT, OPT, etc.). Training in BF16 does not require any extra transformations to the model weights before hosting with DeepSpeed.

• Added support for Hugging Face Diffusers and StableDiffusion models.

• Includes latest versions of DJL Serving, DeepSpeed, Hugging Face Transformers, and CUDA drivers, which bring stability improvement, better error handling, improved model coverage, and latency.

Version updates

• DJL Serving 0.20.0
• DeepSpeed 0.7.5
• Hugging Face Accelerate 0.13.2
• Hugging Face Transformers 4.23.1
• Hugging Face Diffusers 0.7.2
• CUDA 11.6

LMI DLC Release Notes: November 4, 2022

Highlights

• Launched LMI DLCs on AWS. SageMaker and Amazon EC2 support LMI DLCs for inference using model parallelism.

Container URI

• 763104351884.dkr.ecr.region.amazonaws.com/djl-inference:0.19.0-deepspeed0.7.3-cu113

Blogs

• For more information, see Deploy BLOOM-176B and OPT-30B on Amazon SageMaker with large model inference Deep Learning Containers and DeepSpeed.
Update models in production

Deployment guardrails are a set of model deployment options in Amazon SageMaker Inference to update your machine learning models in production. Using the fully managed deployment options, you can control the switch from the current model in production to a new one. Traffic shifting modes in blue/green deployments, such as canary and linear, give you granular control over the traffic shifting process from your current model to the new one during the course of the update. There are also built-in safeguards such as auto-rollbacks that help you catch issues early and automatically take corrective action before they significantly impact production.

Deployment guardrails provide the following benefits:

- **Deployment safety while updating production environments.** A regressive update to a production environment can cause unplanned downtime and business impact, such as increased model latency and high error rates. Deployment guardrails help you mitigate those risks by providing best practices and built-in operational safety guardrails.

- **Fully managed deployment.** SageMaker takes care of setting up and orchestrating these deployments and integrates them with endpoint update mechanisms. You do not need to build and maintain orchestration, monitoring, or rollback mechanisms. You can leverage SageMaker to set up and orchestrate these deployments and focus on leveraging ML for your applications.

- **Visibility.** You can track the progress of your deployment through the DescribeEndpoint API or through Amazon CloudWatch Events (for supported endpoints). To learn more about events in SageMaker, see the Endpoint deployment state change section in Automating Amazon SageMaker with Amazon EventBridge. Note that if your endpoint uses any of the features in the Exclusions page, you cannot use CloudWatch Events.

**Note**

Deployment guardrails only apply to Asynchronous inference and Real-time inference endpoint types.

**How to get started**

We support two types of deployments to update models in production: blue/green deployments and rolling deployments.
**Blue/Green Deployments**: You can shift traffic from your old fleet (the blue fleet) to a new fleet (green fleet) with the updates. Blue/green deployments offer multiple traffic shifting modes. A traffic shifting mode is a configuration that specifies how SageMaker routes endpoint traffic to a new fleet containing your updates. The following traffic shifting modes provide you with different levels of control over the endpoint update process:

- **All At Once Traffic Shifting** shifts all of your endpoint traffic from the blue fleet to the green fleet. Once the traffic shifts to the green fleet, your pre-specified Amazon CloudWatch alarms begin monitoring the green fleet for a set amount of time (the **baking period**). If no alarms trip during the baking period, then SageMaker terminates the blue fleet.

- **Canary Traffic Shifting** shifts one small portion of your traffic (a **canary**) to the green fleet and monitor it for a baking period. If the canary succeeds on the green fleet, then SageMaker shifts the rest of the traffic from the blue fleet to the green fleet before terminating the blue fleet.

- **Linear Traffic Shifting** provides even more customization over the number of traffic-shifting steps and the percentage of traffic to shift for each step. While canary shifting lets you shift traffic in two steps, linear shifting extends this to \( n \) linearly spaced steps.

- **Rolling Deployments**: You can update your endpoint as SageMaker incrementally provisions capacity and shifts traffic to a new fleet in steps of a batch size that you specify. Instances on the new fleet are updated with the new deployment configuration, and if no CloudWatch alarms trip during the baking period, then SageMaker cleans up instances on the old fleet. This option gives you granular control over the instance count or capacity percentage shifted during each step.

You can create and manage your deployment through the **UpdateEndpoint** and **CreateEndpoint** SageMaker API and AWS Command Line Interface commands. See the individual deployment pages for more details on how to set up your deployment. Note that if your endpoint uses any of the features listed in the **Exclusions** page, you cannot use deployment guardrails.

To follow guided examples that shows how to use deployment guardrails, see our example **Jupyter notebooks** for the canary and linear traffic shifting modes.

**Auto-Rollback Configuration and Monitoring**

Amazon CloudWatch alarms are a prerequisite for using baking periods in deployment guardrails. You can only use the auto-rollback functionality in deployment guardrails if you set up CloudWatch alarms that can monitor an endpoint. If any of your alarms trip during the specified monitoring period, SageMaker initiates a complete rollback to the old endpoint to protect your application. If
you do not have any CloudWatch alarms set up to monitor your endpoint, then the auto-rollback functionality does not work during your deployment.

To learn more about Amazon CloudWatch, see What is Amazon CloudWatch? in the Amazon CloudWatch User Guide.

Note

Ensure that your IAM execution role has permission to perform the cloudwatch:DescribeAlarms action on the auto-rollback alarms you specify.

Alarm Examples

To help you get started, we provide the following examples to demonstrate the capabilities of CloudWatch alarms. In addition to using or modifying the following examples, you can create your own alarms and configure the alarms to monitor various metrics on the specified fleets for a certain period of time. To see more SageMaker metrics and dimensions you can add to your alarms, see Monitor Amazon SageMaker with Amazon CloudWatch.

Topics

- Monitor invocation errors on both old and new fleets
- Monitor model latency on the new fleet

Monitor invocation errors on both old and new fleets

The following CloudWatch alarm monitors an endpoint's average error rate. You can use this alarm with any deployment guardrails traffic shifting type to provide overall monitoring on both the old and new fleets. If the alarm trips, then SageMaker initiates a rollback to the old fleet.

Invocation errors coming from both the old fleet and new fleet contribute to the average error rate. If the average error rate exceeds the specified threshold, then the alarm trips. This particular example monitors the 4xx errors (client errors) on both the old and new fleets for the duration of a deployment. You can also monitor the 5xx errors (server errors) by using the metric Invocation5XXErrors.
For this alarm type, if your old fleet trips the alarm during the deployment, SageMaker terminates your deployment. Therefore, if your current production fleet already causes errors, consider using or modifying one of the following examples that only monitors the new fleet for errors.

```json
#Applied deployment type: all types
{
   "AlarmName": "EndToEndDeploymentHighErrorRateAlarm",
   "AlarmDescription": "Monitors the error rate of 4xx errors",
   "MetricName": "Invocation4XXErrors",
   "Namespace": "AWS/SageMaker",
   "Statistic": "Average",
   "Dimensions": [
      {
         "Name": "EndpointName",
         "Value": "<your-endpoint-name>
      },
      {
         "Name": "VariantName",
         "Value": "AllTraffic"
      }
   ],
   "Period": 600,
   "EvaluationPeriods": 2,
   "Threshold": 1,
   "ComparisonOperator": "GreaterThanThreshold",
   "TreatMissingData": "notBreaching"
}
```

In the previous example, note the values for the following fields:

- For `AlarmName` and `AlarmDescription`, enter a name and description you choose for the alarm.
- For `MetricName`, use the value `Invocation4XXErrors` to monitor for 4xx errors on the endpoint.
- For `Namespace`, use the value `AWS/SageMaker`. You can also specify your own custom metric, if applicable.
Monitor model latency on the new fleet

The following CloudWatch alarm example monitors the new fleet's model latency during your deployment. You can use this alarm to monitor only the new fleet and exclude the old fleet. The alarm lasts for the entire deployment. This example gives you comprehensive, end-to-end monitoring of the new fleet and initiates a rollback to the old fleet if the new fleet has any response time issues.

CloudWatch publishes the metrics with the dimension EndpointConfigName:{New-Endpoint-Config} after the new fleet starts receiving traffic, and these metrics last even after the deployment is complete.

You can use the following alarm example with any deployment type.

```json
#Applied deployment type: all types
{
    "AlarmName": "NewEndpointConfigVersionHighModelLatencyAlarm",
    "AlarmDescription": "Monitors the model latency on new fleet",
    "MetricName": "ModelLatency",
    "Namespace": "AWS/SageMaker",
    "Statistic": "Average",
    "Dimensions": [
        {
            "Name": "EndpointName",
            "Value": <your-endpoint-name>
        },
        {
            "Name": "VariantName",
            "Value": "AllTraffic"
        },
        {
```
In the previous example, note the values for the following fields:

- For **MetricName**, use the value `ModelLatency` to monitor the model's response time.
- For **Namespace**, use the value `AWS/SageMaker`. You can also specify your own custom metric, if applicable.
- For the dimension **EndpointName**, use the name of the endpoint you are updating as the value.
- For the dimension **VariantName**, use the value `AllTraffic` to specify all endpoint traffic.
- For the dimension **EndpointConfigName**, the value should refer to the endpoint configuration name for your new or updated endpoint.

**Note**

If you want to monitor your old fleet instead of the new fleet, you can change the dimension **EndpointConfigName** to specify the name of your old fleet's configuration.

### Blue/Green Deployments

When you update your endpoint, Amazon SageMaker automatically uses a blue/green deployment to maximize the availability of your endpoints. In a blue/green deployment, SageMaker provisions a new fleet with the updates (the green fleet). Then, SageMaker shifts traffic from the old fleet (the blue fleet) to the green fleet. Once the green fleet operates smoothly for a set evaluation period (called the baking period), SageMaker terminates the blue fleet. With the additional capabilities in blue/green deployments, you can utilize traffic shifting modes and auto-rollback monitoring to protect your endpoint from significant production impact.

The following list describes the key features of blue/green deployments in SageMaker:
• **Traffic shifting modes.** The traffic shifting modes for deployment guardrails let you control the volume of traffic and number of traffic-shifting steps between the blue fleet and the green fleet. This capability gives you the ability to progressively evaluate the performance of the green fleet without fully committing to a 100% traffic shift.

• **Baking period.** The baking period is a set amount of time to monitor the green fleet before proceeding to the next deployment stage. If any of the pre-specified alarms trip during any baking period, then all endpoint traffic rolls back to the blue fleet. The baking period helps you to build confidence in your update before making the traffic shift permanent.

• **Auto-rollbacks.** You can specify Amazon CloudWatch alarms that SageMaker uses to monitor the green fleet. If an issue with the updated code trips any of the alarms, SageMaker initiates an auto-rollback to the blue fleet in order to maintain availability thereby minimizing risk.

### Traffic Shifting Modes

The various traffic shifting modes in blue/green deployments give you more granular control over traffic shifting between the blue fleet and the green fleet. The available traffic shifting modes for blue/green deployments are all at once, canary, and linear. The following table shows a comparison of the options.

<table>
<thead>
<tr>
<th>Name</th>
<th>What is it?</th>
<th>Pros</th>
<th>Cons</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>All at once</td>
<td>Shifts all of the traffic to the new fleet in a single step.</td>
<td>Minimizes the overall update duration.</td>
<td>Regressive updates affect 100% of the traffic.</td>
<td>Use this option to minimize update time and cost.</td>
</tr>
<tr>
<td>Canary</td>
<td>Traffic shifts in two steps. The</td>
<td>Confines the blast radius</td>
<td>Both fleets are operation</td>
<td>Use this option to balance</td>
</tr>
</tbody>
</table>

⚠️ **Important**

For blue/green deployments that involve multiple stage traffic shifting or baking periods, you are billed for both the fleets for the duration of the update, irrespective of the traffic to the fleet. This is in contrast to blue/green deployments with all at once traffic shifting and no baking periods, where you are only billed for one fleet during the course of the update.
<table>
<thead>
<tr>
<th>Name</th>
<th>What is it</th>
<th>Pros</th>
<th>Cons</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>first (canary) step</td>
<td>shifts a small portion of the traffic followed by the second step, which</td>
<td>of regressive updates to only the canary fleet.</td>
<td>al in parallel for entire deployment.</td>
<td>between minimizing the blast radius of regressive</td>
</tr>
<tr>
<td>step shifts</td>
<td>shifts the remainder of the traffic.</td>
<td></td>
<td></td>
<td>updates and minimizing the time that two fleets are</td>
</tr>
<tr>
<td>Linear</td>
<td>A fixed portion of the traffic shifts in a pre-specified number of</td>
<td>Minimizes the risk of regressive updates by shifting traffic</td>
<td>The update duration and cost are proportional to the number of steps.</td>
<td>operational.</td>
</tr>
<tr>
<td></td>
<td>equally spaced steps.</td>
<td>over several steps.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Get Started**

Once you specify your desired deployment configuration, SageMaker handles provisioning new instances, terminating old instances, and shifting traffic for you. You can create and manage your deployment through the existing `UpdateEndpoint` and `CreateEndpoint` SageMaker API and AWS Command Line Interface commands. Note that if your endpoint uses any of the features listed in the Exclusions page, you cannot use deployment guardrails. See the individual deployment pages for more details on how to set up your deployment:

- **Blue/Green Update with All At Once Traffic Shifting**
- **Blue/Green Update with Canary Traffic Shifting**
- **Blue/Green Update with Linear Traffic Shifting**

To follow guided examples that show how to use deployment guardrails, see our example Jupyter notebooks for the canary and linear traffic shifting modes.
All At Once Traffic Shifting

With all at once traffic shifting, you can quickly roll out an endpoint update using the safety guardrails of a blue/green deployment. You can use this traffic shifting option to minimize the update duration while still taking advantage of the availability guarantees of blue/green deployments. The baking period feature helps you to monitor the performance and functionality of your new instances before terminating your old instances, ensuring that your new fleet is fully operational.

The following diagram shows how all at once traffic shifting manages the old and new fleets.

When you use all at once traffic shifting, SageMaker routes 100% of the traffic to the new fleet (green fleet). Once the green fleet starts receiving traffic, the baking period begins. The baking period is a set amount of time in which pre-specified Amazon CloudWatch alarms monitor the performance of the green fleet. If no alarms trip during the baking period, SageMaker terminates the old fleet (blue fleet). If any alarms trip during the baking period, then an auto-rollback initiates and 100% of the traffic shifts back to the blue fleet.

Prerequisites

Before setting up a deployment with all at once traffic shifting, you must create Amazon CloudWatch alarms to watch metrics from your endpoint. If any of the alarms trip during the baking period, then the traffic rolls back to your blue fleet. To learn how to set up CloudWatch alarms on an endpoint, see the prerequisite page Auto-Rollback Configuration and Monitoring. To learn more about CloudWatch alarms, see Using Amazon CloudWatch alarms in the Amazon CloudWatch User Guide.

Configure All At Once Traffic Shifting

Once you are ready for your deployment and have set up CloudWatch alarms for your endpoint, you can use either the SageMaker UpdateEndpoint API or the update-endpoint command in the AWS Command Line Interface to initiate the deployment.

Topics
The following example shows how you can update your endpoint with all at once traffic shifting using `UpdateEndpoint` in the Amazon SageMaker API.

```python
import boto3
client = boto3.client("sagemaker")
response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    DeploymentConfig={
        "BlueGreenUpdatePolicy": {
            "TrafficRoutingConfiguration": {
                "Type": "ALL_AT_ONCE"
            },
            "TerminationWaitInSeconds": 600,
            "MaximumExecutionTimeoutInSeconds": 1800
        },
        "AutoRollbackConfiguration": {
            "Alarms": [
                {
                    "AlarmName": "<your-cw-alarm>"
                }
            ]
        }
    }
)
```

To configure the all at once traffic shifting option, do the following:

- For `EndpointName`, use the name of the existing endpoint you want to update.
- For `EndpointConfigName`, use the name of the endpoint configuration you want to use.
- Under `DeploymentConfig` and `BlueGreenUpdatePolicy`, in `TrafficRoutingConfiguration`, set the `Type` parameter to `ALL_AT_ONCE`. This specifies that the deployment uses the all at once traffic shifting mode.
• For **TerminationWaitInSeconds**, use 600. This parameter tells SageMaker to wait for the specified amount of time (in seconds) after your green fleet is fully active before terminating the instances in the blue fleet. In this example, SageMaker waits for 10 minutes after the final baking period before terminating the blue fleet.

• For **MaximumExecutionTimeoutInSeconds**, use 1800. This parameter sets the maximum amount of time that the deployment can run before it times out. In the preceding example, your deployment has a limit of 30 minutes to finish.

• In **AutoRollbackConfiguration**, within the **Alarms** field, you can add your CloudWatch alarms by name. Create one **AlarmName**: `<your-cw-alarm>` entry for each alarm you want to use.

---

**How to update an endpoint with an existing blue/green update policy (API)**

When you use the [CreateEndpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateEndpoint.html) API to create an endpoint, you can optionally specify a deployment configuration to reuse for future endpoint updates. You can use the same **DeploymentConfig** options as the previous UpdateEndpoint API example. There are no changes to the CreateEndpoint API behavior. Specifying the deployment configuration does not automatically perform a blue/green update on your endpoint.

The option to use a previous deployment configuration happens when using the [UpdateEndpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/API_UpdateEndpoint.html) API to update your endpoint. When updating your endpoint, you can use the **RetainDeploymentConfig** option to keep the deployment configuration you specified when you created the endpoint.

When calling the [UpdateEndpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/API_UpdateEndpoint.html) API, set **RetainDeploymentConfig** to `True` to keep the **DeploymentConfig** options from your original endpoint configuration.

```python
response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    RetainDeploymentConfig=True
)
```

**How to update an endpoint (CLI)**

If you are using the AWS CLI, the following example shows how to start a blue/green all at once deployment using the `update-endpoint` command.

```bash
update-endpoint
```

---

Blue/Green Deployments 4414
To configure the all at once traffic shifting option, do the following:

- For **endpoint-name**, use the name of the endpoint you want to update.
- For **endpoint-config-name**, use the name of the endpoint configuration you want to use.
- For **deployment-config**, use a `BlueGreenUpdatePolicy` JSON object.

---

**Note**

If you would rather save your JSON object in a file, see [Generating AWS CLI skeleton and input parameters](#) in the [AWS CLI User Guide](#).

---

### Canary Traffic Shifting

With canary traffic shifting, you can test a portion of your endpoint traffic on the new fleet while the old fleet serves the remainder of the traffic. This testing step is a safety guardrail that validates the new fleet's functionality before shifting all of your traffic to the new fleet. You still have the benefits of a blue/green deployment, and the added canary feature lets you ensure that your new (green) fleet can serve inference before letting it handle 100% of the traffic.

The portion of your green fleet that turns on to receive traffic is called the canary, and you can choose the size of this canary. Note that the canary size should be less than or equal to 50% of the new fleet's capacity. Once the baking period finishes and no pre-specified Amazon CloudWatch alarms trip, the rest of the traffic shifts from the old (blue) fleet to the green fleet. Canary traffic shifting provides you with more safety during your deployment since any issues with the updated model only impact the canary.

The following diagram shows how canary traffic shifting manages the distribution of traffic between the blue and green fleets.
Once SageMaker provisions the green fleet, SageMaker routes a portion of the incoming traffic (for example, 25%) to the canary. Then the baking period begins, during which your CloudWatch alarms monitor the performance of the green fleet. During this time, both the blue fleet and green fleet are partially active and receiving traffic. If any of the alarms trip during the baking period, then SageMaker initiates a rollback and all traffic returns to the blue fleet. If none of the alarms trip, then all of the traffic shifts to the green fleet and there is a final baking period. If the final baking period finishes without tripping any alarms, then the green fleet serves all traffic and SageMaker terminates the blue fleet.

**Prerequisites**

Before setting up a deployment with canary traffic shifting, you must create Amazon CloudWatch alarms to monitor metrics from your endpoint. The alarms are active during the baking period, and if any alarms trip, then all endpoint traffic rolls back to the blue fleet. To learn how to set up CloudWatch alarms on an endpoint, see the prerequisite page [Auto-Rollback Configuration and Monitoring](#). To learn more about CloudWatch alarms, see [Using Amazon CloudWatch alarms](#) in the *Amazon CloudWatch User Guide*. 

---

**Deployment Stages**

- **Canary - Create**
- **Canary - CanaryFlipTraffic**
- **Canary - CanaryBake**
- **Canary - FinalFlipTraffic**
- **Canary - FinalBake**
- **Canary - Cleanup**
Configure Canary Traffic Shifting

Once you are ready for your deployment and have set up Amazon CloudWatch alarms for your endpoint, you can use either the Amazon SageMaker UpdateEndpoint API or the update-endpoint command in the AWS CLI to initiate the deployment.

Topics

• How to update an endpoint (API)
• How to update an endpoint with an existing blue/green update policy (API)
• How to update an endpoint (CLI)

How to update an endpoint (API)

The following example of the UpdateEndpoint API shows how you can update an endpoint with canary traffic shifting.

```python
import boto3
client = boto3.client("sagemaker")
response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    DeploymentConfig=
    "BlueGreenUpdatePolicy": {
        "TrafficRoutingConfiguration": {
            "Type": "CANARY",
            "CanarySize": {
                "Type": "CAPACITY_PERCENT",
                "Value": 30
            },
            "WaitIntervalInSeconds": 600
        },
        "TerminationWaitInSeconds": 600,
        "MaximumExecutionTimeoutInSeconds": 1800
    },
    "AutoRollbackConfiguration": {
        "Alarms": [
            {
                "AlarmName": "<your-cw-alarm>",
        ]
    }
```

Blue/Green Deployments
To configure the canary traffic shifting option, do the following:

- For **EndpointName**, use the name of the existing endpoint you want to update.
- For **EndpointConfigName**, use the name of the endpoint configuration you want to use.
- Under **DeploymentConfig** and **BlueGreenUpdatePolicy**, in **TrafficRoutingConfiguration**, set the **Type** parameter to **CANARY**. This specifies that the deployment uses canary traffic shifting.
- In the **CanarySize** field, you can change the size of the canary by modifying the **Type** and **Value** parameters. For **Type**, use **CAPACITY_PERCENT**, meaning the percentage of your green fleet you want to use as the canary, and then set **Value** to 30. In this example, you use 30% of the green fleet's capacity as the canary. Note that the canary size should be equal to or less than 50% of the green fleet's capacity.
- For **WaitIntervalInSeconds**, use 600. The parameter tells SageMaker to wait for the specified amount of time (in seconds) between each interval shift. This interval is the duration of the canary baking period. In the preceding example, SageMaker waits for 10 minutes after the canary shift and then completes the second and final traffic shift.
- For **TerminationWaitInSeconds**, use 600. This parameter tells SageMaker to wait for the specified amount of time (in seconds) after your green fleet is fully active before terminating the instances in the blue fleet. In this example, SageMaker waits for 10 minutes after the final baking period before terminating the blue fleet.
- For **MaximumExecutionTimeoutInSeconds**, use 1800. This parameter sets the maximum amount of time that the deployment can run before it times out. In the preceding example, your deployment has a limit of 30 minutes to finish.
- In **AutoRollbackConfiguration**, within the **Alarms** field, you can add your CloudWatch alarms by name. Create one **AlarmName**: `<your-cw-alarm>` entry for each alarm you want to use.

### How to update an endpoint with an existing blue/green update policy (API)

When you use the [CreateEndpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/api-endpoints.html) API to create an endpoint, you can optionally specify a deployment configuration to reuse for future endpoint updates. You can use the same **DeploymentConfig** options as the previous UpdateEndpoint API example. There are no
changes to the CreateEndpoint API behavior. Specifying the deployment configuration does not automatically perform a blue/green update on your endpoint.

The option to use a previous deployment configuration happens when using the `UpdateEndpoint` API to update your endpoint. When updating your endpoint, you can use the `RetainDeploymentConfig` option to keep the deployment configuration you specified when you created the endpoint.

When calling the `UpdateEndpoint` API, set `RetainDeploymentConfig` to `True` to keep the DeploymentConfig options from your original endpoint configuration.

```python
response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    RetainDeploymentConfig=True
)
```

**How to update an endpoint (CLI)**

If you are using the AWS CLI, the following example shows how to start a blue/green canary deployment using the `update-endpoint` command.

```bash
update-endpoint
--endpoint-name <your-endpoint-name>
--endpoint-config-name <your-config-name>
--deployment-config "BlueGreenUpdatePolicy": {
  "TrafficRoutingConfiguration": {
    "Type": "CANARY",
    "CanarySize": {
      "Type": "CAPACITY_PERCENT",
      "Value": 30,
      "WaitIntervalInSeconds": 600
    },
    "TerminationWaitInSeconds": 600,
    "MaximumExecutionTimeoutInSeconds": 1800,
    "AutoRollbackConfiguration": {
      "Alarms": [{
        "AlarmName": "<your-alarm>
      }]
    }
  }
}
```

To configure the canary traffic shifting option, do the following:

- For `endpoint-name`, use the name of the endpoint you want to update.
- For `endpoint-config-name`, use the name of the endpoint configuration you want to use.
- For `deployment-config`, use a [BlueGreenUpdatePolicy](https://docs.aws.amazon.com/sagemaker/latest/dg/endpoint-blue-green-update.html) JSON object.
Note

If you would rather save your JSON object in a file, see Generating AWS CLI skeleton and input parameters in the AWS CLI User Guide.

Linear Traffic Shifting

Linear traffic shifting enables you to gradually shift traffic from your old fleet (blue fleet) to your new fleet (green fleet). With linear traffic shifting, you can shift traffic in multiple steps, minimizing the chance of a disruption to your endpoint. This blue/green deployment option gives you the most granular control over traffic shifting.

You can choose either the number of instances or the percentage of the green fleet's capacity to activate during each step. Each linear step should only be between 10-50% of the green fleet's capacity. For each step, there is a baking period during which your pre-specified Amazon CloudWatch alarms monitor metrics on the green fleet. Once the baking period finishes and no alarms trip, the active portion of your green fleet continues receiving traffic and a new step begins. If alarms trip during any of the baking periods, 100% of the endpoint traffic rolls back to the blue fleet.

The following diagram shows how linear traffic shifting routes traffic to the blue and green fleets.

Once SageMaker provisions the new fleet, the first portion of the green fleet turns on and receives traffic. SageMaker deactivates the same size portion of the blue fleet, and the baking period begins. If any alarms trip, all of the endpoint traffic rolls back to the blue fleet. If the baking period
finishes, then the next step begins. Another portion of the green fleet activates and receives traffic, part of the blue fleet deactivates, and another baking period begins. The same process repeats until the blue fleet is fully deactivated and the green fleet is fully active and receiving all traffic. If an alarm goes off at any point, SageMaker terminates the shifting process and 100% of the traffic rolls back to the blue fleet.

Prerequisites

Before setting up a deployment with linear traffic shifting, you must create CloudWatch alarms to monitor metrics from your endpoint. The alarms are active during the baking period, and if any alarms trip, then all endpoint traffic rolls back to the blue fleet. To learn how to set up CloudWatch alarms on an endpoint, see the prerequisite page Auto-Rollback Configuration and Monitoring. To learn more about CloudWatch alarms, see Using Amazon CloudWatch alarms in the Amazon CloudWatch User Guide.

Configure Linear Traffic Shifting

Once you are ready for your deployment and have set up CloudWatch alarms for your endpoint, you can use either the Amazon SageMaker UpdateEndpoint API or the update-endpoint command in the AWS CLI to initiate the deployment.

Topics

- How to update an endpoint (API)
- How to update an endpoint with an existing blue/green update policy (API)
- How to update an endpoint (CLI)

How to update an endpoint (API)

The following example of the UpdateEndpoint API shows how you can update an endpoint with linear traffic shifting.

```python
import boto3
client = boto3.client("sagemaker")

response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    DeploymentConfig={
        "BlueGreenUpdatePolicy": {
```
To configure the linear traffic shifting option, do the following:

- For **EndpointName**, use the name of the existing endpoint you want to update.
- For **EndpointConfigName**, use the name of the endpoint configuration you want to use.
- Under **DeploymentConfig** and **BlueGreenUpdatePolicy**, in **TrafficRoutingConfiguration**, set the **Type** parameter to **LINEAR**. This specifies that the deployment uses linear traffic shifting.
- In the **LinearStepSize** field, you can change the size of the steps by modifying the **Type** and **Value** parameters. For **Type**, use **CAPACITY_PERCENT**, meaning the percentage of your green fleet you want to use as the step size, and then set **Value** to 20. In this example, you turn on 20% of the green fleet's capacity for each traffic shifting step. Note that when customizing your linear step size, you should only use steps that are 10-50% of the green fleet's capacity.
- For **WaitIntervalInSeconds**, use 300. The parameter tells SageMaker to wait for the specified amount of time (in seconds) between each traffic shift. This interval is the duration of the baking period between each linear step. In the preceding example, SageMaker waits for 5 minutes between each traffic shift.
- For **TerminationWaitInSeconds**, use 300. This parameter tells SageMaker to wait for the specified amount of time (in seconds) after your green fleet is fully active before terminating the
instances in the blue fleet. In this example, SageMaker waits for 5 minutes after the final baking period before terminating the blue fleet.

- For `MaximumExecutionTimeoutInSeconds`, use `3600`. This parameter sets the maximum amount of time that the deployment can run before it times out. In the preceding example, your deployment has a limit of 1 hour to finish.

- In `AutoRollbackConfiguration`, within the `Alarms` field, you can add your CloudWatch alarms by name. Create one `AlarmName: <your-cw-alarm>` entry for each alarm you want to use.

**How to update an endpoint with an existing blue/green update policy (API)**

When you use the `CreateEndpoint` API to create an endpoint, you can optionally specify a deployment configuration to reuse for future endpoint updates. You can use the same `DeploymentConfig` options as the previous `UpdateEndpoint` API example. There are no changes to the `CreateEndpoint` API behavior. Specifying the deployment configuration does not automatically perform a blue/green update on your endpoint.

The option to use a previous deployment configuration happens when using the `UpdateEndpoint` API to update your endpoint. When updating your endpoint, you can use the `RetainDeploymentConfig` option to keep the deployment configuration you specified when you created the endpoint.

When calling the `UpdateEndpoint` API, set `RetainDeploymentConfig` to `True` to keep the `DeploymentConfig` options from your original endpoint configuration.

```python
response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    RetainDeploymentConfig=True
)
```

**How to update an endpoint (CLI)**

If you are using the AWS CLI, the following example shows how to start a blue/green linear deployment using the `update-endpoint` command.

```bash
update-endpoint
--endpoint-name <your-endpoint-name>
--endpoint-config-name <your-config-name>
```
To configure the linear traffic shifting option, do the following:

- For endpoint-name, use the name of the endpoint you want to update.
- For endpoint-config-name, use the name of the endpoint configuration you want to use.
- For deployment-config, use a BlueGreenUpdatePolicy JSON object.

Note

If you would rather save your JSON object in a file, see Generating AWS CLI skeleton and input parameters in the AWS CLI User Guide.

Rolling Deployments

When you update your endpoint, you can specify a rolling deployment to gradually shift traffic from your old fleet to a new fleet. You can control the size of the traffic shifting steps, as well as specify an evaluation period to monitor the new instances for issues before terminating instances from the old fleet. With rolling deployments, instances on the old fleet are cleaned up after each traffic shift to the new fleet, reducing the amount of additional instances needed to update your endpoint. This is useful especially for accelerated instances that are in high demand.

Rolling deployments gradually replace the previous deployment of your model version with the new version by updating your endpoint in configurable batch sizes. The traffic shifting behavior of rolling deployments is similar to the linear traffic shifting mode in blue/green deployments, but rolling deployments provide you with the benefit of reduced capacity requirements when compared to blue/green deployments. With rolling deployments, fewer instances are active at a time, and you have more granular control over how many instances you want to update in the new fleet. You should consider using a rolling deployment instead of a blue/green deployment if you have large models or a large endpoint with many instances.

The following list describes the key features of rolling deployments in Amazon SageMaker:
• **Baking period.** The baking period is a set amount of time to monitor the new fleet before proceeding to the next deployment stage. If any of the pre-specified alarms trip during any baking period, then all endpoint traffic rolls back to the old fleet. The baking period helps you to build confidence in your update before making the traffic shift permanent.

• **Rolling batch size.** You have granular control over the size of each batch for traffic shifting, or the number of instances you want to update in each batch. This number can range for 5–50% of the size of your fleet. You can specify the batch size as a number of instances or as the overall percentage of your fleet.

• **Auto-rollback.** You can specify Amazon CloudWatch alarms that SageMaker uses to monitor the new fleet. If an issue with the updated code trips any of the alarms, SageMaker initiates an auto-rollback to the old fleet in order to maintain availability, thereby minimizing risk.

---

**Note**

If your endpoint uses any of the features listed in the [Exclusions](#) page, you cannot use rolling deployments.

---

**How it works**

During a rolling deployment, SageMaker provides the infrastructure to shift traffic from the old fleet to the new fleet without having to provision all of the new instances at once. SageMaker uses the following steps to shift traffic:

1. SageMaker provisions the first batch of instances in the new fleet.
2. A portion of traffic is shifted from the old instances to the first batch of new instances.
3. After the baking period, if no Amazon CloudWatch alarms are tripped, then SageMaker cleans up a batch of old instances.
4. SageMaker continues to provision, shift, and clean up instances in batches until the deployment is complete.

If an alarm is tripped during one of the baking periods, then traffic is rolled back to the old fleet in batches of a size that you specify. Alternatively, you can specify the rolling deployment to shift 100% of the traffic back to the old fleet if an alarm is tripped.
The following diagram shows the progression of a successful rolling deployment, as described in the previous steps.

To create a rolling deployment, you only have to specify your desired deployment configuration. Then SageMaker handles provisioning new instances, terminating old instances, and shifting traffic for you. You can create and manage your deployment through the existing UpdateEndpoint and CreateEndpoint SageMaker API and AWS Command Line Interface commands.

Prerequisites

Before setting up a rolling deployment, you must create Amazon CloudWatch alarms to watch metrics from your endpoint. If any of the alarms trip during the baking period, then the traffic begins rolling back to your old fleet. To learn how to set up CloudWatch alarms on an endpoint, see the prerequisite page Auto-Rollback Configuration and Monitoring. To learn more about CloudWatch alarms, see Using Amazon CloudWatch alarms in the Amazon CloudWatch User Guide.

Also, review the Exclusions page to make sure that your endpoint meets the requirements for a rolling deployment.
Determine the rolling batch size

Before updating your endpoint, determine the batch size that you want to use for incrementally shifting traffic to the new fleet.

For rolling deployments, you can specify a batch size that is 5–50% of the capacity of your fleet. If you choose a large batch size, the deployment completes more quickly. However, keep in mind that the endpoint requires more capacity while updating, roughly the batch size overhead. If you choose a smaller batch size, the deployment takes longer, but you use less capacity during the deployment.

Configure a rolling deployment

Once you are ready for your deployment and have set up CloudWatch alarms for your endpoint, you can use the SageMaker UpdateEndpoint API or the update-endpoint command in the AWS Command Line Interface to initiate the deployment.

How to update an endpoint

The following example shows how you can update your endpoint with a rolling deployment using the update_endpoint method of the Boto3 SageMaker client.

To configure a rolling deployment, use the following example and fields:

- For EndpointName, use the name of the existing endpoint you want to update.
- For EndpointConfigName, use the name of the endpoint configuration you want to use.
- In the AutoRollbackConfiguration object, within the Alarms field, you can add your CloudWatch alarms by name. Create one AlarmName: <your-cw-alarm> entry for each alarm you want to use.
- Under DeploymentConfig, for the RollingUpdatePolicy object, specify the following fields:
  - MaximumExecutionTimeoutInSeconds — The time limit for the total deployment. Exceeding this limit causes a timeout. The maximum value you can specify for this field is 28800 seconds, or 8 hours.
  - WaitIntervalInSeconds — The length of the baking period, during which SageMaker monitors alarms for each batch on the new fleet.
  - MaximumBatchSize — Specify the Type of batch you want to use (either instance count or overall percentage of your fleet) and the Value, or the size of each batch.
RollbackMaximumBatchSize — Use this object to specify the rollback strategy in case an alarm trips. Specify the Type of batch you want to use (either instance count or overall percentage of your fleet), and the Value, or the size of each batch. If you don’t specify these fields, or if you set the value to 100% of your endpoint, then SageMaker uses a blue/green rollback strategy and rolls all traffic back to the old fleet when an alarm trips.

```python
import boto3
client = boto3.client("sagemaker")

response = client.update_endpoint(
    EndpointName="<your-endpoint-name>",
    EndpointConfigName="<your-config-name>",
    DeploymentConfig={
        "AutoRollbackConfiguration": {
            "Alarms": [
                {
                    "AlarmName": "<your-cw-alarm>"
                },
            ],
        },
        "RollingUpdatePolicy": {
            "MaximumExecutionTimeoutInSeconds": number,
            "WaitIntervalInSeconds": number,
            "MaximumBatchSize": {
                "Type": "INSTANCE_COUNT" | "CAPACITY_PERCENTAGE" (default),
                "Value": number
            },
            "RollbackMaximumBatchSize": {
                "Type": "INSTANCE_COUNT" | "CAPACITY_PERCENTAGE" (default),
                "Value": number
            }
        }
    }
)
```

After updating your endpoint, you might want to check the status of your rolling deployment and check the health of your endpoint. You can review your endpoint’s status in the SageMaker console, or you can review the status of your endpoint by using the `DescribeEndpoint` API.
In the `VariantStatus` object returned by the `DescribeEndpoint` API, the `Status` field tells you the current deployment or operational status of your endpoint. For more information about the possible statuses and what they mean, see `ProductionVariantStatus`.

If you attempted to do a rolling deployment and the status of your endpoint is `UpdateRollbackFailed`, see the following section for troubleshooting help.

**Failure handling**

If your rolling deployments fail and the auto-rollback fails as well, your endpoint can be left with a status of `UpdateRollbackFailed`. This status means that different endpoint configurations are deployed to the instances behind your endpoint, and your endpoint is in service with a mix of old and new endpoint configurations.

You can make another call to the `UpdateEndpoint` API to return your endpoint to a healthy state. Specify your desired endpoint configuration and deployment configuration (either as a rolling deployment, a blue/green deployment, or neither) to update your endpoint.

You can call the `DescribeEndpoint` API to check the health of your endpoint again, which is returned in the `VariantStatus` object as the `Status` field. If your update is successful, your endpoint's `Status` returns to `InService`.

**Exclusions**

When doing a blue/green or rolling deployment, your new endpoint configuration must have the same variant name as the old endpoint configuration. There are also feature-based exclusions that make your endpoint incompatible with deployment guardrails at this time. If your endpoint uses any of the following features, you cannot use deployment guardrails on your endpoint, and your endpoint will fall back to using a blue/green deployment with all at once traffic shifting and no final baking period:

- Marketplace containers
- Endpoints that use Inf1 (Inferentia-based) instances
- Amazon Elastic Inference endpoints

If you're doing a rolling deployment, there are additional feature-based exclusions:

- Serverless inference endpoints
- Multi-variant inference endpoints
Shadow tests

With Amazon SageMaker you can evaluate any changes to your model serving infrastructure by comparing its performance against the currently deployed infrastructure. This practice is known as shadow testing. Shadow testing can help you catch potential configuration errors and performance issues before they impact end users. With SageMaker, you don't need to invest in building your shadow testing infrastructure, so you can focus on model development.

You can use this capability to validate changes to any component of your production variant, namely the model, the container, or the instance, without any end user impact. It is useful in situations including but not limited to the following:

- You are considering promoting a new model that has been validated offline to production, but want to evaluate operational performance metrics such as latency and error rate before making this decision.
- You are considering changes to your serving infrastructure container, such as patching vulnerabilities or upgrading to newer versions, and want to assess the impact of these changes prior to promotion to production.
- You are considering changing your ML instance and want to evaluate how the new instance would perform with live inference requests.

The SageMaker console provides a guided experience to manage the workflow of shadow testing. You can setup shadow tests for a predefined duration of time, monitor the progress of the test through a live dashboard, clean up upon completion, and act on the results. Select a production variant you want to test against, and SageMaker automatically deploys the new variant in shadow mode and routes a copy of the inference requests to it in real time within the same endpoint. Only the responses of the production variant are returned to the calling application. You can choose to discard or log the responses of the shadow variant for offline comparison. For more information on production and shadow variants, see [Safely validate models in production](#). Note that if your endpoint uses any of the features listed in the [Exclusions](#) page, you cannot use shadow tests.

See [Create a shadow test](#) for instructions on creating a shadow test.

**Create a shadow test**

You can create a shadow test to compare the performance of a shadow variant against a production variant. You can run the test on an existing endpoint that is serving inference requests or you can create a new endpoint on which to run the test.
To create a shadow test you need to specify the following:

- A *production variant* that receives and responds to 100 percent of the incoming inference requests.
- A *shadow variant* that receives a percentage of the incoming requests, replicated from the production variant, but does not return any responses.

For each variant, you can use SageMaker to control the model, instance type, and instance count. You can configure the percentage of incoming requests, known as the traffic sampling percentage, that you want replicated to your shadow variant. SageMaker manages the replication of requests to your shadow variant and you can modify the traffic sampling percentage when your test is scheduled or running. You can also optionally turn on Data Capture to log requests and responses of your production and shadow variants.

**Note**

SageMaker supports a maximum of one shadow variant per endpoint. For an endpoint with a shadow variant, there can be a maximum of one production variant.

You can schedule the test to start at any time and continue for a specified duration. The default duration is 7 days and the maximum is 30 days. After the test is complete, the endpoint reverts to the state it was in prior to starting the test. This ensures that you do not have to manually clean up resources upon the completion of the test.

You can monitor a test that is running through a dashboard in the SageMaker console. The dashboard provides a side by side comparison of invocation metrics and instance metrics between the production and shadow variants, along with a tabular view with relevant metric statistics. This dashboard is also available for completed tests. Once you have reviewed the metrics, you can either choose to promote the shadow variant to be the new production variant or retain the existing production variant. Once you promote the shadow variant, it responds to all incoming requests. For more information, see [Promote a shadow variant](#).

The following procedure describes how to create a shadow test through the SageMaker console. There are variations in the workflow depending on whether you want to use an existing endpoint or to create a new endpoint for the shadow test.

**Topics**

Create a shadow test
Prerequisites

Before creating a shadow test with the SageMaker console, you must have a SageMaker model ready to use. For more information about how to create a SageMaker model, see Deploy models for real-time inference.

You can get started with shadow tests with an existing endpoint with a production variant and a shadow variant, an existing endpoint with only a production variant, or just the SageMaker models you'd like to compare. Shadow tests support creating an endpoint and adding variants before your test begins.

Enter shadow test details

To start creating your shadow test, fill out the Enter shadow test details page by doing the following:

1. Open the SageMaker console.
2. In the left navigation panel, choose Inference, and then choose Shadow tests.
3. Choose Create shadow test.
4. Under Name, enter a name for the test.
5. (Optional) Under Description, enter a description for the test.
6. (Optional) Specify Tags using Key and Value pairs.
7. Choose Next.

Enter shadow test settings

After filling out the Enter shadow test details page, fill out the Enter shadow test settings page. If you already have a SageMaker Inference endpoint and a production variant, follow the Use an existing endpoint workflow. If you don't already have an endpoint, follow the Create a new endpoint workflow.
Use an existing endpoint

If you want to use an existing endpoint for your test, fill out the Enter shadow test settings page by doing the following:

1. Choose a role that has the AmazonSageMakerFullAccess IAM policy attached.
2. Choose Use an existing endpoint, and then choose one of the available endpoints.
3. (Optional) To encrypt the storage volume on your endpoint, either choose an existing KMS key or choose Enter a KMS key ARN from the dropdown list under Encryption key. If you choose the second option, a field to enter the KMS key ARN appears. Enter the KMS key ARN in that field.
4. If you have multiple production variants behind that endpoint, remove the ones you don't want to use for the test. You can remove a model variant by selecting it and then choosing Remove.
5. If you do not already have a shadow variant, add a shadow variant. To add a shadow variant, do the following:
   a. Choose Add.
   b. Choose Shadow variant.
   c. In the Add model dialog box, choose the model you want to use for your shadow variant.
   d. Choose Save.
6. (Optional) In the preceding step, the shadow variant is added with the default settings. To modify these settings, select the shadow variant and choose Edit. The Edit shadow variant dialog box appears. For more information on filling out this dialog box, see Edit a shadow test.
7. In the Schedule section, enter the duration of the test by doing the following:
   a. Choose the box under Duration. A popup calendar appears.
   b. Select the start and end dates from the calendar, or enter the start and end dates in the fields for Start date and End date, respectively.
   c. (Optional) For the fields Start time and End time, enter the start and end times, respectively, in the 24 hour format.
   d. Choose Apply.
The minimum duration is 1 hour, and the maximum duration is 30 days.

8. (Optional) Turn on **Enable data capture** to save inference request and response information from your endpoint to an Amazon S3 bucket, and then enter the location of the Amazon S3 bucket.

9. Choose **Create shadow test**.

Create a new endpoint

If you don't have an existing endpoint, or you want to create a new endpoint for your test, fill out the **Enter shadow test settings** page by doing the following:

1. Choose a role that has the **AmazonSageMakerFullAccess** IAM policy attached.
2. Choose **Create a new endpoint**.
3. Under **Name**, enter a name for the endpoint.
4. Add one production variant and one shadow variant to the endpoint:
   - To add a production variant choose **Add**, and then choose **Production variant**. In the **Add model** dialog box, choose the model you want to use for your production variant, and then choose **Save**.
   - To add a shadow variant choose **Add**, and then choose **Shadow variant**. In the **Add model** dialog box, choose the model you want to use for your shadow variant, and then choose **Save**.
5. (Optional) In the preceding step, the shadow variant is added with the default settings. To modify these settings, select the shadow variant and choose **Edit**. The **Edit shadow variant** dialog box appears. For more information on filling out this dialog box, see **Edit a shadow test**.
6. In the **Schedule** section, enter the duration of the test by doing the following:
   a. Choose the box under **Duration**. A popup calendar appears.
   b. Select the start and end dates from the calendar, or enter the start and end dates under **Start date** and **End date**, respectively.
   c. (Optional) Under **Start time** and **End time**, enter the start and end times, respectively, in the 24 hour format.
   d. Choose **Apply**.
The minimum duration is 1 hour, and the maximum duration is 30 days.

7. (Optional) Turn on **Enable data capture** to save inference request and response information from your endpoint to an Amazon S3 bucket, and then enter the location of the Amazon S3 bucket.

8. Choose **Create shadow test**.

After completing the preceding procedures, you should now have a test scheduled to begin at your specified start date and time. You can view the progress of the test from a dashboard. For more information about viewing your test and the actions you can take, see [View, monitor, and edit shadow tests](#).

**View, monitor, and edit shadow tests**

You can view the statuses of your shadow tests, monitor their progress from a dashboard, and perform actions, such as starting or stopping an test early or deleting an test. The following sections show how you can view and modify your shadow tests using the SageMaker console.

**Topics**
- [View shadow tests](#)
- [Monitor a shadow test](#)
- [Start a shadow test early](#)
- [Complete a shadow test early](#)
- [Delete a shadow test](#)
- [Edit a shadow test](#)

**View shadow tests**

You can view the statuses of all of your shadow tests on the **Shadow tests** page on the SageMaker console.

To view your tests in the console, do the following:

1. Open the [SageMaker console](#).
2. In the navigation panel, choose **Inference**.
3. Choose **Shadow tests** to view the page that lists all of your shadow tests. The page should look like the following screenshot, with all the tests listed under the **Shadow test** section.

You can see the status of a test in the console on the **Shadow tests** page by checking the **Status** field for the test.

The following are the possible statuses for a test:

- **Creating** – SageMaker is creating your test.
- **Created** – SageMaker has finished creating your test, and it will begin at the scheduled time.
- **Updating** – When you make changes to your test, your test shows as updating.
- **Starting** – SageMaker is beginning your test.
- **Running** – Your test is in progress.
- **Stopping** – SageMaker is stopping your test.
- **Completed** – Your test has completed.
- **Cancelled** – When you conclude your test early, it shows as cancelled.
Monitor a shadow test

You can view the details of a shadow test and monitor it while it is in progress or after it has completed. SageMaker presents a live dashboard comparing the operational metrics like model latency, and error rate aggregated, of the production and shadow variants.

To view the details of an individual test in the console, do the following:

1. Select the test you want to monitor from the Shadow test section on the Shadow tests page.
2. From the Actions dropdown list, choose View. An overview page with the details of the test and a metrics dashboard appears.

The overview page has the following three sections.

**Summary**

This section summarizes the progress and status of the test. It also shows the summary statistics of the metric chosen from the Select metric dropdown list in the Metrics subsection. The following screenshot shows this section.

![Screenshot of the Summary section of a shadow test overview page](image_url)
In the preceding screenshot, the **Settings**, and **Details** tabs show the settings that you selected, and the details that you entered when creating the test.

**Analysis**

This section shows a metrics dashboard with separate graphs for the following metrics:

- Invocations
- InvocationsPerInstance
- ModelLatency
- Invocation4XXErrors
- Invocation5XXErrors
- InvocationModelErrors
- CPUUtilization
- MemoryUtilization
- DiskUtilization

The last three metrics monitor the model container runtime resource usage. The rest are CloudWatch metrics that you can use to analyse the performance of your variant. In general, fewer errors indicate a more stable model. A lower latency indicates either a faster model or a faster infrastructure. For more information about CloudWatch metrics, see [SageMaker Endpoint Invocation Metrics](https://docs.aws.amazon.com/sagemaker/latest/dg/endpoint-invocation-metrics.html). The following screenshot shows the metrics dashboard.
Environment

This section shows the variants that you compared in the test. If you are satisfied by the performance of the shadow variant, based on the aforementioned metrics, you can promote the shadow variant to production, by choosing **Deploy shadow variant**. For more details about deploying a shadow variant, see **Promote a shadow variant**. You can also change the traffic sampling percentage, and continue testing, by choosing **Edit traffic**. For more details about editing a shadow variant, see **Edit a shadow test**. The following screenshot shows this section.
Start a shadow test early

You can start your test before its scheduled start time. If the new duration of the test exceeds 30 days, SageMaker automatically sets the end of the test to 30 days after the new start time. This action starts the test immediately. If you want to change the start or end time of the test, see Edit a shadow test.

To immediately start your test, before its scheduled start time, through the console, do the following:

1. Select the test you want to start immediately from the Shadow test section on the Shadow tests page.
2. From the Actions dropdown list, choose Start. The Start shadow test? dialog box appears.
3. Choose Start now.

Complete a shadow test early

You can complete an in-progress test before the end of its scheduled duration. For more information see Complete a shadow test early.

Delete a shadow test

You can delete a test that you no longer need. Deleting your test only deletes the test metadata and not your endpoint, variants, or data captured in Amazon S3. If you want your endpoint to stop running, you must delete your endpoint. For more information about deleting an endpoint, see Delete Endpoints and Resources.

To delete a test through the console, do the following:
1. Select the test you want to delete from the **Shadow test** section on the **Shadow tests** page.
2. From the **Actions** dropdown list, choose **Delete**. The **Delete shadow test** dialog box appears.
3. In the **To confirm deletion, type delete in the field** text box, enter _delete_.
4. Choose **Delete**.

**Edit a shadow test**

You can modify both scheduled and in-progress tests. Before your test starts, you can change the description, the shadow variant configuration, the start date, and the end date of the test. You can also turn on or turn off data capture.

After your test starts, you can only change the description, the traffic sampling percentage for the shadow variant, and the end date.

To edit the details of your test through the console, do the following:

1. Select the test you want to edit from the **Shadow test** section on the **Shadow tests** page.
2. From the **Actions** dropdown list, choose **Edit**. The **Enter shadow test details** page appears.
3. (Optional) Under **Description**, enter a description of your test.
4. Choose **Next**. The **Enter shadow test settings** page appears.
5. (Optional) To edit your shadow variant, do the following:
   
   a. Select the shadow variant and choose **Edit**. The **Edit shadow variant** dialog box appears. If your test has already started, then you can only change the traffic sampling percentage.
   
   b. (Optional) Under **Name**, enter the new name to replace the old name.
   
   c. (Optional) Under **Traffic sample**, enter the new traffic sampling percentage to replace the old traffic sampling percentage.
   
   d. (Optional) Under **Instance type**, select the new instance type from the dropdown list.
   
   e. (Optional) Under **Instance count**, enter the new instance count to replace the old instance count.
   
   f. Choose **Apply**.

You cannot change the model in your shadow variant using the above procedure. If you want to change the model, first remove the shadow variant by selecting it and choosing **Remove**. Then add a new shadow variant.
6. (Optional) To edit the duration of the test, do the following:

   a. Choose the box under **Duration** in the **Schedule** section. A popup calendar appears.

   b. If your test is yet to start, you can change both the start and end dates. Select the new start and end dates from the calendar, or enter the new start and end dates under **Start date** and **End date**, respectively.

      If your test has already started, you can only change the end date. Enter the new end date under **End date**.

   c. (Optional) If your test is yet to start, you can change both the start and end times. Enter the new start and end times under **Start time**, and **End time**, respectively, in the 24 hour format.

      If your test has already started, you can only change the end time. Enter the new end time under **End time**, in the 24 hour format.

   d. Choose **Apply**.

7. (Optional) Turn on or turn off **Enable data capture**.

8. Choose **Update shadow test**.

**Complete a shadow test**

Your test automatically completes at the end of the scheduled duration, or you can stop an in-progress test early. After your test has completed, the test’s status in the **Shadow tests** section on the **Shadow tests** page shows as **Complete**. Then you can review and analyze the final metrics of your test.

You can use the metrics dashboard to decide whether to promote the shadow variant to production. For more information about analyzing the metrics dashboard of your test, see [Monitor a shadow test](#).

For instructions on how to complete your test before the end of its scheduled completion time, see [Complete a shadow test early](#).

For instructions on promoting your shadow variant to production, see [Promote a shadow variant](#).
Complete a shadow test early

One reason you might want to complete an in-progress shadow test is if you’ve decided that the metrics for your shadow variant look good and you want to promote it to production. You might also decide to complete the test if one or more of the variants aren’t performing well.

To complete your test before its scheduled end date, do the following:

1. Select the test you want to mark complete from the **Shadow tests** section on the **Shadow tests** page.
2. From the **Actions** dropdown list, choose **Complete**, and the **Complete shadow test** dialog box appears.
3. In the dialog box, choose one of the following options:
   - **Yes, deploy shadow variant**
   - **No, remove shadow variant**
4. (Optional) In the **Comment** text box, enter your reason for completing the test before its scheduled end time.
5. 1. If you decided to deploy the shadow variant, choose **Complete and proceed to deploy**. The **Deploy shadow variant** page appears. For instructions on how to fill out this page, see [Promote a shadow variant](#).
    2. If you decide to remove the shadow variant, choose **Confirm**.

Promote a shadow variant

If you’ve decided that you want to replace your production variant with your shadow variant, you can update your endpoint and promote your shadow variant to respond to inference requests. This removes your current production variant from production and replaces it with your shadow variant.

If your shadow test is still in-progress, you must first complete your test. To complete your shadow test before its scheduled end, follow the instructions in [Complete a shadow test early](#) before continuing with this section.

When you promote a shadow variant to production, you have the following options for the instance count of the shadow variant.
To promote your shadow variant, do the following:

1. If your test has completed, do the following:
   a. Select the test from the Shadow test section on the Shadow tests page.
   b. From the Actions dropdown list, choose View. The dashboard appears.
   c. Choose Deploy shadow variant in the Environment section. The Deploy shadow variant page appears.

   If your test has not completed, see Complete a shadow test early to complete it.

2. In the Variant settings section, select one of the following options:
   - Retain production settings
   - Retain shadow settings
   - Custom instance settings

   If you selected Custom instance settings, do the following:
   a. Select the instance type from the Instance type dropdown list.
   b. Under Instance count, enter the number of instances.

3. In Enter 'deploy' to confirm deployment text box, enter deploy.
4. Choose **Deploy shadow variant**.

Your SageMaker Inference endpoint is now using the shadow variant as your production variant, and your production variant has been removed from the endpoint.

**Best Practices**

When creating an inference experiment, keep the following information in mind:

- **Traffic sampling percentage** – Sampling 100 percent of the inference requests lets you validate that your shadow variant can handle production traffic when promoted. You may start off with a lower traffic sampling percentage and dial up as you gain confidence in your variant, but it is best practice to ensure that you’ve increased the traffic to 100 percent prior to promotion.

- **Instance type** – Unless you are using shadow variants to evaluate alternate instance types or sizes, we recommend that you use the same instance type, size, and count so that you can be certain that your shadow variant can handle the volume of inference requests after you promote it.

- **Auto scaling** – To ensure that your shadow variant can respond to spikes in the number of inference requests or changes in inference requests patterns, we highly recommend that you configure autoscaling on your shadow variants. To learn how to configure autoscaling, see [Automatically Scale Amazon SageMaker Models](#). If you have configured autoscaling, you can also validate changes to autoscaling policies without causing impact to users.

- **Metrics monitoring** – After you initiate a shadow experiment and have sufficient invocations, monitor the metrics dashboard to ensure that the metrics such as latency and error rate are within acceptable bounds. This helps you catch misconfigurations early and take corrective action. For information about how to monitor the metrics of an in-progress inference experiment, see [View, monitor, and edit shadow tests](#).

**Exclusions**

There are feature-based exclusions that make your endpoint incompatible with shadow tests at this time. If your endpoint uses any of the following features you cannot use shadow tests on your endpoint, and your request to setup shadow tests will lead to validation errors.

- Serverless inference
- Asynchronous inference
• Marketplace containers
• Multiple-containers endpoints
• Multi-model endpoints
• Endpoints that use Inf1 (Inferentia-based) instances
• Amazon Elastic Inference endpoints

Access containers through SSM

Amazon SageMaker allows you to securely connect to the Docker containers on which your models are deployed on for Inference using AWS Systems Manager (SSM). This gives you shell level access to the container so that you can debug the processes running within the container and log commands and responses with Amazon CloudWatch. You can also set up an AWS PrivateLink connection to the ML instances that host your containers for accessing the containers via SSM privately.

⚠️ Warning
Enabling SSM access can impact the performance of your endpoint. We recommend using this feature with your dev or test endpoints and not with the endpoints in production. Also, SageMaker automatically applies security patches, and replaces or terminates faulty endpoint instances within 10 minutes. However for endpoints with SSM enabled production variants, SageMaker delays security patching and replacing or terminating faulty endpoint instances by a day, to allow you to debug.

The following sections detail how you can use this feature.

Allowlist

You have to contact customer support, and get your account allowlisted, to use this feature. You cannot create an endpoint with SSM access enabled, if your account is not allow listed for this access.
Enable SSM access

To enable SSM access for an existing container on an endpoint, update the endpoint with a new endpoint configuration, with the EnableSSMAccess parameter set to true. The following example provides a sample endpoint configuration.

```
{
    "EndpointConfigName": "endpoint-config-name",
    "ProductionVariants": [
        {
            "InitialInstanceCount": 1,
            "InitialVariantWeight": 1.0,
            "InstanceType": "ml.t2.medium",
            "ModelName": "model-name",
            "VariantName": "variant-name",
            "EnableSSMAccess": true,
        }]
}
```

For more information on enabling SSM access, see EnableSSMAccess.

IAM configuration

Endpoint IAM permissions

If you have enabled SSM access for an endpoint instance, SageMaker starts and manages the SSM agent when it initiates the endpoint instance. To allow the SSM agent to communicate with the SSM services, add the following policy to the execution role that the endpoint runs under.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": ["ssmmessages:CreateControlChannel", "ssmmessages:CreateDataChannel", "ssmmessages:OpenControlChannel", "ssmmessages:OpenDataChannel"]
        }
    ]
}
```
User IAM permissions

Add the following policy to give an IAM user SSM session permissions to connect to a SSM target.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "ssm:StartSession",
                "ssm:TerminateSession"
            ],
            "Resource": "*
        }
    ]
}
```

You can restrict the endpoints that an IAM user can connect to, with the following policy. Replace the italicized placeholder text with your own information.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "ssm:StartSession",
            ],
            "Resource": [
                "sagemaker-endpoint-arn"
            ]
        }
    ]
}
```
SSM access with AWS PrivateLink

If your endpoints run within a virtual private cloud (VPC) that is not connected to the public internet, you can use AWS PrivateLink to enable SSM. AWS PrivateLink restricts all network traffic between your endpoint instances, SSM, and Amazon EC2 to the Amazon network. For more information on how to setup SSM access with AWS PrivateLink, see Set up a VPC endpoint for Session Manager.

Logging with Amazon CloudWatch Logs

For SSM access enabled endpoints, you can log errors from the SSM agent with Amazon CloudWatch Logs. For more information on how to log errors with CloudWatch Logs, see Logging session activity. The log is available at the SSM log stream, variant-name/ec2-instance-id/ssm, under the endpoint log group /aws/sagemaker/endpoints/endpoint-name. For more information on how to view the log, see View log data sent to CloudWatch Logs.

Production variants behind your endpoint can have multiple model containers. The log for each model container is recorded in the log stream. Each log is preceded by [sagemaker ssm logs] [container-name], where container-name is either the name that you gave to the container, or the default name, such as container_0, and container_1.

Accessing model containers

To access a model container on your endpoint instance, you need its target ID. The target ID is in one of the following formats:

- sagemaker-endpoint:endpoint-name_variant-name_ec2-instance-id for containers on single container endpoints
- sagemaker-endpoint:endpoint-name_variant-name_ec2-instance-id_container-name for containers on multi-container endpoints

The following example shows how you can use the AWS CLI to access a model container using its target ID.
If you enable logging, as mentioned in Logging with Amazon CloudWatch Logs, you can find the target IDs for all the containers listed at the beginning of the SSM log stream.

Note

- You cannot connect to 1P algorithm containers or containers of models obtained from SageMaker MarketPlace with SSM. However you can connect to deep learning containers (DLCs) provided by AWS or any custom container that you own.
- If you have enabled network isolation for a model container that prevents it from making outbound network calls, you cannot start an SSM session for that container.
- You can only access one container from one SSM session. To access another container, even if it is behind the same endpoint, start a new SSM session with the target ID of that endpoint.

Deploy models with model servers

The following content shows you how to deploy your models on SageMaker using popular model servers, such as TorchServe and Triton.

Deploy models with TorchServe

TorchServe is the recommended model server for PyTorch, preinstalled in the AWS PyTorch Deep Learning Container (DLC). This powerful tool offers customers a consistent and user-friendly experience, delivering high performance in deploying multiple PyTorch models across various AWS instances, including CPU, GPU, Neuron, and Graviton, regardless of the model size or distribution.

TorchServe supports a wide array of advanced features, including dynamic batching, microbatching, model A/B testing, streaming, torch XLA, tensorRT, ONNX and IPEX. Moreover, it seamlessly integrates PyTorch's large model solution, PiPPy, enabling efficient handling of large models. Additionally, TorchServe extends its support to popular open-source libraries like DeepSpeed, Accelerate, Fast Transformers, and more, expanding its capabilities even further. With
TorchServe, AWS users can confidently deploy and serve their PyTorch models, taking advantage of its versatility and optimized performance across various hardware configurations and model types. For more detailed information, you can refer to the PyTorch documentation and TorchServe on GitHub.

The following table lists the AWS PyTorch DLCs supported by TorchServe.

<table>
<thead>
<tr>
<th>Instance type</th>
<th>SageMaker PyTorch DLC link</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU and GPU</td>
<td><a href="#">SageMaker PyTorch containers</a></td>
</tr>
<tr>
<td>Neuron</td>
<td><a href="#">PyTorch Neuron containers</a></td>
</tr>
<tr>
<td>Graviton</td>
<td><a href="#">SageMaker PyTorch Graviton containers</a></td>
</tr>
</tbody>
</table>

The following sections describe the setup to build and test PyTorch DLCs on Amazon SageMaker.

**Getting started**

To get started, ensure that you have the following prerequisites:

1. Ensure that you have access to an AWS account. Set up your environment so that the AWS CLI can access your account through either an AWS IAM user or an IAM role. We recommend using an IAM role. For the purposes of testing in your personal account, you can attach the following managed permissions policies to the IAM role:
   - [AmazonEC2ContainerRegistryFullAccess](#)
   - [AmazonEC2FullAccess](#)
   - [AWSServiceRoleForAmazonEKSNodegroup](#)
   - [AmazonSageMakerFullAccess](#)
   - [AmazonS3FullAccess](#)
2. Locally configure your dependencies, as shown in the following example:

```python
from datetime import datetime
import os
import json
import logging
import time
```
3. Retrieve the PyTorch DLC image, as shown in the following example.

SageMaker PyTorch DLC images are available in all AWS regions. For more information, see the list of DLC container images.

```python
baseimage = sagemaker.image_uris.retrieve(  
    framework="pytorch",  
    region="<region>" ,  
    py_version="py310",  
    image_scope="inference",  
    version="2.0.1",  
    instance_type="ml.g4dn.16xlarge",  
)
```

4. Create a local workspace.

```bash
mkdir -p workspace/
```

**Adding a package**

The following sections describe how to add and preinstall packages to your PyTorch DLC image.

**BYOC use cases**
The following steps outline how to add a package to your PyTorch DLC image. For more information about customizing your container, see Building AWS Deep Learning Containers Custom Images.

1. Suppose you want to add a package to the PyTorch DLC docker image. Create a Dockerfile under the `docker` directory, as shown in the following example:

```
mkdir -p workspace/docker
  cat workspace/docker/Dockerfile

ARG BASE_IMAGE
FROM $BASE_IMAGE

# Install any additional libraries
RUN pip install transformers==4.28.1
```

2. Build and publish the customized docker image by using the following `build_and_push.sh` script.

```
# Download script build_and_push.sh to workspace/docker
  ls workspace/docker
  build_and_push.sh  Dockerfile

  # Build and publish your docker image
  reponame = "torchserve"
  versiontag = "demo-0.1"

  ./build_and_push.sh {reponame} {versiontag} {baseimage} {region} {account}
```

**SageMaker preinstall use cases**

The following example shows you how to preinstall a package to your PyTorch DLC container. You must create a `requirements.txt` file locally under the directory `workspace/code`.

```
mkdir -p workspace/code
  cat workspace/code/requirements.txt

  transformers==4.28.1
```
Create TorchServe model artifacts

In the following example, we use the pre-trained **MNIST model**. We create a directory `workspace/mnist`, implement `mnist_handler.py` by following the **TorchServe custom service instructions**, and configure the model parameters (such as batch size and workers) in `model-config.yaml`. Then, we use the TorchServe tool `torch-model-archiver` to build the model artifacts and upload to Amazon S3.

1. Configure the model parameters in `model-config.yaml`.

   ```bash
   ls -al workspace/mnist-dev
   
mnist.py
   mnist_handler.py
   mnist_cnn.pt
   model-config.yaml
   
   # config the model
   cat workspace/mnist-dev/model-config.yaml
   minWorkers: 1
   maxWorkers: 1
   batchSize: 4
   maxBatchDelay: 200
   responseTimeout: 300
   ```

2. Build the model artifacts by using `torch-model-archiver`.

   ```bash
   
   cd workspace
   torch-model-archiver --model-name mnist --version 1.0 --model-file mnist-dev/mnist.py --serialized-file mnist-dev/mnist_cnn.pt --handler mnist-dev/mnist_handler.py --config-file mnist-dev/model-config.yaml --archive-format no-archive
   
   cd mnist
   mv ../code .
   ```

Deploy models with TorchServe
3. Upload `mnist.tar.gz` to Amazon S3.

```bash
# upload mnist.tar.gz to S3
output_path = f"s3://{bucket_name}/{prefix}/models"
aws s3 cp mnist.tar.gz {output_path}/mnist.tar.gz
```

### Using single model endpoints to deploy with TorchServe

The following example shows you how to create a **single model real-time inference endpoint**, deploy the model to the endpoint, and test the endpoint by using the [Amazon SageMaker Python SDK](https://github.com/aws/sagemaker-python-sdk).

```python
from sagemaker.model import Model
from sagemaker.predictor import Predictor

# create the single model endpoint and deploy it on SageMaker
model = Model(model_data = f'{output_path}/mnist.tar.gz',
               image_uri = baseimage,
               role = role,
               predictor_cls = Predictor,
               name = "mnist",
               sagemaker_session = smsess)

endpoint_name = 'torchserve-endpoint-' + time.strftime("%Y-%m-%d-%H-%M-%S",
              time.gmtime())
predictor = model.deploy(instance_type='ml.g4dn.xlarge',
                          initial_instance_count=1,
                          endpoint_name = endpoint_name,
                          serializer=JSONSerializer(),
                          deserializer=JSONDeserializer())

# test the endpoint
import random
import numpy as np
dummy_data = {"inputs": np.random.rand(16, 1, 28, 28).tolist()}
res = predictor.predict(dummy_data)
```
Multi-model endpoints are a scalable and cost-effective solution to hosting large numbers of models behind one endpoint. They improve endpoint utilization by sharing the same fleet of resources and serving container to host all of your models. They also reduce deployment overhead because SageMaker manages dynamically loading and unloading models, as well as scaling resources based on traffic patterns. Multi-model endpoints are particularly useful for deep learning and generative AI models that require accelerated compute power.

By using TorchServe on SageMaker multi-model endpoints, you can speed up your development by using a serving stack that you are familiar with while leveraging the resource sharing and simplified model management that SageMaker multi-model endpoints provide.

The following example shows you how to create a multi-model endpoint, deploy the model to the endpoint, and test the endpoint by using the Amazon SageMaker Python SDK. Additional details can be found in this notebook example.

```python
from sagemaker.multidatamodel import MultiDataModel
    from sagemaker.model import Model
    from sagemaker.predictor import Predictor

    # create the single model endpoint and deploy it on SageMaker
    model = Model(model_data = f'{output_path}/mnist.tar.gz',
        image_uri = baseimage,
        role = role,
        sagemaker_session = smsess)

    endpoint_name = 'torchserve-endpoint-' + time.strftime("%Y-%m-%d-%H-%M-%S",
    time.gmtime())
    mme = MultiDataModel(
        name = endpoint_name,
        model_data_prefix = output_path,
        model = model,
        sagemaker_session = smsess)

    mme.deploy(
        initial_instance_count = 1,
        instance_type = "ml.g4dn.xlarge",
        serializer=sagemaker.serializers.JSONSerializer(),
        deserializer=sagemaker.deserializers.JSONDeserializer())

    # list models
```
list(mme.list_models())

# create mnist v2 model artifacts
cp mnist.tar.gz mnistv2.tar.gz

# add mnistv2
mme.add_model(mnistv2.tar.gz)

# list models
list(mme.list_models())

predictor = Predictor(endpoint_name=mme.endpoint_name, sagemaker_session=smsess)

# test the endpoint
import random
import numpy as np
dummy_data = {"inputs": np.random.rand(16, 1, 28, 28).tolist()}
res = predictor.predict(date=dummy_data, target_model="mnist.tar.gz")

## Metrics

TorchServe supports both system level and model level metrics. You can enable metrics in either log format mode or Prometheus mode through the environment variable TS_METRICS_MODE. You can use the TorchServe central metrics config file metrics.yaml to specify the types of metrics to be tracked, such as request counts, latency, memory usage, GPU utilization, and more. By referring to this file, you can gain insights into the performance and health of the deployed models and effectively monitor the TorchServe server's behavior in real-time. For more detailed information, see the [TorchServe metrics documentation](https://docs.aws.amazon.com/sagemaker/latest/dg/TorchServe-model-serving.html).

You can access TorchServe metrics logs that are similar to the StatsD format through the Amazon CloudWatch log filter. The following is an example of a TorchServe metrics log:

```
CPUUtilization.Percent:0.0|#Level:Host|#hostname:my_machine_name,timestamp:1682098185
DiskAvailable.Gigabytes:318.0416717529297|#Level:Host|
#hostname:my_machine_name,timestamp:1682098185
```
Deploy models with DJL Serving

DJL Serving is a high performance universal stand-alone model serving solution. It takes a deep learning model, several models, or workflows and makes them available through an HTTP endpoint.

You can use one of the DJL Serving Deep Learning Containers (DLCs) to serve your models on AWS. To learn about the supported model types and frameworks, see the DJL Serving GitHub repository.

DJL Serving offers many features that help you to deploy your models with high performance:

- **Ease of use** – DJL Serving can serve most models without any modifications. You bring your model artifacts, and DJL Serving can host them.
- **Multiple device and accelerator support** – DJL Serving supports deploying models on CPUs, GPUs, and AWS Inferentia.
- **Performance** – DJL Serving runs multithreaded inference in a single Java virtual machine (JVM) to boost throughput.
- **Dynamic batching** – DJL Serving supports dynamic batching to increase throughput.
- **Auto scaling** – DJL Serving automatically scales workers up or down based on the traffic load.
- **Multi-engine support** – DJL Serving can simultaneously host models using different frameworks (for example, PyTorch and TensorFlow).
- **Ensemble and workflow models** – DJL Serving supports deploying complex workflows comprised of multiple models and can execute parts of the workflow on CPUs and other parts on GPUs. Models within a workflow can leverage different frameworks.

The following sections describe how to set up an endpoint with DJL Serving on SageMaker.

**Getting started**

To get started, ensure that you have the following prerequisites:

1. Ensure that you have access to an AWS account. Set up your environment so that the AWS CLI can access your account through either an AWS IAM user or an IAM role. We recommend using an IAM role. For the purposes of testing in your personal account, you can attach the following managed permissions policies to the IAM role:
   - AmazonEC2ContainerRegistryFullAccess
   - AmazonEC2FullAccess
2. Ensure that you have the `docker` client set up on your system.

3. Log in to Amazon Elastic Container Registry and set the following environment variables:

```bash
export ACCOUNT_ID=<your_account_id>
export REGION=<your_region>
aws ecr get-login-password --region $REGION | docker login --username AWS --password-stdin $ACCOUNT_ID.dkr.ecr.$REGION.amazonaws.com
```

4. Pull the docker image.

```bash
docker pull 763104351884.dkr.ecr.us-west-2.amazonaws.com/djl-inference:0.22.1-deepspeed0.9.2-cu118
```

For all of the available DJL Serving container images, see the large model inference containers and the DJL Serving CPU inference containers. When choosing an image from the tables in the preceding links, replace the AWS region in the example URL column with the region you are in. The DLCs are available in the regions listed in the table at the top of the Available Deep Learning Containers Images page.

**Customize your container**

You can add packages to the base DLC images to customize your container. Suppose you want to add a package to the `763104351884.dkr.ecr.us-west-2.amazonaws.com/djl-inference:0.22.1-deepspeed0.9.2-cu118` docker image. You must create a dockerfile with your desired image as the base image, add the required packages, and push the image to Amazon ECR.

To add a package, complete the following steps:

1. Specify instructions for running your desired libraries or packages in the base image's dockerfile.

```bash
FROM 763104351884.dkr.ecr.us-west-2.amazonaws.com/djl-inference:0.22.1-deepspeed0.9.2-cu118
## add custom packages/libraries
```
2. Build the Docker image from the dockerfile. Specify your Amazon ECR repository, the name of the base image, and a tag for the image. If you don't have an Amazon ECR repository, see [Using Amazon ECR with the AWS CLI](https://docs.aws.amazon.com/AmazonECR/latest/userguide/using-ecr-with-aws-cli.html) in the *Amazon ECR User Guide* for instructions on how to create one.

```
docker build -f Dockerfile -t <registry>/<image_name>:<image_tag>
```

3. Push the Docker image to your Amazon ECR repository.

```
docker push $ACCOUNT_ID.dkr.ecr.$REGION.amazonaws.com/<image_name>:<image_tag>
```

You should now have a customized container image that you can use for model serving. For more examples of customizing your container, see [Building AWS Deep Learning Containers Custom Images](https://docs.aws.amazon.com/djl/userguide/lambda.html).

### Prepare your model artifacts

Before deploying your model on SageMaker, you must package your model artifacts in a `.tar.gz` file. DJL Serving accepts the following artifacts in your archive:

- **Model checkpoint**: Files that store your model weights.
- **serving.properties**: A configuration file that you can add for each model. Place `serving.properties` in the same directory as your model file.
- **model.py**: The inference handler code. This is only applicable when using Python mode. If you don't specify `model.py`, djl-serving uses one of the default handlers.

The following is an example of a `model.tar.gz` structure:

```
- model_root_dir # root directory
  - serving.properties
  - model.py # your custom handler file for Python, if you choose not to use the default handlers provided by DJL Serving
  - model binary files # used for Java mode, or if you don't want to use option.model_id and option.s3_url for Python mode
```
DJL Serving supports Java engines powered by DJL or Python engines. Not all of the preceding artifacts are required; the required artifacts vary based on the mode you choose. For example, in Python mode, you only need to specify `option.model_id` in the `serving.properties` file; you don't need to specify the model checkpoint inside LMI containers. In Java mode, you are required to package the model checkpoint. For more details on how to configure `serving.properties` and operate with different engines, see [DJL Serving Operation Modes](#).

### Use single model endpoints to deploy with DJL Serving

After preparing your model artifacts, you can deploy your model to a SageMaker endpoint. This section describes how to deploy a single model to an endpoint with DJL Serving. If you're deploying multiple models, skip this section and go to [Use multi-model endpoints to deploy with DJL Serving](#).

The following example shows you a method to create a model object using the Amazon SageMaker Python SDK. You'll need to specify the following fields:

- **image_uri**: You can either retrieve one of the base DJL Serving images as shown in this example, or you can specify a custom Docker image from your Amazon ECR repository, if you followed the instructions in [Customize your container](#).
- **model_s3_url**: This should an Amazon S3 URI that points to your `.tar.gz` file.
- **model_name**: Specify a name for the model object.

```python
import boto3
import sagemaker
from sagemaker.model import Model
from sagemaker import image_uris, get_execution_role

aws_region = "aws-region"
sagemaker_session =
  sagemaker.Session(boto_session=boto3.Session(region_name=aws_region))
role = get_execution_role()

def create_model(model_name, model_s3_url):
    # Get the DJL DeepSpeed image uri
    image_uri = image_uris.retrieve(
        framework="djl-deepspeed",
        region=sagemaker_session.boto_session.region_name,
        version="0.20.0"
    )
```

Deploy models with DJL Serving
model = Model(
    image_uri=image_uri,
    model_data=model_s3_url,
    role=role,
    name=model_name,
    sagemaker_session=sagemaker_session,
)
return model

After you create the model object, you can deploy the model to an endpoint and make predictions. For an end-to-end example of how to perform these steps, see the tutorial Large model inference with DeepSpeed and DJL Serving.

**Use multi-model endpoints to deploy with DJL Serving**

If you want to deploy multiple models to an endpoint, SageMaker offers multi-model endpoints, which are a scalable and cost-effective solution to deploying large numbers of models. DJL Serving also supports loading multiple models simultaneously and running inference on each of the models concurrently. DJL Serving containers adhere to the SageMaker multi-model endpoints contracts and can be used to deploy multi-model endpoints.

Each individual model artifact needs to be packaged in the same way as described in the previous section Prepare your model artifacts. You can set model-specific configurations in the serving.properties file and model-specific inference handler code in model.py. For a multi-model endpoint, models need to be arranged in the following way:

```
root_dir
|-- model_1.tar.gz
|-- model_2.tar.gz
|-- model_3.tar.gz
   ...
   ...
```

The Amazon SageMaker Python SDK uses the MultiDataModel object to instantiate a multi-model endpoint. The Amazon S3 URI for the root directory should be passed as the model_data_prefix argument to the MultiDataModel constructor.

DJL Serving also provides several configuration parameters to manage model memory requirements, such as required_memory_mb and reserved_memory_mb, that can be configured
for each model in the `serving.properties` file. These parameters are useful to handle out of memory errors more gracefully. For all of the configurable parameters, see `OutofMemory handling in djl-serving`.

The auto scaling feature of DJL Serving makes it easy to ensure that the models are scaled appropriately for incoming traffic. By default, DJL Serving determines the maximum number of workers for a model that can be supported based on the hardware available (such as CPU cores or GPU devices). You can set lower and upper bounds for each model to ensure that a minimum traffic level can always be served, and that a single model does not consume all available resources. You can set the following properties in the `serving.properties` file:

- `gpu.minWorkers`: Minimum number of workers for GPUs.
- `gpu.maxWorkers`: Maximum number of workers for CPUs.
- `cpu.minWorkers`: Minimum number of workers for CPUs.
- `cpu.maxWorkers`: Maximum number of workers for CPUs.

For an end-to-end example of how to deploy a multi-model endpoint on SageMaker using a DJL Serving container, see the example notebook `Multi-Model-Inference-Demo.ipynb`.

**Deploy models with Triton Inference Server**

Triton Inference Server is an open source inference serving software that streamlines AI inference. With Triton, you can deploy any model built with multiple deep learning and machine learning frameworks, including TensorRT, TensorFlow, PyTorch, ONNX, OpenVINO, Python, RAPIDS FIL, and more.

The SageMaker Triton containers help you deploy Triton Inference Server on the SageMaker Hosting platform to serve trained models in production. It supports the different modes in which SageMaker operates. For a list of available Triton Inference Server containers available on SageMaker, see `NVIDIA Triton Inference Containers (SM support only)`.

For end-to-end notebook examples, we recommend taking a look at the `amazon-sagemaker-examples repository`.

**Hosting modes**

The following SageMaker Hosting modes are supported by Triton containers:
• Single model endpoints
  • This is SageMaker's default mode of operation. In this mode, the Triton container can load a single model, or a single ensemble model.
  • The name of the model must be passed as a property of the container environment, which is part of the CreateModel SageMaker API call. The environment variable used to pass in the model name is SAGEMAKER_TRITON_DEFAULT_MODEL_NAME.

• Single model endpoints with ensemble
  • Triton Inference Server supports ensemble, which is a pipeline, or a DAG (directed acyclic graph) of models. While an ensemble technically comprises of multiple models, in the default single model endpoint mode, SageMaker can treat the ensemble proper (the meta-model that represents the pipeline) as the main model to load, and can subsequently load the associated models.
  • The ensemble proper's model name must be used to load the model. It must be passed as a property of the container environment, which is part of the CreateModel SageMaker API call. The environment variable used to pass in the model name is SAGEMAKER_TRITON_DEFAULT_MODEL_NAME.

• Multi-model endpoints
  • In this mode, SageMaker can serve multiple models on a single endpoint. You can use this mode by specifying the environment variable ‘MultiModel’: true as a property of the container environment, which is part of the CreateModel SageMaker API call.
  • By default, no model is loaded when the instance starts. To run an inference request against a particular model, specify the corresponding model's *.tar.gz file as an argument to the TargetModel property of the InvokeEndpoint SageMaker API call.

• Multi-model endpoints with ensemble
  • In this mode, SageMaker functions as described for multi-model endpoints. However, the SageMaker Triton container can load multiple ensemble models, meaning that multiple model pipelines can run on the same instance. SageMaker treats every ensemble as one model, and the ensemble proper of each model can be invoked by specifying the corresponding *.tar.gz archive as the TargetModel.
  • For better memory management during dynamic memory LOAD and UNLOAD, we recommend that you keep the ensemble size small.
Inference payload types

Triton supports two methods of sending an inference payload over the network – json and binary+json (or binary encoded json). The JSON payload in both cases includes the datatype, shape and the actual inference request tensor. The request tensor must be a binary tensor.

With the binary+json format, you must specify the length of the request metadata in the header to allow Triton to correctly parse the binary payload. In the SageMaker Triton container, this is done using a custom Content-Type header: application/vnd.sagemaker-triton.binary+json;json-header-size={}. This is different from using the Inference-Header-Content-Length header on a stand-alone Triton Inference Server because custom headers are not allowed in SageMaker.

Using config.pbtxt to set the model config

For Triton Inference Servers on SageMaker, each model must include a config.pbtxt file that specifies, at a minimum, the following configurations for the model:

- **name**: While this is optional for models running outside of SageMaker, we recommend that you always provide a name for the models to be run in Triton on SageMaker.
- **platform and/or backend**: Setting a backend is essential to specify the type of the model. Some backends have further classification, such as tensorflow_savedmodel or tensorflow_graphdef. Such options can be specified as part of the platform key in addition to the backend key. The most common backends are tensorrt, onnxruntime, tensorflow, pytorch, python, dali, fil, and openvino.
- **input**: Specify three attributes for the input: name, data_type and dims (the shape).
- **output**: Specify three attributes for the output: name, data_type and dims (the shape).
- **max_batch_size**: Set the batch size to a value greater than or equal to 1 that indicates the maximum batch size that Triton should use with the model.

For more details on configuring config.pbtxt, see Triton’s GitHub repository. Triton provides several configurations for tweaking model behavior. Some of the most common and important configuration options are:

- **instance_groups**: Instance groups help with specifying the number and location for a given model. They have the attributes count, kind, and gpus (used when kind is KIND_GPU). The count attribute is equivalent to the number of workers. For regular model serving, each worker
has its own copy of the model. Similarly, in Triton, the **count** specifies the number of model copies per device. For example, if the **instance_group** type is KIND_CPU, then the CPU has count number of model copies.

**Note**

On a GPU instance, the **instance_group** configuration applies per GPU device. For example, count number of model copies are placed on each GPU device unless you explicitly specify which GPU devices should load the model.

- **dynamic_batching** and **sequence_batching**: Dynamic batching is used for stateless models, and sequence batching is used for stateful models (where you want to route a request to the same model instance every time). Batching schedulers enable a per-model queue, which help in increasing throughput, depending on the batching configuration.

- **ensemble**: An ensemble model represents a pipeline of one or more models and the connection of input and output tensors between those models. It can be configured by specifying **platform** as ensemble. The ensemble configuration is just a representation of the model pipeline. On SageMaker, all the models under an ensemble are treated as dependents of the ensemble model and are counted as a single model for SageMaker metrics, such as LoadedModelCount.

**Publishing default Triton metrics to Amazon CloudWatch**

The NVIDIA Triton Inference Container exposes metrics at port 8002 (configurable) for the different models and GPUs that are utilized in the Triton Inference Server. For full details of the default metrics that are available, see the GitHub page for the [Triton Inference Server metrics](https://github.com/triton-inference-server/metrics). These metrics are in Prometheus format and can be scraped using a Prometheus scraper configuration.

Starting with version v23.07 onwards, the SageMaker Triton container supports publishing these metrics to Amazon CloudWatch by specifying a few environment variables. In order to scrape the Prometheus metrics, the SageMaker Triton container leverages the Amazon CloudWatch agent.

The required environment variables that you must specify to collect metrics are as follows:
<table>
<thead>
<tr>
<th>Environment variable</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAGEMAKER_TRITON.AllowMetrics</td>
<td>Specify this option to allow Triton to publish metrics to its Prometheus endpoint.</td>
<td>&quot;true&quot;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_PUBLISH_METRICS_TO_CLOUDWATCH</td>
<td>Specify this option to start the pre-checks necessary to publish metrics to Amazon CloudWatch.</td>
<td>&quot;true&quot;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_CLOUDWATCH_LOG_GROUP</td>
<td>Specify this option to point to the log group to which metrics are written.</td>
<td>&quot;/aws/Smaker/Enpoints/TritonMetrics/SageMakerTwoEnsemblesTest&quot;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_CLOUDWATCH_METRICNAMESPACE</td>
<td>Specify this option to point to the metric namespace where you want to see and plot the metrics.</td>
<td>&quot;/aws/Smaker/Enpoints/TritonMetrics/SageMakerTwoEnsemblesPublicTest&quot;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_METRICS_PORT</td>
<td>Specify this as 8002, or any other port. If SageMaker has not blocked the specified port, it is used. Otherwise, another non-blocked port is chosen automatically.</td>
<td>&quot;8002&quot;</td>
</tr>
</tbody>
</table>

When publishing metrics with Triton on SageMaker, keep in mind the following limitations:

- While you can generate custom metrics through the C-API and Python backend (v23.05 onwards), these are currently not supported for publishing to Amazon CloudWatch.
- In SageMaker multi-model endpoints (MME) mode, Triton runs in an environment that requires model namespace to be enabled because each model (except ensemble models) are treated as if they are in their own model repository. Currently, this creates a limitation for metrics. When model namespace is enabled, Triton does not distinguish the metrics between two models with the same name belonging to different ensembles. As a workaround, make sure that every
model being deployed has a unique name. This also makes it easier to look up your metrics in CloudWatch.

### Environment variables

The following table lists the supported environment variables for Triton on SageMaker.

<table>
<thead>
<tr>
<th>Environment variable</th>
<th>Description</th>
<th>Type</th>
<th>Possible values</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAGEMAKER_MULTI_MODEL</td>
<td>Allows Triton to operate in SageMaker multi-model endpoints mode.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_DEFAULT_MODEL_NAME</td>
<td>Specify the model to be loaded in the SageMaker single model (default) mode. For ensemble mode, specify the name of the ensemble proper.</td>
<td>String</td>
<td><code>&lt;model_name&gt;</code> as specified in config.pbtxt</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_PING_MODE</td>
<td>'ready' is the default mode in SageMaker's single model mode, and 'live' is the default in SageMaker's multi-model endpoints mode.</td>
<td>String</td>
<td>ready, live</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_DISABLE_MODE</td>
<td>In the SageMaker Triton container, this</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
<td>Type</td>
<td>Possible values</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------</td>
<td>------</td>
<td>-----------------</td>
</tr>
<tr>
<td>DEL_NAMES PACING</td>
<td>is set to <code>true</code> by default.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SAGEMAKER_BIND_TO_PORT</td>
<td>While on SageMaker, the default port is 8080. You can customize to a different port in multi-container scenarios.</td>
<td>String</td>
<td><code>&lt;port_number&gt;</code></td>
</tr>
<tr>
<td>SAGEMAKER_SAFE_PORT_RANGE</td>
<td>This is set by the SageMaker platform when using multi-container mode.</td>
<td>String</td>
<td><code>&lt;port_1&gt;</code>–<code>&lt;port_2&gt;</code></td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_ALLOW_GRPC</td>
<td>While SageMaker doesn't support GRPC currently, if you're using Triton in front of a custom reverse proxy, you may choose to enable GRPC.</td>
<td>Boolean</td>
<td><code>true, false</code></td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_GRPC_PORT</td>
<td>The default port for GRPC is 8001, but you can change it.</td>
<td>String</td>
<td><code>&lt;port_number&gt;</code></td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_THREAD_COUNT</td>
<td>You can set the number of default HTTP request handler threads.</td>
<td>String</td>
<td><code>&lt;number&gt;</code></td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
<td>Type</td>
<td>Possible values</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------</td>
<td>---------------------</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_LOG_VERBOSE</td>
<td>true by default on SageMaker, but you can selectively turn this option off.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_LOG_INFO</td>
<td>false by default on SageMaker.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_LOG_WARNING</td>
<td>false by default on SageMaker.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_LOG_ERROR</td>
<td>false by default on SageMaker.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_SHM_DEFAULT_BYTE_SIZE</td>
<td>Specify the shm size for the Python backend, in bytes. The default value is 16 MB but can be increased.</td>
<td>String</td>
<td>&lt;number&gt;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_SHM_GROWTH_BYTE_SIZE</td>
<td>Specify the shm growth size for the Python backend, in bytes. The default value is 1 MB but can be increased to allow greater increments.</td>
<td>String</td>
<td>&lt;number&gt;</td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
<td>Type</td>
<td>Possible values</td>
</tr>
<tr>
<td>----------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------</td>
<td>-----------------</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_TENSORFLOW_VERSION</td>
<td>The default value is 2. Triton no longer supports Tensorflow 2 from Triton v23.04. You can configure this variable for previous versions.</td>
<td>String</td>
<td>&lt;number&gt;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_MODEL_LOAD_GPU_LIMIT</td>
<td>Restrict the maximum GPU memory percentage which is used for model loading, allowing the remainder to be used for the inference requests.</td>
<td>String</td>
<td>&lt;number&gt;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_ALLOW_METRICS</td>
<td>false by default on SageMaker.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_METRICS_PORT</td>
<td>The default port is 8002.</td>
<td>String</td>
<td>&lt;number&gt;</td>
</tr>
<tr>
<td>Environment variable</td>
<td>Description</td>
<td>Type</td>
<td>Possible values</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------</td>
<td>------</td>
<td>-----------------</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_PUBLISH_Metrics_TO_CLOUDWATCH</td>
<td>false by default on SageMaker. Set this variable to true to allow pushing Triton default metrics to Amazon CloudWatch. If this option is enabled, you are responsible for CloudWatch costs when metrics are published to your account.</td>
<td>Boolean</td>
<td>true, false</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_CLOUDWATCH_LOG_GROUP</td>
<td>Required if you've enabled metrics publishing to CloudWatch.</td>
<td>String</td>
<td>&lt;cloudwatch_log_group_name&gt;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_CLOUDWATCH_METRIC_NAMESPACE</td>
<td>Required if you've enabled metrics publishing to CloudWatch.</td>
<td>String</td>
<td>&lt;cloudwatch_metric_namespace&gt;</td>
</tr>
<tr>
<td>SAGEMAKER_TRITON_ADDITIONAL_ARGS</td>
<td>Appends any additional arguments when starting the Triton Server.</td>
<td>String</td>
<td>&lt;additional_args&gt;</td>
</tr>
</tbody>
</table>
Amazon SageMaker Edge Manager provides model management for edge devices so you can optimize, secure, monitor, and maintain machine learning models on fleets of edge devices such as smart cameras, robots, personal computers, and mobile devices.

Why Use Edge Manager?

Many machine learning (ML) use cases require running ML models on a fleet of edge devices, which allows you to get predictions in real-time, preserves the privacy of the end users, and lowers the cost of network connectivity. With the increasing availability of low-power edge hardware designed for ML, it is now possible to run multiple complex neural network models on edge devices.

However, operating ML models on edge devices is challenging, because devices, unlike cloud instances, have limited compute, memory, and connectivity. After the model is deployed, you need to continuously monitor the models, because model drift can cause the quality of model to decay overtime. Monitoring models across your device fleets is difficult because you need to write custom code to collect data samples from your device and recognize skew in predictions. In addition, models are often hard-coded into the application. To update the model, you must rebuild and update the entire application or device firmware, which can disrupt your operations.

With SageMaker Edge Manager, you can optimize, run, monitor, and update machine learning models across fleets of devices at the edge.

How Does it Work?

At a high level, there are five main components in the SageMaker Edge Manager workflow: compiling models with SageMaker Neo, packaging Neo-compiled models, deploying models to your devices, running models on the SageMaker inference engine (Edge Manager agent), and maintaining models on the devices.

Deploy models at the edge with SageMaker Edge Manager
SageMaker Edge Manager uses SageMaker Neo to optimize your models for the target hardware in one click, then to cryptographically sign your models before deployment. Using SageMaker Edge Manager, you can sample model input and output data from edge devices and send it to the cloud for monitoring and analysis, and view a dashboard that tracks and visually reports on the operation of the deployed models within the SageMaker console.

SageMaker Edge Manager extends capabilities that were previously only available in the cloud to the edge, so developers can continuously improve model quality by using Amazon SageMaker Model Monitor for drift detection, then relabel the data with SageMaker Ground Truth and retrain the models in SageMaker.

**How Do I Use SageMaker Edge Manager?**

If you are a first time user of SageMaker Edge Manager, we recommend that you do the following:

1. **Read the Getting Started section** - This section walks you through setting up your first edge packaging job and creating your first fleet.
2. **Explore Edge Manager Jupyter notebook examples** - Example notebooks are stored in the `amazon-sagemaker-examples` GitHub repository in the `sagemaker_edge_manager` folder.

**Getting Started**

This guide demonstrates how to complete the necessary steps to register, deploy, and manage a fleet of devices, and how to satisfy Amazon SageMaker Edge Manager prerequisites.

**Topics**

- Setting Up
- Train, Compile, and Package Your Model
- Create and Register Fleets and Authenticate Devices
Setting Up

Before you begin using SageMaker Edge Manager to manage models on your device fleets, you must first create IAM Roles for both SageMaker and AWS IoT. You will also want to create at least one Amazon S3 bucket where you will store your pre-trained model, the output of your SageMaker Neo compilation job, as well as input data from your edge devices.

Sign up for an AWS account

If you do not have an AWS account, complete the following steps to create one.

To sign up for an AWS account

2. Follow the online instructions.

   Part of the sign-up procedure involves receiving a phone call and entering a verification code on the phone keypad.

   When you sign up for an AWS account, an AWS account root user is created. The root user has access to all AWS services and resources in the account. As a security best practice, assign administrative access to an administrative user, and use only the root user to perform tasks that require root user access.

   AWS sends you a confirmation email after the sign-up process is complete. At any time, you can view your current account activity and manage your account by going to https://aws.amazon.com/ and choosing My Account.

Create an administrative user

After you sign up for an AWS account, secure your AWS account root user, enable AWS IAM Identity Center, and create an administrative user so that you don't use the root user for everyday tasks.

Secure your AWS account root user

1. Sign in to the AWS Management Console as the account owner by choosing Root user and entering your AWS account email address. On the next page, enter your password.
For help signing in by using root user, see Signing in as the root user in the AWS Sign-In User Guide.

2. Turn on multi-factor authentication (MFA) for your root user.

   For instructions, see Enable a virtual MFA device for your AWS account root user (console) in the IAM User Guide.

Create an administrative user

1. Enable IAM Identity Center.

   For instructions, see Enabling AWS IAM Identity Center in the AWS IAM Identity Center User Guide.

2. In IAM Identity Center, grant administrative access to an administrative user.

   For a tutorial about using the IAM Identity Center directory as your identity source, see Configure user access with the default IAM Identity Center directory in the AWS IAM Identity Center User Guide.

Sign in as the administrative user

- To sign in with your IAM Identity Center user, use the sign-in URL that was sent to your email address when you created the IAM Identity Center user.

   For help signing in using an IAM Identity Center user, see Signing in to the AWS access portal in the AWS Sign-In User Guide.

Create roles and storage

SageMaker Edge Manager needs access to your Amazon S3 bucket URI. To facilitate this, create an IAM role that can run SageMaker and has permission to access Amazon S3. Using this role, SageMaker can run under your account and access to your Amazon S3 bucket.

You can create an IAM role by using the IAM console, AWS SDK for Python (Boto3), or AWS CLI. The following is an example of how to create an IAM role, attach the necessary policies with the IAM console, and create an Amazon S3 bucket.

1. Create an IAM role for Amazon SageMaker.
a. Sign in to the AWS Management Console and open the IAM console at https://console.aws.amazon.com/iam/.
b. In the navigation pane of the IAM console, choose Roles, and then choose Create role.
c. For Select type of trusted entity, choose AWS service.
d. Choose the service that you want to allow to assume this role. In this case, choose SageMaker. Then choose Next: Permissions.
   • This automatically creates an IAM policy that grants access to related services such as Amazon S3, Amazon ECR, and CloudWatch Logs.
e. Choose Next: Tags.
f. (Optional) Add metadata to the role by attaching tags as key–value pairs. For more information about using tags in IAM, see Tagging IAM resources.
g. Choose Next: Review.
h. Type in a Role name.
i. If possible, type a role name or role name suffix. Role names must be unique within your AWS account. They are not distinguished by case. For example, you cannot create roles named both PRODROLE and prodrole. Because other AWS resources might reference the role, you cannot edit the name of the role after it has been created.
j. (Optional) For Role description, type a description for the new role.
k. Review the role and then choose Create role.

Note the SageMaker Role ARN, which you use to create a compilation job with SageMaker Neo and a packaging job with Edge Manager. To find out the role ARN using the console, do the following:

i. Go to the IAM console: https://console.aws.amazon.com/iam/
ii. Select Roles.
iii. Search for the role you just created by typing in the name of the role in the search field.
iv. Select the role.
v. The role ARN is at the top of the Summary page.

2. Create an IAM role for AWS IoT.
The AWS IoT IAM role you create is used to authorize your thing objects. You also use the IAM role ARN to create and register device fleets with a SageMaker client object.

Configure an IAM role in your AWS account for the credentials provider to assume on behalf of the devices in your device fleet. Then, attach a policy to authorize your devices to interact with AWS IoT services.

Create a role for AWS IoT either programmatically or with the IAM console, similar to what you did when you created a role for SageMaker.

a. Sign in to the AWS Management Console and open the IAM console at https://console.aws.amazon.com/iam/.
b. In the navigation pane of the IAM console, choose Roles, and then choose Create role.
c. For Select type of trusted entity, choose AWS service.
d. Choose the service that you want to allow to assume this role. In this case, choose IoT. Select IoT as the Use Case.
e. Choose Next: Permissions.
f. Choose Next: Tags.
g. (Optional) Add metadata to the role by attaching tags as key–value pairs. For more information about using tags in IAM, see Tagging IAM resources.
h. Choose Next: Review.
i. Type in a Role name. The role name must start with SageMaker.
j. (Optional) For Role description, type a description for the new role.
k. Review the role and then choose Create role.
l. Once the role is created, choose Roles in the IAM console. Search for the role you created by typing in role name in the Search field.
m. Choose your role.
n. Next, choose Attach Policies.
o. Search for AmazonSageMakerEdgeDeviceFleetPolicy in the Search field. Select AmazonSageMakerEdgeDeviceFleetPolicy.
p. Choose Attach policy.
q. Add the following policy statement to the trust relationship:
A trust policy is a JSON policy document in which you define the principals that you trust to assume the role. For more information about trust policies, see Roles terms and concepts.

r. Note the AWS IoT role ARN. You use the AWS IoT Role ARN to create and register the device fleet. To find the IAM role ARN with the console:

i. Go to the IAM console: https://console.aws.amazon.com/iam/

ii. Choose Roles.

iii. Search for the role you created by typing in the name of the role in the Search field.

iv. Select the role.

v. The role ARN is on the Summary page.

3. **Create an Amazon S3 bucket.**

   SageMaker Neo and Edge Manager access your pre-compiled model and compiled model from an Amazon S3 bucket. Edge Manager also stores sample data from your device fleet in Amazon S3.

   a. Open the Amazon S3 console at https://console.aws.amazon.com/s3/.

   b. Choose Create bucket.

   c. In **Bucket name**, enter a name for your bucket.

   d. In **Region**, choose the AWS Region where you want the bucket to reside.

   e. In **Bucket settings for Block Public Access**, choose the settings that you want to apply to the bucket.
Train, Compile, and Package Your Model

In this section you will create SageMaker and AWS IoT client objects, download a pre-trained machine learning model, upload your model to your Amazon S3 bucket, compile your model for your target device with SageMaker Neo, and package your model so that it can be deployed with the Edge Manager agent.

1. Import libraries and create client objects.

This tutorial uses the AWS SDK for Python (Boto3) to create clients to interact with SageMaker, Amazon S3, and AWS IoT.

Import Boto3, specify your Region, and initialize the client objects you need as shown in the following example:

```python
import boto3
import json
import time

AWS_REGION = 'us-west-2'  # Specify your Region
bucket = 'bucket-name'

sagemaker_client = boto3.client('sagemaker', region_name=AWS_REGION)
iot_client = boto3.client('iot', region_name=AWS_REGION)
```

Define variables and assign them the role ARN you created for SageMaker and AWS IoT as strings:

```python
# Replace with the role ARN you created for SageMaker
sagemaker_role_arn = "arn:aws:iam::<account>:role/*"

# Replace with the role ARN you created for AWS IoT.
# Note: The name must start with 'SageMaker'
iot_role_arn = "arn:aws:iam::<account>:role/SageMaker*"
```

2. Train a machine learning model.
See [Train a Model with Amazon SageMaker](#) for more information on how to train a machine learning model using SageMaker. You can optionally upload your locally trained model directly into an Amazon S3 URI bucket.

If you do not have a model yet, you can use a pre-trained model for the next steps in this tutorial. For example, you can save the MobileNet V2 models from the TensorFlow framework. MobileNet V2 is an image classification model optimized for mobile applications. For more information about MobileNet V2, see the [MobileNet GitHub README](#).

Type the following into your Jupyter Notebook to save the pre-trained MobileNet V2 model:

```python
# Save the MobileNet V2 model to local storage
import tensorflow as tf
model = tf.keras.applications.MobileNetV2()
model.save("mobilenet_v2.h5")
```

**Note**

- If you do not have TensorFlow installed, you can do so by running `pip install tensorflow=2.4`
- Use TensorFlow version 2.4 or lower for this tutorial.

The model will be saved into the `mobilenet_v2.h5` file. Before packaging the model, you will need to first compile your model using SageMaker Neo. See [Supported Frameworks, Devices, Systems, and Architectures](#) to check if your version of TensorFlow (or other framework of choice) is currently supported by SageMaker Neo.

SageMaker Neo requires models to be stored as a compressed TAR file. Repackage it as a compressed TAR file (*.tar.gz):

```python
# Package MobileNet V2 model into a TAR file
import tarfile
tarfile_name='mobilenet-v2.tar.gz'

with tarfile.open(tarfile_name, mode='w:gz') as archive:
```
3. **Upload your model to Amazon S3.**

Once you have a machine learning model, store it in an Amazon S3 bucket. The following example uses an AWS CLI command to upload the model to the Amazon S3 bucket you created earlier in a directory called `models`. Type in the following into your Jupyter Notebook:

```bash
!aws s3 cp mobilenet-v2.tar.gz s3://{bucket}/models/
```

4. **Compile your model with SageMaker Neo.**

Compile your machine learning model with SageMaker Neo for an edge device. You need to know your Amazon S3 bucket URI where you stored the trained model, the machine learning framework you used to train your model, the shape of your model's input, and your target device.

For the MobileNet V2 model, use the following:

```python
framework = 'tensorflow'
target_device = 'jetson_nano'
data_shape = '{"data":[1,3,224,224]}'
```

SageMaker Neo requires a specific model input shape and model format based on the deep learning framework you use. For more information about how to save your model, see [What input data shapes does SageMaker Neo expect?](http://example.com). For more information about devices and frameworks supported by Neo, see [Supported Frameworks, Devices, Systems, and Architectures](http://example.com).

Use the `CreateCompilationJob` API to create a compilation job with SageMaker Neo. Provide a name to the compilation job, the SageMaker Role ARN, the Amazon S3 URI where your model is stored, the input shape of the model, the name of the framework, the Amazon S3 URI where you want SageMaker to store your compiled model, and your edge device target.

```python
# Specify the path where your model is stored
model_directory = 'models'
s3_model_uri = 's3:///{}/{}'.format(bucket, model_directory, tarfile_name)

# Store compiled model in S3 within the 'compiled-models' directory
compilation_output_dir = 'compiled-models'
```
5. **Package your compiled model.**

Packaging jobs take SageMaker Neo–compiled models and make any changes necessary to deploy the model with the inference engine, Edge Manager agent. To package your model, create an edge packaging job with the `create_edge_packaging` API or the SageMaker console.

You need to provide the name that you used for your Neo compilation job, a name for the packaging job, a role ARN (see Setting Up section), a name for the model, a model version, and the Amazon S3 bucket URI for the output of the packaging job. Note that Edge Manager packaging job names are case-sensitive. The following is an example of how to create a packaging job using the API.

```python
edge_packaging_name='edge-packaging-demo'
model_name="sample-model"
model_version="1.1"
```

Define the Amazon S3 URI where you want to store the packaged model.

```python
# Output directory where you want to store the output of the packaging job
packaging_output_dir = 'packaged_models'
packaging_s3_output = 's3://{}/{}'.format(bucket, packaging_output_dir)
```
Use CreateEdgePackagingJob to package your Neo-compiled model. Provide a name for your edge packaging job and the name you provided for your compilation job (in this example, it was stored in the variable `compilation_job_name`). Also provide a name for your model, a version for your model (this is used to help you keep track of what model version you are using), and the S3 URI where you want SageMaker to store the packaged model.

```python
sagemaker_client.create_edge_packaging_job(
    EdgePackagingJobName=edge_packaging_name,
    CompilationJobName=compilation_job_name,
    RoleArn=sagemaker_role_arn,
    ModelName=model_name,
    ModelVersion=model_version,
    OutputConfig={
        "S3OutputLocation": packaging_s3_output
    }
)
```

**Create and Register Fleets and Authenticate Devices**

In this section you will create your AWS IoT thing object, create a device fleet, register your device fleet so it can interact with the cloud, create X.509 certificates to authenticate your devices to AWS IoT Core, associate the role alias with AWS IoT that was generated when you created your fleet, get your AWS account-specific endpoint for the credentials provider, get an official Amazon Root CA file, and upload the Amazon CA file to Amazon S3.

1. **Create AWS IoT things.**

   SageMaker Edge Manager takes advantage of the AWS IoT Core services to facilitate the connection between the edge devices and endpoints in the AWS cloud. You can take advantage of existing AWS IoT functionality after you set up your devices to work with Edge Manager.

   To connect your device to AWS IoT, you need to create AWS IoT *thing objects*, create and register a client certificate with AWS IoT, and create and configure the IAM role for your devices.

   First, create AWS IoT thing objects with the AWS IoT client (`iot_client`) you created earlier with Boto3. The following example shows how to create two thing objects:
2. **Create your device fleet.**

Create a device fleet with the SageMaker client object defined in a previous step. You can also use the SageMaker console to create a device fleet.

```
import time
device_fleet_name="demo-device-fleet" + str(time.time()).split('.')[0]
device_name="sagemaker-edge-demo-device" + str(time.time()).split('.')[0]
```

Specify your IoT role ARN. This lets AWS IoT grant temporary credentials to devices.

```
device_model_directory='device_output'
s3_device_fleet_output = 's3://{}/{}'.format(bucket, device_model_directory)
sagemaker_client.create_device_fleet(
    DeviceFleetName=device_fleet_name,
    RoleArn=iot_role_arn, # IoT Role ARN specified in previous step
    OutputConfig={
        'S3OutputLocation': s3_device_fleet_output
    }
)
```

An AWS IoT role alias is created when you create a device fleet. This role alias is associated with AWS IoT using the `iot_client` object in a later step.

3. **Register your device fleet.**
To interact with the cloud, you need to register your device with SageMaker Edge Manager. In this example, you register a single device with the fleet you created. To register the device, you need to provide a device name and the AWS IoT thing name as shown in the following example:

```python
# Device name should be 36 characters
device_name = "sagemaker-edge-demo-device" + str(time.time()).split('.')[0]

sagemaker_client.register_devices(
    DeviceFleetName=device_fleet_name,
    Devices=[
        {
            "DeviceName": device_name,
            "IotThingName": iot_thing_name
        }
    ]
)
```

4. **Create X.509 certificates.**

After creating the AWS IoT thing object, you must create a X.509 device certificate for your thing object. This certificate authenticates your device to AWS IoT Core.

Use the following to create a private key, public key, and a X.509 certificate file using the AWS IoT client defined (iot_client) earlier.

```python
# Creates a 2048-bit RSA key pair and issues an X.509 # certificate
# using the issued public key.
create_cert = iot_client.create_keys_and_certificate(
    setAsActive=True
)

# Get certificate from dictionary object and save in its own
with open('./device.pem.crt', 'w') as f:
    for line in create_cert['certificatePem'].split('
'): f.write(line)
    f.write('
')

# Get private key from dictionary object and save in its own
with open('./private.pem.key', 'w') as f:
    for line in create_cert['keyPair']['PrivateKey'].split('
'): f.write(line)
```
f.write('n')
# Get a private key from dictionary object and save in its own
with open('./public.pem.key', 'w') as f:
  for line in create_cert['keyPair']['PublicKey'].split('n'):
    f.write(line)
    f.write('n')

5. **Associate the role alias with AWS IoT.**

When you create a device fleet with SageMaker
(sagemaker_client.create_device_fleet()), a role alias is generated for you. An
AWS IoT role alias provides a mechanism for connected devices to authenticate to AWS IoT
using X.509 certificates, and then obtain short-lived AWS credentials from an IAM role that is
associated with an AWS IoT role alias. The role alias allows you to change the role of the device
without having to update the device. Use DescribeDeviceFleet to get the role alias name and ARN.

# Print Amazon Resource Name (ARN) and alias that has access
# to AWS Internet of Things (IoT).
sagemaker_client.describe_device_fleet(DeviceFleetName=device_fleet_name)

# Store iot role alias string in a variable
# Grabs role ARN
full_role_alias_name =
  sagemaker_client.describe_device_fleet(DeviceFleetName=device_fleet_name)
  ['IotRoleAlias']
start_index = full_role_alias_name.find('SageMaker') # Find beginning of role name
role_alias_name = full_role_alias_name[start_index:]

Use the iot_client to facilitate associating the role alias generated from creating the device
fleet with AWS IoT:

role_alias = iot_client.describe_role_alias(
    roleAlias=role_alias_name)

For more information about IAM role alias, see [Role alias allows access to unused services](#).
You created and registered a certificate with AWS IoT earlier for successful authentication of your device. Now, you need to create and attach a policy to the certificate to authorize the request for the security token.

```python
alias_policy = {
    "Version": "2012-10-17",
    "Statement": {
        "Effect": "Allow",
        "Action": "iot:AssumeRoleWithCertificate",
        "Resource": role_alias['roleAliasDescription']['roleAliasArn']
    }
}

policy_name = 'aliaspolicy-+' + str(time.time()).split('.')[0]
aliaspolicy = iot_client.create_policy(policyName=policy_name,
                    policyDocument=json.dumps(alias_policy))

# Attach policy
iot_client.attach_policy(policyName=policy_name,
                    target=create_cert['certificateArn'])
```

6. **Get your AWS account-specific endpoint for the credentials provider.**

Edge devices need an endpoint in order to assume credentials. Obtain your AWS account-specific endpoint for the credentials provider.

```python
# Get the unique endpoint specific to your AWS account that is making the call.
iot_endpoint = iot_client.describe_endpoint(
    endpointType='iot:CredentialProvider'
)

eventpoint="https://{}/role-aliases/{}/
credentials".format(iot_endpoint['endpointAddress'],role_alias_name)
```

7. **Get the official Amazon root CA file and upload it to the Amazon S3 bucket.**

Use the following in your Jupyter Notebook or AWS CLI (if you use your terminal, remove the ‘!’ magic function):

```bash
!wget https://www.amazontrust.com/repository/AmazonRootCA1.pem
```
Use the endpoint to make an HTTPS request to the credentials provider to return a security token. The following example command uses `curl`, but you can use any HTTP client.

```bash
!curl --cert device.pem.crt --key private.pem.key --cacert AmazonRootCA1.pem $endpoint
```

If the certificate is verified, upload the keys and certificate to your Amazon S3 bucket URI:

```bash
!aws s3 cp private.pem.key s3://{bucket}/authorization-files/
!aws s3 cp device.pem.crt s3://{bucket}/authorization-files/
!aws s3 cp AmazonRootCA1.pem s3://{bucket}/authorization-files/
```

Clean your working directory by moving your keys and certificate to a different directory:

```bash
# Optional - Clean up working directory
!mkdir authorization-files
!mv private.pem.key device.pem.crt AmazonRootCA1.pem authorization-files/
```

---

**Download and Set Up Edge Manager**

The Edge Manager agent is an inference engine for your edge devices. Use the agent to make predictions with models loaded onto your edge devices. The agent also collects model metrics and captures data at specific intervals.

In this section you will set up your device with the agent. To do so, first copy a release artifact and signing root certificate from the release bucket locally to your machine. After you unzip the release artifact, upload it to Amazon S3. Next, define and save a configuration file for the agent. A template is provided for you to copy and paste. Finally, copy the release artifacts, configuration file, and credentials to your device.

1. **Download the SageMaker Edge Manager agent.**

   The agent is released in binary format for supported operating systems. This example runs inference on a Jetson Nano which uses a Linux operating system and has an ARM64 architecture. For more information about what operating system and architecture supported devices use, see [Supported Devices, Chip Architectures, and Systems](#).
Fetch the latest version of binaries from the SageMaker Edge Manager release bucket from the us-west-2 Region.

```
!aws s3 ls s3://sagemaker-edge-release-store-us-west-2-linux-armv8/Releases/ | sort -r
```

This returns release artifacts sorted by their version.

<table>
<thead>
<tr>
<th>Version</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRE 1.20210512.96da6cc/</td>
</tr>
<tr>
<td>PRE 1.20210305.a4bc999/</td>
</tr>
<tr>
<td>PRE 1.20201218.81f481f/</td>
</tr>
<tr>
<td>PRE 1.20201207.02d0e97/</td>
</tr>
</tbody>
</table>

The version has the following format: `<MAJOR_VERSION>.<YYYY-MM-DD>.<SHA-7>`. It consists of three components:

- `<MAJOR_VERSION>`: The release version. The release version is currently set to 1.
- `<YYYY-MM-DD>`: The time stamp of the artifact release.
- `<SHA-7>`: The repository commit ID from which the release is built.

Copy the zipped TAR file locally or to your device directly. The following example shows how to copy the latest release artifact at the time this document was released.

```
!aws s3 cp s3://sagemaker-edge-release-store-us-west-2-linux-x64/Releases/1.20201218.81f481f/1.20201218.81f481f.tgz ./
```

Once you have the artifact, unzip the zipped TAR file. The following unzips the TAR file and stores it in a directory called `agent_demo`:

```
!mkdir agent_demo
!tar -xvzf 1.20201218.81f481f.tgz -C ./agent_demo
```

Upload the agent release artifacts to your Amazon S3 bucket. The following code example copies the content within `agent_demo` and uploads it to a directory within your Amazon S3 bucket called `agent_demo`:

```
```
You also need the signing root certificates from the release bucket:

```bash
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-linux-x64/Certificates/us-west-2/us-west-2.pem ./
```

Upload the signing root certificate to your Amazon S3 bucket:

```bash
aws s3 cp us-west-2.pem s3://{bucket}/authorization-files/
```

2. **Define a SageMaker Edge Manager agent configuration file.**

First, define the agent configuration file as follows:

```json
sagemaker_edge_config = {
    "sagemaker_edge_core_device_name": "device_name",
    "sagemaker_edge_core_device_fleet_name": "device_fleet_name",
    "sagemaker_edge_core_capture_data_buffer_size": 30,
    "sagemaker_edge_core_capture_data_push_period_seconds": 4,
    "sagemaker_edge_core_folder_prefix": "demo_capture",
    "sagemaker_edge_core_region": "us-west-2",
    "sagemaker_edge_core_root_certs_path": "agent_demo/certificates",
    "sagemaker_edge_provider_aws_ca_cert_file": "agent_demo/iot-credentials/AmazonRootCA1.pem",
    "sagemaker_edge_provider_aws_cert_file": "agent_demo/iot-credentials/device.pem.crt",
    "sagemaker_edge_provider_aws_cert_pk_file": "agent_demo/iot-credentials/private.pem.key",
    "sagemaker_edge_provider_aws_iot_cred_endpoint": "endpoint",
    "sagemaker_edge_provider_provider": "Aws",
    "sagemaker_edge_provider_s3_bucket_name": bucket,
    "sagemaker_edge_core_capture_data_destination": "Cloud"
}
```

Replace the following:

- "device_name" with the name of your device (this string was stored in an earlier step in a variable named `device_name`).
• "device_fleet_name" with the name of your device fleet (this string was stored an earlier step in a variable named device_fleet_name)

• "endpoint" with your AWS account-specific endpoint for the credentials provider (this string was stored in an earlier step in a variable named endpoint).

Next, save it as a JSON file:

```python
edge_config_file = open("sagemaker_edge_config.json", "w")
json.dump(sagemaker_edge_config, edge_config_file, indent = 6)
edge_config_file.close()
```

Upload the configuration file to your Amazon S3 bucket:

```bash
!aws s3 cp sagemaker_edge_config.json s3://{bucket}/
```

3. Copy the release artifacts, configuration file, and credentials to your device.

The following instructions are performed on the edge device itself.

**Note**

You must first install Python, the AWS SDK for Python (Boto3), and the AWS CLI on your edge device.

Open a terminal on your device. Create a folder to store the release artifacts, your credentials, and the configuration file.

```bash
mkdir agent_demo
cd agent_demo
```

Copy the contents of the release artifacts that you stored in your Amazon S3 bucket to your device:

```bash
# Copy release artifacts
aws s3 cp s3://<bucket-name>/agent_demo/ ./ --recursive
```
(The contents of the release artifact was stored in a directory called `agent_demo` in a previous step). Replace `<bucket-name>` and `agent_demo` with the name of your Amazon S3 bucket and the file path to your release artifacts, respectively.

Go the `/bin` directory and make the binary files executable:

```bash
cd bin
chmod +x sagemaker_edge_agent_binary
chmod +x sagemaker_edge_agent_client_example

cd agent_demo
```

Make a directory to store your AWS IoT credentials and copy your credentials from your Amazon S3 bucket to your edge device (use the same on you define in the variable `bucket`):

```bash
mkdir iot-credentials
cd iot-credentials
aws s3 cp s3://<bucket-name>/authorization-files/AmazonRootCA1.pem ./
aws s3 cp s3://<bucket-name>/authorization-files/device.pem.crt ./
aws s3 cp s3://<bucket-name>/authorization-files/private.pem.key ./
cd ../
```

Make a directory to store your model signing root certificates:

```bash
mkdir certificates
cd certificates
aws s3 cp s3://<bucket-name>/authorization-files/us-west-2.pem ./
cd agent_demo
```

Copy your configuration file to your device:

```bash
#Download config file from S3
aws s3 cp s3://<bucket-name>/sagemaker_edge_config.json ./
```
cd agent_demo

Your agent_demo directory on your edge device should look similar to the following:

```
###agent_demo
|   ### bin
|   |   ### sagemaker_edge_agent_binary
|   |   ### sagemaker_edge_agent_client_example
|   |   ### sagemaker_edge_config.json
|   ### certificates
|   |   ###us-west-2.pem
|   |   ### iot-credentials
|   |   |   ### AmazonRootCA1.pem
|   |   |   ### device.pem.crt
|   |   |   ### private.pem.key
|   |   ### docs
|   |   |   ### api
|   |   |   ### examples
|   |   |   ### ATTRIBUTIONS.txt
|   |   ### LICENSE.txt
|   |   ### RELEASE_NOTES.md
```

**Run Agent**

In this section you will run the agent as a binary using gRPC, and check that both your device and fleet are working and collecting sample data.

1. **Launch the agent.**

   The SageMaker Edge Manager agent can be run as a standalone process in the form of an Executable and Linkable Format (ELF) executable binary or can be linked against as a Dynamic Shared Object (.dll). Running as a standalone executable binary is the preferred mode and is supported on Linux.

   This example uses gRPC to run the agent. gRPC is an open source high-performance Remote Procedure Call (RPC) framework that can run in any environment. For more information about gRPC, see the [gRPC documentation](#).

   To use gRPC, perform the following steps:

b. Generate server and client code using the protocol buffer compiler.

c. Use the Python (or other languages supported by gRPC) gRPC API to write the server for your service.

d. Use the Python (or other languages supported by gRPC) gRPC API to write a client for your service.

The release artifact you downloaded contains a gRPC application ready for you to run the agent. The example is located within the /bin directory of your release artifact. The sagemaker_edge_agent_binary binary executable is in this directory.

To run the agent with this example, provide the path to your socket file (.sock) and JSON .config file:

```
./bin/sagemaker_edge_agent_binary -a /tmp/sagemaker_edge_agent_example.sock -c sagemaker_edge_config.json
```

2. **Check your device.**

Check that your device is connected and sampling data. Making periodic checks, manually or automatically, allows you to check that your device or fleet is working properly.

Provide the name of the fleet to which the device belongs and the unique device identifier. From your local machine, run the following:

```
sagemaker_client.describe_device(
    DeviceName=device_name,
    DeviceFleetName=device_fleet_name
)
```

For the given model, you can see the name, model version, latest sample time, and when the last inference was made.

```
{
    "DeviceName": "sample-device",
    "DeviceFleetName": "demo-device-fleet",
    "IoTThingName": "sample-thing-name-1",
    "RegistrationTime": 1600977370,
```
The timestamp provided by `LatestHeartbeat` indicates the last signal that was received from the device. `LatestSampleTime` and `LatestInference` describe the time stamp of the last data sample and inference, respectively.

3. **Check your fleet.**

Check that your fleet is working with `GetDeviceFleetReport`. Provide the name of the fleet the device belongs to.

```python
sagemaker_client.get_device_fleet_report(
    DeviceFleetName=device_fleet_name
)
```

For a given model, you can see the name, model version, latest sample time, and when the last inference was made, along with the Amazon S3 bucket URI where the data samples are stored.

```json
# Sample output
{
    "DeviceFleetName": "sample-device-fleet",
    "OutputConfig": {
        "S3OutputLocation": "s3://fleet-bucket/package_output",
    },
    "AgentVersions": [{"Version": "1.1", "AgentCount": 2}]
    "DeviceStats": {"Connected": 2, "Registered": 2},
    "Models": [{
        "ModelName": "sample-model",
        "ModelVersion": "1.1",
        "OfflineDeviceCount": 0,
        "ConnectedDeviceCount": 2,
```
Set Up Devices and Fleets

Fleets are collections of logically grouped devices you can use to collect and analyze data. You can use SageMaker Edge Manager to operate machine learning models on a fleet of smart cameras, smart speakers, robots, and other edge devices.

Create a fleet and register your devices either programmatically with the AWS SDK for Python (Boto3) or through the SageMaker console.

Topics
- Create a Fleet
- Register a Device
- Check Status

Create a Fleet

You can create a fleet programatically with the AWS SDK for Python (Boto3) or through the SageMaker console https://console.aws.amazon.com/sagemaker.

Create a Fleet (Boto3)

Use the CreateDeviceFleet API to create a fleet. Specify a name for the fleet, your AWS IoT Role ARN for the RoleArn field, as well as an Amazon S3 URI where you want the device to store sampled data.

You can optionally include a description of the fleet, tags, and an AWS KMS Key ID.

```python
import boto3

# Create SageMaker client so you can interact and manage SageMaker resources
sagemaker_client = boto3.client("sagemaker", region_name="aws-region")

sagemaker_client.create_device_fleet(
    DeviceFleetName="sample-fleet-name",
    RoleArn="role-arn", 
    # Other options...
)
```
An AWS IoT Role Alias is created for you when you create a device fleet. The AWS IoT role alias provides a mechanism for connected devices to authenticate to AWS IoT using X.509 certificates and then obtain short-lived AWS credentials from an IAM role that is associated with the AWS IoT role alias.

Use `DescribeDeviceFleet` to get the role alias name and ARN.

```python
# Print Amazon Resource Name (ARN) and alias that has access
# to AWS Internet of Things (IoT).
sagemaker_client.describe_device_fleet(DeviceFleetName=device_fleet_name)
['IotRoleAlias']
```

Use `DescribeDeviceFleet` API to get a description of fleets you created.

```python
sagemaker_client.describe_device_fleet(
    DeviceFleetName="sample-fleet-name"
)
```

By default, it returns the name of the fleet, the device fleet ARN, the Amazon S3 bucket URI, the IAM role, the role alias created in AWS IoT, a timestamp of when the fleet was created, and a timestamp of when the fleet was last modified.

```json
{ "DeviceFleetName": "sample-fleet-name",
  "IAMRole": "arn:aws:iam::9999999999:role/rolename",
  "Description": "this is a sample fleet",
}
```
Create a Fleet (Console)

You can create a Edge Manager packaging job using the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker.

1. In the SageMaker console, choose **Edge Manager** and then choose **Edge device fleets**.

2. Choose **Create device fleet**.

3. Enter a name for the device fleet in the **Device fleet name** field. Choose **Next**.
4. On the **Output configuration** page, specify the Amazon S3 bucket URI where you want to store sample data from your device fleet. You can optionally add an encryption key as well by electing an existing AWS KMS key from the dropdown list or by entering a key's ARN. Choose **Submit**.
5. Choose the name of your device fleet to be redirected to the device fleet details. This page displays the name of the device fleet, ARN, description (if you provided one), date the fleet was created, last time the fleet was modified, Amazon S3 bucket URI, AWS KMS key ID (if provided), AWS IoT alias (if provided), and IAM role. If you added tags, they appear in the **Device fleet tags** section.

**Register a Device**

⚠️ **Important**

Device registration is required to use any part of SageMaker Edge Manager.

You can create a fleet programmatically with the AWS SDK for Python (Boto3) or through the SageMaker console at [https://console.aws.amazon.com/sagemaker](https://console.aws.amazon.com/sagemaker).

**Register a Device (Boto3)**

To register your device, first create and register an AWS IoT thing object and configure an IAM role. SageMaker Edge Manager takes advantage of the AWS IoT Core services to facilitate the connection between the edge devices and the cloud. You can take advantage of existing AWS IoT functionality after you set up your devices to work with Edge Manager.

To connect your device to AWS IoT you need to create AWS IoT thing objects, create and register a client certificate with AWS IoT, and create and configure IAM role for your devices.
See the [Getting Started Guide](https://console.aws.amazon.com/sagemaker) for an in-depth example or the [Explore AWS IoT Core services in hands-on tutorial](https://console.aws.amazon.com/sagemaker).

Use the `register_devices` API to register your device. Provide the name of the fleet of which you want the devices to be a part, as well as a name for the device. You can optionally add a description to the device, tags, and AWS IoT thing name associated with the device.

```python
sagemaker_client.register_devices(
    DeviceFleetName="sample-fleet-name",
    Devices=[
        {
            "DeviceName": "sample-device-1",
            "IotThingName": "sample-thing-name-1",
            "Description": "Device #1"
        }
    ],
    Tags=[
        {
            "Key": "string",
            "Value": "string"
        }
    ]
)
```

**Register a Device (Console)**


1. In the SageMaker console, choose **Edge Inference** and then choose **Edge devices**.
2. Choose **Register devices**.
3. In the **Device properties** section, enter the name of the fleet the device belongs to under the **Device fleet name** field. Choose **Next**.
4. In the **Device source** section, add your devices one by one. You must include a **Device Name** for each device in your fleet. You can optionally provide a description (in the **Description** field) and an Internet of Things (IoT) object name (in the **IoT name** field). Choose **Submit** once you have added all your devices.

The **Devices** page displays the name of the device you have added, the fleet to which it belongs, when it was registered, the last heartbeat, and the description and AWS IoT name, if you provided one.

Choose a device to view the device's details, including the device name, fleet, ARN, description, IoT Thing name, when the device was registered, and the last heartbeat.

**Check Status**

Check that your device or fleet is connected and sampling data. Making periodic checks, manually or automatically, allows you to check that your device or fleet is working properly.
Use the Amazon S3 console at https://console.aws.amazon.com/s3/ to interactively choose a fleet for a status check. You can also use the AWS SDK for Python (Boto3). The following describes different APIs from Boto3 you can use to check the status of your device or fleet. Use the API that best fits your use case.

- **Check an individual device.**

  To check the status of an individual device, use DescribeDevice API. A list containing one or more models is provided if a models have been deployed to the device.

  ```python
  sagemaker_client.describe_device(
      DeviceName="sample-device-1",
      DeviceFleetName="sample-fleet-name"
  )
  ```

  Running DescribeDevice returns:

  ```json
  {   "DeviceName": "sample-device",   "Description": "this is a sample device",   "DeviceFleetName": "sample-device-fleet",   "IoTThingName": "SampleThing",   "RegistrationTime": 1600977370,   "LatestHeartbeat": 1600977370,   "Models": [     {       "ModelName": "sample-model",       "ModelVersion": "1.1",       "LatestSampleTime": 1600977370,       "LatestInference": 1600977370     }   ] }
  ```

- **Check a fleet of devices.**

  To check the status of the fleet, use the GetDeviceFleetReport API. Provide the name of the device fleet to get a summary of the fleet.

  ```python
  sagemaker_client.get_device_fleet_report(
      DeviceFleetName="sample-fleet-name"
  )
  ```
• **Check for a heartbeat.**

Each device within a fleet periodically generates a signal, or “heartbeat”. The heartbeat can be used to check that the device is communicating with Edge Manager. If the timestamp of the last heartbeat is not being updated, the device may be failing.

Check the last heartbeat with made by a device with the `DescribeDevice` API. Specify the name of the device and the fleet to which the edge device belongs.

```python
sagemaker_client.describe_device(
    DeviceName="sample-device-1",
    DeviceFleetName="sample-fleet-name"
)
```

**Package Model**

SageMaker Edge Manager packaging jobs take Amazon SageMaker Neo–compiled models and make any changes necessary to deploy the model with the inference engine, Edge Manager agent.

**Topics**

• **Prerequisites**

• **Package a Model (Amazon SageMaker Console)**

• **Package a Model (Boto3)**

**Prerequisites**

To package a model, you must do the following:

1. **Compile your machine learning model with SageMaker Neo.**

   If you have not already done so, compile your model with SageMaker Neo. For more information on how to compile your model, see [Compile and Deploy Models with Neo](https://docs.aws.amazon.com/sagemaker/latest/dg/deploy-models-and-edge-manager.html). If you are first-time user of SageMaker Neo, go through [Getting Started with Neo Edge Devices](https://docs.aws.amazon.com/sagemaker/latest/dg/neo-edge-devices.html).

2. **Get the name of your compilation job.**

   Provide the name of the compilation job name you used when you compiled your model with SageMaker Neo. Open the SageMaker console at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/)
and choose **Compilation jobs** to find a list of compilations that have been submitted to your AWS account. The names of submitted compilation jobs are in the **Name** column.

3. **Get your IAM ARN.**

You need an Amazon Resource Name (ARN) of an IAM role that you can use to download and upload the model and contact SageMaker Neo.

Use one of the following methods to get your IAM ARN:

- **Programmatically with the SageMaker Python SDK**

  ```python
  import sagemaker

  # Initialize SageMaker Session object so you can interact with AWS resources
  sess = sagemaker.Session()

  # Get the role ARN
  role = sagemaker.get_execution_role()

  print(role)
  >> arn:aws:iam::<your-aws-account-id>:role/<your-role-name>
  ```

  For more information about using the SageMaker Python SDK, see the [SageMaker Python SDK API](https://docs.aws.amazon.com/sagemaker/latest/dg/).  

- **Using the AWS Identity and Access Management (IAM) console**

  Navigate to the IAM console at [https://console.aws.amazon.com/iam/](https://console.aws.amazon.com/iam/). In the IAM Resources section, choose **Roles** to view a list of roles in your AWS account. Select or create a role that has AmazonSageMakerFullAccess, AWSIoTFullAccess, and AmazonS3FullAccess.

  For more information on IAM, see [What is IAM?](https://docs.aws.amazon.com/)

4. **Have an S3 bucket URI.**

You need to have at least one Amazon Simple Storage Service (Amazon S3) bucket URI to store your Neo-compiled model, the output of the Edge Manager packaging job, and sample data from your device fleet.

Use one of the following methods to create an Amazon S3 bucket:

- **Programmatically with the SageMaker Python SDK**
You can use the default Amazon S3 bucket during a session. A default bucket is created based on the following format: sagemaker-{region}-{aws-account-id}. To create a default bucket with the SageMaker Python SDK, use the following:

```python
import sagemaker

session=sagemaker.create_session()

bucket=session.default_bucket()
```

- **Using the Amazon S3 console**


**Package a Model (Amazon SageMaker Console)**

You can create a SageMaker Edge Manager packaging job using the SageMaker console at [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/). Before continuing, make sure you have satisfied the Prerequisites.

1. In the SageMaker console, choose **Edge Inference** and then choose **Edge packaging jobs**, as shown in the following image.

2. On the **Job properties** page, enter a name for your packaging job under **Edge packaging job name**. Note that Edge Manager packaging job names are case-sensitive. Name your model and give it a version: enter this under **Model name** and **Model version**, respectively.

3. Next, select an **IAM role**. You can chose a role or let AWS create a role for you. You can optionally specify a **resource key ARN** and **job tags**.

4. Choose **Next**.
5. Specify the name of the compilation job you used when compiling your model with SageMaker Neo in the **Compilation job name** field. Choose **Next**.
6. On the **Output configuration** page, enter the Amazon S3 bucket URI in which you want to store the output of the packaging job.

The **Status** column on the **Edge packaging** jobs page should read **IN PROGRESS**. Once the packaging job is complete, the status updates to **COMPLETED**.

Selecting a packaging job directs you to that job's settings. The **Job settings** section displays the job name, ARN, status, creation time, last modified time, duration of the packaging job, and role ARN.

The **Input configuration** section displays the location of the model artifacts, the data input configuration, and the machine learning framework of the model.
The **Output configuration** section displays the output location of the packaging job, the target device for which the model was compiled, and any tags you created.

7. Choose the name of your device fleet to be redirected to the device fleet details. This page displays the name of the device fleet, ARN, description (if you provided one), date the fleet was created, last time the fleet was modified, Amazon S3 bucket URI, AWS KMS key ID (if provided), AWS IoT alias (if provided), and IAM role. If you added tags, they appear in the **Device fleet tags** section.

### Package a Model (Boto3)

You can create a SageMaker Edge Manager packaging job with the AWS SDK for Python (Boto3). Before continuing, make sure you have satisfied the [Prerequisites](#).

To request an edge packaging job, use `CreateEdgePackagingJob`. You need to provide a name to your edge packaging job, the name of your SageMaker Neo compilation job, your role Amazon resource name (ARN), a name for your model, a version for your model, and the Amazon S3 bucket URI where you want to store the output of your packaging job. Note that Edge Manager packaging job names and SageMaker Neo compilation job names are case-sensitive.

```python
# Import AWS SDK for Python (Boto3)
import boto3

# Create Edge client so you can submit a packaging job
sagemaker_client = boto3.client("sagemaker", region_name='aws-region')

sagemaker_client.create_edge_packaging_job(
    EdgePackagingJobName="edge-packaging-name",
    CompilationJobName="neo-compilation-name",
    RoleArn="arn:aws:iam::99999999999:role/rolename",
    ModelName="sample-model-name",
    ModelVersion="model-version",
    OutputConfig={
        "S3OutputLocation": "s3://your-bucket/",
    }
)
```

You can check the status of an edge packaging job using `DescribeEdgePackagingJob` and providing the case-sensitive edge packaging job name:
response = sagemaker_client.describe_edge_packaging_job(
    EdgePackagingJobName="edge-packaging-name")

This returns a dictionary that can be used to poll the status of the packaging job:

# Optional - Poll every 30 sec to check completion status
import time

while True:
    response = sagemaker_client.describe_edge_packaging_job(
        EdgePackagingJobName="edge-packaging-name")

    if response['EdgePackagingJobStatus'] == 'Completed':
        break
    elif response['EdgePackagingJobStatus'] == 'Failed':
        raise RuntimeError('Packaging job failed')
    print('Packaging model...')
    time.sleep(30)
    print('Done!')

For a list of packaging jobs, use ListEdgePackagingJobs. You can use this API to search for a specific packaging job. Provide a partial name to filter packaging job names for NameContains, a partial name for ModelNameContains to filter for jobs in which the model name contains the name you provide. Also specify with which column to sort with SortBy, and by which direction to sort for SortOrder (either Ascending or Descending).

sagemaker_client.list_edge_packaging_jobs(
    "NameContains": "sample",
    "ModelNameContains": "sample",
    "SortBy": "column-name",
    "SortOrder": "Descending"
)

To stop a packaging job, use StopEdgePackagingJob and provide the name of your edge packaging job.

sagemaker_client.stop_edge_packaging_job(
    EdgePackagingJobName="edge-packaging-name"
)
For a full list of Edge Manager APIs, see the Boto3 documentation.

**The Edge Manager Agent**

The Edge Manager agent is an inference engine for your edge devices. Use the agent to make predictions with models loaded onto your edge devices. The agent also collects model metrics and captures data at specific intervals. Sample data is stored in your Amazon S3 bucket.

There are two methods of installing and deploying the Edge Manager agent onto your edge devices:

1. Download the agent as a binary from the Amazon S3 release bucket. For more information, see [Download and Set Up the Edge Manager Agent Manually](#).
2. Use the AWS IoT Greengrass V2 console or the AWS CLI to deploy `aws.greengrass.SageMakerEdgeManager`. See [Create the AWS IoT Greengrass V2 Components](#).

**Download and Set Up the Edge Manager Agent Manually**

Download the Edge Manager agent based on your operating system, architecture, and AWS Region. The agent is periodically updated, so you have the option to choose your agent based on release dates and versions. Once you have the agent, create a JSON configuration file. Specify the device IoT thing name, fleet name, device credentials, and other key-value pairs. See [Running the Edge Manager agent](#) for full a list of keys you must specify in the configuration file. You can run the agent as an executable binary or link against it as a dynamic shared object (DSO).

**How the agent works**

The agent runs on the CPU of your devices. The agent runs inference on the framework and hardware of the target device you specified during the compilation job. For example, if you compiled your model for the Jetson Nano, the agent supports the GPU in the provided Deep Learning Runtime (DLR).

The agent is released in binary format for supported operating systems. Check that your operating system is supported and meets the minimum OS requirement in the following table:

**Linux**

**Version**: Ubuntu 18.04
**Supported Binary Formats:** x86-64 bit (ELF binary) and ARMv8 64 bit (ELF binary)

**Windows**

**Version:** Windows 10 version 1909

**Supported Binary Formats:** x86-32 bit (DLL) and x86-64 bit (DLL)

### Installing the Edge Manager agent

To use the Edge Manager agent, you first must obtain the release artifacts and a root certificate. The release artifacts are stored in an Amazon S3 bucket in the us-west-2 Region. To download the artifacts, specify your operating system (<OS>) and the <VERSION>.

Based on your operating system, replace <OS> with one of the following:

<table>
<thead>
<tr>
<th>Windows 32-bit</th>
<th>Windows 64-bit</th>
<th>Linux x86-64</th>
<th>Linux ARMv8</th>
</tr>
</thead>
<tbody>
<tr>
<td>windows-x86</td>
<td>windows-x64</td>
<td>linux-x64</td>
<td>linux-armv8</td>
</tr>
</tbody>
</table>

The VERSION is broken into three components: `<MAJOR_VERSION>.<YYYY-MM-DD>-<SHA-7>`, where:

- `<MAJOR_VERSION>`: The release version. The release version is currently set to 1.
- `<YYYY-MM-DD>`: The time stamp of the artifacts release.
- `<SHA-7>`: The repository commit ID from which the release is built.

You must provide the `<MAJOR_VERSION>` and the time stamp in YYYY-MM-DD format. We suggest you use the latest artifact release time stamp.

Run the following in your command line to get the latest time stamp. Replace <OS> with your operating system:

```bash
aws s3 ls s3://sagemaker-edge-release-store-us-west-2-<OS>/Releases/ | sort -r
```

For example, if you have a Windows 32-bit OS, run:

```bash
aws s3 ls s3://sagemaker-edge-release-store-us-west-2-windows-x86/Releases/ | sort -r
```
This returns:

```
2020-12-01 23:33:36 0
```

```text
PRE 1.20201218.81f481f/
PRE 1.20201207.02d0e97/
```

The return output in this example shows two release artifacts. The first release artifact file notes that the release version has a major release version of 1, a time stamp of 20201218 (in YYYY-MM-DD format), and a 81f481f SHA-7 commit ID.

**Note**

The preceding command assumes you have configured the AWS Command Line Interface. For more information, about how to configure the settings that the AWS CLI uses to interact with AWS, see [Configuring the AWS CLI](#).

Based on your operating system, use the following commands to install the artifacts:

**Windows 32-bit**

```
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-windows-x86/Releases/<VERSION>/<VERSION>.zip .
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-windows-x86/Releases/<VERSION>/sha256_hex.shasum .
```

**Windows 64-bit**

```
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-windows-x64/Releases/<VERSION>/<VERSION>.zip .
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-windows-x64/Releases/<VERSION>/sha256_hex.shasum .
```

**Linux x86-64**

```
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-linux-x64/Releases/<VERSION>/<VERSION>.tgz .
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-linux-x64/Releases/<VERSION>/sha256_hex.shasum .
```
Linux ARMv8

```bash
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-linux-armv8/
    Releases/<VERSION>/<VERSION>.tgz .
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-linux-armv8/
    Releases/<VERSION>/sha256_hex.shasum .
```

You also must download a root certificate. This certificate validates model artifacts signed by AWS before loading them onto your edge devices.

Replace `<OS>` corresponding to your platform from the list of supported operation systems and replace `<REGION>` with your AWS Region.

```bash
aws s3 cp s3://sagemaker-edge-release-store-us-west-2-<OS>/
    Certificates/<REGION>/<REGION>.pem .
```

**Running the Edge Manager agent**

You can run the SageMaker Edge Manager agent as a standalone process in the form of an Executable and Linkable Format (ELF) executable binary or you can link against it as a dynamic shared object (.dll). Linux supports running it as a standalone executable binary and is the preferred mode. Windows supports running it as a shared object (.dll).

On Linux, we recommend that you run the binary via a service that's a part of your initialization (init) system. If you want to run the binary directly, you can do so in a terminal as shown in the following example. If you have a modern OS, there are no other installations necessary prior to running the agent, since all the requirements are statically built into the executable. This gives you flexibility to run the agent on the terminal, as a service, or within a container.

To run the agent, first create a JSON configuration file. Specify the following key-value pairs:

- **sagemaker_edge_core_device_name**: The name of the device. This device name needs to be registered along with the device fleet in the SageMaker Edge Manager console.
- **sagemaker_edge_core_device_fleet_name**: The name of the fleet to which the device belongs.
- **sagemaker_edge_core_region**: The AWS Region associated with the device, the fleet and the Amazon S3 buckets. This corresponds to the Region where the device is registered and where the Amazon S3 bucket is created (they are expected to be the same). The models themselves can be
compiled with SageMaker Neo in a different Region, this configuration is not related to model
compilation Region.

• **sagemaker_edge_core_root_certs_path**: The absolute folder path to root certificates. This
is used to validate the device with the relevant AWS account.

• **sagemaker_edge_provider_aws_ca_cert_file**: The absolute path to Amazon Root CA
certificate (AmazonRootCA1.pem). This is used to validate the device with the relevant AWS
account. AmazonCA is a certificate owned by AWS.

• **sagemaker_edge_provider_aws_cert_file**: The absolute path to AWS IoT signing root
certificate (*.pem.crt).

• **sagemaker_edge_provider_aws_cert_pk_file**: The absolute path to AWS IoT private key.
(*.pem.key).

• **sagemaker_edge_provider_aws_iot_cred_endpoint**: The AWS IoT credentials endpoint
(identifier.iot.region.amazonaws.com). This endpoint is used for credential validation. See
Connecting devices to AWS IoT for more information.

• **sagemaker_edge_provider_provider**: This indicates the implementation of the provider
interface being used. The provider interface communicates with the end network services for
uploads, heartbeats and registration validation. By default this is set to "Aws". We allow custom
implementations of the provider interface. It can be set to None for no provider or Custom for
custom implementation with the relevant shared object path provided.

• **sagemaker_edge_provider_provider_path**: Provides the absolute path to the provider
implementation shared object. (.so or .dll file). The "Aws" provider .dll or .so file is provided with
the agent release. This field is mandatory.

• **sagemaker_edge_provider_s3_bucket_name**: The name of your Amazon S3 bucket (not the
Amazon S3 bucket URI). The bucket must have a sagemaker string within its name.

• **sagemaker_edge_log_verbose** (Boolean.): Optional. This sets the debug log. Select either
True or False.

• **sagemaker_edge_telemetry_libsystemd_path**: For Linux only, systemd implements
the agent crash counter metric. Set the absolute path of libsystemd to turn on the crash
counter metric. You can find the default libsystemd path can be found by running whereis
libsystemd in the device terminal.

• **sagemaker_edge_core_capture_data_destination**: The destination for uploading
capture data. Choose either "Cloud" or "Disk". The default is set to "Disk". Setting it to
"Disk" writes the input and output tensor(s) and auxiliary data to the local file system at your
preferred location of. When writing to "Cloud" use the Amazon S3 bucket name provided in the `sagemaker_edge_provider_s3_bucket_name` configuration.

- `sagemaker_edge_core_capture_data_disk_path`: Set the absolute path in the local file system, into which capture data files are written when "Disk" is the destination. This field is not used when "Cloud" is specified as the destination.

- `sagemaker_edge_core_folder_prefix`: The parent prefix in Amazon S3 where captured data is stored when you specify "Cloud" as the capture data destination (`sagemaker_edge_core_capture_data_disk_path`). Captured data is stored in a sub-folder under `sagemaker_edge_core_capture_data_disk_path` if "Disk" is set as the data destination.

- `sagemaker_edge_core_capture_data_buffer_size` (Integer value): The capture data circular buffer size. It indicates the maximum number of requests stored in the buffer.

- `sagemaker_edge_core_capture_data_batch_size` (Integer value): The capture data batch size. It indicates the size of a batch of requests that are handled from the buffer. This value must be less than `sagemaker_edge_core_capture_data_buffer_size`. A maximum of half the size of the buffer is recommended for batch size.

- `sagemaker_edge_core_capture_data_push_period_seconds` (Integer value): The capture data push period in seconds. A batch of requests in the buffer is handled when there are batch size requests in the buffer, or when this time period has completed (whichever comes first). This configuration sets that time period.

- `sagemaker_edge_core_capture_data_base64_embed_limit`: The limit for uploading capture data in bytes. Integer value.

Your configuration file should look similar to the following example(with your specific values specified). This example uses the default AWS provider("Aws") and does not specify a periodic upload.

```json
{
    "sagemaker_edge_core_device_name": "device-name",
    "sagemaker_edge_core_device_fleet_name": "fleet-name",
    "sagemaker_edge_core_region": "region",
    "sagemaker_edge_core_root_certs_path": "<Absolute path to root certificates>",
    "sagemaker_edge_provider_provider": "Aws",
    "sagemaker_edge_provider_provider_path": "/path/to/libprovider_aws.so",
    "sagemaker_edge_provider_aws_ca_cert_file": "<Absolute path to Amazon Root CA certificate>/AmazonRootCA1.pem",
}
```
"sagemaker_edge_provider_aws_cert_file": "<Absolute path to AWS IoT signing root certificate>/device.pem.crt",
"sagemaker_edge_provider_aws_cert_pk_file": "<Absolute path to AWS IoT private key.>/private.pem.key",
"sagemaker_edge_provider_aws_iot_cred_endpoint": "https://<AWS IoT Endpoint Address>",
"sagemaker_edge_core_capture_data_destination": "Cloud",
"sagemaker_edge_provider_s3_bucket_name": "sagemaker-bucket-name",
"sagemaker_edge_core_folder_prefix": "Amazon S3 folder prefix",
"sagemaker_edge_core_capture_data_buffer_size": 30,
"sagemaker_edge_core_capture_data_batch_size": 10,
"sagemaker_edge_core_capture_data_push_period_seconds": 4000,
"sagemaker_edge_core_capture_data_base64_embed_limit": 2,
"sagemaker_edge_log_verbose": false
}

The release artifact includes a binary executable called `sagemaker_edge_agent_binary` in the `/bin` directory. To run the binary, use the `-a` flag to create a socket file descriptor (.sock) in a directory of your choosing and specify the path of the agent JSON config file you created with the `-c` flag.

```
./sagemaker_edge_agent_binary -a <ADDRESS_TO_SOCKET> -c <PATH_TO_CONFIG_FILE>
```

The following example shows the code snippet with a directory and file path specified:

```
./sagemaker_edge_agent_binary -a /tmp/sagemaker_edge_agent_example.sock -c sagemaker_edge_config.json
```

In this example, a socket file descriptor named `sagemaker_edge_agent_example.sock` is created in the `/tmp` directory and points to a configuration file that is in the same working directory as the agent called `sagemaker_edge_config.json`.

## Deploy the Model Package and Edge Manager Agent with AWS IoT Greengrass

SageMaker Edge Manager integrates AWS IoT Greengrass version 2 to simplify accessing, maintaining, and deploying the Edge Manager agent and model to your devices. Without AWS IoT Greengrass V2, setting up your devices and fleets to use SageMaker Edge Manager requires you to manually copy the Edge Manager agent from an Amazon S3 release bucket. You use the agent to make predictions with models loaded onto your edge devices. With AWS IoT Greengrass V2 and SageMaker Edge Manager integration, you can use AWS IoT Greengrass V2 components.
Components are pre-built software modules that can connect your edge devices to AWS services or third-party service via AWS IoT Greengrass.

You must install the AWS IoT Greengrass Core software onto your device(s) if you want to use AWS IoT Greengrass V2 to deploy the Edge Manager agent and your model. For more information about device requirements and how to set up your devices, see Setting up AWS IoT Greengrass core devices in the AWS IoT Greengrass documentation.

You use the following three components to deploy the Edge Manager agent:

- **A pre-built public component**: SageMaker maintains the public Edge Manager component.

- **A autogenerated private component**: The private component is autogenerated when you package your machine learning model with the CreateEdgePackagingJob API and specify GreengrassV2Component for the Edge Manager API field PresetDeploymentType.

- **A custom component**: This is the inference application that is responsible for preprocessing and making inferences on your device. You must create this component. See either Create a Hello World custom component in the SageMaker Edge Manager documentation or Create custom AWS IoT Greengrass components in the AWS IoT Greengrass documentation for more information on how to create custom components.

**Prerequisites**

SageMaker Edge Manager uses AWS IoT Greengrass V2 to simplify the deployment of the Edge Manager agent, your machine learning models, and your inference application to your devices with the use of components. To make it easier to maintain your AWS IAM roles, Edge Manager allows you to reuse your existing AWS IoT role alias. If you do not have one yet, Edge Manager generates a role alias as part of the Edge Manager packaging job. You no longer need to associate a role alias generated from the SageMaker Edge Manager packaging job with your AWS IoT role.

Before you start, you must complete the following prerequisites:

1. Install the AWS IoT Greengrass Core software. For detailed information, see Install the AWS IoT Greengrass Core software.

2. Set up AWS IoT Greengrass V2. For more information, see Install AWS IoT Greengrass Core software with manual resource provisioning.
3. Attach the following permission and inline policy to the IAM role created during AWS IoT Greengrass V2 setup.

- Navigate to the IAM console [https://console.aws.amazon.com/iam/](https://console.aws.amazon.com/iam/).
- Search for the role you created by typing in the role name in the **Search** field.
- Choose your role.
- Next, choose **Attach policies**.
- Search for **AmazonSageMakerEdgeDeviceFleetPolicy**.
- Select **AmazonSageMakerFullAccess** (This is an optional step that makes it easier for you to reuse this IAM role in model compilation and packaging).
- Add required permissions to a role's permissions policy, don't attach inline policies to IAM users.

```json
{
    "Version":"2012-10-17",
    "Statement": [
        {
            "Sid":"GreengrassComponentAccess",
            "Effect":"Allow",
            "Action": [
                "greengrass:CreateComponentVersion",
                "greengrass:DescribeComponent"
            ],
            "Resource": "*"
        }
    ]
}
```
- Choose **Attach policy**.
- Choose **Trust relationship**.
- Choose **Edit trust relationship**.

**Note**

- Make sure the AWS IoT thing name is all lowercase and does not contain characters except (optionally) dashes (-).
- The IAM Role must start with **SageMaker**.
4. Create an Edge Manager device fleet. For information on how to create a fleet, see Set Up Devices and Fleets.

5. Register your device with the same name as your AWS IoT thing name created during the AWS IoT Greengrass V2 setup.

6. Create at least one custom private AWS IoT Greengrass component. This component is the application that runs inference on the device. For more information, see Create a Hello World custom component

Note

- The SageMaker Edge Manager and AWS IoT Greengrass integration only works for AWS IoT Greengrass v2.
- Both your AWS IoT thing name and Edge Manager device name must be the same.
- SageMaker Edge Manager does not load local AWS IoT certificates and call the AWS IoT credential provider endpoint directly. Instead, SageMaker Edge Manager uses the AWS
IoT Greengrass v2 TokenExchangeService and it fetches a temporary credential from a TES endpoint.

Create the AWS IoT Greengrass V2 Components

AWS IoT Greengrass uses components, a software module that is deployed to and runs on a AWS IoT Greengrass core device. You need (at a minimum) three components:

1. A public Edge Manager Agent AWS IoT Greengrass component which deploys the Edge Manager agentbinary.

2. A model component that is autogenerated when you package your machine learning model with either the AWS SDK for Python (Boto3) API or with the SageMaker console. For information, see Create an autogenerated component.

3. A private, custom component to implement the Edge Manager agent client application, and do any preprocessing and post-processing of the inference results. For more information about how to create a custom component, see Create an autogenerated component or Create custom AWS IoT Greengrass components.

Create an autogenerated component

Generate the model component with the CreateEdgePackagingJob API and specify GreengrassV2Component for the SageMaker Edge Manager packaging job API field PresetDeploymentType. When you call the CreateEdgePackagingJob API, Edge Manager takes your SageMaker Neo–compiled model in Amazon S3 and creates a model component. The model component is automatically stored in your account. You can view any of your components by navigating to the AWS IoT console https://console.aws.amazon.com/iot/. Select Greengrass and then select Core devices. The page has a list of AWS IoT Greengrass core devices associated with your account. If a model component name is not specified in PresetDeploymentConfig, the default name generated consists of "SagemakerEdgeManager" and the name of your Edge Manager agent packaging job. The following example demonstrates how to specify to Edge Manager to create a AWS IoT Greengrass V2 component with the CreateEdgePackagingJob API.

```python
import sagemaker
import boto3

# Create a SageMaker client object to make it easier to interact with other AWS services.
```
sagemaker_client = boto3.client('sagemaker', region=<YOUR_REGION>)

# Replace with your IAM Role ARN
sagemaker_role_arn = "arn:aws:iam::<account>::role/*"

# Replace string with the name of your already created S3 bucket.
bucket = 'edge-manager-demo-bucket'

# Specify a name for your edge packaging job.
edge_packaging_name = "edge_packag_job_demo"

# Replace the following string with the name you used for the SageMaker Neo compilation job.
compilation_job_name = "getting-started-demo"

# The name of the model and the model version.
model_name = "sample-model"
model_version = "1.1"

# Output directory in S3 where you want to store the packaged model.
packaging_output_dir = 'packaged_models'
packaging_s3_output = 's3://{}/{}'.format(bucket, packaging_output_dir)

# The name you want your Greengrass component to have.
component_name = "SagemakerEdgeManager" + edge_packaging_name

sagemaker_client.create_edge_packaging_job(
    EdgePackagingJobName=edge_packaging_name,
    CompilationJobName=compilation_job_name,
    RoleArn=sagemaker_role_arn,
    ModelName=model_name,
    ModelVersion=model_version,
    OutputConfig={
        "S3OutputLocation": packaging_s3_output,
        "PresetDeploymentType": "GreengrassV2Component",
        "PresetDeploymentConfig": "{"ComponentName": "sample-component-name", "ComponentVersion": "1.0.2"}"
    }
)

You can also create the autogenerated component with the SageMaker console. Follow steps 1-6 in [Package a Model (Amazon SageMaker Console)](https://docs.aws.amazon.com/sagemaker/latest/dg/packaging-edge-models.html)
Enter the Amazon S3 bucket URI where you want to store the output of the packaging job and optional encryption key.

Complete the following to create the model component:

1. Choose **Preset deployment**.
2. Specify the name of the component for the **Component name** field.
3. Optionally, provide a description of the component, a component version, the platform OS, or the platform architecture for the **Component description**, **Component version**, **Platform OS**, and **Platform architecture**, respectively.
4. Choose **Submit**.

**Create a Hello World custom component**

The custom application component is used to perform inference on the edge device. The component is responsible for loading models to SageMaker Edge Manager, invoking the Edge Manager agent for inference, and unloading the model when the component is shut down. Before you create your component, ensure the agent and application can communicate with Edge Manager. To do this, configure **gRPC**. The Edge Manager agent uses methods defined in Protobuf Buffers and the gRPC server to establish communication with the client application on the edge device and the cloud.

To use gRPC, you must:

1. Create a gRPC stub using the .proto file provided when you download the Edge Manager agent from Amazon S3 release bucket.
2. Write client code with the language you prefer.

You do not need to define the service in a .proto file. The service .proto files are included in the compressed TAR file when you download the Edge Manager agent release binary from the Amazon S3 release bucket.

Install gRPC and other necessary tools on your host machine and create the gRPC stubs `agent_pb2_grpc.py` and `agent_pb2.py` in Python. Make sure you have `agent.proto` in your local directory.

```bash
pip install grpcio
```
The preceding code generates the gRPC client and server interfaces from your .proto service definition. In other words, it creates the gRPC model in Python. The API directory contains the Protobuf specification for communicating with the agent.

Next, use the gRPC API to write a client and server for your service (2). The following example script, edge_manager_python_example.py, uses Python to load, list, and unload a yolov3 model to the edge device.

```python
import grpc
from PIL import Image
import agent_pb2
import agent_pb2_grpc
import os

model_path = '<PATH-TO-SagemakerEdgeManager-COMPONENT>'
agent_socket = 'unix:///tmp/aws.greengrass.SageMakerEdgeManager.sock'
agent_channel = grpc.insecure_channel(agent_socket, options=(("grpc.enable_http_proxy", 0),))
agent_client = agent_pb2_grpc.AgentStub(agent_channel)

def list_models():
    return agent_client.ListModels(agent_pb2.ListModelsRequest())

def list_model_tensors(models):
    return {
        model.name: {
            'inputs': model.input_tensor_metadatas,
            'outputs': model.output_tensor_metadatas
        }
        for model in list_models().models
    }
```
def load_model(model_name, model_path):
    load_request = agent_pb2.LoadModelRequest()
    load_request.url = model_path
    load_request.name = model_name
    return agent_client.LoadModel(load_request)

def unload_model(name):
    unload_request = agent_pb2.UnLoadModelRequest()
    unload_request.name = name
    return agent_client.UnLoadModel(unload_request)

def predict_image(model_name, image_path):
    image_tensor = agent_pb2.Tensor()
    image_tensor.byte_data = Image.open(image_path).tobytes()
    image_tensor_metadata = list_model_tensors(list_models())[model_name]['inputs'][0]
    image_tensor.tensor_metadata.name = image_tensor_metadata.name
    image_tensor.tensor_metadata.data_type = image_tensor_metadata.data_type
    for shape in image_tensor_metadata.shape:
        image_tensor.tensor_metadata.shape.append(shape)
    predict_request = agent_pb2.PredictRequest()
    predict_request.name = model_name
    predict_request.tensors.append(image_tensor)
    predict_response = agent_client.Predict(predict_request)
    return predict_response

def main():
    try:
        unload_model('your-model')
    except:
        pass

    print('LoadModel...', end='')
    try:
        load_model('your-model', model_path)
        print('done.')
    except Exception as e:
        print()
        print(e)
        print('Model already loaded!')

    print('ListModels...', end='')
    try:
print(list_models())
print('done. ')

except Exception as e:
    print()
    print(e)
    print('List model failed! ')

print('Unload model... ', end=' ')
try:
    unload_model('your-model')
    print('done. ')
except Exception as e:
    print()
    print(e)
    print('unload model failed! ')

if __name__ == '__main__':
    main()

Ensure model_path points to the name of the AWS IoT Greengrass component containing the model if you use the same client code example.

You can create your AWS IoT Greengrass V2 Hello World component once you have generated your gRPC stubs and you have your Hello World code ready. To do so:

- Upload your edge_manager_python_example.py, agent_pb2_grpc.py, and agent_pb2.py to your Amazon S3 bucket and note down their Amazon S3 path.
- Create a private component in the AWS IoT Greengrass V2 console and define the recipe for your component. Specify the Amazon S3 URI to your Hello World application and gRPC stub in the following recipe.

```---
RecipeFormatVersion: 2020-01-25
ComponentName: com.sagemaker.edgePythonExample
ComponentVersion: 1.0.0
ComponentDescription: Sagemaker Edge Manager Python example
ComponentPublisher: Amazon Web Services, Inc.
ComponentDependencies:
    aws.greengrass.SageMakerEdgeManager:
        VersionRequirement: '>=1.0.0'
        DependencyType: HARD
```
Manifests:
- Platform:
  os: linux
  architecture: "/amd64|x86/"
Lifecycle:
  install: |
    apt-get install python3-pip
    pip3 install grpcio
    pip3 install grpcio-tools
    pip3 install protobuf
    pip3 install Pillow
  run:
    script: |
      python3 {artifacts:path}/edge_manager_python_example.py
Artifacts:
- URI: <code-s3-path>
- URI: <pb2-s3-path>
- URI: <pb2-grpc-s3-path>

For detailed information about creating a Hello World recipe, see Create your first component in the AWS IoT Greengrass documentation.

Deploy the components to your device

Deploy your components with the AWS IoT console or with the AWS CLI.

To deploy your components (console)

Deploy your AWS IoT Greengrass components with the AWS IoT console.

1. In the AWS IoT Greengrass console at https://console.aws.amazon.com/iot/ navigation menu, choose Deployments.
2. On the Components page, on the Public components tab, choose aws.greengrass.SageMakerEdgeManager.
4. From Add to deployment, choose one of the following:
   a. To merge this component to an existing deployment on your target device, choose Add to existing deployment, and then select the deployment that you want to revise.
b. To create a new deployment on your target device, choose Create new deployment. If you have an existing deployment on your device, choosing this step replaces the existing deployment.

5. On the Specify target page, do the following:
   a. Under Deployment information, enter or modify the friendly name for your deployment.
   b. Under Deployment targets, select a target for your deployment, and choose Next. You cannot change the deployment target if you are revising an existing deployment.

6. On the Select components page, under My components, choose:
   - com.<CUSTOM-COMPONENT-NAME>
   - aws.greengrass.SageMakerEdgeManager
   - SagemakerEdgeManager.<YOUR-PACKAGING-JOB>

7. On the Configure components page, choose com.greengrass.SageMakerEdgeManager, and do the following.
   a. Choose Configure component.
   b. Under Configuration update, in Configuration to merge, enter the following configuration.

   ```json
   {
   "DeviceFleetName": "device-fleet-name",
   "BucketName": "DOC-EXAMPLE-BUCKET"
   }
   ``
   Replace device-fleet-name with the name of the edge device fleet that you created, and replace DOC-EXAMPLE-BUCKET with the name of the Amazon S3 bucket that is associated with your device fleet.
   c. Choose Confirm, and then choose Next.

8. On the Configure advanced settings page, keep the default configuration settings, and choose Next.


To deploy your components (AWS CLI)

1. Create a deployment.json file to define the deployment configuration for your SageMaker Edge Manager components. This file should look like the following example.
```json
{
  "targetArn":"targetArn",
  "components": {
    "aws.greengrass.SageMakerEdgeManager": {
      "componentVersion": 1.0.0,
      "configurationUpdate": {
        "merge": {
          "DeviceFleetName": "device-fleet-name",
          "BucketName": "DOC-EXAMPLE-BUCKET"
        }
      }
    }
  }
}
```

- In the `targetArn` field, replace `targetArn` with the Amazon Resource Name (ARN) of the thing or thing group to target for the deployment, in the following format:
  - Thing: `arn:aws:iot:region:account-id:thing/thingName`
  - Thing group: `arn:aws:iot:region:account-id:thinggroup/thingGroupName`

- In the `merge` field, replace `device-fleet-name` with the name of the edge device fleet that you created, and replace `DOC-EXAMPLE-BUCKET` with the name of the Amazon S3 bucket that is associated with your device fleet.

- Replace the component versions for each component with the latest available version.

2. Run the following command to deploy the components on the device:

```
aws greengrassv2 create-deployment \
  --cli-input-json file://path/to/deployment.json
```
The deployment can take several minutes to complete. In the next step, check the component log to verify that the deployment completed successfully and to view the inference results.

For more information about deploying components to individual devices or groups of devices, see [Deploy AWS IoT Greengrass components to devices](#).

### Deploy the Model Package Directly with SageMaker Edge Manager Deployment API

SageMaker Edge Manager provides a deployment API that you can use to deploy models to device targets without AWS IoT Greengrass. It is useful in situations where you want to update models independently of firmware updates or application deployment mechanisms. You can use the API to integrate your edge deployments into a CI/CD workflow to automatically deploy models once you have validated your model for accuracy. The API also has convenient rollback and staged rollout options for you to ensure models work well in a particular environment before wider rollout.

To use the Edge Manager deployment API first compile and package your model. For information on how to compile and package your model, see [Train, Compile, and Package Your Model](#). The following sections of this guide show how you can create edge deployments using SageMaker API, after you have compiled and packaged your models.

**Topics**

- [Create an edge deployment plan](#)
- [Start the edge deployment](#)
- [Check the status of the deployment](#)

### Create an edge deployment plan

You can create an edge deployment plan with the `CreateEdgeDeploymentPlan` API. The deployment plan can have multiple stages. You can configure each stage to rollout the deployment to a subset of edge devices (by percentage, or by device name). You can also configure how rollout failures are handled at each stage.

The following code snippet shows how you can create an edge deployment plan with 1 stage to deploy a compiled and package model to 2 specific edge devices:

```python
import boto3
```
```python
client = boto3.client("sagemaker")

client.create_edge_deployment_plan(
    EdgeDeploymentPlanName="edge-deployment-plan-name",
    DeviceFleetName="device-fleet-name",
    ModelConfigs=[
        {
            "EdgePackagingJobName": "edge-packaging-job-name",
            "ModelHandle": "model-handle"
        }
    ],
    Stages=[
        {
            "StageName": "stage-name",
            "DeviceSelectionConfig": {
                "DeviceSubsetType": "SELECTION",
                "DeviceNames": ["device-name-1", "device-name-2"]
            },
            "DeploymentConfig": {
                "FailureHandlingPolicy": "ROLLBACK_ON_FAILURE"
            }
        }
    ]
)
```

Instead of specific devices, if you want to deploy to the model to a percentage of devices in your fleet, then set the value of DeviceSubsetType to "PERCENTAGE" and replace "DeviceNames": ["device-name-1", "device-name-2"] with "Percentage": desired-percentage in the above example.

Stages can be added after the deployment plan has been created with the CreateEdgeDeploymentStage API, in case you want to start rolling out new stages after validating your test rollout success. For more information about deployment stages see DeploymentStage.

**Start the edge deployment**

After creating the deployment plan and the deployment stages, you can start the deployment with the StartEdgeDeploymentStage API.

```python
client.start_edge_deployment_stage(
```

The Edge Manager Agent
Check the status of the deployment

You can check the status of the edge deployment with the `DescribeEdgeDeploymentPlan` API.

```python
client.describe_edge_deployment_plan(
    EdgeDeploymentPlanName="edge-deployment-plan-name"
)
```

Manage Model

The Edge Manager agent can load multiple models at a time and make inference with loaded models on edge devices. The number of models the agent can load is determined by the available memory on the device. The agent validates the model signature and loads into memory all the artifacts produced by the edge packaging job. This step requires all the required certificates described in previous steps to be installed along with rest of the binary installation. If the model's signature cannot be validated, then loading of the model fails with appropriate return code and reason.

SageMaker Edge Manager agent provides a list of Model Management APIs that implement control plane and data plane APIs on edge devices. Along with this documentation, we recommend going through the sample client implementation which shows canonical usage of the below described APIs.

The proto file is available as a part of the release artifacts (inside the release tarball). In this doc, we list and describe the usage of APIs listed in this proto file.

Note

There is one-to-one mapping for these APIs on Windows release and a sample code for an application implement in C# is shared with the release artifacts for Windows. Below instructions are for running the Agent as a standalone process, applicable for to the release artifacts for Linux.
Extract the archive based on your OS. Where VERSION is broken into three components: 
<MAJOR_VERSION>.<YYYY-MM-DD>-.<SHA-7>. See Installing the Edge Manager agent for information on how to obtain the release version (<MAJOR_VERSION>), time stamp of the release artifact (<YYYY-MM-DD>), and the repository commit ID (SHA-7)

Linux

The zip archive can be extracted with the command:

```
tar -xvzf <VERSION>.tgz
```

Windows

The zip archive can be extracted with the UI or command:

```
unzip <VERSION>.tgz
```

The release artifact hierarchy (after extracting the tar/zip archive) is shown below. The agent proto file is available under api/.

```
0.20201205.7ee4b0b
### bin
  #    ### sagemaker_edge_agent_binary
  #    ### sagemaker_edge_agent_client_example
### docs
### api
  ### agent.proto
### attributions
  #    ### agent.txt
  #    ### core.txt
### examples
### ipc_example
### CMakeLists.txt
### sagemaker_edge_client.cc
### sagemaker_edge_client_example.cc
### sagemaker_edge_client.hh
### sagemaker_edge.proto
### README.md
### shm.cc
### shm.hh
### street_small.bmp
```
Load Model

The Edge Manager agent supports loading multiple models. This API validates the model signature and loads into memory all the artifacts produced by the EdgePackagingJob operation. This step requires all the required certificates to be installed along with rest of the agent binary installation. If the model's signature cannot be validated then this step fails with appropriate return code and error messages in the log.

```plaintext
// perform load for a model
// Note:
// 1. currently only local filesystem paths are supported for loading models.
// 2. multiple models can be loaded at the same time, as limited by available device memory
// 3. users are required to unload any loaded model to load another model.
// Status Codes:
// 1. OK - load is successful
// 2. UNKNOWN - unknown error has occurred
// 3. INTERNAL - an internal error has occurred
// 4. NOT_FOUND - model doesn't exist at the url
// 5. ALREADY_EXISTS - model with the same name is already loaded
// 6. RESOURCE_EXHAUSTED - memory is not available to load the model
// 7. FAILED_PRECONDITION - model is not compiled for the machine.
// rpc LoadModel(LoadModelRequest) returns (LoadModelResponse);
```

Input

```plaintext
//
// request for LoadModel rpc call
```
message LoadModelRequest {
  string url = 1;
  string name = 2;  // Model name needs to match regex "^[a-zA-Z0-9]-*[a-zA-Z0-9]*$"
}

Output

message LoadModelResponse {
  Model model = 1;
}

// Model represents the metadata of a model
// url - url representing the path of the model
// name - name of model
// input_tensor_metadatas - TensorMetadata array for the input tensors
// output_tensor_metadatas - TensorMetadata array for the output tensors
// Note:
// 1. input and output tensor metadata could empty for dynamic models.
message Model {
  string url = 1;
  string name = 2;
  repeated TensorMetadata input_tensor_metadatas = 3;
  repeated TensorMetadata output_tensor_metadatas = 4;
}

Unload Model

Unloads a previously loaded model. It is identified via the model alias which was provided during loadModel. If the alias is not found or model is not loaded then returns error.

// // perform unload for a model // Status Codes:
Input

```
// request for UnLoadModel rpc call
//
message UnLoadModelRequest {
  string name = 1; // Model name needs to match regex "^[a-zA-Z0-9\-*a-zA-Z0-9]*$"
}
```

Output

```
// response for UnLoadModel rpc call
//
message UnLoadModelResponse {}  
```

**List Models**

Lists all the loaded models and their aliases.

```
// lists the loaded models
// Status Codes:
// 1. OK - unload is successful
// 2. UNKNOWN - unknown error has occurred
// 3. INTERNAL - an internal error has occurred
//
rpc ListModels(ListModelsRequest) returns (ListModelsResponse);
```

Input

```
// request for ListModels rpc call
```
Describe Model

Describes a model that is loaded on the agent.

// Status Codes:
// 1. OK - load is successful
// 2. UNKNOWN - unknown error has occurred
// 3. INTERNAL - an internal error has occurred
// 4. NOT_FOUND - model doesn't exist at the url
//
// rpc DescribeModel(DescribeModelRequest) returns (DescribeModelResponse);

Input

//
// request for DescribeModel rpc call
//
message DescribeModelRequest {
  string name = 1;
}

Output

//
// response for DescribeModel rpc call
//
message DescribeModelResponse {
  Model model = 1;
}
Capture Data

Allows the client application to capture input and output tensors in Amazon S3 bucket, and optionally the auxiliary. The client application is expected to pass a unique capture ID along with each call to this API. This can be later used to query status of the capture.

```java
// allows users to capture input and output tensors along with auxiliary data.
// Status Codes:
// 1. OK - data capture successfully initiated
// 2. UNKNOWN - unknown error has occurred
// 3. INTERNAL - an internal error has occurred
// 5. ALREADY_EXISTS - capture initiated for the given capture_id
// 6. RESOURCE_EXHAUSTED - buffer is full cannot accept any more requests.
// 7. OUT_OF_RANGE - timestamp is in the future.
// 8. INVALID_ARGUMENT - capture_id is not of expected format.
//
rpc CaptureData(CaptureDataRequest) returns (CaptureDataResponse);
```

Input

```java
enum Encoding {
    CSV = 0;
    JSON = 1;
    NONE = 2;
    BASE64 = 3;
}

// AuxiliaryData represents a payload of extra data to be capture along with inputs and outputs of inference
// encoding - supports the encoding of the data
// data - represents the data of shared memory, this could be passed in two ways:
// a. send across the raw bytes of the multi-dimensional tensor array
// b. send a SharedMemoryHandle which contains the posix shared memory segment id and
// offset in bytes to location of multi-dimensional tensor array.
//
message AuxiliaryData {
```
```java
string name = 1;
    Encoding encoding = 2;
    oneof data {
        bytes byte_data = 3;
        SharedMemoryHandle shared_memory_handle = 4;
    }
}

//
// Tensor represents a tensor, encoded as contiguous multi-dimensional array.
// tensor_metadata - represents metadata of the shared memory segment
// data_or_handle - represents the data of shared memory, this could be passed in
two ways:
// a. send across the raw bytes of the multi-dimensional tensor array
// b. send a SharedMemoryHandle which contains the posix shared memory segment
// id and offset in bytes to location of multi-dimensional tensor array.
//
message Tensor {
    TensorMetadata tensor_metadata = 1; //optional in the predict request
    oneof data {
        bytes byte_data = 4;
        // will only be used for input tensors
        SharedMemoryHandle shared_memory_handle = 5;
    }
}

//
// request for CaptureData rpc call
//
message CaptureDataRequest {
    string model_name = 1;
    string capture_id = 2; //uuid string
    Timestamp inference_timestamp = 3;
    repeated Tensor input_tensors = 4;
    repeated Tensor output_tensors = 5;
    repeated AuxilaryData inputs = 6;
    repeated AuxilaryData outputs = 7;
}

Output

//
// response for CaptureData rpc call
```
Get Capture Status

Depending on the models loaded the input and output tensors can be large (for many edge devices). Capture to the cloud can be time consuming. So the `CaptureData()` is implemented as an asynchronous operation. A capture ID is a unique identifier that the client provides during capture data call, this ID can be used to query the status of the asynchronous call.

```java
// allows users to query status of capture data operation
// Status Codes:
// 1. OK - data capture successfully initiated
// 2. UNKNOWN - unknown error has occurred
// 3. INTERNAL - an internal error has occurred
// 4. NOT_FOUND - given capture id doesn't exist.
//
// rpc GetCaptureDataStatus(GetCaptureDataStatusRequest) returns (GetCaptureDataStatusResponse);
```

Input

```java
// request for GetCaptureDataStatus rpc call
//
message GetCaptureDataStatusRequest {
  string capture_id = 1;
}
```

Output

```java
enum CaptureDataStatus {
  FAILURE = 0;
  SUCCESS = 1;
  IN_PROGRESS = 2;
  NOT_FOUND = 3;
}

// response for GetCaptureDataStatus rpc call
```
Predict

The `predict` API performs inference on a previously loaded model. It accepts a request in the form of a tensor that is directly fed into the neural network. The output is the output tensor (or scalar) from the model. This is a blocking call.

```
// perform inference on a model.

// Note:
// 1. users can chose to send the tensor data in the protobuf message or
// through a shared memory segment on a per tensor basis, the Predict
// method will handle the decode transparently.
// 2. serializing large tensors into the protobuf message can be quite expensive,
// based on our measurements it is recommended to use shared memory of
// tenors larger than 256KB.
// 3. SMEdge IPC server will not use shared memory for returning output tensors,
// i.e., the output tensor data will always send in byte form encoded
// in the tensors of PredictResponse.
// 4. currently SMEdge IPC server cannot handle concurrent predict calls, all
// these call will be serialized under the hood. this shall be addressed
// in a later release.
// Status Codes:
// 1. OK - prediction is successful
// 2. UNKNOWN - unknown error has occurred
// 3. INTERNAL - an internal error has occurred
// 4. NOT_FOUND - when model not found
// 5. INVALID_ARGUMENT - when tenors types mismatch
//
// rpc Predict(PredictRequest) returns (PredictResponse);
```

Input

```
// request for Predict rpc call
//
message PredictRequest {
```
string name = 1;
repeated Tensor tensors = 2;
}

//
// Tensor represents a tensor, encoded as contiguous multi-dimensional array.
//    tensor_metadata - represents metadata of the shared memory segment
//    data_or_handle - represents the data of shared memory, this could be passed in
two ways:
//                        a. send across the raw bytes of the multi-dimensional
tensor array
//                        b. send a SharedMemoryHandle which contains the posix
shared memory segment
//                            id and offset in bytes to location of multi-
dimensional tensor array.
//
message Tensor {
    TensorMetadata tensor_metadata = 1; //optional in the predict request
    oneof data {
        bytes byte_data = 4;
        // will only be used for input tensors
        SharedMemoryHandle shared_memory_handle = 5;
    }
}

//
// Tensor represents a tensor, encoded as contiguous multi-dimensional array.
//    tensor_metadata - represents metadata of the shared memory segment
//    data_or_handle - represents the data of shared memory, this could be passed in
two ways:
//                        a. send across the raw bytes of the multi-dimensional
tensor array
//                        b. send a SharedMemoryHandle which contains the posix
shared memory segment
//                            id and offset in bytes to location of multi-
dimensional tensor array.
//
message Tensor {
    TensorMetadata tensor_metadata = 1; //optional in the predict request
    oneof data {
        bytes byte_data = 4;
        // will only be used for input tensors
        SharedMemoryHandle shared_memory_handle = 5;
    }
}
// TensorMetadata represents the metadata for a tensor
//    name - name of the tensor
//    data_type - data type of the tensor
//    shape - array of dimensions of the tensor
//
message TensorMetadata {
  string name = 1;
  DataType data_type = 2;
  repeated int32 shape = 3;
}

// SharedMemoryHandle represents a posix shared memory segment
//    offset - offset in bytes from the start of the shared memory segment.
//    segment_id - shared memory segment id corresponding to the posix shared memory segment.
//    size - size in bytes of shared memory segment to use from the offset position.
//
message SharedMemoryHandle {
  uint64 size = 1;
  uint64 offset = 2;
  uint64 segment_id = 3;
}

Output

⚠️ **Note**

The PredictResponse only returns Tensors and not SharedMemoryHandle.

// response for Predict rpc call
//
message PredictResponse {
  repeated Tensor tensors = 1;
}
SageMaker Edge Manager end of life

Starting in April 26, 2024, you can no longer access Amazon SageMaker Edge Manager through the AWS management console, make edge packaging jobs, and manage edge device fleets.

FAQs

Use the following sections to get answers to commonly asked questions about the SageMaker Edge Manager end of life (EOL).

Q: What happens to my Amazon SageMaker Edge Manager after the EOL date?

A: After April 26, 2024, all references to edge packaging jobs, devices, and device fleets are deleted from the Edge Manager service. You can no longer discover or access the Edge Manager service from your AWS console and applications that call on the Edge Manager service APIs no longer work.

Q: Will I be billed for Edge Manager resources remaining in my account after the EOL date?

A: Resources created by Edge Manager, such as edge packages inside Amazon S3 buckets, AWS IoT things, and AWS IAM roles, continue to exist on their respective services after April 26, 2024. To avoid being billed after Edge Manager is no longer supported, delete your resources. For more information on deleting your resources, see Delete Edge Manager resources.

Q: How do I delete my Amazon SageMaker Edge Manager resources?

A: Resources created by Edge Manager, such as edge packages inside Amazon S3 buckets, AWS IoT things, and AWS IAM roles, continue to exist on their respective services after April 26, 2024. To avoid being billed after Edge Manager is no longer supported, delete your resources. For more information on deleting your resources, see Delete Edge Manager resources.

Q: How can I continue deploying models on the edge?

A: We suggest you try one the following machine learning tools. For a cross-platform edge runtime, use ONNX. ONNX is a popular, well-maintained open-source solution that translates your models into instructions that many types of hardware can run, and is compatible with the latest ML frameworks. ONNX can be integrated into your SageMaker workflows as an automated step for your edge deployments.

For edge deployments and monitoring use AWS IoT Greengrass V2. AWS IoT Greengrass V2 has an extensible packaging and deployment mechanism that can fit models and applications at the edge.
You can use the built-in MQTT channels to send model telemetry back for Amazon SageMaker Model Monitor or use the built-in permissions system to send data captured from the model back to Amazon Simple Storage Service (Amazon S3). If you don't or can't use AWS IoT Greengrass V2, we suggest using MQTT and IoT Jobs (C/C++ library) to create a lightweight OTA mechanism to deliver models.

We have prepared [sample code available at this GitHub repository](https://github.com) to help you transition to these suggested tools.

### Delete Edge Manager resources

Resources created by Edge Manager continue to exist after April 26, 2024. To avoid billing, delete these resources.

To delete AWS IoT Greengrass resources, do the following:

1. In the AWS IoT Core console, choose **Greengrass devices** under **Manage**.
2. Choose **Components**.
3. Under **My components**, Edge Manager created components are in the format `SageMakerEdge (EdgePackagingJobName)`. Select the component you want to delete.
4. Then choose **Delete version**.

To delete a AWS IoT role alias, do the following:

1. In the AWS IoT Core console, choose **Security** under **Manage**.
2. Choose **Role aliases**.
3. Edge Manager created role aliases are in the format `SageMakerEdge-{DeviceFleetName}`. Select the role you want to delete.
4. Choose **Delete**.

To delete packaging jobs in Amazon S3 buckets, do the following:

1. In the SageMaker console, choose **Edge Inference**.
2. Choose **Edge packaging jobs**.
3. Select one of the edge packaging jobs. Copy the Amazon S3 URI under **Model artifact** in the **Output configuration** section.
4. In the Amazon S3 console, navigate to the corresponding location, and check if you need to delete the model artifact. To delete the model artifact, select the Amazon S3 object and choose **Delete**.

**Optimize model performance using Neo**

Neo is a capability of Amazon SageMaker that enables machine learning models to train once and run anywhere in the cloud and at the edge.

If you are a first time user of SageMaker Neo, we recommend you check out the [Getting Started with Edge Devices](#) section to get step-by-step instructions on how to compile and deploy to an edge device.

**What is SageMaker Neo?**

Generally, optimizing machine learning models for inference on multiple platforms is difficult because you need to hand-tune models for the specific hardware and software configuration of each platform. If you want to get optimal performance for a given workload, you need to know the hardware architecture, instruction set, memory access patterns, and input data shapes, among other factors. For traditional software development, tools such as compilers and profilers simplify the process. For machine learning, most tools are specific to the framework or to the hardware. This forces you into a manual trial-and-error process that is unreliable and unproductive.

Neo automatically optimizes Gluon, Keras, MXNet, PyTorch, TensorFlow, TensorFlow-Lite, and ONNX models for inference on Android, Linux, and Windows machines based on processors from Ambarella, ARM, Intel, Nvidia, NXP, Qualcomm, Texas Instruments, and Xilinx. Neo is tested with computer vision models available in the model zoos across the frameworks. SageMaker Neo supports compilation and deployment for two main platforms: cloud instances (including Inferentia) and edge devices.

For more information about supported frameworks and cloud instance types you can deploy to, see [Supported Instance Types and Frameworks](#) for cloud instances.

For more information about supported frameworks, edge devices, operating systems, chip architectures, and common machine learning models tested by SageMaker Neo for edge devices, see [Supported Frameworks, Devices, Systems, and Architectures](#) for edge devices.
How it Works

Neo consists of a compiler and a runtime. First, the Neo compilation API reads models exported from various frameworks. It converts the framework-specific functions and operations into a framework-agnostic intermediate representation. Next, it performs a series of optimizations. Then it generates binary code for the optimized operations, writes them to a shared object library, and saves the model definition and parameters into separate files. Neo also provides a runtime for each target platform that loads and executes the compiled model.

You can create a Neo compilation job from either the SageMaker console, the AWS Command Line Interface (AWS CLI), a Python notebook, or the SageMaker SDK. For information on how to compile a model, see Use Neo to Compile a Model. With a few CLI commands, an API invocation, or a few clicks, you can convert a model for your chosen platform. You can deploy the model to a SageMaker endpoint or on an AWS IoT Greengrass device quickly.

Neo can optimize models with parameters either in FP32 or quantized to INT8 or FP16 bit-width.

Topics

- Use Neo to Compile a Model
- Cloud Instances
- Edge Devices
- Troubleshoot Errors

Use Neo to Compile a Model

This section shows how to create, describe, stop, and list compilation jobs. The following options are available in Amazon SageMaker Neo for managing the compilation jobs for machine learning
models: the AWS Command Line Interface, the Amazon SageMaker console, or the Amazon SageMaker SDK.

Topics

- Prepare Model for Compilation
- Compile a Model (AWS Command Line Interface)
- Compile a Model (Amazon SageMaker Console)
- Compile a Model (Amazon SageMaker SDK)

Prepare Model for Compilation

SageMaker Neo requires machine learning models to satisfy specific input data shapes. The input shape required for compilation depends on the deep learning framework you use. Once your model input shape is correctly formatted, save your model according to the requirements below. Once you have a saved model, compress the model artifacts.

Topics

- What input data shapes does SageMaker Neo expect?
- Saving Models for SageMaker Neo

What input data shapes does SageMaker Neo expect?

Before you compile your model, make sure your model is formatted correctly. Neo expects the name and shape of the expected data inputs for your trained model with JSON format or list format. The expected inputs are framework specific.

Below are the input shapes SageMaker Neo expects:

Keras

Specify the name and shape (NCHW format) of the expected data inputs using a dictionary format for your trained model. Note that while Keras model artifacts should be uploaded in NHWC (channel-last) format, DataInputConfig should be specified in NCHW (channel-first) format. The dictionary formats required are as follows:

- For one input: `{'input_1': [1,3,224,224]}`
- For two inputs: `{'input_1': [1,3,224,224], 'input_2': [1,3,224,224]}`
MXNet/ONNX

Specify the name and shape (NCHW format) of the expected data inputs using a dictionary format for your trained model. The dictionary formats required are as follows:

- For one input: `{data':[1,3,1024,1024]}
- For two inputs: `{var1': [1,1,28,28], var2':[1,1,28,28]}

PyTorch

For a PyTorch model, you don't need to provide the name and shape of the expected data inputs if you meet both of the following conditions:

- You created your model definition file by using PyTorch 2.0 or later. For more information about how to create the definition file, see the PyTorch section under Saving Models for SageMaker Neo.
- You are compiling your model for a cloud instance. For more information about the instance types that SageMaker Neo supports, see Supported Instance Types and Frameworks.

If you meet these conditions, SageMaker Neo gets the input configuration from the model definition file (.pt or .pth) that you create with PyTorch.

Otherwise, you must do the following:

Specify the name and shape (NCHW format) of the expected data inputs using a dictionary format for your trained model. Alternatively, you can specify the shape only using a list format. The dictionary formats required are as follows:

- For one input in dictionary format: `{input0':[1,3,224,224]}
- For one input in list format: [[1,3,224,224]]
- For two inputs in dictionary format: `{input0':[1,3,224,224], input1':[1,3,224,224]}
- For two inputs in list format: [[1,3,224,224], [1,3,224,224]]

TensorFlow

Specify the name and shape (NHWC format) of the expected data inputs using a dictionary format for your trained model. The dictionary formats required are as follows:
• For one input: {'input': [1,1024,1024,3]}
• For two inputs: {'data1': [1,28,28,1], 'data2':[1,28,28,1]}

**TFLite**

Specify the name and shape (NHWC format) of the expected data inputs using a dictionary format for your trained model. The dictionary formats required are as follows:

• For one input: {'input':[1,224,224,3]}

**Note**

SageMaker Neo only supports TensorFlow Lite for edge device targets. For a list of supported SageMaker Neo edge device targets, see the SageMaker Neo [Devices](#) page. For a list of supported SageMaker Neo cloud instance targets, see the SageMaker Neo [Supported Instance Types and Frameworks](#) page.

**XGBoost**

An input data name and shape are not needed.

**Saving Models for SageMaker Neo**

The following code examples show how to save your model to make it compatible with Neo. Models must be packaged as compressed tar files (*.tar.gz).

**Keras**

Keras models require one model definition file (.h5).

There are two options for saving your Keras model in order to make it compatible for SageMaker Neo:

1. Export to .h5 format with `model.save("<model-name>", save_format="h5").`
2. Freeze the SavedModel after exporting.

Below is an example of how to export a tf.keras model as a frozen graph (option two):
```python
import os
import tensorflow as tf
from tensorflow.keras.applications.resnet50 import ResNet50
from tensorflow.keras import backend

tf.keras.backend.set_learning_phase(0)
model = tf.keras.applications.ResNet50(weights='imagenet', include_top=False,
    input_shape=(224, 224, 3), pooling='avg')
model.summary()

# Save as a SavedModel
export_dir = 'saved_model/
model.save(export_dir, save_format='tf')

# Freeze saved model
input_node_names = [inp.name.split(':')[0] for inp in model.inputs]
output_node_names = [output.name.split(':')[0] for output in model.outputs]
print("Input names: ", input_node_names)
with tf.Session() as sess:
    loaded = tf.saved_model.load(sess, export_dir=export_dir, tags=["serve"])
    frozen_graph = tf.graph_util.convert_variables_to_constants(sess,

    sess.graph.as_graph_def(),

    tf.io.write_graph(graph_or_graph_def=frozen_graph, logdir=".",
    name="frozen_graph.pb", as_text=False)

import tarfile
tar = tarfile.open("frozen_graph.tar.gz", "w:gz")
tar.add("frozen_graph.pb")
tar.close()
```

⚠️ **Warning**

Do not export your model with the SavedModel class using `model.save(<path>, save_format='tf')`. This format is suitable for training, but it is not suitable for inference.
MXNet

MXNet models must be saved as a single symbol file `*-symbol.json` and a single parameter `*.params` files.

Gluon Models

Define the neural network using the `HybridSequential` Class. This will run the code in the style of symbolic programming (as opposed to imperative programming).

```python
from mxnet import nd, sym
from mxnet.gluon import nn

def get_net():
    net = nn.HybridSequential()  # Here we use the class HybridSequential.
    net.add(nn.Dense(256, activation='relu'),
            nn.Dense(128, activation='relu'),
            nn.Dense(2))
    net.initialize()
    return net

# Define an input to compute a forward calculation.
x = nd.random.normal(shape=(1, 512))
net = get_net()

# During the forward calculation, the neural network will automatically infer
# the shape of the weight parameters of all the layers based on the shape of
# the input.
net(x)

# hybridize model
net.hybridize()
net(x)

# export model
net.export('<model_name>') # this will create model-symbol.json and
# model-0000.params files

import tarfile
tar = tarfile.open('<model_name>.tar.gz', 'w:gz')
for name in ['<model_name>-0000.params', '<model_name>-symbol.json']:
    tar.add(name)
tar.close()
```
Amazon SageMaker

Developer Guide

For more information about hybridizing models, see the MXNet hybridize documentation.
Gluon Model Zoo (GluonCV)
GluonCV model zoo models come pre-hybridized. So you can just export them.
import numpy as np
import mxnet as mx
import gluoncv as gcv
from gluoncv.utils import export_block
import tarfile
net = gcv.model_zoo.get_model('<model_name>', pretrained=True) # For example, choose
<model_name> as resnet18_v1
export_block('<model_name>', net, preprocess=True, layout='HWC')
tar = tarfile.open("<model_name>.tar.gz", "w:gz")
for name in ["<model_name>-0000.params", "<model_name>-symbol.json"]:
tar.add(name)
tar.close()

Non Gluon Models
All non-Gluon models when saved to disk use *-symbol and *.params ﬁles. They are
therefore already in the correct format for Neo.
# Pass the following 3 parameters: sym, args, aux
mx.model.save_checkpoint('<model_name>',0,sym,args,aux) # this will create
<model_name>-symbol.json and <model_name>-0000.params files
import tarfile
tar = tarfile.open("<model_name>.tar.gz", "w:gz")
for name in ["<model_name>-0000.params", "<model_name>-symbol.json"]:
tar.add(name)
tar.close()

PyTorch
PyTorch models must be saved as a deﬁnition ﬁle (.pt or .pth) with input datatype of float32.
Compile Models

4554


To save your model, use the `torch.jit.trace` method followed by the `torch.save` method. This process saves an object to a disk file and by default uses Python pickle (`pickle_module=pickle`) to save the objects and some metadata. Next, convert the saved model to a compressed tar file.

```python
import torchvision
import torch

model = torchvision.models.resnet18(pretrained=True)
model.eval()
inp = torch.rand(1, 3, 224, 224)
model_trace = torch.jit.trace(model, inp)

# Save your model. The following code saves it with the .pth file extension
model_trace.save('model.pth')

# Save as a compressed tar file
import tarfile
with tarfile.open('model.tar.gz', 'w:gz') as f:
    f.add('model.pth')
f.close()
```

If you save your model with PyTorch 2.0 or later, SageMaker Neo derives the input configuration for the model (the name and shape for its input) from the definition file. In that case, you don't need to specify the data input configuration to SageMaker when you compile the model.

If you want to prevent SageMaker Neo from deriving the input configuration, you can set the `_store_inputs` parameter of `torch.jit.trace` to False. If you do this, you must specify the data input configuration to SageMaker when you compile the model.

For more information about the `torch.jit.trace` method, see [TORCH.JIT.TRACE](#) in the PyTorch documentation.

**TensorFlow**

TensorFlow requires one `.pb` or one `.pbtxt` file and a variables directory that contains variables. For frozen models, only one `.pb` or `.pbtxt` file is required.

The following code example shows how to use the `tar` Linux command to compress your model.

Run the following in your terminal or in a Jupyter notebook (if you use a Jupyter notebook, insert the `!` magic command at the beginning of the statement):
# Download SSD_Mobilenet trained model
!wget http://download.tensorflow.org/models/object_detection/ssd_mobilenet_v2_coco_2018_03_29.tar.gz

# unzip the compressed tar file
!tar xvf ssd_mobilenet_v2_coco_2018_03_29.tar.gz

# Compress the tar file and save it in a directory called 'model.tar.gz'
!tar czvf model.tar.gz ssd_mobilenet_v2_coco_2018_03_29/frozen_inference_graph.pb

The command flags used in this example accomplish the following:

- c: Create an archive
- z: Compress the archive with gzip
- v: Display archive progress
- f: Specify the filename of the archive

**Built-In Estimators**

Built-in estimators are either made by framework-specific containers or algorithm-specific containers. Estimator objects for both the built-in algorithm and framework-specific estimator saves the model in the correct format for you when you train the model using the built-in `.fit` method.

For example, you can use `sagemaker.TensorFlow` to define a TensorFlow estimator:

```python
from sagemaker.tensorflow import TensorFlow

estimator = TensorFlow(entry_point='mnist.py',
    role=role,  #param role can be arn of a sagemaker execution
    role
    framework_version='1.15.3',
    py_version='py3',
    training_steps=1000,
    evaluation_steps=100,
    instance_count=2,
    instance_type='ml.c4.xlarge')
```

Then train the model with `.fit` built-in method:
estimator.fit(inputs)

Before finally compiling model with the build in compile_model method:

```python
# Specify output path of the compiled model
output_path = '/'.join(estimator.output_path.split('/')[:-1])

# Compile model
optimized_estimator = estimator.compile_model(target_instance_family='ml_c5',
input_shape={'data':[1, 784]},  # Batch size 1, 3 channels, 224x224 Images.
                               output_path=output_path,
                               framework='tensorflow', framework_version='1.15.3')
```

You can also use the `sagemaker.estimator.Estimator` Class to initialize an estimator object for training and compiling a built-in algorithm with the compile_model method from the SageMaker Python SDK:

```python
import sagemaker
from sagemaker.image_uris import retrieve
sagemaker_session = sagemaker.Session()
aws_region = sagemaker_session.boto_region_name

# Specify built-in algorithm training image
training_image = retrieve(framework='image-classification',
                          region=aws_region, image_scope='training')

training_image = retrieve(framework='image-classification', region=aws_region,
                          image_scope='training')

# Create estimator object for training
estimator = sagemaker.estimator.Estimator(image_uri=training_image,
                                         role=role,  #param role can be arn of a
                                         sagemaker execution role
                                         instance_count=1,
                                         instance_type='ml.p3.8xlarge',
                                         volume_size = 50,
                                         max_run = 360000,
                                         input_mode= 'File',
                                         output_path=s3_training_output_location,
                                         base_job_name='image-classification-training')
```
# Setup the input data_channels to be used later for training.

```python
train_data = sagemaker.inputs.TrainingInput(s3_training_data_location,
                                           content_type='application/x-recordio',
                                           s3_data_type='S3Prefix')

validation_data = sagemaker.inputs.TrainingInput(s3_validation_data_location,
                                                  content_type='application/x-recordio',
                                                  s3_data_type='S3Prefix')

data_channels = {'train': train_data, 'validation': validation_data}
```

# Train model

```python
estimator.fit(inputs=data_channels, logs=True)
```

# Compile model with Neo

```python
optimized_estimator = estimator.compile_model(target_instance_family='ml_c5',
                                              input_shape={'data': [1, 3, 224, 224],
                                                           'softmax_label': [1]},
                                              output_path=s3_compilation_output_location,
                                              framework='mxnet',
                                              framework_version='1.7')
```

For more information about compiling models with the SageMaker Python SDK, see [Compile a Model (Amazon SageMaker SDK)](link).

## Compile a Model (AWS Command Line Interface)

This section shows how to manage Amazon SageMaker Neo compilation jobs for machine learning models using AWS Command Line Interface (CLI). You can create, describe, stop, and list the compilation jobs.

1. **Create a Compilation Job**

   With the [CreateCompilationJob](link) API operation, you can specify the data input format, the S3 bucket in which to store your model, the S3 bucket to which to write the compiled model, and the target hardware device or platform.

   The following table demonstrates how to configure CreateCompilationJob API based on whether your target is a device or a platform.
Device Example

```json
{
    "CompilationJobName": "neo-compilation-job-demo",
    "RoleArn": "arn:aws:iam::<your-account>:role/service-role/AmazonSageMaker-ExecutionRole-yyyymmddThhmmss",
    "InputConfig": {
        "S3Uri": "s3://<your-bucket>/sagemaker.neo-compilation-job-demo-data/train",
        "DataInputConfig": "{data: [1,3,1024,1024]}",
        "Framework": "MXNET"
    },
    "OutputConfig": {
        "S3OutputLocation": "s3://<your-bucket>/sagemaker.neo-compilation-job-demo-data/compile",
        "TargetDevice": "ml_c5"
    },
    "StoppingCondition": {
        "MaxRuntimeInSeconds": 300
    }
}
```

You can optionally specify the framework version you used with the `FrameworkVersion` field if you used the PyTorch framework to train your model and your target device is a ml_* target.

```json
{
    "CompilationJobName": "neo-compilation-job-demo",
    "RoleArn": "arn:aws:iam::<your-account>:role/service-role/AmazonSageMaker-ExecutionRole-yyyymmddThhmmss",
    "InputConfig": {
        "S3Uri": "s3://<your-bucket>/sagemaker.neo-compilation-job-demo-data/train",
        "DataInputConfig": "{data: [1,3,1024,1024]}",
        "Framework": "PYTORCH",
        "FrameworkVersion": "1.6"
    },
    "OutputConfig": {
        "S3OutputLocation": "s3://<your-bucket>/sagemaker.neo-compilation-job-demo-data/compile",
```

Compile Models
# A target device specification example for a ml_c5 instance family
"TargetDevice": "ml_c5",
# When compiling for ml_* instances using PyTorch framework, use the "CompilerOptions" field in
# OutputConfig to provide the correct data type ("dtype") of the model’s input. Default assumed is "float32"
"CompilerOptions": "{'dtype': 'long'}"
},
"StoppingCondition": {
    "MaxRuntimeInSeconds": 300
}
}

## Notes:

- If you saved your model by using PyTorch version 2.0 or later, the `DataInputConfig` field is optional. SageMaker Neo gets the input configuration from the model definition file that you create with PyTorch. For more information about how to create the definition file, see the PyTorch section under Saving Models for SageMaker Neo.
- This API field is only supported for PyTorch.

### Platform Example

```
{
    "CompilationJobName": "neo-test-compilation-job",
    "RoleArn": "arn:aws:iam::<your-account>:role/service-role/AmazonSageMaker-ExecutionRole-yyyymmddThhmmss",
    "InputConfig": {
        "S3Uri": "s3://<your-bucket>/sagemaker/neo-compilation-job-demo-data/train",
        "DataInputConfig": "{'data': [1,3,1024,1024]}",
        "Framework": "MXNET"
    },
    "OutputConfig": {
        "S3OutputLocation": "s3://<your-bucket>/sagemaker/neo-compilation-job-demo-data/compile",
        # A target platform configuration example for a p3.2xlarge instance
        "TargetPlatform": {
```
Amazon SageMaker

"Os": "LINUX",
"Arch": "X86_64",
"Accelerator": "NVIDIA"
},
"CompilerOptions": "{'cuda-ver': '10.0', 'trt-ver': '6.0.1', 'gpu-code': 'sm_70'}"
},
"StoppingCondition": {
"MaxRuntimeInSeconds": 300
}
}

Note

For the OutputConfig API operation, the TargetDevice and TargetPlatform API operations are mutually exclusive. You have to choose one of the two options.

To find the JSON string examples of DataInputConfig depending on frameworks, see What input data shapes Neo expects.

For more information about setting up the configurations, see the InputConfig, OutputConfig, and TargetPlatform API operations in the SageMaker API reference.

2. After you configure the JSON file, run the following command to create the compilation job:

aws sagemaker create-compilation-job \
--cli-input-json file://job.json \
--region us-west-2

# You should get CompilationJobArn

3. Describe the compilation job by running the following command:

aws sagemaker describe-compilation-job \
--compilation-job-name $JOB_NM \
--region us-west-2

4. Stop the compilation job by running the following command:

aws sagemaker stop-compilation-job \

5. List the compilation job by running the following command:

```bash
aws sagemaker list-compilation-jobs \
--region us-west-2
```

### Compile a Model (Amazon SageMaker Console)

You can create an Amazon SageMaker Neo compilation job in the Amazon SageMaker console.

1. In the Amazon SageMaker console, choose **Compilation jobs**, and then choose **Create compilation job**.

2. On the **Create compilation job** page, under **Job name**, enter a name. Then select an **IAM role**.
3. If you don’t have an IAM role, choose **Create a new role**.

4. On the **Create an IAM role** page, choose **Any S3 bucket**, and choose **Create role**.
5. Non PyTorch Frameworks

Within the Input configuration section, enter the full path of the Amazon S3 bucket URI that contains your model artifacts in the Location of model artifacts input field. Your model artifacts must be in a compressed tarball file format (.tar.gz).

For the Data input configuration field, enter the JSON string that specifies the shape of the input data.

For Machine learning framework, choose the framework of your choice.
To find the JSON string examples of input data shapes depending on frameworks, see [What input data shapes Neo expects](#).

**PyTorch Framework**

Similar instructions apply for compiling PyTorch models. However, if you trained with PyTorch and are trying to compile the model for `ml_*` (except `ml_inf`) target, you can optionally specify the version of PyTorch you used.
Input configuration

Amazon SageMaker needs to know where model artifacts are stored, what the shape of the data matrix is, and which machine learning framework to use. Learn more

Location of model artifacts

Amazon SageMaker needs the path to the model artifacts in Amazon S3. To find the path, look in your Amazon S3 directories.

s3://demo-test-neo-model/resnet50.tar.gz

To find a path, go to Amazon S3

Data input configuration

Amazon SageMaker needs to know what the shape of the data matrix is.

{"input" : [1,3,224,224]}

Machine learning framework

Choose the machine learning framework that your model was trained in.

PyTorch

Framework version

Choose the machine learning framework version that your model was trained in.

1.4

1.5

1.6

Notes

- If you saved your model by using PyTorch version 2.0 or later, the Data input configuration field is optional. SageMaker Neo gets the input configuration from the model definition file that you create with PyTorch. For more information about how to create the definition file, see the PyTorch section under Saving Models for SageMaker Neo.

- When compiling for ml_* instances using PyTorch framework, use Compiler options field in Output Configuration to provide the correct data type (dtype) of the model's input. The default is set to "float32".

To find the JSON string examples of input data shapes depending on frameworks, see What input data shapes Neo expects.
Warning

If you specify a Amazon S3 bucket URI path that leads to .pth file, you will receive the following error after starting compilation: ClientError: InputConfiguration: Unable to untar input model. Please confirm the model is a tar.gz file.

6. Go to the Output configuration section. Choose where you want to deploy your model. You can deploy your model to a Target device or a Target platform. Target devices include cloud and edge devices. Target platforms refer to specific OS, architecture, and accelerators you want your model to run on.

For S3 Output location, enter the path to the S3 bucket where you want to store the model. You can optionally add compiler options in JSON format under the Compiler options section.
7. Check the status of the compilation job when started. This status of the job can be found at the top of the **Compilation Job** page, as shown in the following screenshot. You can also check the status of it in the **Status** column.
8. Check the status of the compilation job when completed. You can check the status in the **Status** column as shown in the following screenshot.

![Compilation jobs screenshot](image)

**Compile a Model (Amazon SageMaker SDK)**

You can use the `compile_model` API in the [Amazon SageMaker SDK for Python](https://sagemaker-developer-guide.readthedocs.io/en/stable/) to compile a trained model and optimize it for specific target hardware. The API should be invoked on the estimator object used during model training.

**Note**

You must set `MMS_DEFAULT_RESPONSE_TIMEOUT` environment variable to 500 when compiling the model with MXNet or PyTorch. The environment variable is not needed for TensorFlow.

The following is an example of how you can compile a model using the `trained_model_estimator` object:

```python
# Replace the value of expected_trained_model_input below and
# specify the name & shape of the expected inputs for your trained model
# in json dictionary form
expected_trained_model_input = {'data':[1, 784]}

# Replace the example target_instance_family below to your preferred
# target_instance_family
compiled_model = trained_model_estimator.compile_model(target_instance_family='ml_c5',
```
input_shape=expected_trained_model_input,
output_path='insert s3 output path',
env={'MMS_DEFAULT_RESPONSE_TIMEOUT': '500'})

The code compiles the model, saves the optimized model at output_path, and creates a SageMaker model that can be deployed to an endpoint. Sample notebooks of using the SDK for Python are provided in the Neo Model Compilation Sample Notebooks section.

Cloud Instances

Amazon SageMaker Neo provides compilation support for popular machine learning frameworks such as TensorFlow, PyTorch, MXNet, and more. You can deploy your compiled model to cloud instances and AWS Inferentia instances. For a full list of supported frameworks and instances types, see Supported Instances Types and Frameworks.

You can compile your model in one of three ways: through the AWS CLI, the SageMaker Console, or the SageMaker SDK for Python. See, Use Neo to Compile a Model for more information. Once compiled, your model artifacts are stored in the Amazon S3 bucket URI you specified during the compilation job. You can deploy your compiled model to cloud instances and AWS Inferentia instances using the SageMaker SDK for Python, AWS SDK for Python (Boto3), AWS CLI, or the AWS console.

If you deploy your model using AWS CLI, the console, or Boto3, you must select a Docker image Amazon ECR URI for your primary container. See Neo Inference Container Images for a list of Amazon ECR URIs.

Topics

- Supported Instance Types and Frameworks
- Deploy a Model
- Request Inferences from a Deployed Service
- Inference Container Images

Supported Instance Types and Frameworks

Amazon SageMaker Neo supports popular deep learning frameworks for both compilation and deployment. You can deploy your model to cloud instances, AWS Inferentia instance types, or Amazon Elastic Inference accelerators.
The following describes frameworks SageMaker Neo supports and the target cloud instances you can compile and deploy to. For information on how to deploy your compiled model to a cloud or Inferentia instance, see [Deploy a Model with Cloud Instances](#). For information on how to deploy your compiled model with Elastic Inference accelerators, see [Use EI on Amazon SageMaker Hosted Endpoints](#).

### Cloud Instances

SageMaker Neo supports the following deep learning frameworks for CPU and GPU cloud instances:

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>MXNet</td>
<td>1.8.0</td>
<td>Supports 1.8.0 or earlier</td>
<td>Image Classification, Object Detection, Semantic Segmentation, Pose Estimation, Activity Recognition</td>
<td>One symbol file (.json) and one parameter file (.params)</td>
<td>GluonCV v0.8.0</td>
</tr>
<tr>
<td>ONNX</td>
<td>1.7.0</td>
<td>Supports 1.7.0 or earlier</td>
<td>Image Classification, SVM</td>
<td>One model file (.onnx)</td>
<td></td>
</tr>
<tr>
<td>Keras</td>
<td>2.2.4</td>
<td>Supports 2.2.4 or earlier</td>
<td>Image Classification</td>
<td>One model definition file (.h5)</td>
<td></td>
</tr>
<tr>
<td>PyTorch</td>
<td>1.4, 1.5, 1.6, 1.7, 1.8, 1.12, 1.13, or 2.0</td>
<td>Supports 1.4, 1.5, 1.6, 1.7</td>
<td>Image Classification</td>
<td>One model definition file (.pt or .pth)</td>
<td></td>
</tr>
<tr>
<td>-----------</td>
<td>-------------------</td>
<td>---------------</td>
<td>--------</td>
<td>-------------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>TensorFlow</td>
<td>1.15.3 or 2.9</td>
<td>Supports 1.15.3 and 2.9</td>
<td>Image Classification</td>
<td>For saved models, one .pb or one .pbtxt file and a variables directory that contains variables</td>
<td></td>
</tr>
<tr>
<td>1.8, 1.12, 1.13, and 2.0</td>
<td>Versions 1.13 and 2.0 support Object Detection, Vision Transformer, and HuggingFace</td>
<td></td>
<td>with input dtype of float32</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------</td>
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<td>---------------</td>
<td>--------</td>
<td>--------------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>XGBoost</td>
<td>1.3.3</td>
<td>Supports 1.3.3 or earlier</td>
<td>Decision Trees</td>
<td>One XGBoost model file (.model) where the number of nodes in a tree is less than 2^31</td>
<td></td>
</tr>
</tbody>
</table>

**Note**

"Model Version" is the version of the framework used to train and export the model.

**Instance Types**

You can deploy your SageMaker compiled model to one of the cloud instances listed below:

<table>
<thead>
<tr>
<th>Instance</th>
<th>Compute Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml_c4</td>
<td>Standard</td>
</tr>
<tr>
<td>ml_c5</td>
<td>Standard</td>
</tr>
<tr>
<td>ml_m4</td>
<td>Standard</td>
</tr>
<tr>
<td>ml_m5</td>
<td>Standard</td>
</tr>
<tr>
<td>ml_p2</td>
<td>Accelerated computing</td>
</tr>
</tbody>
</table>
For information on the available vCPU, memory, and price per hour for each instance type, see Amazon SageMaker Pricing.

### Note

When compiling for `ml_*` instances using PyTorch framework, use **Compiler options** field in **Output Configuration** to provide the correct data type (dtype) of the model's input. The default is set to "float32".

### AWS Inferentia

SageMaker Neo supports the following deep learning frameworks for Inf1:

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>MXNet</td>
<td>1.5 or 1.8</td>
<td>Supports 1.8, 1.5 and earlier</td>
<td>Image Classification, Object Detection, Semantic Segmentation, Pose Estimation</td>
<td>One symbol file (.json) and one parameter file (.params)</td>
<td>GluonCV v0.8.0</td>
</tr>
<tr>
<td>-----------</td>
<td>-------------------</td>
<td>---------------</td>
<td>--------</td>
<td>-------------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>PyTorch</td>
<td>1.7, 1.8 or 1.9</td>
<td>Supports 1.9 and earlier</td>
<td>Image Classification</td>
<td>One model definition file (.pt or .pth) with input dtype of float32</td>
<td>n, Activity Recognition</td>
</tr>
<tr>
<td>TensorFlow</td>
<td>1.15 or 2.5</td>
<td>Supports 2.5, 1.15 and earlier</td>
<td>Image Classification</td>
<td>For saved models, one .pb or one .pbtxt file and a variables directory that contains variables For frozen models, only one .pb or .pbtxt file</td>
<td></td>
</tr>
</tbody>
</table>

**Note**

“Model Version” is the version of the framework used to train and export the model.
You can deploy your SageMaker Neo-compiled model to AWS Inferentia-based Amazon EC2 Inf1 instances. AWS Inferentia is Amazon's first custom silicon chip designed to accelerate deep learning. Currently, you can use the ml_inf1 instance to deploy your compiled models.

**AWS Inferentia2 and AWS Trainium**

Currently, you can deploy your SageMaker Neo-compiled model to AWS Inferentia2-based Amazon EC2 Inf2 instances (in US East (Ohio) Region), and to AWS Trainium-based Amazon EC2 Trn1 instances (in US East (N. Virginia) Region). For more information about supported models on these instances, see [Model Architecture Fit Guidelines](#) in the AWS Neuron documentation, and the examples in the [Neuron Github repository](#).

**Amazon Elastic Inference**

SageMaker Neo supports the following deep learning frameworks for Elastic Inference:

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>TensorFlow</td>
<td>2.3.2</td>
<td>Supports 2.3</td>
<td>Image Classification, Object Detection, Semantic Segmentation, Pose Estimation, Activity Recognition</td>
<td>For saved models, one .pb or one .pbtxt file and a variables directory that contains variables. For frozen models, only one .pb or .pbtxt file.</td>
</tr>
</tbody>
</table>

You can deploy your SageMaker Neo-compiled model to an Elastic Inference Accelerator. For more information, see [Use EI on Amazon SageMaker Hosted Endpoints](#).
Deploy a Model

To deploy an Amazon SageMaker Neo-compiled model to an HTTPS endpoint, you must configure and create the endpoint for the model using Amazon SageMaker hosting services. Currently, developers can use Amazon SageMaker APIs to deploy modules on to ml.c5, ml.c4, ml.m5, ml.m4, ml.p3, ml.p2, and ml.inf1 instances.

For Inferentia and Trainium instances, models need to be compiled specifically for those instances. Models compiled for other instance types are not guaranteed to work with Inferentia or Trainium instances.

For Elastic Inference accelerators, models need to be compiled specifically for ml_eia2 devices. For information on how to deploy your compiled model to an Elastic Inference accelerator, see Use El on Amazon SageMaker Hosted Endpoints.

When you deploy a compiled model, you need to use the same instance for the target that you used for compilation. This creates a SageMaker endpoint that you can use to perform inferences. You can deploy a Neo-compiled model using any of the following: Amazon SageMaker SDK for Python, SDK for Python (Boto3), AWS Command Line Interface, and the SageMaker console.

Note

For deploying a model using AWS CLI, the console, or Boto3, see Neo Inference Container Images to select the inference image URI for your primary container.

Topics

- Prerequisites
- Deploy a Compiled Model Using SageMaker SDK
- Deploy a Compiled Model Using Boto3
- Deploy a Compiled Model Using the AWS CLI
- Deploy a Compiled Model Using the Console
Prerequisites

Note
Follow the instructions in this section if you compiled your model using AWS SDK for Python (Boto3), AWS CLI, or the SageMaker console.

To create a SageMaker Neo-compiled model, you need the following:

1. A Docker image Amazon ECR URI. You can select one that meets your needs from this list.
2. An entry point script file:
   a. For PyTorch and MXNet models:
      
      *If you trained your model using SageMaker*, the training script must implement the functions described below. The training script serves as the entry point script during inference. In the example detailed in [MNIST Training, Compilation and Deployment with MXNet Module and SageMaker Neo](#), the training script (`mnist.py`) implements the required functions.

      *If you did not train your model using SageMaker*, you need to provide an entry point script (`inference.py`) file that can be used at the time of inference. Based on the framework—MXNet or PyTorch—the inference script location must conform to the SageMaker Python SDK [Model Directory Structure for MxNet](https://docs.aws.amazon.com/sagemaker/latest/dg/neo-model-deployment-model-definition.html) or [Model Directory Structure for PyTorch](https://docs.aws.amazon.com/sagemaker/latest/dg/neo-model-deployment-model-definition.html).

      When using Neo Inference Optimized Container images with **PyTorch** and **MXNet** on CPU and GPU instance types, the inference script must implement the following functions:

      - `model_fn`: Loads the model. (Optional)
      - `input_fn`: Converts the incoming request payload into a numpy array.
      - `predict_fn`: Performs the prediction.
      - `output_fn`: Converts the prediction output into the response payload.
      - Alternatively, you can define `transform_fn` to combine `input_fn`, `predict_fn`, and `output_fn`. 
The following are examples of `inference.py` script within a directory named code (code/inference.py) for **PyTorch and MXNet (Gluon and Module)**. The examples first load the model and then serve it on image data on a GPU:

### MXNet Module

```python
import numpy as np
import json
import mxnet as mx
import neomx  # noqa: F401
from collections import namedtuple

Batch = namedtuple('Batch', ['data'])

# Change the context to mx.cpu() if deploying to a CPU endpoint
ctx = mx.gpu()

def model_fn(model_dir):
    # The compiled model artifacts are saved with the prefix 'compiled'
sym, arg_params, aux_params = mx.model.load_checkpoint('compiled', 0)
mod = mx.mod.Module(symbol=sym, context=ctx, label_names=None)
exe = mod.bind(for_training=False,
               data_shapes=[('data', (1,3,224,224))],
               label_shapes=mod._label_shapes)
mod.set_params(arg_params, aux_params, allow_missing=True)

    # Run warm-up inference on empty data during model load (required for GPU)
data = mx.nd.empty((1,3,224,224), ctx=ctx)
mod.forward(Batch([data]))
return mod

def transform_fn(mod, image, input_content_type, output_content_type):
    # pre-processing
    decoded = mx.image.imdecode(image)
    resized = mx.image.resize_short(decoded, 224)
cropped, crop_info = mx.image.center_crop(resized, (224, 224))
normalized = mx.image.color_normalize(cropped.astype(np.float32) / 255,
                                      mean=mx.nd.array([0.485, 0.456, 0.406]),
                                      std=mx.nd.array([0.229, 0.224, 0.225]))
```
transposed = normalized.transpose((2, 0, 1))
batchified = transposed.expand_dims(axis=0)
casted = batchified.astype(dtype='float32')
processed_input = casted.as_in_context(ctx)

# prediction/inference
mod.forward(Batch([processed_input]))

# post-processing
prob = mod.get_outputs()[0].asnumpy().tolist()
prob_json = json.dumps(prob)
return prob_json, output_content_type

MXNet Gluon

import numpy as np
import json
import mxnet as mx
import neomx  # noqa: F401

# Change the context to mx.cpu() if deploying to a CPU endpoint
ctx = mx.gpu()

def model_fn(model_dir):
    # The compiled model artifacts are saved with the prefix 'compiled'
    block = mx.gluon.nn.SymbolBlock.imports('compiled-symbol.json',
    ['data'], 'compiled-0000.params', ctx=ctx)

    # Hybridize the model & pass required options for Neo: static_alloc=True
    & static_shape=True
    block.hybridize(static_alloc=True, static_shape=True)

    # Run warm-up inference on empty data during model load (required for
    GPU)
    data = mx.nd.empty((1,3,224,224), ctx=ctx)
    warm_up = block(data)
    return block

def input_fn(image, input_content_type):
    # pre-processing
    decoded = mx.image.imdecode(image)
    resized = mx.image.resize_short(decoded, 224)
cropped, crop_info = mx.image.center_crop(resized, (224, 224))
normalized = mx.image.color_normalize(cropped.astype(np.float32) / 255,
    mean=mx.nd.array([0.485, 0.456, 0.406]),
    std=mx.nd.array([0.229, 0.224, 0.225]))
transposed = normalized.transpose((2, 0, 1))
batchified = transposed.expand_dims(axis=0)
casted = batchified.astype(dtype='float32')
processed_input = casted.as_in_context(ctx)
return processed_input

def predict_fn(processed_input_data, block):
    # prediction/inference
    prediction = block(processed_input_data)
    return prediction

def output_fn(prediction, output_content_type):
    # post-processing
    prob = prediction.asnumpy().tolist()
    prob_json = json.dumps(prob)
    return prob_json, output_content_type

PyTorch 1.4 and Older

import os
import torch
import torch.nn.parallel
import torch.optim
import torch.utils.data
import torch.utils.data.distributed
import torchvision.transforms as transforms
from PIL import Image
import io
import json
import pickle

def model_fn(model_dir):
    """Load the model and return it.
    Providing this function is optional.
    There is a default model_fn available which will load the model
    compiled using SageMaker Neo. You can override it here."""
Keyword arguments:
model_dir -- the directory path where the model artifacts are present

# The compiled model is saved as "compiled.pt"
model_path = os.path.join(model_dir, 'compiled.pt')
with torch.neon.config(model_dir=model_dir, neo_runtime=True):
    model = torch.jit.load(model_path)
    device = torch.device("cuda" if torch.cuda.is_available() else "cpu")
    model = model.to(device)

# We recommend that you run warm-up inference during model load
sample_input_path = os.path.join(model_dir, 'sample_input.pkl')
with open(sample_input_path, 'rb') as input_file:
    model_input = pickle.load(input_file)
if torch.is_tensor(model_input):
    model_input = model_input.to(device)
model(model_input)
elif isinstance(model_input, tuple):
    model_input = (inp.to(device) for inp in model_input if torch.is_tensor(inp))
    model(*model_input)
else:
    print("Only supports a torch tensor or a tuple of torch tensors")
    return model

def transform_fn(model, request_body, request_content_type, response_content_type):
    """Run prediction and return the output.
The function
- Pre-processes the input request
- Runs prediction
- Post-processes the prediction output.
    """
    # preprocess
    decoded = Image.open(io.BytesIO(request_body))
    preprocess = transforms.Compose([
        transforms.Resize(256),
        transforms.CenterCrop(224),
        transforms.ToTensor(),
        transforms.Normalize(
            mean=[
            ]
        )
    )
    return preprocess(decoded)
import os
import torch
import torch.nn.parallel
import torch.optim
import torch.utils.data
import torch.utils.data.distributed
import torchvision.transforms as transforms
from PIL import Image
import io
import json
import pickle

def model_fn(model_dir):
    
    # Load the model and return it.
    Providing this function is optional.
    There is a default_model_fn available, which will load the model compiled using SageMaker Neo. You can override the default here. The model_fn only needs to be defined if your model needs extra steps to load, and can otherwise be left undefined.

    Keyword arguments:
    model_dir -- the directory path where the model artifacts are present

    # The compiled model is saved as "model.pt"
    model_path = os.path.join(model_dir, 'model.pt')
    device = torch.device("cuda" if torch.cuda.is_available() else "cpu")
    model = torch.jit.load(model_path, map_location=device)
model = model.to(device)

return model

def transform_fn(model, request_body, request_content_type, response_content_type):
    """Run prediction and return the output.
    The function
    1. Pre-processes the input request
    2. Runs prediction
    3. Post-processes the prediction output.
    """
    # preprocess
    decoded = Image.open(io.BytesIO(request_body))
    preprocess = transforms.Compose([
        transforms.Resize(256),
        transforms.CenterCrop(224),
        transforms.ToTensor(),
        transforms.Normalize(
            mean=[
                0.485, 0.456, 0.406],
            std=[
                0.229, 0.224, 0.225]),
    ])
    normalized = preprocess(decoded)
    batchified = normalized.unsqueeze(0)

    # predict
    device = torch.device("cuda" if torch.cuda.is_available() else "cpu")
    batchified = batchified.to(device)
    output = model.forward(batchified)
    return json.dumps(output.cpu().numpy().tolist()), response_content_type

b. **For inf1 instances or onnx, xgboost, keras container images**

For all other Neo Inference-optimized container images, or inferentia instance types, the entry point script must implement the following functions for Neo Deep Learning Runtime:

- neo_preprocess: Converts the incoming request payload into a numpy array.
• neo_postprocess: Converts the prediction output from Neo Deep Learning Runtime into the response body.

Note
The preceding two functions do not use any of the functionalities of MXNet, PyTorch, or TensorFlow.

For examples of how to use these functions, see Neo Model Compilation Sample Notebooks.

c. For TensorFlow models

If your model requires custom pre- and post-processing logic before data is sent to the model, then you must specify an entry point script inference.py file that can be used at the time of inference. The script should implement either a either a pair of input_handler and output_handler functions or a single handler function.

Note
Note that if handler function is implemented, input_handler and output_handler are ignored.

The following is a code example of inference.py script that you can put together with the compile model to perform custom pre- and post-processing on an image classification model. The SageMaker client sends the image file as an application/x-image content type to the input_handler function, where it is converted to JSON. The converted image file is then sent to the Tensorflow Model Server (TFX) using the REST API.

```python
import json
import numpy as np
import json
import io
from PIL import Image

def input_handler(data, context):
```
Pre-process request input before it is sent to TensorFlow Serving REST API

Args:
data (obj): the request data, in format of dict or string
context (Context): an object containing request and configuration details

Returns:
(dict): a JSON-serializable dict that contains request body and headers

```python
f = data.read()
f = io.BytesIO(f)
image = Image.open(f).convert('RGB')
batch_size = 1
image = np.asarray(image.resize((512, 512)))
image = np.concatenate([image[np.newaxis, :, :]] * batch_size)
body = json.dumps({"signature_name": "serving_default", "instances":
image.tolist()})
return body
```

def output_handler(data, context):
    """Post-process TensorFlow Serving output before it is returned to the client.

    Args:
data (obj): the TensorFlow serving response
context (Context): an object containing request and configuration details

    Returns:
(bytes, string): data to return to client, response content type
    """
    if data.status_code != 200:
        raise ValueError(data.content.decode('utf-8'))

    response_content_type = context.accept_header
    prediction = data.content
    return prediction, response_content_type
```

If there is no custom pre- or post-processing, the SageMaker client converts the file image to JSON in a similar way before sending it over to the SageMaker endpoint.

For more information, see the [Deploying to TensorFlow Serving Endpoints in the SageMaker Python SDK](#).
3. The Amazon S3 bucket URI that contains the compiled model artifacts.

**Deploy a Compiled Model Using SageMaker SDK**

You must satisfy the [prerequisites](#) section if the model was compiled using AWS SDK for Python (Boto3), AWS CLI, or the Amazon SageMaker console. Follow one of the following use cases to deploy a model compiled with SageMaker Neo based on how you compiled your model.

**Topics**

- [If you compiled your model using the SageMaker SDK](#)
- [If you compiled your model using MXNet or PyTorch](#)
- [If you compiled your model using Boto3, SageMaker console, or the CLI for TensorFlow](#)

**If you compiled your model using the SageMaker SDK**

The `sagemaker.Model` object handle for the compiled model supplies the `deploy()` function, which enables you to create an endpoint to serve inference requests. The function lets you set the number and type of instances that are used for the endpoint. You must choose an instance for which you have compiled your model. For example, in the job compiled in the job compiled in [Compile a Model (Amazon SageMaker SDK)](#) section, this is `ml_c5`.

```python
predictor = compiled_model.deploy(initial_instance_count = 1, instance_type = 'ml.c5.4xlarge')

# Print the name of newly created endpoint
print(predictor.endpoint_name)
```

**If you compiled your model using MXNet or PyTorch**

Create the SageMaker model and deploy it using the `deploy()` API under the framework-specific Model APIs. For MXNet, it is `MXNetModel` and for PyTorch, it is `PyTorchModel`. When you are creating and deploying an SageMaker model, you must set MMS_DEFAULT_RESPONSE_TIMEOUT environment variable to 500 and specify the `entry_point` parameter as the inference script (`inference.py`) and the `source_dir` parameter as the directory location (`code`) of the inference script. To prepare the inference script (`inference.py`) follow the Prerequisites step.

The following example shows how to use these functions to deploy a compiled model using the SageMaker SDK for Python:
MXNet

```python
from sagemaker.mxnet import MXNetModel

# Create SageMaker model and deploy an endpoint
sm_mxnet_compiled_model = MXNetModel(
    model_data='insert S3 path of compiled MXNet model archive',
    role='AmazonSageMaker-ExecutionRole',
    entry_point='inference.py',
    source_dir='code',
    framework_version='1.8.0',
    py_version='py3',
    image_uri='insert appropriate ECR Image URI for MXNet',
    env={'MMS_DEFAULT_RESPONSE_TIMEOUT': '500'},
)

# Replace the example instance_type below to your preferred instance_type
predictor = sm_mxnet_compiled_model.deploy(initial_instance_count = 1, instance_type = 'ml.p3.2xlarge')

# Print the name of newly created endpoint
print(predictor.endpoint_name)
```

PyTorch 1.4 and Older

```python
from sagemaker.pytorch import PyTorchModel

# Create SageMaker model and deploy an endpoint
sm_pytorch_compiled_model = PyTorchModel(
    model_data='insert S3 path of compiled PyTorch model archive',
    role='AmazonSageMaker-ExecutionRole',
    entry_point='inference.py',
    source_dir='code',
    framework_version='1.4.0',
    py_version='py3',
    image_uri='insert appropriate ECR Image URI for PyTorch',
    env={'MMS_DEFAULT_RESPONSE_TIMEOUT': '500'},
)

# Replace the example instance_type below to your preferred instance_type
predictor = sm_pytorch_compiled_model.deploy(initial_instance_count = 1, instance_type = 'ml.p3.2xlarge')
```
PyTorch 1.5 and Newer

```python
from sagemaker.pytorch import PyTorchModel

# Create SageMaker model and deploy an endpoint
sm_pytorch_compiled_model = PyTorchModel(
    model_data='insert S3 path of compiled PyTorch model archive',
    role='AmazonSageMaker-ExecutionRole',
    entry_point='inference.py',
    source_dir='code',
    framework_version='1.5',
    py_version='py3',
    image_uri='insert appropriate ECR Image URI for PyTorch',
)

# Replace the example instance_type below to your preferred instance_type
predictor = sm_pytorch_compiled_model.deploy(initial_instance_count = 1,
                                              instance_type = 'ml.p3.2xlarge')

# Print the name of newly created endpoint
print(predictor.endpoint_name)
```

**Note**

The `AmazonSageMakerFullAccess` and `AmazonS3ReadOnlyAccess` policies must be attached to the `AmazonSageMaker-ExecutionRole` IAM role.

**If you compiled your model using Boto3, SageMaker console, or the CLI for TensorFlow**

Construct a `TensorFlowModel` object, then call `deploy`:

```python
role='AmazonSageMaker-ExecutionRole'
model_path='S3 path for model file'
framework_image='inference container arn'
tf_model = TensorFlowModel(model_data=model_path,
                           framework_version='1.15.3',
```

Cloud Instances 4589
See [Deploying directly from model artifacts](#) for more information.

You can select a Docker image Amazon ECR URI that meets your needs from [this list](#).

For more information on how to construct a `TensorFlowModel` object, see the [SageMaker SDK](#).

**Note**

Your first inference request might have high latency if you deploy your model on a GPU. This is because an optimized compute kernel is made on the first inference request. We recommend that you make a warm-up file of inference requests and store that alongside your model file before sending it off to a TFX. This is known as “warming up” the model.

The following code snippet demonstrates how to produce the warm-up file for image classification example in the [prerequisites](#) section:

```python
import tensorflow as tf
from tensorflow_serving.apis import classification_pb2
from tensorflow_serving.apis import inference_pb2
from tensorflow_serving.apis import model_pb2
from tensorflow_serving.apis import predict_pb2
from tensorflow_serving.apis import prediction_log_pb2
import numpy as np

with tf.python_io.TFRecordWriter("tf_serving_warmup_requests") as writer:
    img = np.random.uniform(0, 1, size=[224, 224, 3]).astype(np.float32)
    img = np.expand_dims(img, axis=0)
    test_data = np.repeat(img, 1, axis=0)
    request = predict_pb2.PredictRequest()
    request.model_spec.name = 'compiled_models'
    request.model_spec.signature_name = 'serving_default'
    request.inputs['Placeholder:0'].CopyFrom(tf.compat.v1.make_tensor_proto(test_data,
        shape=test_data.shape, dtype=tf.float32))
    log = prediction_log_pb2.PredictionLog(
```
predict_log=prediction_log_pb2.PredictLog(request=request))
writer.write(log.SerializeToString())

For more information on how to “warm up” your model, see the TensorFlow TFX page.

Deploy a Compiled Model Using Boto3

You must satisfy the prerequisites section if the model was compiled using AWS SDK for Python (Boto3), AWS CLI, or the Amazon SageMaker console. Follow the steps below to create and deploy a SageMaker Neo-compiled model using Amazon Web Services SDK for Python (Boto3).

Topics

- Deploy the Model

Deploy the Model

After you have satisfied the prerequisites, use the create_model, create_endpoint_config, and create_endpoint APIs.

The following example shows how to use these APIs to deploy a model compiled with Neo:

```python
import boto3
client = boto3.client('sagemaker')

# create sagemaker model
create_model_api_response = client.create_model(
    ModelName='my-sagemaker-model',
    PrimaryContainer={
        'Image': '<insert the ECR Image URI>',
        'ModelDataUrl': 's3://path/to/model/artifact/model.tar.gz',
        'Environment': {}
    },
    ExecutionRoleArn='ARN for AmazonSageMaker-
    ExecutionRole'
)

print("create_model API response", create_model_api_response)

# create sagemaker endpoint config
create_endpoint_config_api_response = client.create_endpoint_config(
```
```

EndpointConfigName='sagemaker-neomxnet-endpoint-configuration',
ProductionVariants=[
  {
    'VariantName': <provide your variant name>,
    'ModelName': 'my-sagemaker-model',
    'InitialInstanceCount': 1,
    'InstanceType': <provide your instance type here>
  },
  
  
}

print ("create_endpoint_config API response", create_endpoint_config_api_response)

# create sagemaker endpoint
create_endpoint_api_response = client.create_endpoint(
  EndpointName='provide your endpoint name',
  EndpointConfigName=<insert your endpoint config name>,
)

print ("create_endpoint API response", create_endpoint_api_response)
```

**Note**

The AmazonSageMakerFullAccess and AmazonS3ReadOnlyAccess policies must be attached to the AmazonSageMaker-ExecutionRole IAM role.

For full syntax of `create_model`, `create_endpoint_config`, and `create_endpoint` APIs, see `create_model`, `create_endpoint_config`, and `create_endpoint`, respectively.

If you did not train your model using SageMaker, specify the following environment variables:

**MXNet and PyTorch**

```
"Environment": {
  "SAGEMAKER_PROGRAM": "inference.py",
  "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code",
  "SAGEMAKER_CONTAINER_LOG_LEVEL": "20",
```

Cloud Instances
If you trained your model using SageMaker, specify the environment variable `SAGEMAKER_SUBMIT_DIRECTORY` as the full Amazon S3 bucket URI that contains the training script.

**Deploy a Compiled Model Using the AWS CLI**

You must satisfy the [prerequisites](#) section if the model was compiled using AWS SDK for Python (Boto3), AWS CLI, or the Amazon SageMaker console. Follow the steps below to create and deploy a SageMaker Neo-compiled model using the AWS CLI.

**Topics**

- [Deploy the Model](#)

**Deploy the Model**

After you have satisfied the [prerequisites](#), use the create-model, create-endpoint-config, and create-endpoint AWS CLI commands. The following steps explain how to use these commands to deploy a model compiled with Neo:

**Create a Model**

From [Neo Inference Container Images](#), select the inference image URI and then use create-model API to create a SageMaker model. You can do this with two steps:
1. Create a `create_model.json` file. Within the file, specify the name of the model, the image URI, the path to the `model.tar.gz` file in your Amazon S3 bucket, and your SageMaker execution role:

```json
{
  "ModelName": "insert model name",
  "PrimaryContainer": {
    "Image": "insert the ECR Image URI",
    "ModelDataUrl": "insert S3 archive URL",
    "Environment": {"See details below"}
  },
  "ExecutionRoleArn": "ARN for AmazonSageMaker-ExecutionRole"
}
```

If you trained your model using SageMaker, specify the following environment variable:

```json
"Environment": {
  "SAGEMAKER_SUBMIT_DIRECTORY" : "[Full S3 path for *.tar.gz file containing the training script]"
}
```

If you did not train your model using SageMaker, specify the following environment variables:

**MXNet and PyTorch**

```json
"Environment": {
  "SAGEMAKER_PROGRAM": "inference.py",
  "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code",
  "SAGEMAKER_CONTAINER_LOG_LEVEL": "20",
  "SAGEMAKER_REGION": "insert your region",
  "MMS_DEFAULT_RESPONSE_TIMEOUT": "500"
}
```

**TensorFlow**

```json
"Environment": {
  "SAGEMAKER_PROGRAM": "inference.py",
  "SAGEMAKER_SUBMIT_DIRECTORY": "/opt/ml/model/code",
  "SAGEMAKER_CONTAINER_LOG_LEVEL": "20",
  "SAGEMAKER_REGION": "insert your region"
}
```
Note

The AmazonSageMakerFullAccess and AmazonS3ReadonlyAccess policies must be attached to the AmazonSageMaker-ExecutionRole IAM role.

2. Run the following command:

```
aws sagemaker create-model --cli-input-json file://create_model.json
```

For the full syntax of the create-model API, see `create-model`.

Create an Endpoint Configuration

After creating a SageMaker model, create the endpoint configuration using the `create-endpoint-config` API. To do this, create a JSON file with your endpoint configuration specifications. For example, you can use the following code template and save it as `create_config.json`:

```
{
   "EndpointConfigName": "<provide your endpoint config name>",
   "ProductionVariants": [
      {
         "VariantName": "<provide your variant name>",
         "ModelName": "my-sagemaker-model",
         "InitialInstanceCount": 1,
         "InstanceType": "<provide your instance type here>",
         "InitialVariantWeight": 1.0
      }
   ]
}
```

Now run the following AWS CLI command to create your endpoint configuration:

```
aws sagemaker create-endpoint-config --cli-input-json file://create_config.json
```

For the full syntax of the create-endpoint-config API, see `create-endpoint-config`. 
Create an Endpoint

After you have created your endpoint configuration, create an endpoint using the create-endpoint API:

```
aws sagemaker create-endpoint --endpoint-name '<provide your endpoint name>' --endpoint-config-name '<insert your endpoint config name>'
```

For the full syntax of the create-endpoint API, see `create-endpoint`.

Deploy a Compiled Model Using the Console

You must satisfy the `prerequisites` section if the model was compiled using AWS SDK for Python (Boto3), the AWS CLI, or the Amazon SageMaker console. Follow the steps below to create and deploy a SageMaker Neo-compiled model using the SageMaker console `https://console.aws.amazon.com/SageMaker`.

Topics

- Deploy the Model

Deploy the Model

After you have satisfied the `prerequisites`, use the following steps to deploy a model compiled with Neo:

1. Choose **Models**, and then choose **Create models** from the **Inference** group. On the **Create model** page, complete the **Model name**, **IAM role**, and **VPC** fields (optional), if needed.
2. To add information about the container used to deploy your model, choose **Add container** container, then choose **Next**. Complete the **Container input options**, **Location of inference code image**, and **Location of model artifacts**, and optionally, **Container host name**, and **Environmental variables** fields.
3. To deploy Neo-compiled models, choose the following:

- **Container input options**: Choose **Provide model artifacts and inference image**.

- **Location of inference code image**: Choose the inference image URI from [Neo Inference Container Images](#), depending on the AWS Region and kind of application.

- **Location of model artifact**: Enter the Amazon S3 bucket URI of the compiled model artifact generated by the Neo compilation API.

- **Environment variables**:
  - Leave this field blank for **SageMaker XGBoost**.
• If you trained your model using SageMaker, specify the environment variable `SAGEMAKER_SUBMIT_DIRECTORY` as the Amazon S3 bucket URI that contains the training script.

• If you did not train your model using SageMaker, specify the following environment variables:

<table>
<thead>
<tr>
<th>Key</th>
<th>Values for MXNet and PyTorch</th>
<th>Values TensorFlow</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAGEMAKER_PROGRAM</td>
<td>inference.py</td>
<td>inference.py</td>
</tr>
<tr>
<td>SAGEMAKER_SUBMIT_DIRECTORY</td>
<td>/opt/ml/model/code</td>
<td>/opt/ml/model/code</td>
</tr>
<tr>
<td>SAGEMAKER_CONTAINER</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SAGEMAKER_CONTAINER</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>SAGEMAKER_LOG_LEVEL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MMS_DEFAULT_RESPONSE</td>
<td>500</td>
<td>Leave this field blank for TF</td>
</tr>
</tbody>
</table>

4. Confirm that the information for the containers is accurate, and then choose **Create model**. On the **Create model landing page**, choose **Create endpoint**.
5. In **Create and configure endpoint** diagram, specify the **Endpoint name**. For **Attach endpoint configuration**, choose **Create a new endpoint configuration**.

6. In **New endpoint configuration** page, specify the **Endpoint configuration name**.
7. Choose **Edit** next to the name of the model and specify the correct **Instance type** on the **Edit Production Variant** page. It is imperative that the **Instance type** value match the one specified in your compilation job.
8. Choose Save.

9. On the New endpoint configuration page, choose Create endpoint configuration, and then choose Create endpoint.

Request Inferences from a Deployed Service

If you have followed instructions in Deploy a Model, you should have a SageMaker endpoint set up and running. Regardless of how you deployed your Neo-compiled model, there are three ways you can submit inference requests:

Topics

- Request Inferences from a Deployed Service (Amazon SageMaker SDK)
- Request Inferences from a Deployed Service (Boto3)
- Request Inferences from a Deployed Service (AWS CLI)

Request Inferences from a Deployed Service (Amazon SageMaker SDK)

Use the following the code examples to request inferences from your deployed service based on the framework you used to train your model. The code examples for the different frameworks are similar. The main difference is that TensorFlow requires application/json as the content type.
PyTorch and MXNet

If you are using **PyTorch v1.4 or later** or **MXNet 1.7.0 or later** and you have an Amazon SageMaker endpoint InService, you can make inference requests using the `predictor` package of the SageMaker SDK for Python.

**Note**

The API varies based on the SageMaker SDK for Python version:

- For version 1.x, use the `RealTimePredictor` and `Predict` API.
- For version 2.x, use the `Predictor` and the `Predict` API.

The following code example shows how to use these APIs to send an image for inference:

**SageMaker Python SDK v1.x**

```python
from sagemaker.predictor import RealTimePredictor

endpoint = 'insert name of your endpoint here'

# Read image into memory
payload = None
with open("image.jpg", 'rb') as f:
    payload = f.read()

predictor = RealTimePredictor(endpoint=endpoint, content_type='application/x-image')
inference_response = predictor.predict(data=payload)
print (inference_response)
```

**SageMaker Python SDK v2.x**

```python
from sagemaker.predictor import Predictor

endpoint = 'insert name of your endpoint here'

# Read image into memory
payload = None
with open("image.jpg", 'rb') as f:
    payload = f.read()
```
predictor = Predictor(endpoint)
inference_response = predictor.predict(data=payload)
print (inference_response)

TensorFlow

The following code example shows how to use the SageMaker Python SDK API to send an image for inference:

```python
from sagemaker.predictor import Predictor
from PIL import Image
import numpy as np
import json

endpoint = 'insert the name of your endpoint here'

# Read image into memory
image = Image.open(input_file)
batch_size = 1
image = np.asarray(image.resize((224, 224)))
image = image / 128 - 1
image = np.concatenate([image[np.newaxis, :, :]] * batch_size)
body = json.dumps({'instances': image.tolist()})

predictor = Predictor(endpoint)
inference_response = predictor.predict(data=body)
print(inference_response)
```

Request Inferences from a Deployed Service (Boto3)

You can submit inference requests using SageMaker SDK for Python (Boto3) client and `invoke_endpoint()` API once you have an SageMaker endpoint InService. The following code example shows how to send an image for inference:

PyTorch and MXNet

```python
import boto3
import json

endpoint = 'insert name of your endpoint here'
```
runtime = boto3.Session().client('sagemaker-runtime')

# Read image into memory
with open(image, 'rb') as f:
    payload = f.read()
# Send image via InvokeEndpoint API
response = runtime.invoke_endpoint(EndpointName=endpoint, ContentType='application/x-image', Body=payload)

# Unpack response
result = json.loads(response['Body'].read().decode())

TensorFlow

For TensorFlow submit an input with application/json for the content type.

from PIL import Image
import numpy as np
import json
import boto3

client = boto3.client('sagemaker-runtime')
input_file = 'path/to/image'
image = Image.open(input_file)
batch_size = 1
image = np.asarray(image.resize((224, 224)))
image = image / 128 - 1
image = np.concatenate([image[np.newaxis, :, :]] * batch_size)
body = json.dumps({'instances': image.tolist()})

ioc_predictor_endpoint_name = 'insert name of your endpoint here'
content_type = 'application/json'
ioc_response = client.invoke_endpoint(
    EndpointName=ioc_predictor_endpoint_name,
    Body=body,
    ContentType=content_type
)

XGBoost

For an XGBoost application, you should submit a CSV text instead:

import boto3
import json
endpoint = 'insert your endpoint name here'

runtime = boto3.Session().client('sagemaker-runtime')

csv_text = '1,-1.0,1.0,1.5,2.6'
# Send CSV text via InvokeEndpoint API
response = runtime.invoke_endpoint(EndpointName=endpoint, ContentType='text/csv',
    Body=csv_text)
# Unpack response
result = json.loads(response['Body'].read().decode())

Note that BYOM allows for a custom content type. For more information, see runtime_InvokeEndpoint.

Request Inferences from a Deployed Service (AWS CLI)

Inference requests can be made with the sagemaker-runtime invoke-endpoint once you have an Amazon SageMaker endpoint InService. You can make inference requests with the AWS Command Line Interface (AWS CLI). The following example shows how to send an image for inference:

```bash
aws sagemaker-runtime invoke-endpoint --endpoint-name 'insert name of your endpoint here' --body fileb://image.jpg --content-type=application/x-image output_file.txt
```

An output_file.txt with information about your inference requests is made if the inference was successful.

For TensorFlow submit an input with application/json as the content type.

```bash
aws sagemaker-runtime invoke-endpoint --endpoint-name 'insert name of your endpoint here' --body fileb://input.json --content-type=application/json output_file.txt
```

Inference Container Images

SageMaker Neo now provides inference image URI information for ml_* targets. For more information see DescribeCompilationJob.

Based on your use case, replace the highlighted portion in the inference image URI template provided below with appropriate values.
Amazon SageMaker XGBoost

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/xgboost-neo:latest
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Keras

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-neo-keras:fx_version-instance_type-py3
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Replace `fx_version` with 2.2.4.

Replace `instance_type` with either cpu or gpu.

MXNet

CPU or GPU instance types

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-inference-mxnet:fx_version-instance_type-py3
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Replace `fx_version` with 1.8.0.

Replace `instance_type` with either cpu or gpu.

Inferentia1

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-neo-mxnet:fx_version-instance_type-py3
```

Replace `aws_region` with either us-east-1 or us-west-2.

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.
Replace `fx_version` with 1.5.1.

Replace `instance_type` with `inf`.

**ONNX**

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-neo-onnx:fx_version-
instance_type-py3
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Replace `fx_version` with 1.5.0.

Replace `instance_type` with either `cpu` or `gpu`.

**PyTorch**

**CPU or GPU instance types**

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-inference-
pytorch:fx_version-instance_type-py3
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Replace `fx_version` with 1.4, 1.5, 1.6, 1.7, 1.8, 1.12, 1.13, or 2.0.

Replace `instance_type` with either `cpu` or `gpu`.

**Inferentia**

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-neo-
pytorch:fx_version-instance_type-py3
```

Replace `aws_region` with either `us-east-1` or `us-west-2`.

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Replace `fx_version` with 1.5.1.
Replace `instance_type` with `inf`.

Inferentia2 and Trainium1

```
```

Replace `aws_region` with `us-east-2` for Inferentia2, and `us-east-1` for Trainium1.

**TensorFlow**

**CPU or GPU instance types**

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-inference-tensorflow:fx_version-instance_type-py3
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used.

Replace `fx_version` with 1.15.3 or 2.9.

Replace `instance_type` with either `cpu` or `gpu`.

Inferentia1

```
aws_account_id.dkr.ecr.aws_region.amazonaws.com/sagemaker-neo-tensorflow:fx_version-instance_type-py3
```

Replace `aws_account_id` from the table at the end of this page based on the `aws_region` you used. Note that for instance type `inf` only `us-east-1` and `us-west-2` are supported.

Replace `fx_version` with 1.15.0.

Replace `instance_type` with `inf`.

Inferentia2 and Trainium1

```
```

Replace `aws_region` with `us-east-2` for Inferentia2, and `us-east-1` for Trainium1.
The following table maps `aws_account_id` with `aws_region`. Use this table to find the correct inference image URI you need for your application.

<table>
<thead>
<tr>
<th>aws_account_id</th>
<th>aws_region</th>
</tr>
</thead>
<tbody>
<tr>
<td>785573368785</td>
<td>us-east-1</td>
</tr>
<tr>
<td>007439368137</td>
<td>us-east-2</td>
</tr>
<tr>
<td>710691900526</td>
<td>us-west-1</td>
</tr>
<tr>
<td>301217895009</td>
<td>us-west-2</td>
</tr>
<tr>
<td>802834080501</td>
<td>eu-west-1</td>
</tr>
<tr>
<td>205493899709</td>
<td>eu-west-2</td>
</tr>
<tr>
<td>254080097072</td>
<td>eu-west-3</td>
</tr>
<tr>
<td>601324751636</td>
<td>eu-north-1</td>
</tr>
<tr>
<td>966458181534</td>
<td>eu-south-1</td>
</tr>
<tr>
<td>746233611703</td>
<td>eu-central-1</td>
</tr>
<tr>
<td>110948597952</td>
<td>ap-east-1</td>
</tr>
<tr>
<td>763008648453</td>
<td>ap-south-1</td>
</tr>
<tr>
<td>941853720454</td>
<td>ap-northeast-1</td>
</tr>
<tr>
<td>151534178276</td>
<td>ap-northeast-2</td>
</tr>
<tr>
<td>925152966179</td>
<td>ap-northeast-3</td>
</tr>
<tr>
<td>324986816169</td>
<td>ap-southeast-1</td>
</tr>
<tr>
<td>355873309152</td>
<td>ap-southeast-2</td>
</tr>
<tr>
<td>474822919863</td>
<td>cn-northwest-1</td>
</tr>
<tr>
<td>472730292857</td>
<td>cn-north-1</td>
</tr>
</tbody>
</table>
Edge Devices

Amazon SageMaker Neo provides compilation support for popular machine learning frameworks. You can deploy your Neo-compiled edge devices such as the Raspberry Pi 3, Texas Instruments' Sitara, Jetson TX1, and more. For a full list of supported frameworks and edge devices, see Supported Frameworks, Devices, Systems, and Architectures.

You must configure your edge device so that it can use AWS services. One way to do this is to install DLR and Boto3 to your device. To do this, you must set up the authentication credentials. See Boto3 AWS Configuration for more information. Once your model is compiled and your edge device is configured, you can download the model from Amazon S3 to your edge device. From there, you can use the Deep Learning Runtime (DLR) to read the compiled model and make inferences.

For first-time users, we recommend you check out the Getting Started guide. This guide walks you through how to set up your credentials, compile a model, deploy your model to a Raspberry Pi 3, and make inferences on images.

Topics

- Supported Frameworks, Devices, Systems, and Architectures
- Deploy Models
- Getting Started with Neo on Edge Devices
Supported Frameworks, Devices, Systems, and Architectures

Amazon SageMaker Neo supports common machine learning frameworks, edge devices, operating systems, and chip architectures. Find out if Neo supports your framework, edge device, OS, and chip architecture by selecting one of the topics below.

You can find a list of models that have been tested by the Amazon SageMaker Neo Team in the Tested Models section.

Note

- Ambarella devices require additional files to be included within the compressed TAR file before it is sent for compilation. For more information, see Troubleshoot Ambarella Errors.
- TIM-VX (libtim-vx.so) is required for i.MX 8M Plus. For information on how to build TIM-VX, see the TIM-VX GitHub repository.

Topics

- Supported Frameworks
- Supported Devices, Chip Architectures, and Systems
- Tested Models

Supported Frameworks

Amazon SageMaker Neo supports the following frameworks.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>MXNet</td>
<td>1.8</td>
<td>Supports 1.8 or earlier</td>
<td>Image Classification, Object Detection, Semantic</td>
<td>One symbol file (.json) and one parameter file (.params)</td>
<td>GluonCV v0.8.0</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------------</td>
<td>---------------</td>
<td>------------------------------</td>
<td>--------------------------------------</td>
<td>----------------------------------------------</td>
</tr>
<tr>
<td>ONNX</td>
<td>1.7</td>
<td>Supports 1.7 or earlier</td>
<td>Image Classification, SVM</td>
<td>One model file (.onnx)</td>
<td></td>
</tr>
<tr>
<td>Keras</td>
<td>2.2</td>
<td>Supports 2.2 or earlier</td>
<td>Image Classification</td>
<td>One model definition file (.h5)</td>
<td></td>
</tr>
<tr>
<td>PyTorch</td>
<td>1.7, 1.8</td>
<td>Supports 1.7, 1.8 or earlier</td>
<td>Image Classification, Object Detection</td>
<td>One model definition file (.pth)</td>
<td></td>
</tr>
<tr>
<td>TensorFlow</td>
<td>1.15, 2.4, 2.5 (only for ml.inf1.* instances)</td>
<td>Supports 1.15, 2.4, 2.5 (only for ml.inf1.* instances) or earlier</td>
<td>Image Classification, Object Detection</td>
<td>*For saved models, one .pb or one .pbtxt file and a variables directory that contains variables *For frozen models, only one .pb or .pbtxt file</td>
<td></td>
</tr>
<tr>
<td>--------------</td>
<td>-------------------</td>
<td>---------------</td>
<td>----------------------------</td>
<td>---------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>TensorFlow-Lite</td>
<td>1.15</td>
<td>Supports 1.15 or earlier</td>
<td>Image Classification, Object Detection</td>
<td>One model definition flatbuffer file (.tflite)</td>
<td></td>
</tr>
<tr>
<td>XGBoost</td>
<td>1.3</td>
<td>Supports 1.3 or earlier</td>
<td>Decision Trees</td>
<td>One XGBoost model file (.model) where the number of nodes in a tree is less than $2^{31}$</td>
<td></td>
</tr>
<tr>
<td>DARKNET</td>
<td></td>
<td></td>
<td>Image Classification, Object Detection (Yolo model is not supported)</td>
<td>One config (.cfg) file and one weights (.weights) file</td>
<td></td>
</tr>
</tbody>
</table>

**Supported Devices, Chip Architectures, and Systems**

Amazon SageMaker Neo supports the following devices, chip architectures, and operating systems.

**Devices**

You can select a device using the dropdown list in the Amazon SageMaker console or by specifying the TargetDevice in the output configuration of the CreateCompilationJob API.

You can choose from one of the following edge devices:
<table>
<thead>
<tr>
<th>Device List</th>
<th>System on a Chip (SoC)</th>
<th>Operating System</th>
<th>Architecture</th>
<th>Accelerator</th>
<th>Compiler Options Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>aisage</td>
<td></td>
<td>Linux</td>
<td>ARM64</td>
<td>Mali</td>
<td></td>
</tr>
<tr>
<td>amba_cv2</td>
<td>CV2</td>
<td>Arch Linux</td>
<td>ARM64</td>
<td>cvflow</td>
<td></td>
</tr>
<tr>
<td>amba_cv22</td>
<td>CV22</td>
<td>Arch Linux</td>
<td>ARM64</td>
<td>cvflow</td>
<td></td>
</tr>
<tr>
<td>amba_cv25</td>
<td>CV25</td>
<td>Arch Linux</td>
<td>ARM64</td>
<td>cvflow</td>
<td></td>
</tr>
<tr>
<td>coreml</td>
<td></td>
<td>iOS, macOS</td>
<td></td>
<td></td>
<td>{&quot;class_labels&quot;: &quot;imagenet_labels_1000.txt&quot;}</td>
</tr>
<tr>
<td>imx8qm</td>
<td>NXP imx8</td>
<td>Linux</td>
<td>ARM64</td>
<td></td>
<td></td>
</tr>
<tr>
<td>imx8mplus</td>
<td>i.MX 8M Plus</td>
<td>Linux</td>
<td>ARM64</td>
<td>NPU</td>
<td></td>
</tr>
<tr>
<td>jacinto_tda4vm</td>
<td>TDA4VM</td>
<td>Linux</td>
<td>ARM</td>
<td>TDA4VM</td>
<td></td>
</tr>
<tr>
<td>jetson_nano</td>
<td></td>
<td>Linux</td>
<td>ARM64</td>
<td>NVIDIA</td>
<td>{&quot;gpu-code&quot;: &quot;sm_53&quot;, 'trt-ver': '5.0.6', 'cuda-ver': '10.0'}</td>
</tr>
</tbody>
</table>

For TensorFlow2, Edge Devices 4615
### Device List

<table>
<thead>
<tr>
<th>Device List</th>
<th>System on a Chip (SoC)</th>
<th>Operating System</th>
<th>Architecture</th>
<th>Accelerator</th>
<th>Compiler Options Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>jetson_tx1</td>
<td>Linux</td>
<td>ARM64</td>
<td>NVIDIA</td>
<td></td>
<td>{'JETPACK_VERSION': '4.6', 'gpu_code': 'sm_72'}</td>
</tr>
<tr>
<td>jetson_tx2</td>
<td>Linux</td>
<td>ARM64</td>
<td>NVIDIA</td>
<td></td>
<td>{'gpu-code': 'sm_53', 'trt-ver': '6.0.1', 'cuda-ver': '10.0'}</td>
</tr>
<tr>
<td>Device List</td>
<td>System on a Chip (SoC)</td>
<td>Operating System</td>
<td>Architecture</td>
<td>Accelerator</td>
<td>Compiler Options Example</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------</td>
<td>------------------</td>
<td>--------------</td>
<td>-------------</td>
<td>--------------------------</td>
</tr>
<tr>
<td>jetson_xavier</td>
<td></td>
<td>Linux</td>
<td>ARM64</td>
<td>NVIDIA</td>
<td>{'gpu-code': 'sm_72', 'trt-ver': '5.1.6', 'cuda-ver': '10.0'}</td>
</tr>
<tr>
<td>qcs605</td>
<td></td>
<td>Android</td>
<td>ARM64</td>
<td>Mali</td>
<td>{'ANDROID_PLATFORM': 27}</td>
</tr>
<tr>
<td>qcs603</td>
<td></td>
<td>Android</td>
<td>ARM64</td>
<td>Mali</td>
<td>{'ANDROID_PLATFORM': 27}</td>
</tr>
<tr>
<td>rasp3b</td>
<td>ARM A56</td>
<td>Linux</td>
<td>ARM_EABIHF</td>
<td></td>
<td>{'mattr': ['+neon']}</td>
</tr>
<tr>
<td>rasp4b</td>
<td>ARM A72</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rk3288</td>
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### System on a Chip (SoC) Operating System Architecture Accelerator Compiler Options Example

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<th>Architecture</th>
<th>Accelerator</th>
<th>Compiler Options Example</th>
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For more information about JSON key-value compiler options for each target device, see the CompilerOptions field in the `OutputConfig API` data type.

### Systems and Chip Architectures

The following look-up tables provide information regarding available operating systems and architectures for Neo model compilation jobs.

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### Tested Models

The following collapsible sections provide information about machine learning models that were tested by the Amazon SageMaker Neo team. Expand the collapsible section based on your framework to check if a model was tested.

#### Note

This is not a comprehensive list of models that can be compiled with Neo.

See [Supported Frameworks](#) and [SageMaker Neo Supported Operators](#) to find out if you can compile your model with SageMaker Neo.
## DarkNet

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TensorFlow-Lite (INT8)

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Deploy Models

You can deploy the compute module to resource-constrained edge devices by: downloading the compiled model from Amazon S3 to your device and using DLR, or you can use AWS IoT Greengrass.

Before moving on, make sure your edge device must be supported by SageMaker Neo. See, Supported Frameworks, Devices, Systems, and Architectures to find out what edge devices are supported. Make sure that you specified your target edge device when you submitted the compilation job, see Use Neo to Compile a Model.

Deploy a Compiled Model (DLR)

DLR is a compact, common runtime for deep learning models and decision tree models. DLR uses the TVM runtime, Treelite runtime, NVIDIA TensorRT™, and can include other hardware-specific runtimes. DLR provides unified Python/C++ APIs for loading and running compiled models on various devices.

You can install latest release of DLR package using the following pip command:

```
pip install dlr
```

For installation of DLR on GPU targets or non-x86 edge devices, please refer to Releases for prebuilt binaries, or Installing DLR for building DLR from source. For example, to install DLR for Raspberry Pi 3, you can use:

```
```

Deploy a Model (AWS IoT Greengrass)

AWS IoT Greengrass extends cloud capabilities to local devices. It enables devices to collect and analyze data closer to the source of information, react autonomously to local events, and communicate securely with each other on local networks. With AWS IoT Greengrass, you can perform machine learning inference at the edge on locally generated data using cloud-trained models. Currently, you can deploy models on to all AWS IoT Greengrass devices based on ARM Cortex-A, Intel Atom, and Nvidia Jetson series processors. For more information on deploying a Lambda inference application to perform machine learning inferences with AWS IoT Greengrass, see How to configure optimized machine learning inference using the AWS Management Console.
Getting Started with Neo on Edge Devices

This guide to getting started with Amazon SageMaker Neo shows you how to compile a model, set up your device, and make inferences on your device. Most of the code examples use Boto3. We provide commands using AWS CLI where applicable, as well as instructions on how to satisfy prerequisites for Neo.

⚠️ Note

You can run the following code snippets on your local machine, within a SageMaker notebook, within SageMaker Studio, or (depending on your edge device) on your edge device. The setup is similar; however, there are two main exceptions if you run this guide within a SageMaker notebook instance or SageMaker Studio session:

- You do not need to install Boto3.
- You do not need to add the ‘AmazonSageMakerFullAccess’ IAM policy

This guide assumes you are running the following instructions on your edge device.

Prerequisites

1. **Install Boto3**

   If you are running these commands on your edge device, you must install the AWS SDK for Python (Boto3). Within a Python environment (preferably a virtual environment), run the following locally on your edge device's terminal or within a Jupyter notebook instance:

   Terminal

   ```bash
   pip install boto3
   ```

   Jupyter Notebook

   ```bash
   !pip install boto3
   ```

2. **Set Up AWS Credentials**

   You need to set up Amazon Web Services credentials on your device in order to run SDK for Python (Boto3). By default, the AWS credentials should be stored in the file `~/.aws/`
credentials on your edge device. Within the credentials file, you should see two environment variables: `aws_access_key_id` and `aws_secret_access_key`.

In your terminal, run:

```
$ more ~/.aws/credentials

[default]
aws_access_key_id = YOUR_ACCESS_KEY
aws_secret_access_key = YOUR_SECRET_KEY
```

The [AWS General Reference Guide](https://docs.aws.amazon.com/general/latest/securityguide/) has instructions on how to get the necessary `aws_access_key_id` and `aws_secret_access_key`. For more information on how to set up credentials on your device, see the Boto3 documentation.

3. **Set up an IAM Role and attach policies.**

Neo needs access to your S3 bucket URI. Create an IAM role that can run SageMaker and has permission to access the S3 URI. You can create an IAM role either by using SDK for Python (Boto3), the console, or the AWS CLI. The following example illustrates how to create an IAM role using SDK for Python (Boto3):

```python
import boto3

AWS_REGION = 'aws-region'

# Create an IAM client to interact with IAM
iam_client = boto3.client('iam', region_name=AWS_REGION)
role_name = 'role-name'
```

For more information on how to create an IAM role with the console, AWS CLI, or through the AWS API, see [Creating an IAM user in your AWS account](https://docs.aws.amazon.com/console/).  

Create a dictionary describing the IAM policy you are attaching. This policy is used to create a new IAM role.

```python
policy = {
    'Statement': [
        {
            'Action': 'sts:AssumeRole',
            'Effect': 'Allow',
```
Create a new IAM role using the policy you defined above:

```python
import json

new_role = iam_client.create_role(
    AssumeRolePolicyDocument=json.dumps(policy),
    Path='/',
    RoleName=role_name
)
```

You need to know what your Amazon Resource Name (ARN) is when you create a compilation job in a later step, so store it in a variable as well.

```python
role_arn = new_role['Role']['Arn']
```

Now that you have created a new role, attach the permissions it needs to interact with Amazon SageMaker and Amazon S3:

```python
iam_client.attach_role_policy(
    RoleName=role_name,
    PolicyArn='arn:aws:iam::aws:policy/AmazonSageMakerFullAccess'
)

iam_client.attach_role_policy(
    RoleName=role_name,
    PolicyArn='arn:aws:iam::aws:policy/AmazonS3FullAccess'
)
```

4. **Create an Amazon S3 bucket to store your model artifacts**

   SageMaker Neo will access your model artifacts from Amazon S3

   Boto3

   ```python
   # Create an S3 client
   s3_client = boto3.client('s3', region_name=AWS_REGION)
   ```
# Name buckets
bucket='name-of-your-bucket'

# Check if bucket exists
if boto3.resource('s3').Bucket(bucket) not in boto3.resource('s3').buckets.all():
    s3_client.create_bucket(
        Bucket=bucket,
        CreateBucketConfiguration={
            'LocationConstraint': AWS_REGION
        }
    )
else:
    print(f'Bucket {bucket} already exists. No action needed.')

CLI

aws s3 mb s3://'name-of-your-bucket' --region specify-your-region

# Check your bucket exists
aws s3 ls s3://'name-of-your-bucket'/

5. **Train a machine learning model**

See [Train a Model with Amazon SageMaker](Train a Model with Amazon SageMaker) for more information on how to train a machine learning model using Amazon SageMaker. You can optionally upload your locally trained model directly into an Amazon S3 URI bucket.

**Note**

Make sure the model is correctly formatted depending on the framework you used. See [What input data shapes does SageMaker Neo expect?](What input data shapes does SageMaker Neo expect?)

If you do not have a model yet, use the curl command to get a local copy of the coco_ssd_mobilenet model from TensorFlow's website. The model you just copied is an object detection model trained from the COCO dataset. Type the following into your Jupyter notebook:

```python
model_zip_filename = './coco_ssd_mobilenet_v1_1.0.zip'
```
Note that this particular example was packaged in a .zip file. Unzip this file and repackage it as a compressed tarfile (.tar.gz) before using it in later steps. Type the following into your Jupyter notebook:

```python
# Extract model from zip file
!unzip -u {model_zip_filename}

model_filename = 'detect.tflite'
model_name = model_filename.split('.')[0]

# Compress model into .tar.gz so SageMaker Neo can use it
model_tar = model_name + '.tar.gz'
!tar -czf {model_tar} {model_filename}
```

6. **Upload trained model to an S3 bucket**

Once you have trained your machine learning mode, store it in an S3 bucket.

**Boto3**

```python
# Upload model
s3_client.upload_file(Filename=model_filename, Bucket=bucket, 
Key=model_filename)
```

**CLI**

Replace `your-model-filename` and `your-S3-bucket` with the name of your S3 bucket.

```
aws s3 cp your-model-filename s3://your-S3-bucket
```

**Step 1: Compile the Model**

Once you have satisfied the [Prerequisites](#), you can compile your model with Amazon SageMaker Neo. You can compile your model using the AWS CLI, the console or the [Amazon Web Services SDK for Python (Boto3)](https://boto3.amazonaws.com/v1/documentation/api/latest/index.html), see [Use Neo to Compile a Model](#). In this example, you will compile your model with Boto3.
To compile a model, SageMaker Neo requires the following information:

1. **The Amazon S3 bucket URI where you stored the trained model.**

   If you followed the prerequisites, the name of your bucket is stored in a variable named `bucket`. The following code snippet shows how to list all of your buckets using the AWS CLI:

   ```bash
   aws s3 ls
   ```

   For example:

   ```bash
   $ aws s3 ls
   2020-11-02 17:08:50 bucket
   ```

2. **The Amazon S3 bucket URI where you want to save the compiled model.**

   The code snippet below concatenates your Amazon S3 bucket URI with the name of an output directory called `output`:

   ```python
   s3_output_location = f's3://{bucket}/output'
   ```

3. **The machine learning framework you used to train your model.**

   Define the framework you used to train your model.

   ```python
   framework = 'framework-name'
   ```

   For example, if you wanted to compile a model that was trained using TensorFlow, you could either use `tflite` or `tensorflow`. Use `tflite` if you want to use a lighter version of TensorFlow that uses less storage memory.

   ```python
   framework = 'tflite'
   ```

   For a complete list of Neo-supported frameworks, see [Supported Frameworks, Devices, Systems, and Architectures](#).

4. **The shape of your model's input.**
Neo requires the name and shape of your input tensor. The name and shape are passed in as key-value pairs. `value` is a list of the integer dimensions of an input tensor and `key` is the exact name of an input tensor in the model.

```python
data_shape = '{"name": [tensor-shape]}'
```

For example:

```python
data_shape = '{"normalized_input_image_tensor":[1, 300, 300, 3]}'
```

⚠️ **Note**

Make sure the model is correctly formatted depending on the framework you used. See *What input data shapes does SageMaker Neo expect?* The key in this dictionary must be changed to the new input tensor’s name.

5. **Either the name of the target device to compile for or the general details of the hardware platform**

```python
target_device = 'target-device-name'
```

For example, if you want to deploy to a Raspberry Pi 3, use:

```python
target_device = 'rasp3b'
```

You can find the entire list of supported edge devices in *Supported Frameworks, Devices, Systems, and Architectures*.

Now that you have completed the previous steps, you can submit a compilation job to Neo.

```python
# Create a SageMaker client so you can submit a compilation job
sagemaker_client = boto3.client('sagemaker', region_name=AWS_REGION)

# Give your compilation job a name
compilation_job_name = 'getting-started-demo'
prefix = f'{compilation_job_name}-{}-
print(f'Compilation job for {compilation_job_name} started')
```
response = sagemaker_client.create_compilation_job(
    CompilationJobName=compilation_job_name,
    RoleArn=role_arn,
    InputConfig={
        'S3Uri': s3_input_location,
        'DataInputConfig': data_shape,
        'Framework': framework.upper()
    },
    OutputConfig={
        'S3OutputLocation': s3_output_location,
        'TargetDevice': target_device
    },
    StoppingCondition={
        'MaxRuntimeInSeconds': 900
    }
)

# Optional - Poll every 30 sec to check completion status
import time
while True:
    response =
    sagemaker_client.describe_compilation_job(CompilationJobName=compilation_job_name)
    if response['CompilationJobStatus'] == 'COMPLETED':
        break
    elif response['CompilationJobStatus'] == 'FAILED':
        raise RuntimeError('Compilation failed')
    print('Compiling ...')
    time.sleep(30)
print('Done!')

If you want additional information for debugging, include the following print statement:

print(response)

If the compilation job is successful, your compiled model is stored in the output Amazon S3 bucket you specified earlier (s3_output_location). Download your compiled model locally:

object_path = f'output/{model}-{target_device}.tar.gz'
neo_compiled_model = f'compiled-{model}.tar.gz'
s3_client.download_file(bucket, object_path, neo_compiled_model)
Step 2: Set Up Your Device

You will need to install packages on your edge device so that your device can make inferences. You will also need to either install AWS IoT Greengrass core or Deep Learning Runtime (DLR). In this example, you will install packages required to make inferences for the coco_ssd_mobilenet object detection algorithm and you will use DLR.

1. Install additional packages

In addition to Boto3, you must install certain libraries on your edge device. What libraries you install depends on your use case.

For example, for the coco_ssd_mobilenet object detection algorithm you downloaded earlier, you need to install NumPy for data manipulation and statistics, PIL to load images, and Matplotlib to generate plots. You also need a copy of TensorFlow if you want to gauge the impact of compiling with Neo versus a baseline.

```
!pip3 install numpy pillow tensorflow matplotlib
```

2. Install inference engine on your device

To run your Neo-compiled model, install the Deep Learning Runtime (DLR) on your device. DLR is a compact, common runtime for deep learning models and decision tree models. On x86_64 CPU targets running Linux, you can install the latest release of the DLR package using the following pip command:

```
!pip install dlr
```

For installation of DLR on GPU targets or non-x86 edge devices, refer to Releases for prebuilt binaries, or Installing DLR for building DLR from source. For example, to install DLR for Raspberry Pi 3, you can use:

```
```
Step 3: Make Inferences on Your Device

In this example, you will use Boto3 to download the output of your compilation job onto your edge device. You will then import DLR, download an example images from the dataset, resize this image to match the model’s original input, and then you will make a prediction.

1. **Download your compiled model from Amazon S3 to your device and extract it from the compressed tarfile.**

   ```python
   # Download compiled model locally to edge device
   object_path = f'output/{model_name}-{target_device}.tar.gz'
   neo_compiled_model = f'compiled-{model_name}.tar.gz'
   s3_client.download_file(bucket_name, object_path, neo_compiled_model)

   # Extract model from .tar.gz so DLR can use it
   !mkdir ./dlr_model # make a directory to store your model (optional)
   !tar -xzvf ./compiled-detect.tar.gz --directory ./dlr_model
   ```

2. **Import DLR and an initialized DLRModel object.**

   ```python
   import dlr
   device = 'cpu'
   model = dlr.DLRModel('./dlr_model', device)
   ```

3. **Download an image for inferencing and format it based on how your model was trained.**

   For the coco_ssd_mobilenet example, you can download an image from the COCO dataset and then reform the image to 300x300:

   ```python
   from PIL import Image

   # Download an image for model to make a prediction
   input_image_filename = './input_image.jpg'
   !curl https://farm9.staticflickr.com/8325/8077197378_79efb4805e_z.jpg --output {input_image_filename}

   # Format image so model can make predictions
   resized_image = image.resize((300, 300))

   # Model is quantized, so convert the image to uint8
   x = np.array(resized_image).astype('uint8')
   ```
4. **Use DLR to make inferences.**

   Finally, you can use DLR to make a prediction on the image you just downloaded:

   ```python
   out = model.run(x)
   ```

   For more examples using DLR to make inferences from a Neo-compiled model on an edge device, see the [neo-ai-dlr Github repository](https://github.com/neo-ai-dlr).

**Troubleshoot Errors**

This section contains information about how to understand and prevent common errors, the error messages they generate, and guidance on how to resolve these errors. Before moving on, ask yourself the following questions:

- **Did you encounter an error before you deployed your model?** If yes, see [Troubleshoot Neo Compilation Errors](#).

- **Did you encounter an error after you compiled your model?** If yes, see [Troubleshoot Neo Inference Errors](#).

- **Did you encounter an error trying to compile your model for Ambarella devices?** If yes, see [Troubleshoot Ambarella Errors](#).

**Error Classification Types**

This list classifies the *user errors* you can receive from Neo. These include access and permission errors and load errors for each of the supported frameworks. All other errors are *system errors*.

**Client permission error**

Neo passes the errors for these straight through from the dependent service.

- *Access Denied* when calling sts:AssumeRole
- *Any 400* error when calling Amazon S3 to download or upload a client model
- *PassRole* error
Load error

Assuming that the Neo compiler successfully loaded .tar.gz from Amazon S3, check whether the tarball contains the necessary files for compilation. The checking criteria is framework-specific:

- **TensorFlow**: Expects only protobuf file (*.pb or *.pbtxt). For saved models, expects one variables folder.
- **Pytorch**: Expect only one pytorch file (*.pth).
- **MXNET**: Expect only one symbol file (*.json) and one parameter file (*.params).
- **XGBoost**: Expect only one XGBoost model file (*.model). The input model has size limitation.

Compilation error

Assuming that the Neo compiler successfully loaded .tar.gz from Amazon S3, and that the tarball contains necessary files for compilation. The checking criteria is:

- **OperatorNotImplemented**: An operator has not been implemented.
- **OperatorAttributeNotImplemented**: The attribute in the specified operator has not been implemented.
- **OperatorAttributeRequired**: An attribute is required for an internal symbol graph, but it is not listed in the user input model graph.
- **OperatorAttributeValueNotValid**: The value of the attribute in the specific operator is not valid.

Topics

- Troubleshoot Neo Compilation Errors
- Troubleshoot Neo Inference Errors
- Troubleshoot Ambarella Errors

Troubleshoot Neo Compilation Errors

This section contains information about how to understand and prevent common compilation errors, the error messages they generate, and guidance on how to resolve these errors.

Topics

- How to Use This Page
How to Use This Page

Attempt to resolve your error by going through these sections in the following order:

1. Check that the input of your compilation job satisfies the input requirements. See [What input data shapes does SageMaker Neo expect?](#).
2. Check common **framework-specific errors**.
3. Check if your error is an **infrastructure error**.
4. Check your **compilation log**.

Framework-Related Errors

**Keras**

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<th>Solution</th>
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<td>Check your h5 file is in the Amazon S3 URI you specified. <em>Or</em> Check that the <strong>h5 file is correctly formatted</strong>.</td>
</tr>
<tr>
<td>InputConfiguration: Multiple h5 files provided, <code>&lt;model path&gt;</code>, when only one is allowed</td>
<td>Check you are only providing one h5 file.</td>
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<tr>
<td>ClientError: InputConfiguration: Unable to load provided Keras model. Error: 'sample_weight_mode'</td>
<td>Check the Keras version you specified is supported. See, supported frameworks for <a href="#">cloud instances</a> and <a href="#">edge devices</a>.</td>
</tr>
</tbody>
</table>
## Troubleshoot Errors

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>ClientError: InputConfiguration: Input input has wrong shape in Input Shape dictionary. Input shapes should be provided in NCHW format.</td>
<td>Check that your model input follows NCHW format. See <a href="#">What input data shapes does SageMaker Neo expect?</a></td>
</tr>
</tbody>
</table>

### MXNet

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>ClientError: InputConfiguration: Only one parameter file is allowed for MXNet model. Please make sure the framework you select is correct.</td>
<td>SageMaker Neo will select the first parameter file given for compilation.</td>
</tr>
</tbody>
</table>

### TensorFlow

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>InputConfiguration: Exactly one .pb file is allowed for TensorFlow models.</td>
<td>Make sure you only provide one .pb or .pbtxt file.</td>
</tr>
<tr>
<td>InputConfiguration: Exactly one .pb or .pbtxt file is allowed for TensorFlow models.</td>
<td>Make sure you only provide one .pb or .pbtxt file.</td>
</tr>
<tr>
<td>ClientError: InputConfiguration: TVM cannot convert &lt;model zoo&gt; model. Please make sure the framework you selected is correct. The following operators are not implemented: {&lt;operator name&gt;}</td>
<td>Check the operator you chose is supported. See <a href="#">SageMaker Neo Supported Frameworks and Operators</a></td>
</tr>
</tbody>
</table>
## PyTorch

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
</table>
| InputConfiguration: We are unable to extract DataInputConfig from the model due to *input_config_derivation_error*. Please override by providing a DataInputConfig during compilation job creation. | Do either of the following:  
- Specify the name and shape of the expected inputs by providing a DataInputConfig definition in your compilation request.  
- Investigate the error in Amazon CloudWatch Logs. Check the /aws/sagemaker/CompilationJobs log group and look for a log stream named *compilationJobName/model-info-extraction*. |

## Infrastructure-Related Errors

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>ClientError: InputConfiguration: S3 object does not exist. Bucket: &lt;bucket&gt;, Key: &lt;bucket key&gt;</td>
<td>Check the Amazon S3 URI your provided.</td>
</tr>
<tr>
<td>ClientError: InputConfiguration: Bucket &lt;bucket name&gt; is in region &lt;region name&gt; which is different from AWS Sagemaker service region &lt;service region&gt;</td>
<td>Create an Amazon S3 bucket that is in the same region as the service.</td>
</tr>
</tbody>
</table>
## Error and Solution Table

<table>
<thead>
<tr>
<th>Error</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>ClientError: InputConfiguration: Unable to untar input model. Please confirm the model is a tar.gz file</td>
<td>Check that your model in Amazon S3 is compressed into a tar.gz file.</td>
</tr>
</tbody>
</table>

## Check your compilation log

2. Select the region you created the compilation job from the Region dropdown list in the top right.
3. In the navigation pane of the Amazon CloudWatch, choose Logs. Select Log groups.
4. Search for the log group called /aws/sagemaker/CompilationJobs. Select the log group.
5. Search for the logstream named after the compilation job name. Select the log stream.

## Troubleshoot Neo Inference Errors

This section contains information about how to prevent and resolve some of the common errors you might encounter upon deploying and/or invoking the endpoint. This section applies to PyTorch 1.4.0 or later and MXNet v1.7.0 or later.

- Make sure the first inference (warm-up inference) on a valid input data is done in model_fn(), if you defined a model_fn in your inference script, otherwise the following error message may be seen on the terminal when predict API is called:

  ```
  An error occurred (ModelError) when calling the InvokeEndpoint operation: Received server error (0) from <users-sagemaker-endpoint> with message "Your invocation timed out while waiting for a response from container model. Review the latency metrics for each container in Amazon CloudWatch, resolve the issue, and try again."
  ```

- Make sure that the environment variables in the following table are set. If they are not set, the following error message might show up:

  **On the terminal:**
An error occurred (ModelError) when calling the InvokeEndpoint operation: Received server error (503) from <users-sagemaker-endpoint> with message "{ "code": 503, "type": "InternalServerException", "message": "Prediction failed" } ".

In CloudWatch:

W-9001-model-stdout com.amazonaws.ml.mms.wlm.WorkerLifeCycle - AttributeError: 'NoneType' object has no attribute 'transform'

<table>
<thead>
<tr>
<th>Key</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAGEMAKER_PROGRAM</td>
<td>inference.py</td>
</tr>
<tr>
<td>SAGEMAKER_SUBMIT_DIRECTORY</td>
<td>/opt/ml/model/code</td>
</tr>
<tr>
<td>SAGEMAKER_CONTAINER_LOG_LEVEL</td>
<td>20</td>
</tr>
<tr>
<td>SAGEMAKER_REGION</td>
<td>&lt;your region&gt;</td>
</tr>
</tbody>
</table>

- Make sure that the MMS_DEFAULT_RESPONSE_TIMEOUT environment variable is set to 500 or a higher value while creating the Amazon SageMaker model; otherwise, the following error message may be seen on the terminal:

An error occurred (ModelError) when calling the InvokeEndpoint operation: Received server error (0) from <users-sagemaker-endpoint> with message "Your invocation timed out while waiting for a response from container model. Review the latency metrics for each container in Amazon CloudWatch, resolve the issue, and try again."

Troubleshoot Ambarella Errors

SageMaker Neo requires models to be packaged in a compressed TAR file (*.tar.gz). Ambarella devices require additional files to be included within the compressed TAR file before it is sent for compilation. Include the following files within your compressed TAR file if you want to compile a model for Ambarella targets with SageMaker Neo:

- A trained model using a framework supported by SageMaker Neo
- A JSON configuration file
Calibration images

For example, the contents of your compressed TAR file should look similar to the following example:

```plaintext
###amba_config.json
###calib_data
|    ### data1
|    ### data2
|    ### .
|    ### .
|    ### .
|    ### data500
###mobilenet_v1_1.0_0224_frozen.pb
```

The directory is configured as follows:

- `amba_config.json`: Configuration file
- `calib_data`: Folder containing calibration images
- `mobilenet_v1_1.0_0224_frozen.pb`: TensorFlow model saved as a frozen graph

For information about frameworks supported by SageMaker Neo, see Supported Frameworks.

**Setting up the Configuration File**

The configuration file provides information required by the Ambarella toolchain to compile the model. The configuration file must be saved as a JSON file and the name of the file must end with `config.json`. The following chart shows the contents of the configuration file.

<table>
<thead>
<tr>
<th>Key</th>
<th>Description</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>inputs</td>
<td>Dictionary mapping input layers to attribute.</td>
<td>`{inputs:{&quot;data&quot;:{.. ..},&quot;data1&quot;:{...}}}</td>
</tr>
<tr>
<td>&quot;data&quot;</td>
<td>Input layer name. Note: &quot;data&quot; is an example of the name you can use to label the input layer.</td>
<td>&quot;data&quot;</td>
</tr>
<tr>
<td><strong>Key</strong></td>
<td><strong>Description</strong></td>
<td><strong>Example</strong></td>
</tr>
<tr>
<td>------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>------------------------------------------</td>
</tr>
<tr>
<td>shape</td>
<td>Describes the shape of the input to the model. This follows the same conventions that SageMaker Neo uses.</td>
<td>&quot;shape&quot;: &quot;1,3,224,224&quot;</td>
</tr>
<tr>
<td>filepath</td>
<td>Relative path to the directory containing calibration images. These can be binary or image files like JPG or PNG.</td>
<td>&quot;filepath&quot;: &quot;calib_data/&quot;</td>
</tr>
<tr>
<td>colorformat</td>
<td>Color format that model expects. This will be used while converting images to binary. Supported values: [RGB, BGR]. Default is RGB.</td>
<td>&quot;colorformat&quot;: &quot;RGB&quot;</td>
</tr>
<tr>
<td>mean</td>
<td>Mean value to be subtracted from the input. Can be a single value or a list of values. When the mean is given as a list the number of entries must match the channel dimension of the input.</td>
<td>&quot;mean&quot;: 128.0</td>
</tr>
<tr>
<td>scale</td>
<td>Scale value to be used for normalizing the input. Can be a single value or a list of values. When the scale is given as a list, the number of entries must match the channel dimension of the input.</td>
<td>&quot;scale&quot;: 255.0</td>
</tr>
</tbody>
</table>

The following is a sample configuration file:
Calibration Images

Quantize your trained model by providing calibration images. Quantizing your model improves the performance of the CVFlow engine on an Ambarella System on a Chip (SoC). The Ambarella toolchain uses the calibration images to determine how each layer in the model should be quantized to achieve optimal performance and accuracy. Each layer is quantized independently to INT8 or INT16 formats. The final model has a mix of INT8 and INT16 layers after quantization.

How many images should you use?

It is recommended that you include between 100–200 images that are representative of the types of scenes the model is expected to handle. The model compilation time increases linearly to the number of calibration images in the input file.

What are the recommended image formats?

Calibration images can be in a raw binary format or image formats such as JPG and PNG.

Your calibration folder can contain a mixture of images and binary files. If the calibration folder contains both images and binary files, the toolchain first converts the images to binary files. Once the conversion is complete, it uses the newly generated binary files along with the binary files that were originally in the folder.

Can I convert the images into binary format first?

Yes. You can convert the images to the binary format with open-source packages such as OpenCV or PIL. Crop and resize the images so they satisfy the input layer of your trained model.
Mean and Scale

You can specify mean and scaling pre-processing options to the Amberalla toolchain. These operations are embedded into the network and are applied during inference on each input. Do not provide processed data if you specify the mean or scale. More specifically, do not provide data you have subtracted the mean from or have applied scaling to.

Check your compilation log

For information on checking compilation log for Ambarella devices, see Check your compilation log.

Use Amazon SageMaker Elastic Inference (EI)

Starting April 15, 2023, AWS will not onboard new customers to Amazon Elastic Inference (EI), and will help current customers migrate their workloads to options that offer better price and performance. After April 15, 2023, new customers will not be able to launch instances with Amazon EI accelerators in Amazon SageMaker, Amazon ECS, or Amazon EC2.

Machine learning (ML) on AWS helps you innovate faster with the most comprehensive set of ML services and infrastructure made available in a low-cost, pay as-you-go usage model. AWS continuously delivers better performing and lower cost infrastructure for ML inference workloads. AWS launched Amazon Elastic Inference (EI) in 2018 to enable customers to attach low-cost GPU-powered acceleration to Amazon EC2, Amazon SageMaker instances, or Amazon Elastic Container Service (ECS) tasks to reduce the cost of running deep learning inference by up to 75% compared to standalone GPU based instances such as Amazon EC2 P4d and Amazon EC2 G5. In 2019, AWS launched AWS Inferentia, Amazon's first custom silicon designed to accelerate deep learning workloads by providing high performance inference in the cloud. Amazon EC2 Inf1 instances based on AWS Inferentia chips deliver up 2.3x higher throughput and up to 70% lower cost per inference than comparable current generation GPU-based Amazon EC2 instances. With the availability of new accelerated compute options such as AWS Inferentia and Amazon EC2 G5 instances, the benefit of attaching a fractional GPU to a CPU host instance using Amazon EI has diminished. For example, customers hosting models on Amazon EI who move to ml.inf1.xlarge instances can get up to 56% in cost savings and 2x performance improvement.

Customers can use Amazon SageMaker Inference Recommender to help them choose the best alternative instances to Amazon EI for deploying their ML models.

Frequently asked questions
1. Why is Amazon encouraging customers to move workloads from Amazon Elastic Inference (EI) to newer hardware acceleration options such as AWS Inferentia?

Customers get better performance at a much better price than Amazon EI with new hardware accelerator options such as AWS Inferentia for their inference workloads. AWS Inferentia is designed to provide high performance inference in the cloud, to drive down the total cost of inference, and to make it easy for developers to integrate machine learning into their business applications. To enable customers to benefit from such newer generation hardware accelerators, we will not onboard new customers to Amazon EI after April 15, 2023.

2. Which AWS services are impacted by the move to stop onboarding new customers to Amazon Elastic Inference (EI)?

This announcement will affect Amazon EI accelerators attached to any Amazon EC2, Amazon SageMaker instances, or Amazon Elastic Container Service (ECS) tasks. In Amazon SageMaker, this applies to both endpoints and notebook kernels using Amazon EI accelerators.

3. Will I be able to create a new Amazon Elastic Inference (EI) accelerator after April 15, 2023?

No, if you are a new customer and have not used Amazon EI in the past 30 days, then you will not be able create a new Amazon EI instance in your AWS account after April 15, 2023. However, if you have used an Amazon EI accelerator at least once in the past 30 days, you can attach a new Amazon EI accelerator to your instance.

4. How do I evaluate alternative instance options for my current Amazon SageMaker Inference Endpoints?

Amazon SageMaker Inference Recommender can help you identify cost-effective deployments to migrate existing workloads from Amazon Elastic Inference (EI) to an appropriate ML instance supported by SageMaker.

5. How do I change the instance type for my existing endpoint in Amazon SageMaker?

You can change the instance type for your existing endpoint by doing the following:

1. First, create a new EndpointConfig that uses the new instance type. If you have an autoscaling policy, delete the existing autoscaling policy.
2. Call UpdateEndpoint while specifying your newly created EndpointConfig.
3. Wait for your endpoint to change status to InService. This will take approximately 10-15 minutes.
4. Finally, if you need autoscaling for your new endpoint, create a new autoscaling policy for this new endpoint and ProductionVariant.

6. How do I change the instance type for my existing Amazon SageMaker Notebook Instance using Amazon Elastic Inference (EI)?

Choose Notebook instances in the SageMaker console, and then choose the Notebook Instance you want to update. Make sure the Notebook Instance has a Stopped status. Finally, you can choose Edit and change your instance type. Make sure that, when your Notebook Instance starts up, you select the right kernel for your new instance.

7. Is there a specific instance type which is a good alternative to Amazon Elastic Inference (EI)?

Every machine learning workload is unique. We recommend using Amazon SageMaker Inference Recommender to help you identify the right instance type for your ML workload, performance requirements, and budget. AWS Inferentia, specifically inf1.xlarge, is the best high performance and low-cost alternative for Amazon EI customers.

**Migrate from Amazon Elastic Inference to other instances**

The following information can help you migrate your SageMaker-hosted endpoints from instances that use Amazon Elastic Inference accelerators to other instances. The advice varies depending on your framework.

**PyTorch**

If you're migrating from PyTorch, use the following guidelines.

1. **Choose the right instance type**

Every machine learning workload is unique. We recommend using Amazon SageMaker Inference Recommender to help you identify the right instance type for your ML workload, performance requirements, and budget. AWS Inferentia, specifically inf1.xlarge, is the best high performance and low-cost alternative for Amazon Elastic Inference customers.

In our load testing with Inference Recommender, g4dn.xlarge instances performed better than m5.large instances with eia.2large attached. With Amazon Elastic Inference, you have to pay the additional cost of the ML instance to which the accelerator is attached. Amazon Elastic Inference also only supports PyTorch 1.5 and TensorFlow 2.3. If you migrate to ml.g4dn instances, you can use the latest versions of PyTorch 1.11 and TensorFlow 2.9. Additionally, ml.g4dn and
AWS Inferentia are available in all AWS Regions, whereas Amazon Elastic Inference is only available in 6 Regions. Both AWS Inferentia and ml.g4dn offer better performance at lower price for most ML inference workloads.

2. Modify `inference.py`

Modify your `inference.py` file to remove any Elastic Inference-specific required changes and use default handlers. Based on different user cases, you might have different input and output handlers, but the main changes you must make are in the model loading handler functions `model_fn` and `predict_fn`. Remove the Elastic Inference-specific predict handler `predict_fn` and restore the model loading handler `model_fn` to the default format. The following example shows how to do this, with the parts you should remove from `inference.py` commented out:

```python
from __future__ import print_function
import os
import torch
import torch.nn as nn
import torch.nn.functional as F
import numpy as np

def model_fn(model_dir, context):
    model = {customer_model}
    # if torch.__version__ in VERSIONS_USE_NEW_API:
    #     import torcheia
    #     loaded_model = loaded_model.eval()
    #     loaded_model = torcheia.jit.attach_eia(loaded_model, 0)
    with open(os.path.join(model_dir, 'model.pth'), 'rb') as f:
        model.load_state_dict(torch.load(f))
    return model

# def predict_fn(input_data, model):
#     logger.info("Performing EIA inference with Torch JIT context with input of size {}\n".format(input_data.shape))
#     device = torch.device("cuda" if torch.cuda.is_available() else "cpu")
#     input_data = input_data.to(device)
```

Migrate from Amazon Elastic Inference to other instances
#     with torch.no_grad():
#        if torch.__version__ in VERSIONS_USE_NEW_API:
#            import torcheia
#            torch._C._jit_set_profiling_executor(False)
#            with torch.jit.optimized_execution(True):
#                return model.forward(input_data)
#        else:
#            with torch.jit.optimized_execution(True, {"target_device": "eia:0"}):
#                return model(input_data)

def predict_fn(input_data, model):
    return model(input_data)

### 3. Create a model

Create a new model that points to your modified inference.py file. You can keep the
inference.py file locally and point to it by specifying source_dir and entry_point or tar the
inference.py file into the model tarball. The following example shows the former case:

```python
from sagemaker.pytorch import PyTorchModel

pytorch = PyTorchModel(
    model_data={model_data_url},
    role=role,
    entry_point="inference.py",
    source_dir="code",
    framework_version="1.5.1",
    py_version="py3",
    sagemaker_session=sagemaker_session,
)
```

### 4. Deploy the model to the endpoint and invoke it

You can use one of the following options for deploying your model after making the preceding
changes.

**Option 1: Deploy from scratch**

You can deploy the model to a new endpoint with a recommended instance from the **Accelerated
Computing** category, such as G4.

```python
predictor = pytorch.deploy(
```
### Option 2: Update the existing endpoint

Complete the following steps to update your existing endpoint:

1. **Call CreateEndpointConfig** to create a new EndpointConfig that uses the new instance type. If you have an autoscaling policy, delete the existing autoscaling policy.

   ```python
   endpoint_config_response = sagemaker_client.create_endpoint_config(
       EndpointConfigName=endpoint_config_name,
       ProductionVariants=[
           {
               "VariantName": "variant1",  # The name of the production variant.
               "ModelName": model_name,  # The name of new created model
               "InstanceType": instance_type,  # Specify the right-sized instance type.
               "InitialInstanceCount": 1  # Number of instances to launch initially.
           }
       ]
   )
   ```

2. **Call UpdateEndpoint** and specify your newly created EndpointConfig.

   ```python
   endpoint_config_response = sagemaker_client.update_endpoint(
       EndpointConfigName=endpoint_config_name,  # The name of the new endpoint config just created
       EndpointName=endpoint_name  # The name of the existing endpoint you want to update
   )
   ```

3. **Wait for your endpoint to change status to InService.** This takes approximately 10–15 minutes.

4. **Finally, if you need autoscaling for your new endpoint,** create a new autoscaling policy for your new endpoint and ProductionVariant.
**TensorFlow**

If you're migrating from TensorFlow, use the following guidelines.

1. **Choose the right instance type**

Refer to the 1. Choose the right instance type guidance in the [PyTorch section](#).

2. **Deploy the model to the endpoint and invoke it**

You can use one of the following options for deploying your model.

*Option 1: Deploy from scratch*

You can migrate from Elastic Inference by re-deploying the model to a new endpoint by removing the accelerator_type field and specifying a right-sized instance type from the Accelerated Computing category, such as G4. In the following example, the commented out line causes you to deploy without using an Elastic Inference accelerator.

```python
predictor = tensorflow_model.deploy(
    ...
    instance_type="ml.g4dn.2xlarge"
    # instance_type="ml.c5.xlarge",
    # accelerator_type="ml.eia1.medium"
    ...
)
```

*Option 2: Update the existing endpoint*

Refer to the Option 2. Update the existing endpoint guidance in Step 4 of the [PyTorch section](#).

**MXNet**

If you're migrating from MXNet, use the following guidelines.

1. **Choose the right instance type**

Refer to the 1. Choose the right instance type guidance in the [PyTorch section](#).

2. **Deploy the model to the endpoint and invoke it**

You can use one of the following options for deploying your model.
Option 1: Deploy from scratch

You can migrate from Elastic Inference by re-deploying the model to a new endpoint by removing the accelerator_type field and specifying a right-sized instance type from the Accelerated Computing category, such as G4. In the following example, the commented out line causes you to deploy without using an Elastic Inference accelerator.

```python
predictor = mxnet_model.deploy(
    ...
    # instance_type="ml.c5.xlarge",
    instance_type="ml.g4dn.2xlarge"
    ...
)
```

Option 2: Update the existing endpoint

Refer to the Option 2: Update the existing endpoint guidance in Step 4 of the PyTorch section.

Topics
- How EI Works
- Choose an EI Accelerator Type
- Use EI in a SageMaker Notebook Instance
- Use EI on a Hosted Endpoint
- Frameworks that Support EI
- Use EI with SageMaker Built-in Algorithms
- EI Sample Notebooks
- Set Up to Use EI
- Attach EI to a Notebook Instance
- Use EI on Amazon SageMaker Hosted Endpoints

How EI Works

Amazon Elastic Inference accelerators are network attached devices that work along with SageMaker instances in your endpoint to accelerate your inference calls. Elastic Inference accelerates inference by allowing you to attach fractional GPUs to any SageMaker instance. You can select the client instance to run your application and attach an Elastic Inference accelerator to use
the right amount of GPU acceleration for your inference needs. Elastic Inference helps you lower your cost when not fully utilizing your GPU instance for inference. We recommend trying Elastic Inference with your model using different CPU instances and accelerator sizes.

The following EI accelerator types are available. You can configure your endpoints or notebook instances with any EI accelerator type.

In the table, the throughput in teraflops (TFLOPS) is listed for both single-precision floating-point (F32) and half-precision floating-point (F16) operations. The memory in GB is also listed.

<table>
<thead>
<tr>
<th>Accelerator Type</th>
<th>F32 Throughput in TFLOPS</th>
<th>F16 Throughput in TFLOPS</th>
<th>Memory in GB</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml.eia2.medium</td>
<td>1</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>ml.eia2.large</td>
<td>2</td>
<td>16</td>
<td>4</td>
</tr>
<tr>
<td>ml.eia2.xlarge</td>
<td>4</td>
<td>32</td>
<td>8</td>
</tr>
<tr>
<td>ml.eia1.medium</td>
<td>1</td>
<td>8</td>
<td>1</td>
</tr>
<tr>
<td>ml.eia1.large</td>
<td>2</td>
<td>16</td>
<td>2</td>
</tr>
<tr>
<td>ml.eia1.xlarge</td>
<td>4</td>
<td>32</td>
<td>4</td>
</tr>
</tbody>
</table>

**Choose an EI Accelerator Type**

Consider the following factors when choosing an accelerator type for a hosted model:

- Models, input tensors and batch sizes influence the amount of accelerator memory you need. Start with an accelerator type that provides at least as much memory as the file size of your trained model. Factor in that a model might use significantly more memory than the file size at runtime.

- Demands on CPU compute resources, main system memory, and GPU-based acceleration and accelerator memory vary significantly between different kinds of deep learning models. The latency and throughput requirements of the application also determine the amount of compute and acceleration you need. Thoroughly test different configurations of instance types and EI accelerator sizes to make sure you choose the configuration that best fits the performance needs of your application.
For more information on selecting an EI accelerator, see:

- Amazon Elastic Inference Overview
- Choosing an Instance and Accelerator Type for Your Model
- Optimizing costs in Amazon Elastic Inference with TensorFlow

**Use EI in a SageMaker Notebook Instance**

Typically, you build and test machine learning models in a SageMaker notebook before you deploy them for production. You can attach EI to your notebook instance when you create the notebook instance. You can set up an endpoint that is hosted locally on the notebook instance by using the local mode supported by TensorFlow, MXNet, and PyTorch estimators and models in the Amazon SageMaker Python SDK to test inference performance. Elastic Inference enabled PyTorch is not currently supported on notebook instances. For instructions on how to attach EI to a notebook instance and set up a local endpoint for inference, see Attach EI to a Notebook Instance. There are also Elastic Inference-enabled SageMaker Notebook Jupyter kernels for Elastic Inference-enabled versions of TensorFlow and Apache MXNet. For information about using SageMaker notebook instances, see Use Amazon SageMaker Notebook Instances

**Use EI on a Hosted Endpoint**

When you are ready to deploy your model for production to provide inferences, you create a SageMaker hosted endpoint. You can attach EI to the instance where your endpoint is hosted to increase its performance at providing inferences. For instructions on how to attach EI to a hosted endpoint instance, see Use EI on Amazon SageMaker Hosted Endpoints.

**Frameworks that Support EI**

Amazon Elastic Inference is designed to be used with AWS enhanced versions of TensorFlow, Apache MXNet, or PyTorch machine learning frameworks. These enhanced versions of the frameworks are automatically built into containers when you use the Amazon SageMaker Python SDK, or you can download them as binary files and import them in your own Docker containers.

You can download the EI-enabled TensorFlow binary files from the public amazonei-tensorflow Amazon S3 bucket to the TensorFlow serving containers. For more information about building a container that uses the EI-enabled version of TensorFlow, see Amazon Elastic Inference with TensorFlow in SageMaker.
You can download the EI-enabled MXNet binary files from the public amazonei-apachemxnet Amazon S3 bucket to the MXNet serving containers. For more information about building a container that uses the EI-enabled version of MXNet, see Amazon Elastic Inference with MXNet in SageMaker.

You can download the Elastic Inference enabled binary for PyTorch. For more information about building a container that uses the EI-enabled version of PyTorch, see Amazon Elastic Inference with PyTorch in SageMaker.

To use Elastic Inference in a hosted endpoint, you can choose any of the following frameworks depending on your needs:

- SageMaker Python SDK - Deploy TensorFlow models
- SageMaker Python SDK - Deploy MXNet models
- SageMaker Python SDK - Deploy PyTorch models

If you need to create a custom container for deploying your model that is complex and requires extensions to a framework that the SageMaker pre-built containers do not support, use the low-level AWS SDK for Python (Boto 3).

### Use EI with SageMaker Built-in Algorithms

Currently, the Image Classification - MXNet and Object Detection - MXNet built-in algorithms support EI. For an example that uses the Image Classification algorithm with EI, see End-to-End Multiclass Image Classification Example.

### EI Sample Notebooks

The following Sample notebooks provide examples of using EI in SageMaker:

- Using Amazon Elastic Inference with MXNet on Amazon SageMaker
- Using Amazon Elastic Inference with MXNet on an Amazon SageMaker Notebook Instance
- Using Amazon Elastic Inference with Neo-compiled TensorFlow model on SageMaker
- Using Amazon Elastic Inference with a pre-trained TensorFlow Serving model on SageMaker

### Set Up to Use EI

Use the instructions in this topic only if one of the following applies to you:
• You want to use a customized role or permission policy.
• You want to use a VPC for your hosted model or notebook instance.

Note
If you already have an execution role that has the AmazonSageMakerFullAccess managed policy attached (this is true for any IAM role that you create when you create a notebook instance, training job, or model in the console) and you are not connecting to an EI model or notebook instance in a VPC, you do not need to make any of these changes to use EI in Amazon SageMaker.

Topics
• Set Up Required Permissions
• Use a Custom VPC to Connect to EI

Set Up Required Permissions
To use EI in SageMaker, the role that you use to open a notebook instance or create a deployable model must have a policy with the required permissions attached. You can attach the AmazonSageMakerFullAccess managed policy, which contains the required permissions, to the role, or you can add a custom policy that has the required permissions. For information about creating an IAM role, see Creating a Role for an AWS Service (Console) in the AWS Identity and Access Management User Guide. For information about attaching a policy to a role, see Adding and Removing IAM Policies.

Add these permissions specifically for connecting EI in an IAM policy.

```json
{
    "Effect": "Allow",
    "Action": [
        "elastic-inference:Connect",
        "ec2:DescribeVpcEndpoints"
    ],
    "Resource": "*
}
```
The following IAM policy is the complete list of required permissions to use EI in SageMaker.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "elastic-inference:Connect",
                "ec2:DescribeVpcEndpoints"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:*"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "ecr:GetAuthorizationToken",
                "ecr:GetDownloadUrlForLayer",
                "ecr:BatchGetImage",
                "ecr:BatchCheckLayerAvailability",
                "cloudwatch:PutMetricData",
                "cloudwatch:PutMetricAlarm",
                "cloudwatch:DescribeAlarms",
                "cloudwatch:DeleteAlarms",
                "ec2:CreateNetworkInterface",
                "ec2:CreateNetworkInterfacePermission",
                "ec2:DeleteNetworkInterface",
                "ec2:DeleteNetworkInterfacePermission",
                "ec2:DescribeNetworkInterfaces",
                "ec2:DescribeVpcs",
                "ec2:DescribeDhcpOptions",
                "ec2:DescribeSubnets",
                "ec2:DescribeSecurityGroups",
                "application-autoscaling:DeleteScalingPolicy",
                "application-autoscaling:DeleteScheduledAction",
                "application-autoscaling:DeregisterScalableTarget",
                "application-autoscaling:DescribeScalableTargets",
```
"application-autoscaling:DescribeScalingActivities",
"application-autoscaling:DescribeScalingPolicies",
"application-autoscaling:DescribeScheduledActions",
"application-autoscaling:PutScalingPolicy",
"application-autoscaling:PutScheduledAction",
"application-autoscaling:RegisterScalableTarget",
"logs:CreateLogGroup",
"logs:CreateLogStream",
"logs:DescribeLogStreams",
"logs:GetLogEvents",
"logs:PutLogEvents"
],
"Resource": "*"
},
{
  "Effect": "Allow",
  "Action": [
    "s3:GetObject",
    "s3:PutObject",
    "s3:DeleteObject"
  ],
  "Resource": [
    "arn:aws:s3:::*SageMaker*",
    "arn:aws:s3:::*Sagemaker*",
    "arn:aws:s3:::*sagemaker*
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:GetObject"
  ],
  "Resource": "*",
  "Condition": {
    "Set Up to Use EI": 4667
  }
}
Use a Custom VPC to Connect to EI

To use EI with SageMaker in a VPC, you need to create and configure two security groups, and set up a PrivateLink VPC interface endpoint. EI uses VPC interface endpoint to communicate with SageMaker endpoints in your VPC. The security groups you create are used to connect to the VPC interface endpoint.

Set up Security Groups to Connect to EI

To use EI within a VPC, you need to create two security groups:
• A security group to control access to the VPC interface endpoint that you will set up for EI.
• A security group that allows SageMaker to call into the first security group.

To configure the two security groups

1. Create a security group with no outbound connections. You will attach this to the VPC endpoint interface you create in the next section.
2. Create a second security group with no inbound connections, but with an outbound connection to the first security group.
3. Edit the first security group to allow inbound connections only to the second security group and all outbound connections.

For more information about VPC security groups, see Security Groups for Your VPC in the Amazon Virtual Private Cloud User Guide.

Set up a VPC Interface Endpoint to Connect to EI

To use EI with SageMaker in a custom VPC, you need to set up a VPC interface endpoint (PrivateLink) for the EI service.

• Set up a VPC interface endpoint (PrivateLink) for the EI. Follow the instructions at Creating an Interface Endpoint. In the list of services, choose com.amazonaws.<region>.elastic-inference.runtime. For Security group, make sure you select the first security group you created in the previous section to the endpoint.
• When you set up the interface endpoint, choose all of the Availability Zones where EI is available. EI fails if you do not set up at least two Availability Zones. For information about VPC subnets, see VPCs and Subnets.

Attach EI to a Notebook Instance

To test and evaluate inference performance using EI, you can attach EI to a notebook instance when you create or update a notebook instance. You can then use EI in local mode to host a model at an endpoint hosted on the notebook instance. You should test various sizes of notebook instances and EI accelerators to evaluate the configuration that works best for your use case.
Set Up to Use EI

To use EI locally in a notebook instance, create a notebook instance with an EI instance.

To create a notebook instance with an EI instance

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the navigation pane, choose Notebook instances.
3. Choose Create notebook instance.
4. For Notebook instance name, provide a unique name for your notebook instance.
5. For notebook instance type, choose a CPU instance such as ml.t2.medium.
6. For Elastic Inference (EI), choose an instance from the list, such as ml.eia2.medium.
7. For IAM role, choose an IAM role that has the required permissions to use SageMaker and EI.
8. (Optional) For VPC - Optional, if you want the notebook instance to use a VPC, choose one from the available list. Otherwise, leave it as No VPC. If you use a VPC follow the instructions at Use a Custom VPC to Connect to EI.
9. (Optional) For Lifecycle configuration - optional, either leave it as No configuration or choose a lifecycle configuration. For more information, see Customize a Notebook Instance Using a Lifecycle Configuration Script.
10. (Optional) For Encryption key - optional, Optional) If you want SageMaker to use an AWS Key Management Service (AWS KMS) key to encrypt data in the ML storage volume attached to the notebook instance, specify the key.
11. (Optional) For Volume Size In GB - optional, leave the default value of 5.
12. (Optional) For Tags, add tags to the notebook instance. A tag is a label you assign to help manage your notebook instances. A tag consists of a key and a value, both of which you define.
13. Choose Create Notebook Instance.

After you create your notebook instance with EI attached, you can create a Jupyter notebook and set up an EI endpoint that is hosted locally on the notebook instance.

Topics

• Use EI in Local Mode in SageMaker
Use EI in Local Mode in SageMaker

To use EI locally in an endpoint hosted on a notebook instance, use local mode with the Amazon SageMaker Python SDK versions of either the TensorFlow, MXNet, or PyTorch estimators or models. For more information about local mode support in the SageMaker Python SDK, see https://github.com/aws/sagemaker-python-sdk#sagemaker-python-sdk-overview.

Topics

- Use EI in Local Mode with SageMaker TensorFlow Estimators and Models
- Use EI in Local Mode with SageMaker Apache MXNet Estimators and Models
- Use EI in Local Mode with SageMaker PyTorch Estimators and Models

Use EI in Local Mode with SageMaker TensorFlow Estimators and Models

To use EI with TensorFlow in local mode, specify local for instance_type and local_sagemaker_notebook for accelerator_type when you call the deploy method of an estimator or a model object. For more information about Amazon SageMaker Python SDK TensorFlow estimators and models, see https://sagemaker.readthedocs.io/en/stable/frameworks/tensorflow/index.html.

The following code shows how to use local mode with an estimator object. To call the deploy method, you must have previously either:

- Trained the model by calling the fit method of an estimator.
- Pass a model artifact when you initialize the model object.

```python
# Deploys the model to a local endpoint
tf_predictor = tf_model.deploy(initial_instance_count=1,
                               instance_type='local',
                               accelerator_type='local_sagemaker_notebook')
```

Use EI in Local Mode with SageMaker Apache MXNet Estimators and Models

To use EI with MXNet in local mode, specify local for instance_type and local_sagemaker_notebook for accelerator_type when you call the deploy method of an estimator or a model object. For more information about Amazon SageMaker Python SDK MXNet estimators and models, see https://sagemaker.readthedocs.io/en/stable/frameworks/mxnet/index.html.
The following code shows how to use local mode with an estimator object. You must have previously called the `fit` method of the estimator to train the model.

```python
# Deploys the model to a local endpoint
mxnet_predictor = mxnet_estimator.deploy(initial_instance_count=1,
                                          instance_type='local',
                                          accelerator_type='local_sagemaker_notebook')
```

For a complete example of using EI in local mode with MXNet, see the sample notebook at https://sagemaker-examples.readthedocs.io/en/latest/sagemaker-python-sdk/mxnet_mnist/mxnet_mnist_elastic_inference_local.html.

**Use EI in Local Mode with SageMaker PyTorch Estimators and Models**

To use EI with PyTorch in local mode, when you call the `deploy` method of an estimator or a model object, specify `local` for `instance_type` and `local_sagemaker_notebook` for `accelerator_type`. For more information about Amazon SageMaker Python SDK PyTorch estimators and models, see SageMaker PyTorch Estimators and Models.

The following code shows how to use local mode with an estimator object. You must have previously called the `fit` method of the estimator to train the model.

```python
# Deploys the model to a local endpoint
pytorch_predictor = pytorch_estimator.deploy(initial_instance_count=1,
                                             instance_type='local',
                                             accelerator_type='local_sagemaker_notebook')
```

**Use EI on Amazon SageMaker Hosted Endpoints**

To use Elastic Inference (EI) in Amazon SageMaker with a hosted endpoint for real-time inference, specify an EI accelerator when you create the deployable model to be hosted at that endpoint. You can do this in one of the following ways:

- Use the Amazon SageMaker Python SDK versions of either the TensorFlow, MXNet, or PyTorch and the SageMaker pre-built containers for TensorFlow, MXNet, and PyTorch
- Build your own container, and use the low-level SageMaker API (Boto 3). You will need to import the EI-enabled version of either TensorFlow, MXNet, or PyTorch from the provided Amazon S3 locations into your container, and use one of those versions to write your training script.
Use either the Image Classification - MXNet or Object Detection - MXNet build-in algorithms, and use the AWS SDK for Python (Boto3) to run your training job and create your deployable model and hosted endpoint.

Topics

- Use EI with a SageMaker TensorFlow Container
- Use EI with a SageMaker MXNet Container
- Use EI with a SageMaker PyTorch Container
- Use EI with Your Own Container

Use EI with a SageMaker TensorFlow Container


SageMaker provides default model training and inference code for your convenience. For custom file formats, you might need to implement custom model training and inference code.

Use an Estimator Object

To use an estimator object with EI, when you use the deploy method, include the accelerator_type input argument. The estimator returns a predictor object, which we call its deploy method, as shown in the example code.

```python
# Deploy an estimator using EI (using the accelerator_type input argument)
predictor = estimator.deploy(initial_instance_count=1,
                             instance_type='ml.m4.xlarge',
                             accelerator_type='ml.eia2.medium')
```

Use a Model Object

To use a model object with EI, when you use the deploy method, include the accelerator_type input argument. The estimator returns a predictor object, which we call its deploy method, as shown in the example code.

```python
# Deploy a model using EI (using the accelerator_type input argument)
```
predictor = model.deploy(initial_instance_count=1,
                        instance_type='ml.m4.xlarge',
                        accelerator_type='ml.eia2.medium')

Use EI with a SageMaker MXNet Container

To use MXNet with EI in SageMaker, you need to call the deploy method of either the `Estimator` or `Model` objects. You then specify an accelerator type using the `accelerator_type` input argument. For information about using MXNet in the Amazon SageMaker Python SDK, see https://sagemaker.readthedocs.io/en/stable/frameworks/mxnet/index.html

For your convenience, SageMaker provides default model training and inference code. For custom file formats, you might need to write custom model training and inference code.

Use an Estimator Object

To use an estimator object with EI, when you use the deploy method, include the `accelerator_type` input argument. The estimator returns a predictor object, which we call its deploy method, as shown in the example code.

```python
# Deploy an estimator using EI (using the accelerator_type input argument)
predictor = estimator.deploy(initial_instance_count=1,
                              instance_type='ml.m4.xlarge',
                              accelerator_type='ml.eia2.medium')
```

Use a Model Object

To use a model object with EI, when you use the deploy method, include the `accelerator_type` input argument. The estimator returns a predictor object, which we call its deploy method, as shown in the example code.

```python
# Deploy a model using EI (using the accelerator_type input argument)
predictor = model.deploy(initial_instance_count=1,
                          instance_type='ml.m4.xlarge',
                          accelerator_type='ml.eia2.medium')
```

Use EI with a SageMaker PyTorch Container

To use PyTorch with EI in SageMaker, you need to call the deploy method of either the `Estimator` or `Model` objects. You then specify an accelerator type using the `accelerator_type` input argument.
argument. For information about using PyTorch in the Amazon SageMaker Python SDK, see SageMaker PyTorch Estimators and Models.

For your convenience, SageMaker provides default model training and inference code. For custom file formats, you might need to write custom model training and inference code.

### Use an Estimator Object

To use an estimator object with EI, when you use the deploy method, include the `accelerator_type` input argument. The estimator returns a predictor object, which we call its deploy method, as shown in this example code.

```python
# Deploy an estimator using EI (using the accelerator_type input argument)
predictor = estimator.deploy(initial_instance_count=1,
                             instance_type='ml.m4.xlarge',
                             accelerator_type='ml.eia2.medium')
```

### Use a Model Object

To use a model object with EI, when you use the deploy method, include the `accelerator_type` input argument. The model returns a predictor object, which we call its deploy method, as shown in this example code.

```python
# Deploy a model using EI (using the accelerator_type input argument)
predictor = model.deploy(initial_instance_count=1,
                          instance_type='ml.m4.xlarge',
                          accelerator_type='ml.eia2.medium')
```

### Use EI with Your Own Container

To use EI with a model in a custom container that you build, use the low-level AWS SDK for Python (Boto 3). download and import the AWS EI-enabled versions of TensorFlow, Apache MXNet, or PyTorch machine learning frameworks, and write your training script using those frameworks.

### Import the EI Version of TensorFlow, MXNet, or PyTorch into Your Docker Container

To use EI with your own container, you need to import either the Amazon EI TensorFlow Serving library, the Amazon EI Apache MXNet library, or the Elastic Inference enabled PyTorch library into your container. The EI-enabled versions of TensorFlow and MXNet are currently available as binary files stored in Amazon S3 locations. You can download the EI-enabled binary for TensorFlow
from the Amazon S3 bucket at console.aws.amazon.com/s3/buckets/amazonei-tensorflow. For information about building a container that uses the EI-enabled version of TensorFlow, see https://github.com/aws/sagemaker-tensorflow-container#building-the-sagemaker-elastic-inference-tensorflow-serving-container. You can download the EI-enabled binary for Apache MXNet from the public Amazon S3 bucket at console.aws.amazon.com/s3/buckets/amazonei-apachemxnet. For information about building a container that uses the EI-enabled version of MXNet, see https://github.com/aws/sagemaker-mxnet-container#building-the-sagemaker-elastic-inference-mxnet-container. You can download the Elastic Inference enabled binary for PyTorch. For information about building a container that uses the Elastic Inference enabled version of PyTorch, see Building your image.

Create an EI Endpoint with AWS SDK for Python (Boto 3)

To create an endpoint by using AWS SDK for Python (Boto 3), you first create an endpoint configuration. The endpoint configuration specifies one or more models (called production variants) that you want to host at the endpoint. To attach EI to one or more of the production variants hosted at the endpoint, you specify one of the EI instance types as the AcceleratorType field for that ProductionVariant. You then pass that endpoint configuration when you create the endpoint.

Create an Endpoint Configuration

To use EI, you need to specify an accelerator type in the endpoint configuration.

```python
# Create Endpoint Configuration
from time import gmtime, strftime

endpoint_config_name = 'ImageClassificationEndpointConfig-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
print(endpoint_config_name)
create_endpoint_config_response = sagemaker.create_endpoint_config(
    EndpointConfigName = endpoint_config_name,
    ProductionVariants=[[
        'InstanceType':'ml.m4.xlarge',
        'InitialInstanceCount':1,
        'ModelName':model_name,
        'VariantName':'AllTraffic',
        'AcceleratorType':'ml.eia2.medium']])

print("Endpoint Config Arn: " + create_endpoint_config_response['EndpointConfigArn'])
```
Create an Endpoint

After you create an endpoint configuration with an accelerator type, you can create an endpoint.

```python
endpoint_name = 'ImageClassificationEndpoint-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
endpoint_response = sagemaker.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name)
```

After creating the endpoint, you can invoke it using the `invoke_endpoint` method in a Boto3 runtime object, as you would any other endpoint.

Best practices

The following topics provide guidance on best practices for deploying machine learning models in Amazon SageMaker.

Topics

- Best practices for deploying models on SageMaker Hosting Services
- Monitor Security Best Practices
- Low latency real-time inference with AWS PrivateLink
- Migrate inference workload from x86 to AWS Graviton
- Troubleshoot Amazon SageMaker model deployments
- Inference cost optimization best practices
- Best practices to minimize interruptions during GPU driver upgrades
- Best practices for endpoint security and health with Amazon SageMaker

Best practices for deploying models on SageMaker Hosting Services

When hosting models using SageMaker hosting services, consider the following:

- Typically, a client application sends requests to the SageMaker HTTPS endpoint to obtain inferences from a deployed model. You can also send requests to this endpoint from your Jupyter notebook during testing.
- You can deploy a model trained with SageMaker to your own deployment target. To do that, you need to know the algorithm-specific format of the model artifacts that were generated by
model training. For more information about output formats, see the section corresponding to the algorithm you are using in Common Data Formats for Training.

- You can deploy multiple variants of a model to the same SageMaker HTTPS endpoint. This is useful for testing variations of a model in production. For example, suppose that you've deployed a model into production. You want to test a variation of the model by directing a small amount of traffic, say 5%, to the new model. To do this, create an endpoint configuration that describes both variants of the model. You specify the ProductionVariant in your request to the CreateEndPointConfig. For more information, see ProductionVariant.

- You can configure a ProductionVariant to use Application Auto Scaling. For information about configuring automatic scaling, see Automatically Scale Amazon SageMaker Models.

- You can modify an endpoint without taking models that are already deployed into production out of service. For example, you can add new model variants, update the ML Compute instance configurations of existing model variants, or change the distribution of traffic among model variants. To modify an endpoint, you provide a new endpoint configuration. SageMaker implements the changes without any downtime. For more information see, UpdateEndpoint and UpdateEndpointWeightsAndCapacities.

- Changing or deleting model artifacts or changing inference code after deploying a model produces unpredictable results. If you need to change or delete model artifacts or change inference code, modify the endpoint by providing a new endpoint configuration. Once you provide the new endpoint configuration, you can change or delete the model artifacts corresponding to the old endpoint configuration.

- If you want to get inferences on entire datasets, consider using batch transform as an alternative to hosting services. For information, see Use Batch Transform

**Deploy Multiple Instances Across Availability Zones**

Create robust endpoints when hosting your model. SageMaker endpoints can help protect your application from Availability Zone outages and instance failures. If an outage occurs or an instance fails, SageMaker automatically attempts to distribute your instances across Availability Zones. For this reason, we strongly recommend that you deploy multiple instances for each production endpoint.

If you are using an Amazon Virtual Private Cloud (VPC), configure the VPC with at least two Subnets, each in a different Availability Zone. If an outage occurs or an instance fails, Amazon SageMaker automatically attempts to distribute your instances across Availability Zones.
In general, to achieve more reliable performance, use more small Instance Types in different Availability Zones to host your endpoints.

**Deploy inference components for high availability.** In addition to the above recommendation for instance numbers, to achieve 99.95% availability, ensure that your inference components are configured to have more than two copies. In addition, in your managed auto scaling policy, set the minimum number of instances to two as well.

**Monitor Security Best Practices**


**Low latency real-time inference with AWS PrivateLink**

Amazon SageMaker provides low latency for real-time inferences while maintaining high availability and resiliency using multi-AZ deployment. The application latency is made up of two primary components: infrastructure or overhead latency and model inference latency. Reduction of overhead latency opens up new possibilities such as deploying more complex, deep, and accurate models or splitting monolithic applications into scalable and maintainable microservice modules. You can reduce the latency for real-time inferences with SageMaker using an AWS PrivateLink deployment. With AWS PrivateLink, you can privately access all SageMaker API operations from your Virtual Private Cloud (VPC) in a scalable manner by using interface VPC endpoints. An interface VPC endpoint is an elastic network interface in your subnet with private IP addresses that serves as an entry point for all SageMaker API calls.

By default, a SageMaker endpoint with 2 or more instances is deployed in at least 2 AWS Availability Zones (AZs) and instances in any AZ can process invocations. This results in one or more AZ “hops” that contribute to the overhead latency. An AWS PrivateLink deployment with the privateDNSEnabled option set as true alleviates this by achieving two objectives:

- It keeps all inference traffic within your VPC.
- It keeps invocation traffic in the same AZ as the client that originated it when using SageMaker Runtime. This avoids the “hops” between AZs reducing the overhead latency.
The following sections of this guide demonstrate how you can reduce the latency for real-time inferences with AWS PrivateLink deployment.

**Topics**
- [Deploy AWS PrivateLink](#)
- [Deploy SageMaker endpoint in a VPC](#)
- [Invoke the SageMaker endpoint](#)

**Deploy AWS PrivateLink**

To deploy AWS PrivateLink, first create an interface endpoint for the VPC from which you connect to the SageMaker endpoints. Please follow the steps in [Access an AWS service using an interface VPC endpoint](#) to create the interface endpoint. While creating the endpoint, select the following settings in the console interface:

- Select the **Enable DNS name** checkbox under **Additional Settings**
- Select the appropriate security groups and the subnets to be used with the SageMaker endpoints.

Also make sure that the VPC has DNS hostnames turned on. For more information on how to change DNS attributes for your VPC, see [View and update DNS attributes for your VPC](#).

**Deploy SageMaker endpoint in a VPC**

To achieve low overhead latency, create a SageMaker endpoint using the same subnets that you specified when deploying AWS PrivateLink. These subnets should match the AZs of your client application, as shown in the following code snippet.

```python
model_name = '<the-name-of-your-model>'

vpc = 'vpc-0123456789abcdef0'
subnet_a = 'subnet-0123456789abcdef0'
subnet_b = 'subnet-0123456789abcdef1'
security_group = 'sg-0123456789abcdef0'

create_model_response = sagemaker_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = sagemaker_role,
    PrimaryContainer = {
```
The aforementioned code snippet assumes that you have followed the steps in Before you begin.

**Invoke the SageMaker endpoint**

Finally, specify the SageMaker Runtime client and invoke the SageMaker endpoint as shown in the following code snippet.

```python
endpoint_name = '<endpoint-name>'

runtime_client = boto3.client('sagemaker-runtime')
response = runtime_client.invoke_endpoint(EndpointName=endpoint_name,
                                       ContentType='text/csv',
                                       Body=payload)
```

For more information on endpoint configuration, see [Deploy models for real-time inference](#).

**Migrate inference workload from x86 to AWS Graviton**

**AWS Graviton** is a series of ARM-based processors designed by AWS. They are more energy efficient than x86-based processors and offer a compelling price-performance ratio. Amazon SageMaker offers Graviton-based instances so that you can take advantage of these advanced processors for your inference needs.

You can migrate your existing inference workloads from x86-based instances to Graviton-based instances, by using either ARM compatible container images or multi-architecture container images. This guide assumes that you are either using [AWS Deep Learning container images](#), or your own ARM compatible container images. For more information on building your own images, check [Building your image](#).

At a high level, migrating inference workload from x86-based instances to Graviton-based instances is a four-step process:

- Migrate inference workload from x86 to AWS Graviton
- 4681
1. Push container images to Amazon Elastic Container Registry (Amazon ECR), an AWS managed container registry.

2. Create a SageMaker Model.

3. Create an endpoint configuration.

4. Create an endpoint.

The following sections of this guide provide more details regarding the above steps. Replace the *user placeholder text* in the code examples with your own information.

**Topics**

- [Push container images to Amazon ECR](#)
- [Create a SageMaker Model](#)
- [Create an endpoint configuration](#)
- [Create an endpoint](#)

**Push container images to Amazon ECR**

You can push your container images to Amazon ECR with the AWS CLI. When using an ARM compatible image, verify that it supports ARM architecture:

```
docker inspect deep-learning-container-uri
```

The response "Architecture": "arm64" indicates that the image supports ARM architecture. You can push it to Amazon ECR with the `docker push` command. For more information, check [Pushing a Docker image](#).

Multi-architecture container images are fundamentally a set of container images supporting different architectures or operating systems, that you can refer to by a common manifest name. If you are using multi-architecture container images, then in addition to pushing the images to Amazon ECR, you will also have to push a manifest list to Amazon ECR. A manifest list allows for the nested inclusion of other image manifests, where each included image is specified by architecture, operating system and other platform attributes. The following example creates a manifest list, and pushes it to Amazon ECR.

1. Create a manifest list.
2. Annotate the manifest list, so that it correctly identifies which image is for which architecture.

```bash
docker manifest annotate --arch arm64 aws-account-id.dkr.ecr.aws-region.amazonaws.com/my-repository \
aws-account-id.dkr.ecr.aws-account-id.amazonaws.com/my-repository:arm64
```

3. Push the manifest.

```bash
docker manifest push aws-account-id.dkr.ecr.aws-region.amazonaws.com/my-repository
```

For more information on creating and pushing manifest lists to Amazon ECR, check [Introducing multi-architecture container images for Amazon ECR](https://docs.aws.amazon.com/goto/WebAPI/sagemaker/latest/userguide/Introducing-multiple-image-container-supported.html) and [Pushing a multi-architecture image](https://docs.aws.amazon.com/goto/WebAPI/sagemaker/latest/userguide/Pushing-multiple-image-container-supported.html).

**Create a SageMaker Model**

Create a SageMaker Model by calling the [CreateModel](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateModel.html) API.

```python
import boto3
from sagemaker import get_execution_role

aws_region = "aws-region"
sagemaker_client = boto3.client("sagemaker", region_name=aws_region)

role = get_execution_role()

sagemaker_client.create_model(
    ModelName = "model-name",
    PrimaryContainer = {
        "Image": "deep-learning-container-uri",
        "ModelDataUrl": "model-s3-location",
        "Environment": {
            "SAGEMAKER_PROGRAM": "inference.py",
        }
    }
)"
"SAGEMAKER_SUBMIT_DIRECTORY": "inference-script-s3-location",
"SAGEMAKER_CONTAINER_LOG_LEVEL": "20",
"SAGEMAKER_REGION": aws_region,
}

Create an endpoint configuration

Create an endpoint configuration by calling the CreateEndpointConfig API. For a list of Graviton-based instances, check Compute optimized instances.

```python
sagemaker_client.create_endpoint_config(
    EndpointConfigName = "endpoint-config-name",
    ProductionVariants = [
        {
            "VariantName": "variant-name",
            "ModelName": "model-name",
            "InitialInstanceCount": 1,
            "InstanceType": "ml.c7g.xlarge", # Graviton-based instance
        }
    ]
)
```

Create an endpoint

Create an endpoint by calling the CreateEndpoint API.

```python
sagemaker_client.create_endpoint(
    EndpointName = "endpoint-name",
    EndpointConfigName = "endpoint-config-name"
)
```

Troubleshoot Amazon SageMaker model deployments

If you encounter an issue when deploying machine learning models in Amazon SageMaker, see the following guidance.
Detection Errors in the Active CPU Count

If you deploy a SageMaker model with a Linux Java Virtual Machine (JVM), you might encounter detection errors that prevent using available CPU resources. This issue affects some JVMs that support Java 8 and Java 9, and most that support Java 10 and Java 11. These JVMs implement a mechanism that detects and handles the CPU count and the maximum memory available when running a model in a Docker container, and, more generally, within Linux taskset commands or control groups (cgroups). SageMaker deployments take advantage of some of the settings that the JVM uses for managing these resources. Currently, this causes the container to incorrectly detect the number of available CPUs.

SageMaker doesn't limit access to CPUs on an instance. However, the JVM might detect the CPU count as 1 when more CPUs are available for the container. As a result, the JVM adjusts all of its internal settings to run as if only 1 CPU core is available. These settings affect garbage collection, locks, compiler threads, and other JVM internals that negatively affect the concurrency, throughput, and latency of the container.

For an example of the misdetection, in a container configured for SageMaker that is deployed with a JVM that is based on Java8_191 and that has four available CPUs on the instance, run the following command to start your JVM:

```
java -XX:+UnlockDiagnosticVMOptions -XX:+PrintActiveCpus -version
```

This generates the following output:

```
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: determined by OSContainer: 1
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: determined by OSContainer: 1
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: determined by OSContainer: 1
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: determined by OSContainer: 1
active_processor_count: determined by OSContainer: 1
openjdk version "1.8.0_191"
```
Many of the JVMs affected by this issue have an option to disable this behavior and reestablish full access to all of the CPUs on the instance. Disable the unwanted behavior and establish full access to all instance CPUs by including the `-XX:-UseContainerSupport` parameter when starting Java applications. For example, run the `java` command to start your JVM as follows:

```
java -XX:-UseContainerSupport -XX:+UnlockDiagnosticVMOptions -XX:+PrintActiveCpus -version
```

This generates the following output:

```
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: sched_getaffinity processor count: 4
active_processor_count: sched_getaffinity processor count: 4
openjdk version "1.8.0_191"
OpenJDK Runtime Environment (build 1.8.0_191-8u191-b12-2ubuntu0.16.04.1-b12)
OpenJDK 64-Bit Server VM (build 25.191-b12, mixed mode)
```

Check whether the JVM used in your container supports the `-XX:-UseContainerSupport` parameter. If it does, always pass the parameter when you start your JVM. This provides access to all of the CPUs in your instances.

You might also encounter this issue when indirectly using a JVM in SageMaker containers. For example, when using a JVM to support SparkML Scala. The `-XX:-UseContainerSupport` parameter also affects the output returned by the Java `Runtime.getRuntime().availableProcessors()` API.

### Issues with deploying a model.tar.gz file

When you deploy a model using a `model.tar.gz` file, the model tarball should not include any symlinks. Symlinks cause the model creation to fail. Also, we recommend that you do not include any unnecessary files in the tarball.

### Primary container did not pass ping health checks

If your primary container fails ping health checks with the following error message, it indicates that there is an issue with your container or script:
The primary container for production variant beta did not pass the ping health check. Please check CloudWatch Logs logs for this endpoint.

To troubleshoot this issue, you should check the CloudWatch Logs logs for the endpoint in question to see if there are any errors or issues that are preventing the container from responding to /ping or /invocations. The logs may provide an error message that could point to the issue. Once you have identified the error and failure reason you should resolve the error.

It is also good practice to test the model deployment locally before creating an endpoint.

- Use local mode in the SageMaker SDK to imitate the hosted environment by deploying the model to a local endpoint. For more information, see Local Mode.
- Use vanilla docker commands to test the container responds to /ping and /invocations. For more information, see local_test.

**Inference cost optimization best practices**

The following content provides techniques and considerations for optimizing the cost of endpoints. You can use these recommendations to optimize the cost for both new and existing endpoints.

**Best practices**

To optimize your SageMaker Inference costs, follow these best practices.

**Pick the best inference option for the job.**

SageMaker offers 4 different inference options to provide the best inference option for the job. You may be able to save on costs by picking the inference option that best matches your workload.

- Use **real-time inference** for low latency workloads with predictable traffic patterns that need to have consistent latency characteristics and are always available. You pay for using the instance.
- Use **serverless inference** for synchronous workloads that have a spiky traffic pattern and can accept variations in the p99 latency. Serverless inference automatically scales to meet your workload traffic so you don’t pay for any idle resources. You only pay for the duration of the inference request. The same model and containers can be used with both real-time and serverless inference so you can switch between these two modes if your needs change.
- Use **asynchronous inference** for asynchronous workloads that process up to 1 GB of data (such as text corpus, image, video, and audio) that are latency insensitive and cost sensitive. With
asynchronous inference, you can control costs by specifying a fixed number of instances for the optimal processing rate instead of provisioning for the peak. You can also scale down to zero to save additional costs.

- Use **batch inference** for workloads for which you need inference for a large set of data for processes that happen offline (that is, you don't need a persistent endpoint). You pay for the instance for the duration of the batch inference job.

**Opt in to a SageMaker Savings Plan.**

- If you have a consistent usage level across all SageMaker services, you can opt in to a SageMaker Savings Plan to help reduce your costs by up to 64%.

- [Amazon SageMaker Savings Plans](https://aws.amazon.com/sagemaker/savings-plans/) provide a flexible pricing model for Amazon SageMaker, in exchange for a commitment to a consistent amount of usage (measured in $/hour) for a one-year or three-year term. These plans automatically apply to eligible SageMaker ML instance usages including SageMaker Studio Classic Notebook, SageMaker On-Demand Notebook, SageMaker Processing, SageMaker Data Wrangler, SageMaker Training, SageMaker Real-Time Inference, and SageMaker Batch Transform regardless of instance family, size, or Region. For example, you can change usage from a CPU ml.c5.xlarge instance running in US East (Ohio) to a ml.Inf1 instance in US West (Oregon) for inference workloads at any time and automatically continue to pay the Savings Plans price.

**Optimize your model to run better.**

- Unoptimized models can lead to longer run times and use more resources. You may choose to use more or bigger instances to improve performance; however, this leads to higher costs.

- By optimizing your models to be more performant, you may be able to lower costs by using fewer or smaller instances while keeping the same or better performance characteristics. You can use [SageMaker Neo](https://aws.amazon.com/sagemaker/neo/) with SageMaker Inference to automatically optimize models. For more details and samples, see Optimize model performance using Neo.

**Use the most optimal instance type and size for real-time inference.**

- SageMaker Inference has over 70 instance types and sizes that can be used to deploy ML models including AWS Inferentia and Graviton chipsets that are optimized for ML. Choosing the right instance for your model helps ensure you have the most performant instance at the lowest cost for your models.
• By using **Inference Recommender**, you can quickly compare different instances to understand the performance of the model and the costs. With these results, you can choose the instance to deploy with the best return on investment.

**Improve efficiency and costs by combining multiple endpoints into a single endpoint for real-time inference.**

• Costs can quickly add up when you deploy multiple endpoints, especially if the endpoints don’t fully utilize the underlying instances. To understand if the instance is under-utilized, check the utilization metrics (CPU, GPU, etc) in Amazon CloudWatch for your instances. If you have more than one of these endpoints, you can combine the models or containers on these multiple endpoints into a single endpoint.

• Using **Multi-model endpoints** (MME) or **Multi-container endpoints** (MCE), you can deploy multiple ML models or containers in a single endpoint to share the instance across multiple models or containers and improve your return on investment. To learn more, see this [Save on inference costs by using Amazon SageMaker multi-model endpoints](https://aws.amazon.com/blogs/machine-learning/save-on-inference-costs-by-using-amazon-sagemaker-multi-model-endpoints/) or [Deploy multiple serving containers on a single instance using Amazon SageMaker multi-container endpoints](https://aws.amazon.com/blogs/machine-learning/deploy-multiple-serving-containers-on-a-single-instance-using-amazon-sagemaker-multi-container-endpoints/) on the AWS Machine Learning blog.

**Set up autoscaling to match your workload requirements for real-time and asynchronous inference.**

• Without autoscaling, you need to provision for peak traffic or risk model unavailability. Unless the traffic to your model is steady throughout the day, there will be excess unused capacity. This leads to low utilization and wasted resources.

• **Autoscaling** is an out-of-the-box feature that monitors your workloads and dynamically adjusts the capacity to maintain steady and predictable performance at the possible lowest cost. When the workload increases, autoscaling brings more instances online. When the workload decreases, autoscaling removes unnecessary instances, helping you reduce your compute cost. To learn more, see [Configuring autoscaling inference endpoints in Amazon SageMaker](https://aws.amazon.com/blogs/machine-learning/configuring-autoscaling-inference-endpoints-in-amazon-sagemaker/) on the AWS Machine Learning blog.

**Best practices to minimize interruptions during GPU driver upgrades**

SageMaker Model Deployment upgrades GPU drivers on the ML instances for Real-time, Batch, and Asynchronous Inference options over time to provide customers access to improvements from the
driver providers. Below you can see the GPU version supported for each Inference option. Different driver versions can change how your model interacts with the GPUs. Below are some strategies to help you understand how your application works with different driver versions.

**Current versions and supported instance families**

Amazon SageMaker Inference supports the following drivers and instance families:

<table>
<thead>
<tr>
<th>Service</th>
<th>GPU</th>
<th>Driver version</th>
<th>Instance types</th>
</tr>
</thead>
<tbody>
<tr>
<td>Real-time</td>
<td>NVIDIA</td>
<td>470.57.02</td>
<td>ml.p2.<em>, ml.p3.</em>, ml.p4d.<em>, ml.p4de.</em>, ml.g4dn.<em>, ml.g5.</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>535.54.03</td>
<td>ml.p5.*</td>
</tr>
<tr>
<td>Batch</td>
<td>NVIDIA</td>
<td>470.57.02</td>
<td>ml.p2.<em>, ml.p3.</em>, ml.p4d.<em>, ml.p4de.</em>, ml.g4dn.<em>, ml.g5.</em></td>
</tr>
<tr>
<td>Asynchronous Inference</td>
<td>NVIDIA</td>
<td>470.57.02</td>
<td>ml.p2.<em>, ml.p3.</em>, ml.p4d.<em>, ml.p4de.</em>, ml.g4dn.<em>, ml.g5.</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>535.54.03</td>
<td>ml.p5.*</td>
</tr>
</tbody>
</table>

**Troubleshoot your model container with GPU capabilities**

If you encounter an issue when running your GPU workload, see the following guidance:

**GPU card detection failure or NVIDIA initialization error**

Run the `nvidia-smi` (NVIDIA System Management Interface) command from within the Docker container. If the NVIDIA System Management Interface detects a GPU detection error or NVIDIA initialization error, it will return the following error message:

Failed to initialize NVML: Driver/library version mismatch

Based on your use case, follow these best practices to resolve the failure or error:
• Follow the best practice recommendation described in the If you bring your own (BYO) model containers dropdown.

• Follow the best practice recommendation described in the If you use a CUDA compatibility layer dropdown.

Refer to the NVIDIA System Management Interface page on the NVIDIA website for more information.

CannotStartContainerError

If your GPU instance uses NVIDIA driver versions that are not compatible with the CUDA version in the Docker container, then deploying an endpoint will fail with the following error message:

Failure reason CannotStartContainerError. Please ensure the model container for variant <variant_name> starts correctly when invoked with 'docker run <image> serve'

Based on your use case, follow these best practices to resolve the failure or error:

• Follow the best practice recommendation described in the The driver my container depends on is greater than the version on the ML GPU instances dropdown.

• Follow the best practice recommendation described in the If you use a CUDA compatibility layer dropdown.

Best practices for working with mismatched driver versions

The following provides information on how to update your GPU driver:

The driver my container depends on is lower than the version on the ML GPU instance

No action is required. NVIDIA provides backwards compatibility.

The driver my container depends on is greater than the version on the ML GPU instances

If it is a minor version difference, no action is required. NVIDIA provides minor version forward compatibility.

If it is a major version difference, the CUDA Compatibility Package will need to be installed. Please refer to CUDA Compatibility Package in the NVIDIA documentation.
Important

The CUDA Compatibility Package is not backwards compatible so it needs to be disabled if the driver version on the instance is greater than the CUDA Compatibility Package version.

If you bring your own (BYO) model containers

Ensure no NVIDIA driver packages are bundled in the image which could cause conflict with on host NVIDIA driver version.

If you use a CUDA compatibility layer

To verify if the platform Nvidia driver version supports the CUDA Compatibility Package version installed in the model container, see the [CUDA documentation](#). If the platform Nvidia driver version does not support the CUDA Compatibility Package version, you can disable or remove the CUDA Compatibility Package from the model container image. If the CUDA compatibility libs version is supported by the latest Nvidia driver version, we suggest that you enable the CUDA Compatibility Package based on the detected Nvidia driver version for future compatibility by adding the code snippet below into the container start up shell script (at the ENTRYPOINT script).

The script demonstrates how to dynamically switch the use of the CUDA Compatibility Package based on the detected Nvidia driver version on the deployed host for your model container. When SageMaker releases a newer Nvidia driver version, the installed CUDA Compatibility Package can be turned off automatically if the CUDA application is supported natively on the new driver.

```bash
#!/bin/bash

verlte() {
    [ "$1" = "$2" ] && return 1 || [ "$2" = "\`echo -e "$1\n$2" | sort -V | head -n1\`" ]
}

if [ -f /usr/local/cuda/compat/libcuda.so.1 ]; then
    cat /usr/local/cuda/version.txt
    CUDA_COMPAT_MAX_DRIVER_VERSION=$(readlink /usr/local/cuda/compat/libcuda.so.1 |cut -d'.' -f 3-)
    echo "CUDA compat package requires Nvidia driver 
    ${CUDA_COMPAT_MAX_DRIVER_VERSION}"
    NVIDIA_DRIVER_VERSION=$(sed -n 's/^NVRM.*Kernel Module */\([0-9.]*\).*/\1/p' /proc/driver/nvidia/version 2>/dev/null || true)
```
echo "Current installed Nvidia driver version is ${NVIDIA_DRIVER_VERSION}" 
if [ $(verlte $CUDA_COMPAT_MAX_DRIVER_VERSION $NVIDIA_DRIVER_VERSION) ]; then 
  echo "Setup CUDA compatibility libs path to LD_LIBRARY_PATH" 
  export LD_LIBRARY_PATH=/usr/local/cuda/compat:$LD_LIBRARY_PATH 
  echo $LD_LIBRARY_PATH 
else 
  echo "Skip CUDA compat libs setup as newer Nvidia driver is installed" 
  fi 
else 
  echo "Skip CUDA compat libs setup as package not found" 
  fi

Best practices for endpoint security and health with Amazon SageMaker

To address the latest security issues, Amazon SageMaker automatically patches endpoints to the latest and most secure software. However, if you incorrectly modify your endpoint dependencies, Amazon SageMaker can't automatically patch your endpoints or replace your unhealthy instances. To ensure your endpoints remain eligible for automatic updates, apply the following best practices.

Don't delete resources while your endpoints use them

Avoid deleting any of the following resources if you have existing endpoints that use them:

- The model definition that you create with the `CreateModel` action in the Amazon SageMaker API.
- Any model artifacts that you specify for the `ModelDataUrl` parameter.
- The IAM role and permissions that you specify for the `ExecutionRoleArn` parameter.

Reminder

In the model definition that your endpoint uses, ensure that the IAM role that you specified has the correct permissions. For more information about the required permissions for Amazon SageMaker endpoints, see [CreateModel API: Execution Role Permissions](#).

- The inference images that you specify for the `Image` parameter, if you use your own inference code.
Reminder

If you use the private registry feature, ensure that Amazon SageMaker can access the private registry as long as you're using the endpoint.

- The Amazon VPC subnets and security groups that you specify for the `VpcConfig` parameter.
- The endpoint configuration that you create with the `CreateEndpointConfig` action in the Amazon SageMaker API.
- Any KMS keys or Amazon S3 buckets that you specify in the endpoint configuration.

Reminder

Ensure you don’t disable these KMS keys.

Follow these procedures to update your endpoints

When you update your Amazon SageMaker endpoints, use any of the following procedures that apply to your needs.

To update your model definition settings

1. Create a new model definition with your updated settings by using the `CreateModel` action in the Amazon SageMaker API.
2. Create a new endpoint configuration that uses the new model definition. To do this, use the `CreateEndpointConfig` action in the Amazon SageMaker API.
3. Update your endpoint with the new endpoint configuration so that your updated model definition settings take effect.
4. (Optional) Delete the old endpoint configuration if you're not using it with any other endpoints. You can also delete the resources that you specified in the model definition if you're not using them with any other endpoints. These resources include model artifacts in Amazon S3 and inference images.

To update your endpoint configuration

1. Create a new endpoint configuration with your updated settings.
2. Update your endpoint with the new configuration so that your updates take effect.

3. (Optional) Delete the old endpoint configuration if you're not using it with any other endpoints. You can also delete the resources that you specified in the model definition if you're not using them with any other endpoints. These resources include model artifacts in Amazon S3 and inference images.

Whenever you create a new model definition or endpoint configuration, we recommend that you use a unique name. If you want to update these resources and retain their original names, use the following procedures.

**To update your model settings and retain the original model name**

1. Delete the existing model definition. At this point, any endpoint that uses the model is broken, but you fix this in the following steps.

2. Create the model definition again with your updated settings, and use the same model name.

3. Create a new endpoint configuration that uses the updated model definition.

4. Update your endpoint with the new endpoint configuration so that your updates take effect.

**To update your endpoint configuration and retain the original configuration name**

1. Delete the existing endpoint configuration.

2. Create a new endpoint configuration with your updated settings, and use the original name.

3. Update your endpoint with the new configuration so that your updates take effect.

**Supported features**

Amazon SageMaker offers the following four options to deploy models for inference.

- Real-time inference for inference workloads with real-time, interactive, low latency requirements.
- Batch transform for offline inference with large datasets.
- Asynchronous inference for near-real-time inference with large inputs that require longer preprocessing times.
- Serverless inference for inference workloads that have idle periods between traffic spurts.
The following table summarizes the core platform features that are supported by each inference option. It does not show features that can be provided by frameworks, custom Docker containers, or through chaining different AWS services.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Real-time inference</th>
<th>Batch transform</th>
<th>Asynchronous inference</th>
<th>Serverless inference</th>
<th>Docker containers</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Autoscaling support</strong></td>
<td>✓</td>
<td>N/A</td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>GPU support</strong></td>
<td>✓¹</td>
<td>✓¹</td>
<td>✓¹</td>
<td></td>
<td>1P, pre-built, BYOC</td>
</tr>
<tr>
<td><strong>Single model</strong></td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Multi-model endpoint</strong></td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td>k-NN, XGBoost, Linear Learner, RCF, TensorFlow, Apache MXNet, PyTorch, scikit-learn ²</td>
</tr>
<tr>
<td><strong>Multi-container endpoint</strong></td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
</tr>
<tr>
<td><strong>Serial inference pipeline</strong></td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
</tr>
<tr>
<td><strong>Inference Recommender</strong></td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
</tr>
</tbody>
</table>

1P: pre-built, BYOC: Bring Your Own Container
<table>
<thead>
<tr>
<th>Feature</th>
<th>Real-time inference</th>
<th>Batch transform</th>
<th>Asynchronous inference</th>
<th>Serverless inference</th>
<th>Docker containers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Private link support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Data capture/Model monitor support</strong></td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td><strong>DLCs supported</strong></td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
<td>N/A</td>
</tr>
<tr>
<td>Protocols supported</td>
<td>HTTP(S)</td>
<td>HTTP(S)</td>
<td>HTTP(S)</td>
<td>HTTP(S)</td>
<td>N/A</td>
</tr>
<tr>
<td>Payload size</td>
<td>&lt; 6 MB</td>
<td>≤ 100 MB</td>
<td>≤ 1 GB</td>
<td>≤ 4 MB</td>
<td></td>
</tr>
<tr>
<td>HTTP chunked encoding</td>
<td>Framework dependent, 1P not supported</td>
<td>N/A</td>
<td>Framework dependent, 1P not supported</td>
<td>Framework dependent, 1P not supported</td>
<td>N/A</td>
</tr>
<tr>
<td>Request timeout</td>
<td>&lt; 60 seconds</td>
<td>Days</td>
<td>&lt; 1 hour</td>
<td>&lt; 60 seconds</td>
<td>N/A</td>
</tr>
<tr>
<td>Deployment guardrails: blue/green deployments</td>
<td>✓</td>
<td>N/A</td>
<td>✓</td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Deployment guardrails: rolling deployments</td>
<td>✓</td>
<td>N/A</td>
<td>✓</td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Feature</td>
<td>Real-time inference</td>
<td>Batch transform</td>
<td>Asynchronous inference</td>
<td>Serverless inference</td>
<td>Docker containers</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>---------------------</td>
<td>-----------------</td>
<td>------------------------</td>
<td>----------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>Shadow testing</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Scale to zero</td>
<td></td>
<td>N/A</td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
</tr>
<tr>
<td>Scale to zero</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Market place model packages support</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Virtual private cloud support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Multiple production variants support</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Network isolation</td>
<td>✓</td>
<td></td>
<td>✓</td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Model parallel serving support</td>
<td>✓³</td>
<td>✓</td>
<td>✓³</td>
<td>✓³</td>
<td>✓³</td>
</tr>
<tr>
<td>Volume encryption</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
</tr>
<tr>
<td>Customer AWS KMS</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
</tr>
<tr>
<td>d instance support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td>N/A</td>
</tr>
</tbody>
</table>
With SageMaker, you can deploy a single model, or multiple models behind a single inference endpoint for real-time inference. The following table summarizes the core features supported by various hosting options that come with real-time inference.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Single model endpoints</th>
<th>Multi-model endpoints</th>
<th>Serial inference pipeline</th>
<th>Multi-container endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autoscaling support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>GPU support</td>
<td>✓¹</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Single model</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Multi-model endpoints</td>
<td></td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
</tr>
<tr>
<td>Multi-container endpoints</td>
<td>✓</td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Serial inference pipeline</td>
<td>✓</td>
<td>✓</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Inference Recommender</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Private link support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Data capture/Model monitor support</td>
<td>✓</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Supported features
<table>
<thead>
<tr>
<th>Feature</th>
<th>Single model endpoints</th>
<th>Multi-model endpoints</th>
<th>Serial inference pipeline</th>
<th>Multi-container endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td>DLCs supported</td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
<td>k-NN, XGBoost, Linear Learner, RCF, TensorFlow, Apache MXNet, PyTorch, scikit-learn</td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
<td>1P, pre-built, Extend pre-built, BYOC</td>
</tr>
<tr>
<td>Protocols supported</td>
<td>HTTP(S)</td>
<td>HTTP(S)</td>
<td>HTTP(S)</td>
<td>HTTP(S)</td>
</tr>
<tr>
<td>Payload size</td>
<td>&lt; 6 MB</td>
<td>&lt; 6 MB</td>
<td>&lt; 6 MB</td>
<td>&lt; 6 MB</td>
</tr>
<tr>
<td>Request timeout</td>
<td>&lt; 60 seconds</td>
<td>&lt; 60 seconds</td>
<td>&lt; 60 seconds</td>
<td>&lt; 60 seconds</td>
</tr>
<tr>
<td><strong>Deployment guardrails:</strong></td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>blue/green deployments</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Deployment guardrails:</strong></td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>rolling deployments</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shadow testing</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Market place model packages</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>support</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Virtual private cloud support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Multiple production variants</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>support</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Supported features
<table>
<thead>
<tr>
<th>Feature</th>
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<th>Multi-model endpoints</th>
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<th>Multi-container endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Network isolation</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Model parallel serving support</td>
<td>✓ ³</td>
<td></td>
<td>✓ ³</td>
<td></td>
</tr>
<tr>
<td>Volume encryption</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Customer AWS KMS</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>d instance support</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>inf1 support</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1 Availability of the Amazon EC2 instance types depends on the AWS Region. For availability of instances specific to AWS, see Amazon SageMaker Pricing.

2 To use any other framework or algorithm, use the SageMaker Inference toolkit to build a container that supports multi-model endpoints.

3 With SageMaker, you can deploy large models (up to 500 GB) for inference. You can configure the container health check and download timeout quotas, up to 60 minutes. This will allow you to have more time to download and load your model and associated resources. For more information, see SageMaker endpoint parameters for large model inference. You can use SageMaker compatible large model Inference containers. You can also use third-party model parallelization libraries, such as Triton with FasterTransformer and DeepSpeed. You have to ensure that they are compatible with SageMaker.

## Resources

Use the following resources for troubleshooting and reference, answerings FAQs, and learning more about Amazon SageMaker.
Blogs, example notebooks, and additional resources

The following sections contain examples and additional resources for you to learn more about Amazon SageMaker.

Blogs and case studies

See the following table for lists of blogs and case studies for various features within SageMaker Inference. You can use the blogs to help put together solutions that work for your use case.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Resources</th>
</tr>
</thead>
<tbody>
<tr>
<td>Real-Time Inference</td>
<td>• Getting started with deploying real-time models on Amazon SageMaker</td>
</tr>
<tr>
<td></td>
<td>• Deploy BLOOM-176B and OPT-30B on Amazon SageMaker with large model</td>
</tr>
<tr>
<td></td>
<td>inference Deep Learning Containers and DeepSpeed</td>
</tr>
<tr>
<td></td>
<td>• Creating a machine learning-powered REST API with Amazon API Gateway</td>
</tr>
<tr>
<td></td>
<td>mapping templates and Amazon SageMaker</td>
</tr>
<tr>
<td>Autoscaling</td>
<td>• Configuring autoscaling inference endpoints in Amazon SageMaker</td>
</tr>
<tr>
<td>Serverless Inference</td>
<td>• Amazon SageMaker Serverless Inference</td>
</tr>
<tr>
<td></td>
<td>– Machine Learning Inference without Worrying about Servers</td>
</tr>
<tr>
<td></td>
<td>• Host Hugging Face transformer models using Amazon SageMaker Serverless</td>
</tr>
<tr>
<td></td>
<td>Inference</td>
</tr>
<tr>
<td>Feature</td>
<td>Resources</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>---------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| Asynchronous Inference        | • Run computer vision inference on large videos with Amazon SageMaker asynchronous endpoints  
                                | • Build a predictive maintenance solution with Amazon Kinesis, AWS Glue, and Amazon SageMaker  
                                | • Improve high-value research with Hugging Face and Amazon SageMaker Asynchronous Inference endpoints |
| Batch Transform               | • Associating prediction results with input data using Amazon SageMaker Batch Transform |
| Multi-Model Endpoints         | • Save on inference costs by using Amazon SageMaker multi-model endpoints  
                                | • Run multiple deep learning models on GPU with Amazon SageMaker multi-model endpoints  
                                | • How to scale machine learning inference for multi-tenant SaaS use cases  
<pre><code>                            | • Run and optimize multi-model inference with Amazon SageMaker multi-model endpoints |
</code></pre>
<p>| Serial Inference Pipelines    | • Design patterns for serial inference on Amazon SageMaker             |
| Multi-Container Endpoints     | • Cost efficient ML inference with multi-framework models on Amazon SageMaker |</p>
<table>
<thead>
<tr>
<th>Feature</th>
<th>Resources</th>
</tr>
</thead>
<tbody>
<tr>
<td>Running Model Ensembles</td>
<td>• Run ensemble ML models on Amazon SageMaker</td>
</tr>
<tr>
<td>Inference Recommender</td>
<td>• SageMaker Inference Recommender example notebook&lt;br&gt;• SageMaker Inference Recommender for HuggingFace BERT Sentiment Analysis example notebook&lt;br&gt;• Achieve hyperscale performance for model serving using NVIDIA Triton Inference Server on Amazon SageMaker</td>
</tr>
<tr>
<td>Advanced model hosting blog series</td>
<td>• Part 1: Common design patterns for building ML application on Amazon SageMaker&lt;br&gt;• Part 2: Getting started with deploying real time models on SageMaker&lt;br&gt;• Part 3: Run and optimize multi-model inference with Amazon SageMaker multi-model endpoints&lt;br&gt;• Part 4: Design patterns for serial inference on Amazon SageMaker&lt;br&gt;• Part 5: Cost efficient ML inference with multi-framework models on Amazon SageMaker&lt;br&gt;• Part 6: Best practices in testing and updating models on SageMaker&lt;br&gt;• Part 7: Run ensemble ML models on Amazon SageMaker</td>
</tr>
</tbody>
</table>
Example notebooks

See the following table for example notebooks that can help you learn more about SageMaker Inference.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Example notebooks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inference Recommender</td>
<td>• <a href="https://example.com">SageMaker Inference Recommender example notebook</a></td>
</tr>
<tr>
<td></td>
<td>• <a href="https://example.com">SageMaker Inference Recommender for HuggingFace BERT Sentiment Analysis example notebook</a></td>
</tr>
<tr>
<td>Optimize large language models (LLMs) for SageMaker</td>
<td><a href="https://example.com">Generative AI LLMs workshop</a></td>
</tr>
</tbody>
</table>

Additional resources

For more information about each SageMaker Inference option in detail, you can watch the following video.

Deploy ML models for inference at high performance and low cost

Troubleshooting and reference

You can use the following resources and reference documentation to understand best practices when using SageMaker Inference and to troubleshooting issues with model deployments:

- For troubleshooting model deployments, see [Troubleshoot Amazon SageMaker model deployments](https://example.com).
- For model deployment best practices, see [Best practices](https://example.com).
- For reference information about the size of storage volumes provided for different sizes of hosting instances, see [Host instance storage volumes](https://example.com).
- For reference information about SageMaker limits and quotas, see [Amazon SageMaker endpoints and quotas](https://example.com).
- For frequently asked questions about SageMaker, see [Model Hosting FAQs](https://example.com).
Model Hosting FAQs

Refer to the following FAQ items for answers to commonly asked questions about SageMaker Inference Hosting.

General Hosting

The following FAQ items answer common general questions for SageMaker Inference.

Q: What deployment options does Amazon SageMaker provide?

A: After you build and train models, Amazon SageMaker provides four options to deploy them so you can start making predictions. Real-Time Inference is suitable for workloads with millisecond latency requirements, payload sizes up to 6 MB, and processing times of up to 60 seconds. Batch Transform is ideal for offline predictions on large batches of data that are available up front. Asynchronous Inference is designed for workloads that do not have sub-second latency requirements, payload sizes up to 1 GB, and processing times of up to 15 minutes. With Serverless Inference, you can quickly deploy machine learning models for inference without having to configure or manage the underlying infrastructure, and you pay only for the compute capacity used to process inference requests, which is ideal for intermittent workloads.

Q: How do I choose a model deployment option in SageMaker?

A: The following diagram can help you choose a SageMaker Hosting model deployment option.
The preceding diagram walks you through the following decision process. If you want to process requests in batches, you might want to choose Batch Transform. Otherwise, if you want to receive inference for each request to your model, you might want to choose Asynchronous Inference, Serverless Inference, or Real-Time Inference. You can choose Asynchronous Inference if you have long processing times or large payloads and want to queue requests. You can choose Serverless Inference if your workload has unpredictable or intermittent traffic. You can choose Real-Time Inference if you have sustained traffic and need lower and consistent latency for your requests.

Q: I've heard SageMaker Inference is expensive. What's the best way to optimize my cost when hosting models?

A: To optimize your costs with SageMaker Inference, you should choose the right hosting option for your use case. You can also use Inference features such as Amazon SageMaker Savings Plans, model optimization with SageMaker Neo, Multi-Model Endpoints and Multi-Container Endpoints, or autoscaling. For tips on how to optimize your Inference costs, see Inference cost optimization best practices.
Q: Why should I use Amazon SageMaker Inference Recommender?

A: You should use Amazon SageMaker Inference Recommender if you need recommendations for the right endpoint configuration to improve performance and reduce costs. Previously, data scientists who wanted to deploy their models had to run manual benchmarks to select the right endpoint configuration. First, they had to select the right machine learning instance type out of more than 70 available instance types based on the resource requirements of their models and sample payloads, and then optimize the model to account for differing hardware. Then, they had to conduct extensive load tests to validate that latency and throughput requirements were met and that the costs were low. Inference Recommender eliminates this complexity by helping you do the following:

- Get started in minutes with an instance recommendation.
- Conduct load tests across instance types to get recommendations on your endpoint configuration within hours.
- Automatically tune container and model server parameters as well as perform model optimizations for a given instance type.

Q: What is a model server?

A: SageMaker endpoints are HTTP REST endpoints that use a containerized web server, which includes a model server. These containers are responsible for loading up and serving requests for a machine learning model. They implement a web server that responds to /invocations and /ping on port 8080.

Common model servers include TensorFlow Serving, TorchServe and Multi Model Server. SageMaker framework containers have these model servers built in.

Q: What is Bring Your Own Container with Amazon SageMaker?

A: Everything in SageMaker Inference is containerized. SageMaker provides managed containers for popular frameworks such as TensorFlow, SKlearn, and HuggingFace. For a comprehensive updated list of those images, see Available Images.

Sometimes there are custom frameworks for which you might need to build a container. This approach is known as Bring Your Own Container or BYOC. With the BYOC approach, you provide the Docker image to set up your framework or library. Then, you push the image to Amazon Elastic Container Registry (Amazon ECR) so that you can use the image with SageMaker. For an example of a BYOC approach, see Overview of Containers for Amazon SageMaker.
Alternatively, instead of building an image from scratch, you can extend a container. You can take one of the base images that SageMaker provides and add your dependencies on top of it in your Dockerfile.

**Q: Do I need to train my models on SageMaker to host them on SageMaker endpoints?**

A: SageMaker offers the capacity to bring your own trained framework model that you've trained outside of SageMaker and deploy it on any of the SageMaker hosting options.

SageMaker requires you to package the model in a `model.tar.gz` file and have a specific directory structure. Each framework has its own model structure (see the following question for example structures). For more information, see the SageMaker Python SDK documentation for TensorFlow, PyTorch, and MXNet.

While you can choose from prebuilt framework images such as TensorFlow, PyTorch, and MXNet to host your trained model, you can also build your own container to host your trained models on SageMaker endpoints. For a walkthrough, see the example Jupyter notebook **Building your own algorithm container**.

**Q: How should I structure my model if I want to deploy on SageMaker but not train on SageMaker?**

A: SageMaker requires your model artifacts to be compressed in a `.tar.gz` file, or a *tarball*. SageMaker automatically extracts this `.tar.gz` file into the `/opt/ml/model/` directory in your container. The tarball shouldn't contain any symlinks or unnecessary files. If you are making use of one of the framework containers, such as TensorFlow, PyTorch, or MXNet, the container expects your TAR structure to be as follows:

**TensorFlow**

```
model.tar.gz/
|---[model_version_number]/
|   |---variables
|   |---saved_model.pb
code/
|---inference.py
|---requirements.txt
```

**PyTorch**

```
model.tar.gz/
```
Q: When invoking a SageMaker endpoint, I can provide a ContentType and Accept MIME Type. Which one is used to identify the data type being sent and received?

A: ContentType is the MIME type of the input data in the request body (the MIME type of the data you are sending to your endpoint). The model server uses the ContentType to determine if it can handle the type provided or not.

Accept is the MIME type of the inference response (the MIME type of the data your endpoint returns). The model server uses the Accept type to determine if it can handle returning the type provided or not.

Common MIME types include text/csv, application/json, and application/jsonlines.

Q: What are the supported data formats for SageMaker Inference?

A: SageMaker passes any request onto the model container without modification. The container must contain the logic to deserialize the request. For information about the formats defined for built-in algorithms, see Common Data Formats for Inference. If you are building your own container or using a SageMaker Framework container, you can include the logic to accept a request format of your choice.

Similarly, SageMaker also returns the response without modification, and then the client must deserialize the response. In case of the built-in algorithms, they return responses in specific formats. If you are building your own container or using a SageMaker Framework container, you can include the logic to return a response in the format you choose.
Q: How do I invoke my endpoint with binary data such as videos or images?

Use the **Invoke Endpoint** API call to make inference against your endpoint.

When passing your input as a payload to the InvokeEndpoint API, you must provide the correct type of input data that your model expects. When passing a payload in the InvokeEndpoint API call, the request bytes are forwarded directly to the model container. For example, for an image, you may use application/jpeg for the ContentType, and make sure that your model can perform inference on this type of data. This applies for JSON, CSV, video, or any other type of input with which you may be dealing.

Another factor to consider is payload size limits. In terms of real-time and serverless endpoints, the payload limit is 6 MB. You can split your video into multiple frames and invoke the endpoint with each frame individually. Alternatively, if your use case permits, you can send the whole video in the payload using an asynchronous endpoint, which supports up to 1 GB payloads.

For an example that showcases how to run computer vision inference on large videos with Asynchronous Inference, see this blog post.

**Real-Time Inference**

The following FAQ items answer common questions for SageMaker Real-Time Inference.

Q: How do I create a SageMaker endpoint?

A: You can create a SageMaker endpoint through AWS-supported tooling such as the AWS SDKs, the SageMaker Python SDK, the AWS Management Console, AWS CloudFormation, and the AWS Cloud Development Kit (AWS CDK).

There are three key entities in endpoint creation: a SageMaker model, a SageMaker endpoint configuration, and a SageMaker endpoint. The SageMaker model points towards the model data and image you are using. The endpoint configuration defines your production variants, which might include the instance type and instance count. You can then use either the `create_endpoint` API call or the `.deploy()` call for SageMaker to create an endpoint using the metadata from your model and endpoint configuration.

Q: Do I need to use the SageMaker Python SDK to create/invoke endpoints?

A: No, you can use the various AWS SDKs (see Invoke/Create for available SDKs) or even call the corresponding web APIs directly.
Q: What is the difference between Multi-Model Endpoints (MME) and Multi Model Server (MMS)?

A: A Multi-Model Endpoint is a Real-Time Inference option that SageMaker provides. With Multi-Model Endpoints, you can host thousands of models behind one endpoint. Multi Model Server is an open-source framework for serving machine learning models. It provides the HTTP front-end and model management capabilities required by multi-model endpoints to host multiple models within a single container, load models into and unload models out of the container dynamically, and perform inference on a specified loaded model.

Q: What are the different model deployment architectures supported by Real-Time Inference?

A: SageMaker Real-Time Inference supports various model deployment architecture such as Multi-Model Endpoints, Multi-Container Endpoints, and Serial Inference Pipelines.

Multi-Model Endpoints (MME) – MME allows customers to deploy 1000s of hyper-personalized models in a cost effective way. All the models are deployed on a shared-resource fleet. MME works best when the models are of similar size and latency and belong to the same ML framework. These endpoints are ideal for when you have don’t need to call the same model at all times. You can dynamically load respective models onto the SageMaker endpoint to serve your request.

Multi-Container Endpoints (MCE) – MCE allows customers to deploy 15 different containers with diverse ML frameworks and functionalities with no cold starts while only using one SageMaker endpoint. You can directly invoke these containers. MCE is best for when you want to keep all the models in memory.

Serial Inference Pipelines (SIP) – You can use SIP to chain together 2-15 containers on a single endpoint. SIP is mostly suitable for combining preprocessing and model inference in one endpoint and for low latency operations.

Serverless Inference

The following FAQ items answer common questions for Amazon SageMaker Serverless Inference.

Q: What is Amazon SageMaker Serverless Inference?

A: Serverless Inference is a purpose-built serverless model serving option that makes it easy to deploy and scale ML models. Serverless Inference endpoints automatically start compute resources and scale them in and out depending on traffic, eliminating the need for you to choose instance type, run provisioned capacity, or manage scaling. You can optionally specify the memory...
requirements for your serverless endpoint. You pay only for the duration of running the inference code and the amount of data processed, not for idle periods.

**Q: Why should I use Serverless Inference?**

A: Serverless Inference simplifies the developer experience by eliminating the need to provision capacity up front and manage scaling policies. Serverless Inference can scale instantly from tens to thousands of inferences within seconds based on the usage patterns, making it ideal for ML applications with intermittent or unpredictable traffic. For example, a chatbot service used by a payroll processing company experiences an increase in inquiries at the end of the month while traffic is intermittent for the rest of the month. Provisioning instances for the entire month in such scenarios is not cost-effective, as you end up paying for idle periods.

Serverless Inference helps address these types of use cases by providing you automatic and fast scaling out of the box without the need for you to forecast traffic up front or manage scaling policies. Additionally, you pay only for the compute time to run your inference code and for data processing, making it ideal for workloads with intermittent traffic.

**Q: How do I choose the right memory size for my serverless endpoint?**

A: Your serverless endpoint has a minimum RAM size of 1024 MB (1 GB), and the maximum RAM size you can choose is 6144 MB (6 GB). The memory sizes you can choose are 1024 MB, 2048 MB, 3072 MB, 4096 MB, 5120 MB, or 6144 MB. Serverless Inference auto-assigns compute resources proportional to the memory you select. If you choose a larger memory size, your container has access to more vCPUs.

Choose your endpoint's memory size according to your model size. Generally, the memory size should be at least as large as your model size. You may need to benchmark in order to choose the right memory selection for your model based on your latency SLAs. The memory size increments have different pricing; see the [Amazon SageMaker pricing page](https://aws.amazon.com/sagemaker/pricing/) for more information.

**Batch Transform**

The following FAQ items answer common questions for SageMaker Batch Transform.

**Q: How does Batch Transform split my data?**

A: For specific file formats such as CSV, RecordIO and TFRecord, SageMaker can split your data into single-record or multi-record mini batches and send this as a payload to your model container. When the value of `BatchStrategy` is `MultiRecord`, SageMaker sends the maximum number of
records in each request, up to the MaxPayloadInMB limit. When the value of BatchStrategy is SingleRecord, SageMaker sends individual records in each request.

**Q: What is the maximum timeout for Batch Transform and payload limit for a single record?**

**A:** The maximum timeout for Batch Transform is 3600 seconds. The [maximum payload size](https://docs.aws.amazon.com/sagemaker/latest/dg/batch-transform.html) for a record (per mini batch) is 100 MB.

**Q: How do I speed up a Batch Transform job?**

**A:** If you are using the [CreateTransformJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateTransformJob.html) API, you can reduce the time it takes to complete batch transform jobs by using optimal values for parameters such as MaxPayloadInMB, MaxConcurrentTransforms, or BatchStrategy. The ideal value for MaxConcurrentTransforms is equal to the number of compute workers in the batch transform job. If you are using the SageMaker console, you can specify these optimal parameter values in the Additional configuration section of the Batch transform job configuration page. SageMaker automatically finds the optimal parameter settings for built-in algorithms. For custom algorithms, provide these values through an execution-parameters endpoint.

**Q: What are the data formats natively supported in Batch Transform?**

**A:** Batch Transform supports CSV and JSON.

**Asynchronous Inference**

The following FAQ items answer common general questions for SageMaker Asynchronous Inference.

**Q: What is Amazon SageMaker Asynchronous Inference?**

**A:** Asynchronous Inference queues incoming requests and processes them asynchronously. This option is ideal for requests with large payload sizes or long processing times that need to be processed as they arrive. Optionally, you can configure auto-scaling settings to scale down the instance count to zero when not actively processing requests.

**Q: How do I scale my endpoints to 0 when there’s no traffic?**

**A:** Amazon SageMaker supports automatic scaling (autoscaling) your asynchronous endpoint. Autoscaling dynamically adjusts the number of instances provisioned for a model in response to changes in your workload. Unlike other hosted models SageMaker supports, with Asynchronous Inference you can also scale down your asynchronous endpoints instances to zero. Requests that
are received when there are zero instances are queued for processing once the endpoint scales up. For more information, see [Autoscale an asynchronous endpoint](#).

Amazon SageMaker Serverless Inference also automatically scales down to zero. You won’t see this because SageMaker manages scaling your serverless endpoints, but if you are not experiencing any traffic, the same infrastructure applies.
Implement MLOps

Amazon SageMaker supports features to implement machine learning models in production environments with continuous integration and deployment. The following topics give information about how to set up MLOps infrastructure when using SageMaker.

Topics

- Why Should You Use MLOps?
- SageMaker Experiments
- SageMaker Workflows
- Amazon SageMaker ML Lineage Tracking
- Register and Deploy Models with Model Registry
- Model Deployment in SageMaker
- SageMaker Model Monitor
- Automate MLOps with SageMaker Projects
- Amazon SageMaker MLOps FAQ

Why Should You Use MLOps?

As you move from running individual artificial intelligence and machine learning (AI/ML) projects to using AI/ML to transform your business at scale, the discipline of ML Operations (MLOps) can help. MLOps accounts for the unique aspects of AI/ML projects in project management, CI/CD, and quality assurance, helping you improve delivery time, reduce defects, and make data science more productive. MLOps refers to a methodology that is built on applying DevOps practices to machine learning workloads. For a discussion of DevOps principles, see the white paper Introduction to DevOps on AWS. To learn more about implementation using AWS services, see Practicing CI/CD on AWS and Infrastructure as Code.

Like DevOps, MLOps relies on a collaborative and streamlined approach to the machine learning development lifecycle where the intersection of people, process, and technology optimizes the end-to-end activities required to develop, build, and operate machine learning workloads.

MLOps focuses on the intersection of data science and data engineering in combination with existing DevOps practices to streamline model delivery across the machine learning development lifecycle. MLOps is the discipline of integrating ML workloads into release management, CI/CD, and
operations. MLOps requires the integration of software development, operations, data engineering, and data science.

**Challenges with MLOps**

Although MLOps can provide valuable tools to help you scale your business, you might face certain issues as you integrate MLOps into your machine learning workloads.

**Project management**

- ML projects involve data scientists, a relatively new role, and one not often integrated into cross-functional teams. These new team members often speak a very different technical language than product owners and software engineers, compounding the usual problem of translating business requirements into technical requirements.

**Communication and collaboration**

- Building visibility on ML projects and enabling collaboration across different stakeholders such as data engineers, data scientists, ML engineers, and DevOps is becoming increasingly important to ensure successful outcomes.

**Everything is code**

- Use of production data in development activities, longer experimentation lifecycles, dependencies on data pipelines, retraining deployment pipelines, and unique metrics in evaluating the performance of a model.
- Models often have a lifecycle independent of the applications and systems integrating with those models.
- The entire end-to-end system is reproducible through versioned code and artifacts. DevOps projects use Infrastructure-as-Code (IaC) and Configuration-as-Code (CaC) to build environments, and Pipelines-as-Code (PaC) to ensure consistent CI/CD patterns. The pipelines have to integrate with Big Data and ML training workflows. That often means that the pipeline is a combination of a traditional CI/CD tool and another workflow engine. There are important policy concerns for many ML projects, so the pipeline may also need to enforce those policies. Biased input data produces biased results, an increasing concern for business stakeholders.

**CI/CD**

Challenges with MLOps
In MLOps, the source data is a first-class input, along with source code. That's why MLOps calls for versioning the source data and initiating pipeline runs when the source or inference data changes.

Pipelines must also version the ML models, along with inputs and other outputs, in order to provide for traceability.

Automated testing must include proper validation of the ML model during build phases and when the model is in production.

Build phases may include model training and retraining, a time-consuming and resource-intensive process. Pipelines must be granular enough to only perform a full training cycle when the source data or ML code changes, not when related components change.

Because machine learning code is typically a small part of an overall solution, a deployment pipeline may also incorporate the additional steps required to package a model for consumption as an API by other applications and systems.

Monitoring and logging

The feature engineering and model training phases needed to capture model training metrics as well as model experiments. Tuning an ML model requires manipulating the form of the input data as well as algorithm hyperparameters, and systematically capture those experiments. Experiment tracking helps data scientists work more effectively and gives a reproducible snapshot of their work.

Deployed ML models require monitoring of the data passed to the model for inference, along with the standard endpoint stability and performance metrics. The monitoring system must also capture the quality of model output, as evaluated by an appropriate ML metric.

Benefits of MLOps

Adopting MLOps practices gives you faster time-to-market for ML projects by delivering the following benefits.

- **Productivity:** Providing self-service environments with access to curated data sets lets data engineers and data scientists move faster and waste less time with missing or invalid data.
- **Repeatability:** Automating all the steps in the MLDC helps you ensure a repeatable process, including how the model is trained, evaluated, versioned, and deployed.
- **Reliability**: Incorporating CI/CD practices allows for the ability to not only deploy quickly but with increased quality and consistency.

- **Auditability**: Versioning all inputs and outputs, from data science experiments to source data to trained model, means that we can demonstrate exactly how the model was built and where it was deployed.

- **Data and model quality**: MLOps lets us enforce policies that guard against model bias and track changes to data statistical properties and model quality over time.

**SageMaker Experiments**

ML model building requires many iterations of training as you tune the algorithm, model architecture, and parameters to achieve high prediction accuracy. You can track the inputs and outputs across these training iterations to improve repeatability of trials and collaboration within your team using Amazon SageMaker Experiments. You can also track parameters, metrics, datasets, and other artifacts related to your model training jobs. SageMaker Experiments offers a single interface where you can visualize your in-progress training jobs, share experiments within your team, and deploy models directly from an experiment.

To learn about SageMaker Experiments, see [Manage Machine Learning with Amazon SageMaker Experiments](#).

**SageMaker Workflows**

As you scale your machine learning (ML) operations, you can use Amazon SageMaker fully managed workflow services to implement continuous integration and deployment (CI/CD) practices for your ML lifecycle. With the SageMaker Pipelines SDK, you choose and integrate pipeline steps into a unified solution that automates the model-building process from data preparation to model deployment. For Kubernetes based architectures, you can install SageMaker Operators on your Kubernetes cluster to create SageMaker jobs natively using the Kubernetes API and command-line Kubernetes tools such as kubectl. With SageMaker components for Kubeflow pipelines, you can create and monitor native SageMaker jobs from your Kubeflow Pipelines. The job parameters, status, and outputs from SageMaker are accessible from the Kubeflow Pipelines UI. Lastly, if you want to schedule non-interactive batch runs of your Jupyter notebook, use the notebook-based workflows service to initiate standalone or regular runs on a schedule you define.

In summary, SageMaker offers the following workflow technologies:
Amazon SageMaker Model Building Pipelines: Tool for building and managing ML pipelines.

Kubernetes Orchestration: SageMaker custom operators for your Kubernetes cluster and components for Kubeflow Pipelines.

SageMaker Notebook Jobs: On demand or scheduled non-interactive batch runs of your Jupyter notebook.

You can also leverage other services that integrate with SageMaker to build your workflow. Options include the following services:

- **Airflow Workflows**: SageMaker APIs to export configurations for creating and managing Airflow workflows.
- **AWS Step Functions**: Multi-step ML workflows in Python that orchestrate SageMaker infrastructure without having to provision your resources separately.

For more information on managing SageMaker training and inference, see Amazon SageMaker Python SDK Workflows.

**Topics**

- Amazon SageMaker Model Building Pipelines
- Kubernetes Orchestration
- SageMaker Notebook Jobs

**Amazon SageMaker Model Building Pipelines**

Amazon SageMaker Model Building Pipelines is a tool for building machine learning pipelines that take advantage of direct SageMaker integration. Because of this integration, you can create a pipeline and set up SageMaker Projects for orchestration using a tool that handles much of the step creation and management for you. You can build the pipeline using the SageMaker Python SDK, or you can author the pipeline using the SageMaker Pipeline Definition JSON Schema.

SageMaker Pipelines provides the following advantages over other AWS workflow offerings:

**SageMaker Integration**
SageMaker Pipelines is integrated directly with SageMaker, so you don't need to interact with any other AWS services. You also don't need to manage any resources because SageMaker Pipelines is a fully managed service, which means that it creates and manages resources for you.

**SageMaker Python SDK Integration**

Because SageMaker Pipelines is integrated with the SageMaker Python SDK, you can create your pipelines programmatically using a high-level Python interface that you might already be familiar with. To view the SageMaker Python SDK API reference, see [Pipelines](#). For SageMaker Python SDK code examples, see [Amazon SageMaker Model Building Pipelines](#).

**SageMaker Studio Integration**

SageMaker Studio offers an environment to manage the end-to-end SageMaker Pipelines experience. Using Studio, you can bypass the AWS console for your entire workflow management. For more information on managing SageMaker Pipelines from SageMaker Studio, see [View, Track, and Execute SageMaker Pipelines in SageMaker Studio](#).

**Data Lineage Tracking**

With SageMaker Pipelines you can track the history of your data within the pipeline execution. Amazon SageMaker ML Lineage Tracking lets you analyze where the data came from, where it was used as an input, and the outputs that were generated from it. For example, you can view the models created from an individual dataset, and you can view the datasets that went into creating an individual model. For more information, see [Amazon SageMaker ML Lineage Tracking](#).

**Step Reuse**

With SageMaker Pipelines, you can designate steps for caching. When a step is cached, it is indexed for reuse later if the same step is executed again. As a result, you can reuse the output from previous step executions of the same step in the same pipeline without having to run the step again. For more information on step caching, see [Caching Pipeline Steps](#).

**Topics**

- [SageMaker Pipelines Overview](#)
- [Create and Manage SageMaker Pipelines](#)
SageMaker Pipelines Overview

An Amazon SageMaker Model Building Pipelines pipeline is a series of interconnected steps that are defined using the Pipelines SDK. You can also build your pipeline without the SDK using the pipeline definition JSON schema. This pipeline definition encodes a pipeline using a directed acyclic graph (DAG) that can be exported as a JSON definition. This DAG gives information on the requirements for and relationships between each step of your pipeline. The structure of a pipeline's DAG is determined by the data dependencies between steps. These data dependencies are created when the properties of a step's output are passed as the input to another step. The following image is an example of a pipeline DAG:

The following topics describe fundamental SageMaker Pipelines concepts. For a tutorial describing the implementation of these concepts, see Create and Manage SageMaker Pipelines.
Pipeline Structure and Execution

Topics

- Pipeline Structure
- Pipeline Execution using Parallelism Configuration

Pipeline Structure

An Amazon SageMaker Model Building Pipelines instance is composed of a name, parameters, and steps. Pipeline names must be unique within an (account, region) pair. All parameters used in step definitions must be defined in the pipeline. Pipeline steps listed automatically determine their order of execution by their data dependencies on one another. The SageMaker Pipelines service resolves the relationships between steps in the data dependency DAG to create a series of steps that the execution completes. The following is an example of a pipeline structure.
from sagemaker.workflow.pipeline import Pipeline

pipeline_name = f"AbalonePipeline"
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[
        processing_instance_type,
        processing_instance_count,
        training_instance_type,
        model_approval_status,
        input_data,
        batch_data,
    ],
    steps=[step_process, step_train, step_eval, step_cond],
)

Pipeline Execution using Parallelism Configuration

By default, a pipeline performs all steps that are available to run in parallel. You can control this behavior by using the `ParallelismConfiguration` property when creating or updating a pipeline, as well as when starting or retrying a pipeline execution.

Parallelism configurations are applied per execution. For example, if two executions are started they can each run a maximum of 50 steps concurrently, for a total of 100 concurrently running steps. Also, `ParallelismConfiguration(s)` specified when starting, retrying or updating an execution take precedence over parallelism configurations defined in the pipeline.

Example Creating a pipeline execution with `ParallelismConfiguration`

```
pipeline = Pipeline(
    name="myPipeline",
    steps=[step_process, step_train]
)

pipeline.create(role, parallelism_config={"MaxParallelExecutionSteps": 50})
```

IAM Access Management

The following sections describe the AWS Identity and Access Management (IAM) requirements for Amazon SageMaker Model Building Pipelines. For an example of how you can implement these permissions, see Prerequisites.
 Topics

- Pipeline Role Permissions
- Pipeline Step Permissions
- Customize access management for SageMaker Pipelines jobs
- Service Control Policies with Pipelines

Pipeline Role Permissions

Your pipeline requires an IAM pipeline execution role that is passed to SageMaker Pipelines when you create a pipeline. The role for the SageMaker instance that is creating the pipeline must have the `iam:PassRole` permission for the pipeline execution role in order to pass it. For more information on IAM roles, see IAM Roles.

Your pipeline execution role requires the following permissions:

- To pass any role to a SageMaker job within a pipeline, the `iam:PassRole` permission for the role that is being passed.
- Create and Describe permissions for each of the job types in the pipeline.
- Amazon S3 permissions to use the `JsonGet` function. You control access to your Amazon S3 resources using resource-based policies and identity-based policies. A resource-based policy is applied to your Amazon S3 bucket and grants SageMaker Pipelines access to the bucket. An identity-based policy gives your pipeline the ability to make Amazon S3 calls from your account. For more information on resource-based policies and identity-based policies, see Identity-based policies and resource-based policies.

```json
{
   "Action": [
      "s3:GetObject"
   ],
   "Resource": "arn:aws:s3:::<your-bucket-name>/*",
   "Effect": "Allow"
}
```

Pipeline Step Permissions

SageMaker Pipelines include steps that run SageMaker jobs. In order for the pipeline steps to run these jobs, they require an IAM role in your account that provides access for the needed resource.
This role is passed to the SageMaker service principal by your pipeline. For more information on IAM roles, see IAM Roles.

By default, each step takes on the pipeline execution role. You can optionally pass a different role to any of the steps in your pipeline. This ensures that the code in each step does not have the ability to impact resources used in other steps unless there is a direct relationship between the two steps specified in the pipeline definition. You pass these roles when defining the processor or estimator for your step. For examples of how to include these roles in these definitions, see the SageMaker Python SDK documentation.

Customize access management for SageMaker Pipelines jobs

You can further customize your IAM policies so selected members in your organization can run any or all pipeline steps. For example, you can give certain users permission to create training jobs, and another group of users permission to create processing jobs, and all of your users permission to run the remaining steps. To use this feature, you select a custom string which prefixes your job name. Your admin prepends the permitted ARNs with the prefix while your data scientist includes this prefix in pipeline instantiations. Because the IAM policy for permitted users contains a job ARN with the specified prefix, subsequent jobs of your pipeline step have necessary permissions to proceed. Job prefixing is off by default—you must toggle on this option in your Pipeline class to use it.

For jobs with prefixing turned off, the job name is formatted as shown and is a concatenation of fields described in the following table:

```
pipelines-<executionId>-<stepNamePrefix>-<entityToken>-<failureCount>
```

<table>
<thead>
<tr>
<th>Field</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>pipelines</td>
<td>A static string always prepended. This string identifies the pipeline orchestration service as the job's source.</td>
</tr>
<tr>
<td>executionId</td>
<td>A randomized buffer for the running instance of the pipeline.</td>
</tr>
<tr>
<td>stepNamePrefix</td>
<td>The user-specified step name (given in the name argument</td>
</tr>
<tr>
<td>Field</td>
<td>Definition</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------------------------------------------</td>
</tr>
<tr>
<td>entityToken</td>
<td>A randomized token to ensure idempotency of the step entity.</td>
</tr>
<tr>
<td>failureCount</td>
<td>The current number of retries attempted to complete the job.</td>
</tr>
</tbody>
</table>

In this case, no custom prefix is prepended to the job name, and the corresponding IAM policy must match this string.

For users who turn on job prefixing, the underlying job name takes the following form, with the custom prefix specified as `MyBaseJobName`:

```
<MyBaseJobName>-<executionId>-<entityToken>-<failureCount>
```

The custom prefix replaces the static pipelines string to help you narrow the selection of users who can run the SageMaker job as a part of a pipeline.

**Prefix length restrictions**

The job names have internal length constraints specific to individual pipeline steps. This constraint also limits the length of the allowed prefix. The prefix length requirements are as follows:

<table>
<thead>
<tr>
<th>Pipeline step</th>
<th>Prefix length</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>TrainingStep</code>, <code>ModelStep</code>, <code>TransformStep</code>, <code>ProcessingStep</code>, <code>ClarifyCheckStep</code>, <code>QualityCheckStep</code>, <code>RegisterModelStep</code></td>
<td>38</td>
</tr>
<tr>
<td><code>TuningStep</code>, <code>AutoML</code></td>
<td>6</td>
</tr>
</tbody>
</table>
Apply job prefixes to an IAM policy

Your admin creates IAM policies allowing users of specific prefixes to create jobs. The following example policy permits data scientists to create training jobs if they use the MyBaseJobName prefix.

```json
{
    "Action": "sagemaker:CreateTrainingJob",
    "Effect": "Allow",
    "Resource": [
        "arn:aws:sagemaker:region:account-id:*/MyBaseJobName-*"
    ]
}
```

Apply job prefixes to pipeline instantiations

You specify your prefix with the *base_job_name argument of the job instance class.

Note

You pass your job prefix with the *base_job_name argument to the job instance before creating a pipeline step. This job instance contains the necessary information for the job to run as a step in a pipeline. This argument varies depending upon the job instance used. The following list shows which argument to use for each pipeline step type:

- **base_job_name** for the **Estimator** (TrainingStep), **Processor** (ProcessingStep), and **AutoML** (AutoMLStep) classes
- **tuning_base_job_name** for the **Tuner** class (TuningStep)
- **transform_base_job_name** for the **Transformer** class (TransformStep)
- **base_job_name** of **CheckJobConfig** for the **QualityCheckStep** (Quality Check) and **ClarifyCheckstep** (Clarify Check) classes
- For the **Model** class, the argument used depends on if you run create or register on your model before passing the result to **ModelStep**
  - If you call create, the custom prefix comes from the name argument when you construct your model (i.e., Model(name=))
If you call `register`, the custom prefix comes from the `model_package_name` argument of your call to `register` (i.e., `my_model.register(model_package_name=)`)

The following example shows how to specify a prefix for a new training job instance.

```python
# Create a job instance
xgb_train = Estimator(
    image_uri=image_uri,
    instance_type="ml.m5.xlarge",
    instance_count=1,
    output_path=model_path,
    role=role,
    subnets=['subnet-0ab12c34567de89f0'],
    base_job_name="MyBaseJobName",
    security_group_ids=['sg-1a2bbcc3bd4444e55'],
    tags = [ ... ]
    encrypt_inter_container_traffic=True,
)

# Attach your job instance to a pipeline step
step_train = TrainingStep(
    name="TestTrainingJob",
    estimator=xgb_train,
    inputs={
        "train": TrainingInput(...),
        "validation": TrainingInput(...)
    }
)
```

Job prefixing is off by default. To opt into this feature, use the `use_custom_job_prefix` option of `PipelineDefinitionConfig` as shown in the following snippet:

```python
from sagemaker.workflow.pipeline_definition_config import PipelineDefinitionConfig

# Create a definition configuration and toggle on custom prefixing
definition_config = PipelineDefinitionConfig(use_custom_job_prefix=True);

# Create a pipeline with a custom prefix
pipeline = Pipeline(
    name="MyJobPrefixedPipeline",
 ```
Create and run your pipeline. The following example creates and runs a pipeline, and also demonstrates how you can turn off job prefixing and rerun your pipeline.

```python
pipeline.create(role_arn=sagemaker.get_execution_role())

# Optionally, call definition() to confirm your prefixed job names are in the built JSON
pipeline.definition()
pipeline.start()

# To run a pipeline without custom-prefixes, toggle off use_custom_job_prefix, update the pipeline
# via upsert() or update(), and start a new run
definition_config = PipelineDefinitionConfig(use_custom_job_prefix=False)
pipeline.pipeline_definition_config = definition_config
pipeline.update()
execution = pipeline.start()

Similarly, you can toggle the feature on for existing pipelines and start a new run which uses job prefixes.

```python
definition_config = PipelineDefinitionConfig(use_custom_job_prefix=True)
pipeline.pipeline_definition_config = definition_config
pipeline.update()
execution = pipeline.start()
```

Finally, you can view your custom-prefixed job by calling `list_steps` on the pipeline execution.

```python
steps = execution.list_steps()

prefixed_training_job_name = steps['PipelineExecutionSteps'][0]['Metadata']
['TrainingJob'] ['Arn']
```

**Service Control Policies with Pipelines**

Service control policies (SCPs) are a type of organization policy that you can use to manage permissions in your organization. SCPs offer central control over the maximum available
permissions for all accounts in your organization. By using SageMaker Pipelines within your organization, you can ensure that data scientists manage your pipeline executions without having to interact with the AWS console.

If you're using a VPC with your SCP that restricts access to Amazon S3, you need to take steps to allow your pipeline to access other Amazon S3 resources.

To allow SageMaker Pipelines to access Amazon S3 outside of your VPC with the JsonGet function, update your organization's SCP to ensure that the role using SageMaker Pipelines can access Amazon S3. To do this, create an exception for roles that are being used by the SageMaker Pipelines executor via the pipeline execution role using a principal tag and condition key.

**To allow SageMaker Pipelines to access Amazon S3 outside of your VPC**

1. Create a unique tag for your pipeline execution role following the steps in [Tagging IAM users and roles](#).
2. Grant an exception in your SCP using the `Aws:PrincipalTag` IAM condition key for the tag you created. For more information, see [Creating, updating, and deleting service control policies](#).

**Cross-Account Support for SageMaker Pipelines**

You can use cross-account support for Amazon SageMaker Model Building Pipelines to share pipeline entities across AWS accounts and access shared pipelines through direct API calls.

**Set up cross-account pipeline sharing**

SageMaker uses [AWS Resource Access Manager](https://aws.amazon.com/ram/) (AWS RAM) to help you securely share your pipeline entities across accounts.

**Create a resource share**

1. Select *Create a resource share* through the [AWS RAM console](https://aws.amazon.com/ram/).
2. When specifying resource share details, choose the SageMaker Pipelines resource type and select one or more pipelines that you want to share. When you share a pipeline with any other account, all of its executions are also shared implicitly.
3. Associate permissions with your resource share. Choose either the default read-only permission policy or the extended pipeline execution permission policy. For more detailed information, see [Permission policies for SageMaker Pipelines resources](#).
**Note**

If you select the extended pipeline execution policy, note that any start, stop, and retry commands called by shared accounts use resources in the AWS account that shared the pipeline.

4. Use AWS account IDs to specify the accounts to which you want to grant access to your shared resources.

5. Review your resource share configuration and select **Create resource share**. It may take a few minutes for the resource share and principal associations to complete.

For more information, see [Sharing your AWS resources](https://docs.aws.amazon.com/ram/latest/userguide/using-share-invitations.html) in the *AWS Resource Access Manager User Guide*.

**Get responses to your resource share invitation**

Once the resource share and principal associations are set, the specified AWS accounts receive an invitation to join the resource share. The AWS accounts must accept the invite to gain access to any shared resources.

For more information on accepting a resource share invite through AWS RAM, see [Using shared AWS resources](https://docs.aws.amazon.com/ram/latest/userguide/using-share-invitations.html) in the *AWS Resource Access Manager User Guide*.

**Permission policies for SageMaker Pipelines resources**

When creating your resource share, choose one of two supported permission policies to associate with the SageMaker pipeline resource type. Both policies grant access to any selected pipeline and all of its executions.

**Default read-only permissions**

The AWSRAMDefaultPermissionSageMakerPipeline policy allows the following read-only actions:

```
"sagemaker:DescribePipeline"
"sagemaker:DescribePipelineDefinitionForExecution"
"sagemaker:DescribePipelineExecution"
"sagemaker:ListPipelineExecutions"
"sagemaker:ListPipelineExecutionSteps"
"sagemaker:ListPipelineParametersForExecution"
```
Extended pipeline execution permissions

The AWSRAMPermissionSageMakerPipelineAllowExecution policy includes all of the read-only permissions from the default policy and also allows shared accounts to start, stop, and retry pipeline executions.

Note

Be mindful of AWS resource usage when using the extended pipeline execution permission policy. With this policy, shared accounts are allowed to start, stop, and retry pipeline executions. Any resources used for shared pipeline executions are consumed by the owner account.

The extended pipeline execution permission policy allows the following actions:

- "sagemaker:DescribePipeline"
- "sagemaker:DescribePipelineDefinitionForExecution"
- "sagemaker:DescribePipelineExecution"
- "sagemaker:ListPipelineExecutions"
- "sagemaker:ListPipelineExecutionSteps"
- "sagemaker:ListPipelineParametersForExecution"
- "sagemaker:StartPipelineExecution"
- "sagemaker:StopPipelineExecution"
- "sagemaker:RetryPipelineExecution"
- "sagemaker:Search"

Access shared pipeline entities through direct API calls

Once cross-account pipeline sharing is set up, you can call the following SageMaker API actions using a pipeline ARN:

Note

You can only call API commands if they are included in the permissions associated with your resource share. If you select the AWSRAMPermissionSageMakerPipelineAllowExecution policy, then the start, stop, and retry commands use resources in the AWS account that shared the pipeline.
Pipeline Parameters

You can introduce variables into your pipeline definition using parameters. You can reference parameters that you define throughout your pipeline definition. Parameters have a default value, which you can override by specifying parameter values when starting a pipeline execution. The default value must be an instance matching the parameter type. All parameters used in step definitions must be defined in your pipeline definition. Amazon SageMaker Model Building Pipelines supports the following parameter types:

- ParameterString – Representing a string parameter.
- ParameterInteger – Representing an integer parameter.
- ParameterFloat – Representing a float parameter.
- ParameterBoolean – Representing a Boolean Python type.

Parameters take the following format:

```python
<parameter> = <parameter_type>(
    name="<parameter_name>",
    default_value=<default_value>
)
```

The following example shows a sample parameter implementation.

```python
from sagemaker.workflow.parameters import (ParameterInteger, ParameterString,
                                        )
```
You pass the parameter when creating your pipeline as shown in the following example.

```python
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[
        processing_instance_count
    ],
    steps=[step_process]
)
```

You can also pass a parameter value that differs from the default value to a pipeline execution, as shown in the following example.

```python
execution = pipeline.start(
    parameters=dict(
        ProcessingInstanceCount="2",
        ModelApprovalStatus="Approved"
    )
)
```

You can manipulate parameters with SageMaker Python SDK functions like `sagemaker.workflow.functions.Join`. For more information on parameters, see SageMaker Pipelines Parameters.

For known limitations of SageMaker Pipelines Parameters, see Limitations - Parameterization in the Amazon SageMaker Python SDK.

**Pipeline Steps**

SageMaker Pipelines are composed of steps. These steps define the actions that the pipeline takes and the relationships between steps using properties.

**Topics**
• **Step Types**
• **Step Properties**
• **Step Parallelism**
• **Data Dependency Between Steps**
• **Custom Dependency Between Steps**
• **Use a Custom Image in a Step**

### Step Types

The following describes the requirements of each step type and provides an example implementation of the step. These are not functional implementations because they don't provide the resource and inputs needed. For a tutorial that implements these steps, see [Create and Manage SageMaker Pipelines](#).

⚠️ **Note**

You can also create a step from your local machine learning code by converting it to a SageMaker Pipelines step with the `@step` decorator. For more information, see [@step decorator](#).

Amazon SageMaker Model Building Pipelines support the following step types:

• **Processing**
• **Training**
• **Tuning**
• **AutoML**
• **Model**
• **CreateModel**
• **RegisterModel**
• **Transform**
• **Condition**
• **Callback**
• **Lambda**
You can create a step from local machine learning code using the `@step` decorator. After you test your code, you can convert the function to a SageMaker pipeline step by annotating it with the `@step` decorator. SageMaker Pipelines creates and runs a pipeline when you pass the output of the `@step`-decorated function as a step to your pipeline. You can also create a multi-step DAG pipeline that includes one or more `@step`-decorated functions as well as traditional SageMaker pipeline steps. For more details about how to create a step with `@step` decorator, see Lift-and-shift Python code with the `@step` decorator.

**Processing Step**

Use a processing step to create a processing job for data processing. For more information on processing jobs, see Process Data and Evaluate Models.

A processing step requires a processor, a Python script that defines the processing code, outputs for processing, and job arguments. The following example shows how to create a `ProcessingStep` definition.

```python
from sagemaker.sklearn.processing import SKLearnProcessor
sklearn_processor = SKLearnProcessor(framework_version='1.0-1',
                                 role=<role>,
                                 instance_type='ml.m5.xlarge',
                                 instance_count=1)

from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker.workflow.steps import ProcessingStep
inputs = [
             ProcessingInput(source=<input_data>, destination="/opt/ml/processing/input"),
         ]
```
`outputs = [
    ProcessingOutput(output_name="train", source="/opt/ml/processing/train"),
    ProcessingOutput(output_name="validation", source="/opt/ml/processing/validation"),
    ProcessingOutput(output_name="test", source="/opt/ml/processing/test")
]

step_process = ProcessingStep(
    name="AbaloneProcess",
    step_args = sklearn_processor.run(inputs=inputs, outputs=outputs,
        code="abalone/preprocessing.py")
)

Pass runtime parameters

The following example shows how to pass runtime parameters from a PySpark processor to a ProcessingStep.

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.spark.processing import PySparkProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker.workflow.steps import ProcessingStep

pipeline_session = PipelineSession()

pyspark_processor = PySparkProcessor(
    framework_version='2.4',
    role=<role>,
    instance_type='ml.m5.xlarge',
    instance_count=1,
    sagemaker_session=pipeline_session,
)

step_args = pyspark_processor.run(
    inputs=[ProcessingInput(source=<input_data>, destination="/opt/ml/processing/input"),],
    outputs=[
        ProcessingOutput(output_name="train", source="/opt/ml/processing/train"),
        ProcessingOutput(output_name="validation", source="/opt/ml/processing/validation"),
        ProcessingOutput(output_name="test", source="/opt/ml/processing/test")
    ],
    code="preprocess.py",
    arguments=None,
)```
step_process = ProcessingStep(
    name="AbaloneProcess",
    step_args=step_args,
)

For more information on processing step requirements, see the
sagemaker.workflow.steps.ProcessingStep documentation. For an in-depth example, see Define a
Processing Step for Feature Engineering in the Orchestrate Jobs to Train and Evaluate Models with
Amazon SageMaker Pipelines example notebook.

Training Step

You use a training step to create a training job to train a model. For more information on training
jobs, see Train a Model with Amazon SageMaker.

A training step requires an estimator, as well as training and validation data inputs. The following
example shows how to create a TrainingStep definition. For more information on training step
requirements, see the sagemaker.workflow.steps.TrainingStep documentation.

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.inputs import TrainingInput
from sagemaker.workflow.steps import TrainingStep
from sagemaker.xgboost.estimator import XGBoost

pipeline_session = PipelineSession()

xgb_estimator = XGBoost(..., sagemaker_session=pipeline_session)

step_args = xgb_estimator.fit(
    inputs={
        "train": TrainingInput(
            s3_data=step_process.properties.ProcessingOutputConfig.Outputs["train"].S3Output.S3Uri,
            content_type="text/csv"
        ),
        "validation": TrainingInput(
```
Tuning Step

You use a tuning step to create a hyperparameter tuning job, also known as hyperparameter optimization (HPO). A hyperparameter tuning job runs multiple training jobs, each one producing a model version. For more information on hyperparameter tuning, see Perform Automatic Model Tuning with SageMaker.

The tuning job is associated with the SageMaker experiment for the pipeline, with the training jobs created as trials. For more information, see Experiments Integration.

A tuning step requires a HyperparameterTuner and training inputs. You can retrain previous tuning jobs by specifying the warm_start_config parameter of the HyperparameterTuner. For more information on hyperparameter tuning and warm start, see Run a Warm Start Hyperparameter Tuning Job.

You use the get_top_model_s3_uri method of the sagemaker.workflow.steps.TuningStep class to get the model artifact from one of the top-performing model versions. For a notebook that shows how to use a tuning step in a SageMaker pipeline, see sagemaker-pipelines-tuning-step.ipynb.

⚠️ Important

Tuning steps were introduced in Amazon SageMaker Python SDK v2.48.0 and Amazon SageMaker Studio Classic v3.8.0. You must update Studio Classic before you use a tuning step or the pipeline DAG doesn't display. To update Studio Classic, see Shut down and Update SageMaker Studio Classic.
```python
from sagemaker.workflow.pipeline_context import PipelineSession

from sagemaker.tuner import HyperparameterTuner
from sagemaker.inputs import TrainingInput
from sagemaker.workflow.steps import TuningStep

tuner = HyperparameterTuner(..., sagemaker_session=PipelineSession())

step_tuning = TuningStep(
    name = "HPTuning",
    step_args = tuner.fit(inputs=TrainingInput(s3_data="s3://my-bucket/my-data"))
)
```

**Get the best model version**

The following example shows how to get the best model version from the tuning job using the `get_top_model_s3_uri` method. At most, the top 50 performing versions are available ranked according to `HyperParameterTuningJobObjective`. The `top_k` argument is an index into the versions, where `top_k=0` is the best-performing version and `top_k=49` is the worst-performing version.

```python
best_model = Model(
    image_uri=image_uri,
    model_data=step_tuning.get_top_model_s3_uri(
        top_k=0,
        s3_bucket=sagemaker_session.default_bucket()
    ),
    ...)
```

For more information on tuning step requirements, see the [sagemaker.workflow.steps.TuningStep](https://docs.aws.amazon.com/sagemaker/latest/dg/automl_tuning.html) documentation.

**AutoML Step**

Use the [AutoML](https://docs.aws.amazon.com/sagemaker/latest/dg/automl.html) API to create an AutoML job to automatically train a model. For more information on AutoML jobs, see [Automate model development with Amazon SageMaker Autopilot](https://docs.aws.amazon.com/sagemaker/latest/dg/autopilot-quickstart.html).

**Note**

Currently, the AutoML step supports only **ensembling training mode**.
The following example shows how to create a definition using AutoMLStep.

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.workflow.automl_step import AutoMLStep

pipeline_session = PipelineSession()

auto_ml = AutoML(...,
    role="<role>",
    target_attribute_name="my_target_attribute_name",
    mode="ENSEMBLING",
    sagemaker_session=pipeline_session)

input_training = AutoMLInput(
    inputs="s3://my-bucket/my-training-data",
    target_attribute_name="my_target_attribute_name",
    channel_type="training",
)

input_validation = AutoMLInput(
    inputs="s3://my-bucket/my-validation-data",
    target_attribute_name="my_target_attribute_name",
    channel_type="validation",
)

step_args = auto_ml.fit(
    inputs=[input_training, input_validation]
)

step_automl = AutoMLStep(
    name="AutoMLStep",
    step_args=step_args,
)
```

Get the best model version

The AutoML step automatically trains several model candidates. You can get the model with the best objective metric from the AutoML job using the `get_best_auto_ml_model` method and an IAM role to access model artifacts as follows.

```python
best_model = step_automl.get_best_auto_ml_model(role=<role>)
```

For more information, see the AutoML step in the SageMaker Python SDK.
Model Step

Use a ModelStep to create or register a SageMaker model. For more information on ModelStep requirements, see the [sagemaker.workflow.model_step.ModelStep](https://docs.aws.amazon.com/sagemaker/latest/dg/model-step.html) documentation.

Create a model

You can use a ModelStep to create a SageMaker model. A ModelStep requires model artifacts and information about the SageMaker instance type that you need to use to create the model. For more information on SageMaker models, see [Train a Model with Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/train-a-model.html).

The following example shows how to create a ModelStep definition.

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.model import Model
from sagemaker.workflow.model_step import ModelStep

text = TrainingStep(...)
model = Model(
    image_uri=pytorch_estimator.training_image_uri(),
    model_data=text.train.properties.ModelArtifacts.S3ModelArtifacts,
    sagemaker_session=PipelineSession(),
    role=role,
)

model_step = ModelStep(
    name="MyModelCreationStep",
    step_args=model.create(instance_type="ml.m5.xlarge"),
)
```

Register a model

You can use a ModelStep to register a sagemaker.model.Model or a sagemaker.pipeline.PipelineModel with the Amazon SageMaker model registry. A PipelineModel represents an inference pipeline, which is a model composed of a linear sequence of containers that process inference requests. For more information about how to register a model, see [Register and Deploy Models with Model Registry](https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry.html).

The following example shows how to create a ModelStep that registers a PipelineModel.

```python
import time
from sagemaker.workflow.pipeline_context import PipelineSession
```
```python
from sagemaker.sklearn import SKLearnModel
from sagemaker.xgboost import XGBoostModel

pipeline_session = PipelineSession()

code_location = 's3://{0}/{1}/code'.format(bucket_name, prefix)

sklearn_model = SKLearnModel(
    model_data=processing_step.properties.ProcessingOutputConfig.Outputs['model'].S3Output.S3Uri,
    entry_point='inference.py',
    source_dir='sklearn_source_dir/',
    code_location=code_location,
    framework_version='1.0-1',
    role=role,
    sagemaker_session=pipeline_session,
    py_version='py3'
)

xgboost_model = XGBoostModel(
    model_data=training_step.properties.ModelArtifacts.S3ModelArtifacts,
    entry_point='inference.py',
    source_dir='xgboost_source_dir/',
    code_location=code_location,
    framework_version='0.90-2',
    py_version='py3',
    sagemaker_session=pipeline_session,
    role=role
)

from sagemaker.workflow.model_step import ModelStep
from sagemaker import PipelineModel

pipeline_model = PipelineModel(
    models=[sklearn_model, xgboost_model],
    role=role, sagemaker_session=pipeline_session,
)

register_model_step_args = pipeline_model.register(
    content_types=['application/json'],
    response_types=['application/json'],
    inference_instances=['ml.t2.medium', 'ml.m5.xlarge'],
    transform_instances=['ml.m5.xlarge'],
    model_package_group_name='sipgroup',
)```
CreateModel Step

⚠️ Important

We recommend using `Model Step` to create models as of v2.90.0 of the SageMaker Python SDK. `CreateModelStep` will continue to work in previous versions of the SageMaker Python SDK, but is no longer actively supported.

You use a `CreateModel` step to create a SageMaker model. For more information on SageMaker models, see [Train a Model with Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/ nephrology.html).

A create model step requires model artifacts and information about the SageMaker instance type that you need to use to create the model. The following example shows how to create a `CreateModel` step definition. For more information on `CreateModel` step requirements, see the `sagemaker.workflow.steps.CreateModelStep` documentation.

```python
from sagemaker.workflow.steps import CreateModelStep

step_create_model = CreateModelStep(
    name="AbaloneCreateModel",
    model=best_model,
    inputs=inputs
)
```

RegisterModel Step

⚠️ Important

We recommend using `Model Step` to register models as of v2.90.0 of the SageMaker Python SDK. `RegisterModel` will continue to work in previous versions of the SageMaker Python SDK, but is no longer actively supported.

```python
step_model_registration = ModelStep(
    name="AbaloneRegisterModel",
    step_args=register_model_step_args,
)
```
You use a RegisterModel step to register a `sagemaker.model.Model` or a `sagemaker.pipeline.PipelineModel` with the Amazon SageMaker model registry. A PipelineModel represents an inference pipeline, which is a model composed of a linear sequence of containers that process inference requests.

For more information about how to register a model, see [Register and Deploy Models with Model Registry](#). For more information on RegisterModel step requirements, see the `sagemaker.workflow.step_collections.RegisterModel` documentation.

The following example shows how to create a RegisterModel step that registers a PipelineModel.

```python
import time
from sagemaker.sklearn import SKLearnModel
from sagemaker.xgboost import XGBoostModel
code_location = 's3://{0}/{1}/code'.format(bucket_name, prefix)
sklearn_model = SKLearnModel(model_data=processing_step.properties.ProcessingOutputConfig.Outputs['model'].S3Output.S3Uri, entry_point='inference.py', source_dir='sklearn_source_dir/', code_location=code_location, framework_version='1.0-1', role=role, sagemaker_session=sagemaker_session, py_version='py3')
xgboost_model = XGBoostModel(model_data=training_step.properties.ModelArtifacts.S3ModelArtifacts, entry_point='inference.py', source_dir='xgboost_source_dir/', code_location=code_location, framework_version='0.90-2', py_version='py3', sagemaker_session=sagemaker_session, role=role)

from sagemaker.workflow.step_collections import RegisterModel
from sagemaker import PipelineModel
pipeline_model = PipelineModel(models=[sklearn_model, xgboost_model], role=role, sagemaker_session=sagemaker_session)
```
```python
step_register = RegisterModel(
    name="AbaloneRegisterModel",
    model=pipeline_model,
    content_types=["application/json"],
    response_types=["application/json"],
    inference_instances=["ml.t2.medium", "ml.m5.xlarge"],
    transform_instances=["ml.m5.xlarge"],
    model_package_group_name='sipgroup',
)
```

If `model` isn't provided, the register model step requires an estimator as shown in the following example.

```python
from sagemaker.workflow.step_collections import RegisterModel

step_register = RegisterModel(
    name="AbaloneRegisterModel",
    estimator=xgb_train,
    model_data=step_train.properties.ModelArtifacts.S3ModelArtifacts,
    content_types=["text/csv"],
    response_types=["text/csv"],
    inference_instances=["ml.t2.medium", "ml.m5.xlarge"],
    transform_instances=["ml.m5.xlarge"],
    model_package_group_name=model_package_group_name,
    approval_status=model_approval_status,
    model_metrics=model_metrics
)
```

### Transform Step

You use a transform step for batch transformation to run inference on an entire dataset. For more information about batch transformation, see [Run Batch Transforms with Inference Pipelines](#).

A transform step requires a transformer and the data on which to run batch transformation. The following example shows how to create a `Transform` step definition. For more information on `Transform` step requirements, see the [sagemaker.workflow.steps.TransformStep](#) documentation.

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.transformer import Transformer
```

---

Amazon SageMaker Model Building Pipelines 4747
from sagemaker.inputs import TransformInput
from sagemaker.workflow.steps import TransformStep

transformer = Transformer(..., sagemaker_session=PipelineSession())

step_transform = TransformStep(
    name="AbaloneTransform",
    step_args=transformer.transform(data="s3://my-bucket/my-data"),
)

Condition Step

You use a condition step to evaluate the condition of step properties to assess which action should be taken next in the pipeline.

A condition step requires a list of conditions, a list of steps to run if the condition evaluates to true, and a list of steps to run if the condition evaluates to false. The following example shows how to create a ConditionStep definition.

Limitations

- SageMaker Pipelines doesn’t support the use of nested condition steps. You can’t pass a condition step as the input for another condition step.
- A condition step can’t use identical steps in both branches. If you need the same step functionality in both branches, duplicate the step and give it a different name.

from sagemaker.workflow.conditions import ConditionLessThanOrEqualTo
from sagemaker.workflow.condition_step import ConditionStep
from sagemaker.workflow.functions import JsonGet

cond_lte = ConditionLessThanOrEqualTo(
    left=JsonGet(
        step_name=step_eval.name,
        property_file=evaluation_report,
        json_path="regression_metrics.mse.value"
    ),
    right=6.0
)

step_cond = ConditionStep(
    name="AbaloneMSECond",
)
conditions=[cond_lte],
    if_steps=[step_register, step_create_model, step_transform],
    else_steps=[]
)

For more information on ConditionStep requirements, see the `sagemaker.workflow.condition_step.ConditionStep` API reference. For more information on supported conditions, see *Amazon SageMaker Model Building Pipelines - Conditions* in the SageMaker Python SDK documentation.

### Callback Step

You use a Callback step to incorporate additional processes and AWS services into your workflow that aren't directly provided by Amazon SageMaker Model Building Pipelines. When a Callback step runs, the following procedure occurs:

- SageMaker Pipelines sends a message to a customer-specified Amazon Simple Queue Service (Amazon SQS) queue. The message contains a SageMaker Pipelines–generated token and a customer-supplied list of input parameters. After sending the message, SageMaker Pipelines waits for a response from the customer.
- The customer retrieves the message from the Amazon SQS queue and starts their custom process.
- When the process finishes, the customer calls one of the following APIs and submits the SageMaker Pipelines–generated token:
  - `SendPipelineExecutionStepSuccess`, along with a list of output parameters
  - `SendPipelineExecutionStepFailure`, along with a failure reason
- The API call causes SageMaker Pipelines to either continue the pipeline process or fail the process.

For more information on Callback step requirements, see the `sagemaker.workflow.callback_step.CallbackStep` documentation. For a complete solution, see *Extend SageMaker Pipelines to include custom steps using callback steps*.

⚠️ **Important**

Callback steps were introduced in Amazon SageMaker Python SDK v2.45.0 and Amazon SageMaker Studio Classic v3.6.2. You must update Studio Classic before you use a
Callback step or the pipeline DAG doesn't display. To update Studio Classic, see [Shut down and Update SageMaker Studio Classic](#).

The following sample demonstrates an implementation of the preceding procedure.

```python
from sagemaker.workflow.callback_step import CallbackStep

step_callback = CallbackStep(
    name="MyCallbackStep",
    sqs_queue_url="https://sqs.us-east-2.amazonaws.com/012345678901/MyCallbackQueue",
    inputs=[...],
    outputs=[...]
)

callback_handler_code = 'import boto3
import json
def handler(event, context):
    sagemaker_client=boto3.client("sagemaker")
    for record in event["Records"]:
        payload=json.loads(record["body"])
        token=payload["token"]
        # Custom processing
        # Call SageMaker to complete the step
        sagemaker_client.send_pipeline_execution_step_success(
            CallbackToken=token,
            OutputParameters=[...]
        )
,'```

**Note**

Output parameters for `CallbackStep` should not be nested. For example, if you use a nested dictionary as your output parameter, then the dictionary is treated as a single string (ex. `{"output1": "{\"nested_output1\":\"my-output\"}"}`). If you provide a
Stopping behavior

A pipeline process doesn't stop while a Callback step is running.

When you call `StopPipelineExecution` on a pipeline process with a running Callback step, SageMaker Pipelines sends an additional Amazon SQS message to the specified SQS queue. The body of the SQS message contains a **Status** field, which is set to **Stopping**. The following shows an example SQS message body.

```
{
  "token": "26vcYbeWsZ",
  "pipelineExecutionArn": "arn:aws:sagemaker:us-east-2:012345678901:pipeline/callback-pipeline/execution/7pinimwddh3a",
  "arguments": {
    "number": 5,
    "stringArg": "some-arg",
    "inputData": "s3://sagemaker-us-west-2-012345678901/abalone/abalone-dataset.csv"
  },
  "status": "Stopping"
}
```

You should add logic to your Amazon SQS message consumer to take any needed action (for example, resource cleanup) upon receipt of the message, followed by a call to `SendPipelineExecutionStepSuccess` or `SendPipelineExecutionStepFailure`.

Only when SageMaker Pipelines receives one of these calls does it stop the pipeline process.

**Lambda Step**

You use a Lambda step to run an AWS Lambda function. You can run an existing Lambda function, or SageMaker can create and run a new Lambda function. For a notebook that shows how to use a Lambda step in a SageMaker pipeline, see [sagemaker-pipelines-lambda-step.ipynb](#).

⚠️ **Important**

Lambda steps were introduced in Amazon SageMaker Python SDK v2.51.0 and Amazon SageMaker Studio Classic v3.9.1. You must update Studio Classic before you use a Lambda.
SageMaker provides the `sagemaker.lambda_helper.Lambda` class to create, update, invoke, and delete Lambda functions. Lambda has the following signature.

```python
Lambda(
    function_arn,       # Only required argument to invoke an existing Lambda function
    function_name,
    execution_role_arn,
    zipped_code_dir,    # Specify either zipped_code_dir and s3_bucket, OR script
    s3_bucket,          # S3 bucket where zipped_code_dir is uploaded
    script,             # Path of Lambda function script
    handler,            # Lambda handler specified as "lambda_script.lambda_handler"
    timeout,            # Maximum time the Lambda function can run before the lambda step fails
...
)
```

The `sagemaker.workflow.lambda_step.LambdaStep` class has a `lambda_func` argument of type `Lambda`. To invoke an existing Lambda function, the only requirement is to supply the Amazon Resource Name (ARN) of the function to `function_arn`. If you don't supply a value for `function_arn`, you must specify `handler` and one of the following:

- `zipped_code_dir` – The path of the zipped Lambda function
- `s3_bucket` – Amazon S3 bucket where `zipped_code_dir` is to be uploaded
- `script` – The path of the Lambda function script file

The following example shows how to create a Lambda step definition that invokes an existing Lambda function.

```python
from sagemaker.workflow.lambda_step import LambdaStep
from sagemaker.lambda_helper import Lambda

step_lambda = LambdaStep(
    name="ProcessingLambda",
    lambda_func=Lambda(
        # The following arguments are required to create a Lambda function:
        function_name="ProcessingLambda",
        execution_role_arn="arn:aws:iam::123456789012:role/sagemaker-execution-role",
        zipped_code_dir="/var/task",
        s3_bucket="my-s3-bucket",
        script="/var/task/lambda.py",
        handler="lambda.py:lambda_handler",
        timeout=300,
    ),
)
```
The following example shows how to create a Lambda step definition that creates and invokes a Lambda function using a Lambda function script.

```python
from sagemaker.workflow.lambda_step import LambdaStep
from sagemaker.lambda_helper import Lambda

step_lambda = LambdaStep(
    name="ProcessingLambda",
    lambda_func=Lambda(
        function_name="split-dataset-lambda",
        execution_role_arn=execution_role_arn,
        script="lambda_script.py",
        handler="lambda_script.lambda_handler",
        ...
    ),
    inputs={
        s3_bucket = s3_bucket,
        data_file = data_file
    },
    outputs=[
        "train_file", "test_file"
    ]
)
```

**Inputs and outputs**

If your Lambda function has inputs or outputs, these must also be defined in your Lambda step.
When defining the Lambda step, inputs must be a dictionary of key-value pairs. Each value of the inputs dictionary must be a primitive type (string, integer, or float). Nested objects are not supported. If left undefined, the inputs value defaults to None.

The outputs value must be a list of keys. These keys refer to a dictionary defined in the output of the Lambda function. Like inputs, these keys must be primitive types, and nested objects are not supported.

**Timeout and stopping behavior**

The Lambda class has a timeout argument that specifies the maximum time that the Lambda function can run. The default value is 120 seconds with a maximum value of 10 minutes. If the Lambda function is running when the timeout is met, the Lambda step fails; however, the Lambda function continues to run.

A pipeline process can't be stopped while a Lambda step is running because the Lambda function invoked by the Lambda step can't be stopped. If you attempt to stop the process while the Lambda function is running, the pipeline waits for the Lambda function to finish or until the timeout is hit, whichever occurs first, and then stops. If the Lambda function finishes, the pipeline process status is Stopped. If the timeout is hit the pipeline process status is Failed.

**ClarifyCheck Step**

You can use the ClarifyCheck step to conduct baseline drift checks against previous baselines for bias analysis and model explainability. You can then generate and register your baselines with the model.register() method and pass the output of that method to Model Step using step_args. These baselines for drift check can be used by Amazon SageMaker Model Monitor for your model endpoints so that you don’t need to do a baseline suggestion separately. The ClarifyCheck step can also pull baselines for drift check from the model registry. The ClarifyCheck step leverages the Amazon SageMaker Clarify prebuilt container that provides a
range of model monitoring capabilities, including constraint suggestion and constraint validation against a given baseline. For more information, see [Getting Started with a SageMaker Clarify Container](#).

**Configuring the ClarifyCheck step**

You can configure the ClarifyCheck step to conduct only one of the following check types each time it’s used in a pipeline.

- Data bias check
- Model bias check
- Model explainability check

You do this by setting the `clarify_check_config` parameter with one of the following check type values:

- `DataBiasCheckConfig`
- `ModelBiasCheckConfig`
- `ModelExplainabilityCheckConfig`

The ClarifyCheck step launches a processing job that runs the SageMaker Clarify prebuilt container and requires dedicated configurations for the check and the processing job. `ClarifyCheckConfig` and `CheckJobConfig` are helper functions for these configurations that are aligned with how the SageMaker Clarify processing job computes for checking model bias, data bias, or model explainability. For more information, see [Run SageMaker Clarify Processing Jobs for Bias Analysis and Explainability](#).

**Controlling step behaviors for drift check**

The ClarifyCheck step requires the following two boolean flags to control its behavior:

- `skip_check`: This parameter indicates if the drift check against the previous baseline is skipped or not. If it is set to False, the previous baseline of the configured check type must be available.
- `register_new_baseline`: This parameter indicates if a newly calculated baseline can be accessed though step property `BaselineUsedForDriftCheckConstraints`. If it is set to False, the previous baseline of the configured check type also must be available. This can be accessed through the `BaselineUsedForDriftCheckConstraints` property.
For more information, see [Baseline calculation, drift detection and lifecycle with ClarifyCheck and QualityCheck steps in Amazon SageMaker Model Building Pipelines](https://docs.aws.amazon.com/sagemaker/latest/dg/model-building-pipelines.html).

**Working with baselines**

You can optionally specify the `model_package_group_name` to locate the existing baseline and the ClarifyCheck step pulls the DriftCheckBaselines on the latest approved model package in the model package group. Or, you can provide a previous baseline through the `supplied_baseline_constraints` parameter. If you specify both the `model_package_group_name` and the `supplied_baseline_constraints`, the ClarifyCheck step uses the baseline specified by the `supplied_baseline_constraints` parameter.

For more information on using the ClarifyCheck step requirements, see the `sagemaker.workflow.steps.ClarifyCheckStep` in the [*Amazon SageMaker SageMaker SDK for Python*](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-sdk.html). For an Amazon SageMaker Studio Classic notebook that shows how to use ClarifyCheck step in SageMaker Pipelines, see [sagemaker-pipeline-model-monitor-clarify-steps.ipynb](https://github.com/awslabs/sagemaker-pipeline-model-monitor-clarify-steps.ipynb).

**Example Create a ClarifyCheck step for data bias check**

```python
from sagemaker.workflow.check_job_config import CheckJobConfig
from sagemaker.workflow.clarify_check_step import DataBiasCheckConfig, ClarifyCheckStep
from sagemaker.workflow.execution_variables import ExecutionVariables

check_job_config = CheckJobConfig(
    role=role,
    instance_count=1,
    instance_type="ml.c5.xlarge",
    volume_size_in_gb=120,
    sagemaker_session=sagemaker_session,
)

data_bias_data_config = DataConfig(
    s3_data_input_path=step_process.properties.ProcessingOutputConfig.Outputs["train"].S3Output.S3Uri,
    s3_output_path=Join(on='/', values=['s3:/', your_bucket, base_job_prefix, ExecutionVariables.PIPELINE_EXECUTION_ID, 'databiascheckstep']),
    label=0,
    dataset_type="text/csv",
    s3_analysis_config_output_path=data_bias_analysis_cfg_output_path,
)
```
data_bias_config = BiasConfig(
    label_values_or_threshold=[15.0], facet_name=[8], facet_values_or_threshold=[[0.5]]
)

data_bias_check_config = DataBiasCheckConfig(
    data_config=data_bias_data_config,
    data_bias_config=data_bias_config,
)

data_bias_check_step = ClarifyCheckStep(
    name="DataBiasCheckStep",
    clarify_check_config=data_bias_check_config,
    check_job_config=check_job_config,
    skip_check=False,
    register_new_baseline=False,
    supplied_baseline_constraints="s3://sagemaker-us-west-2-111122223333/baseline/analysis.json",
    model_package_group_name="MyModelPackageGroup"
)

QualityCheck Step

You can use the QualityCheck step to conduct baseline suggestions and drift checks against a previous baseline for data quality or model quality in a pipeline. You can then generate and register your baselines with the model.register() method and pass the output of that method to Model Step using step_args. Model Monitor can use these baselines for drift check for your model endpoints so that you don't need to do a baseline suggestion separately. The QualityCheck step can also pull baselines for drift check from the model registry. The QualityCheck step leverages the Amazon SageMaker Model Monitor prebuilt container, which has a range of model monitoring capabilities including constraint suggestion, statistics generation, and constraint validation against a baseline. For more information, see Amazon SageMaker Model Monitor prebuilt container.

Configuring the QualityCheck step

You can configure the QualityCheck step to conduct only one of the following check types each time it’s used in a pipeline.

- Data quality check
- Model quality check
You do this by setting the `quality_check_config` parameter with one of the following check type values:

- `DataQualityCheckConfig`
- `ModelQualityCheckConfig`

The `QualityCheck` step launches a processing job that runs the Model Monitor prebuilt container and requires dedicated configurations for the check and the processing job. The `QualityCheckConfig` and `CheckJobConfig` are helper functions for these configurations that are aligned with how Model Monitor creates a baseline for the model quality or data quality monitoring. For more information on the Model Monitor baseline suggestions, see [Create a Baseline](#) and [Create a Model Quality Baseline](#).

**Controlling step behaviors for drift check**

The `QualityCheck` step requires the following two Boolean flags to control its behavior:

- `skip_check`: This parameter indicates if the drift check against the previous baseline is skipped or not. If it is set to `False`, the previous baseline of the configured check type must be available.
- `register_new_baseline`: This parameter indicates if a newly calculated baseline can be accessed through step properties `BaselineUsedForDriftCheckConstraints` and `BaselineUsedForDriftCheckStatistics`. If it is set to `False`, the previous baseline of the configured check type must also be available. These can be accessed through the `BaselineUsedForDriftCheckConstraints` and `BaselineUsedForDriftCheckStatistics` properties.

For more information, see [Baseline calculation, drift detection and lifecycle with ClarifyCheck and QualityCheck steps in Amazon SageMaker Model Building Pipelines](#).

**Working with baselines**

You can specify a previous baseline directly through the `supplied_baseline_statistics` and `supplied_baseline_constraints` parameters, or you can simply specify the `model_package_group_name` and the `QualityCheck` step pulls the `DriftCheckBaselines` on the latest approved model package in the model package group. When you specify the `model_package_group_name`, `supplied_baseline_constraints`, and `supplied_baseline_statistics`, the `QualityCheck` step uses the baseline specified by
supplied_baseline_constraints and supplied_baseline_statistics on the check type of the QualityCheck step you are running.

For more information on using the QualityCheck step requirements, see the sagemaker.workflow.steps.QualityCheckStep in the Amazon SageMaker SageMaker SDK for Python. For an Amazon SageMaker Studio Classic notebook that shows how to use QualityCheck step in SageMaker Pipelines, see sagemaker-pipeline-model-monitor-clarify-steps.ipynb.

Example Create a QualityCheck step for data quality check

```python
from sagemaker.workflow.check_job_config import CheckJobConfig
from sagemaker.workflow.quality_check_step import DataQualityCheckConfig, QualityCheckStep
from sagemaker.workflow.execution_variables import ExecutionVariables

check_job_config = CheckJobConfig(
    role=role,
    instance_count=1,
    instance_type="ml.c5.xlarge",
    volume_size_in_gb=120,
    sagemaker_session=sagemaker_session,
)

data_quality_check_config = DataQualityCheckConfig(
    baseline_dataset=step_process.properties.ProcessingOutputConfig.Outputs["train"].S3Output.S3Uri,
    dataset_format=DatasetFormat.csv(header=False, output_columns_position="START"),
    output_s3_uri=Join(on='/', values=['s3:/', your_bucket, base_job_prefix, ExecutionVariables.PIPELINE_EXECUTION_ID, 'dataqualitycheckstep'])
)

data_quality_check_step = QualityCheckStep(
    name="DataQualityCheckStep",
    skip_check=False,
    register_new_baseline=False,
    quality_check_config=data_quality_check_config,
    check_job_config=check_job_config,
    supplied_baseline_statistics="s3://sagemaker-us-west-2-555555555555/baseline/statistics.json",
    supplied_baseline_constraints="s3://sagemaker-us-west-2-555555555555/baseline/constraints.json",
    model_package_group_name="MyModelPackageGroup"
)```
EMR Step

You can use the Amazon SageMaker Model Building Pipelines EMR step to process Amazon EMR steps on a running Amazon EMR cluster or have the pipeline create and manage an Amazon EMR cluster for you. For more information about Amazon EMR, see Getting started with Amazon EMR.

The EMR step requires that EMRStepConfig include the location of the JAR file to be used by the Amazon EMR cluster and any arguments to be passed. You also provide the Amazon EMR cluster ID if you want to run the step on a running EMR cluster, or the cluster configuration if you want the EMR step to run on a cluster that it creates, manages, and terminates for you. The following sections include examples and links to sample notebooks demonstrating both methods.

Note

- EMR steps require that the role passed to your pipeline has additional permissions. You should attach the AWS managed policy: AmazonSageMakerPipelinesIntegrations to your pipeline role, or ensure that the role includes the permissions in that policy.
- EMR step is not supported on EMR serverless, nor on Amazon EMR on EKS.
- If you process an EMR step on a running cluster, you can only use a cluster that is in one of the following states: STARTING, BOOTSTRAPPING, RUNNING, or WAITING.
- If you process EMR steps on a running cluster, you can have at most 256 EMR steps in a PENDING state on an EMR cluster. EMR steps submitted beyond this limit result in pipeline execution failure. You may consider using Retry Policy for Pipeline Steps.
- You can specify either cluster ID or cluster configuration, but not both.
- The EMR step relies on Amazon EventBridge to monitor changes in the EMR step or cluster state. If you process your Amazon EMR job on a running cluster, the EMR step uses the SageMakerPipelineExecutionEMRStepStatusUpdateRule rule to monitor EMR step state. If you process your job on a cluster that the EMR step creates for you, the step uses the SageMakerPipelineExecutionEMRClusterStatusRule rule to monitor changes in cluster state. If you see either of these EventBridge rules in your AWS account, do not delete them or else your EMR step may not complete.

Launch a new job on a running Amazon EMR cluster
If you want to launch a new job on a running Amazon EMR cluster, you pass the cluster ID as a string to the `cluster_id` argument of `EMRStep`. The following example demonstrates this procedure.

```python
from sagemaker.workflow.emr_step import EMRStep, EMRStepConfig

emr_config = EMRStepConfig(
    jar="jar-location",  # required, path to jar file used
    args=["--verbose", "--force"],  # optional list of arguments to pass to the jar
    main_class="com.my.Main1",  # optional main class, this can be omitted if jar above
    properties=[  # optional list of Java properties that are set when the step runs
        {  # "key": "mapred.tasktracker.map.tasks.maximum",
            "key": "mapred.tasktracker.map.tasks.maximum",
            "value": "2"
            },
        {  # "key": "mapreduce.map.sort.spill.percent",
            "key": "mapreduce.map.sort.spill.percent",
            "value": "0.90"
            },
        {  # "key": "mapreduce.tasktracker.reduce.tasks.maximum",
            "key": "mapreduce.tasktracker.reduce.tasks.maximum",
            "value": "5"
        }  # required
    ]
)

step_emr = EMRStep(
    name="EMRSampleStep",  # required
    cluster_id="j-1ABCDEFG2HIJK",  # include cluster_id to use a running cluster
    step_config=emr_config,  # required
    display_name="My EMR Step",
    description="Pipeline step to execute EMR job"
)
```

For a sample notebook that guides you through a complete example, see `SageMaker Pipelines EMR Step With Running EMR Cluster`.

**Launch a new job on a new Amazon EMR cluster**
If you want to launch a new job on a new cluster that EMRStep creates for you, provide your cluster configuration as a dictionary with the same structure as a `RunJobFlow` request. However, do not include the following fields in your cluster configuration:

- [Name]
- [Steps]
- [AutoTerminationPolicy]
- [Instances][KeepJobFlowAliveWhenNoSteps]
- [Instances][TerminationProtected]

All other `RunJobFlow` arguments are available for use in your cluster configuration. For details about the request syntax, see `RunJobFlow`.

The following example passes a cluster configuration to an EMR step definition, which prompts the step to launch a new job on a new EMR cluster. The EMR cluster configuration in this example includes specifications for primary and core EMR cluster nodes. For more information about Amazon EMR node types, see [Understand node types: primary, core, and task nodes](#).

```python
from sagemaker.workflow.emr_step import EMRStep, EMRStepConfig

emr_step_config = EMRStepConfig(
    jar="jar-location", # required, path to jar file used
    args=["--verbose", "--force"], # optional list of arguments to pass to the jar
    main_class="com.my.Main1", # optional main class, this can be omitted if jar above
    has a manifest
    properties=[ # optional list of Java properties that are set when the step runs
        
        "key": "mapred.tasktracker.map.tasks.maximum",
        "value": "2"
    ],
    
    "key": "mapreduce.map.sort.spill.percent",
    "value": "0.90"
],

"key": "mapreduce.tasktracker.reduce.tasks.maximum",
"value": "5"}
)```
```python
# include your cluster configuration as a dictionary
emr_cluster_config = {
    "Applications": [  
    
    "Name": "Spark",
    
    ],

    "Instances": [
    "InstanceGroups": [

    "InstanceRole": "MASTER",
    "InstanceCount": 1,
    "InstanceType": "m5.2xlarge"

    ],

    "InstanceRole": "CORE",
    "InstanceCount": 2,
    "InstanceType": "m5.2xlarge"

    ]

    },

    "BootstrapActions": [],

    "ReleaseLabel": "emr-6.6.0",

    "JobFlowRole": "job-flow-role",

    "ServiceRole": "service-role"

}

emr_step = EMRStep(
    name="emr-step",
    cluster_id=None,
    display_name="emr_step",
    description="MyEMRStepDescription",
    step_config=emr_step_config,
    cluster_config=emr_cluster_config
)
```

For a sample notebook that guides you through a complete example, see [SageMaker Pipelines EMR Step With Cluster Lifecycle Management](#).
# Notebook Job Step

Use a NotebookJobStep to run your SageMaker Notebook Job non-interactively as a pipeline step. For more information about SageMaker Notebook Jobs, see [SageMaker Notebook Jobs](#).

A NotebookJobStep requires at minimum an input notebook, image URI and kernel name. For more information about Notebook Job step requirements and other parameters you can set to customize your step, see [sagemaker.workflow.steps.NotebookJobStep](#).

The following example uses minimum arguments to define a NotebookJobStep.

```python
from sagemaker.workflow.notebook_job_step import NotebookJobStep

notebook_job_step = NotebookJobStep(
    input_notebook=input_notebook,
    image_uri=image_uri,
    kernel_name=kernel_name
)
```

Your NotebookJobStep pipeline step is treated as a SageMaker notebook job, so you can track the execution status in the Studio Classic UI notebook job dashboard if you include specific tags with the `tags` argument. For more details about tags to include, see [View your notebook jobs in the Studio UI dashboard](#).

Also, if you schedule your notebook job using the SageMaker Python SDK, you can only specify certain images to run your notebook job. For more information, see [Image constraints for SageMaker Python SDK notebook jobs](#).

## Fail Step

You use a FailStep to stop an Amazon SageMaker Model Building Pipelines execution when a desired condition or state is not achieved and to mark that pipeline's execution as failed. The FailStep also allows you to enter a custom error message, indicating the cause of the pipeline's execution failure.

> **Note**
> When a FailStep and other pipeline steps execute concurrently, the pipeline does not terminate until all concurrent steps are completed.
Limitations for using FailStep

- You cannot add a FailStep to the DependsOn list of other steps. For more information, see [Custom Dependency Between Steps](#).
- Other steps cannot reference the FailStep. It is always the last step in a pipeline's execution.
- You cannot retry a pipeline execution ending with a FailStep.

You can create the FailStep ErrorMessage in the form of a static text string. Alternatively, you can also use [Pipeline Parameters](#) a Join operation, or other step properties to create a more informative error message.

**Example**

The following example code snippet uses a FailStep with an ErrorMessage configured with Pipeline Parameters and a Join operation.

```python
from sagemaker.workflow.fail_step import FailStep
from sagemaker.workflow.functions import Join
from sagemaker.workflow.parameters import ParameterInteger

mse_threshold_param = ParameterInteger(name="MseThreshold", default_value=5)
step_fail = FailStep(
    name="AbaloneMSEFail",
    error_message=Join(
        on=" ", values=["Execution failed due to MSE >", mse_threshold_param]
    ),
)
```

**Step Properties**

The properties attribute is used to add data dependencies between steps in the pipeline. These data dependencies are then used by SageMaker Pipelines to construct the DAG from the pipeline definition. These properties can be referenced as placeholder values and are resolved at runtime.

The properties attribute of a SageMaker Pipelines step matches the object returned by a Describe call for the corresponding SageMaker job type. For each job type, the Describe call returns the following response object:

- ProcessingStep – [DescribeProcessingJob](#)
• TrainingStep – DescribeTrainingJob
• TransformStep – DescribeTransformJob

To check which properties are referrable for each step type during data dependency creation, see Data Dependency - Property Reference in the Amazon SageMaker Python SDK.

**Step Parallelism**

When a step does not depend on any other step, it is run immediately upon pipeline execution. However, executing too many pipeline steps in parallel can quickly exhaust available resources. Control the number of concurrent steps for a pipeline execution with ParallelismConfiguration.

The following example uses ParallelismConfiguration to set the concurrent step limit to five.

```python
pipeline.create(
    parallelism_config=ParallelismConfiguration(5),
)
```

**Data Dependency Between Steps**

You define the structure of your DAG by specifying the data relationships between steps. To create data dependencies between steps, pass the properties of one step as the input to another step in the pipeline. The step receiving the input isn't started until after the step providing the input finishes running.

A data dependency uses JsonPath notation in the following format. This format traverses the JSON property file, which means you can append as many `<property>` instances as needed to reach the desired nested property in the file. For more information on JsonPath notation, see the JsonPath repo.

```plaintext
<step_name>.properties.<property>.<property>
```

The following shows how to specify an Amazon S3 bucket using the ProcessingOutputConfig property of a processing step.

```python
step_process.properties.ProcessingOutputConfig.Outputs["train_data"].S3Output.S3Uri
```
To create the data dependency, pass the bucket to a training step as follows.

```python
from sagemaker.workflow.pipeline_context import PipelineSession

sklearn_train = SKLearn(..., sagemaker_session=PipelineSession())

step_train = TrainingStep(
    name="CensusTrain",
    step_args=sklearn_train.fit(inputs=TrainingInput(
        s3_data=step_process.properties.ProcessingOutputConfig.Outputs['train_data'].S3Output.S3Uri
    )))
)
```

To check which properties are referrable for each step type during data dependency creation, see Data Dependency - Property Reference in the Amazon SageMaker Python SDK.

### Custom Dependency Between Steps

When you specify a data dependency, SageMaker Pipelines provides the data connection between the steps. Alternatively, one step can access the data from a previous step without directly using SageMaker Pipelines. In this case, you can create a custom dependency that tells SageMaker Pipelines not to start a step until after another step has finished running. You create a custom dependency by specifying a step's DependsOn attribute.

As an example, the following defines a step C that starts only after both step A and step B finish running.

```json
{
    'Steps': [
        {'Name':'A', ...},
        {'Name':'B', ...},
        {'Name':'C', 'DependsOn': ['A', 'B']
    ]
}
```

SageMaker Pipelines throws a validation exception if the dependency would create a cyclic dependency.

The following example creates a training step that starts after a processing step finishes running.
processing_step = ProcessingStep(...)  
training_step = TrainingStep(...)  

training_step.add_depends_on([processing_step])

The following example creates a training step that doesn't start until two different processing steps finish running.

processing_step_1 = ProcessingStep(...)  
processing_step_2 = ProcessingStep(...)  

training_step = TrainingStep(...)  

training_step.add_depends_on([processing_step_1, processing_step_2])

The following provides an alternate way to create the custom dependency.

training_step.add_depends_on([processing_step_1])  
training_step.add_depends_on([processing_step_2])

The following example creates a training step that receives input from one processing step and waits for a different processing step to finish running.

processing_step_1 = ProcessingStep(...)  
processing_step_2 = ProcessingStep(...)  

training_step = TrainingStep(...)  

training_step.add_depends_on([processing_step_2])

The following example shows how to retrieve a string list of the custom dependencies of a step.

custom_dependencies = training_step.depends_on
Use a Custom Image in a Step

You can use any of the available SageMaker Deep Learning Container images when you create a step in your pipeline.

You can also use your own container with pipeline steps. Because you can’t create an image from within Amazon SageMaker Studio Classic, you must create your image using another method before using it with Amazon SageMaker Model Building Pipelines.

To use your own container when creating the steps for your pipeline, include the image URI in the estimator definition. For more information on using your own container with SageMaker, see Using Docker Containers with SageMaker.

Lift-and-shift Python code with the @step decorator

The @step decorator is a feature that converts your local machine learning (ML) code into one or more pipeline steps. You can write your ML function as you would for any ML project. Once tested locally or as a training job using the @remote decorator, you can convert the function to a SageMaker pipeline step by adding a @step decorator. You can then pass the output of the @step-decorated function call as a step to SageMaker Pipelines to create and run a pipeline. You can chain a series of functions with the @step decorator to create a multi-step directed acyclic graph (DAG) pipeline as well.

The setup to use the @step decorator is the same as the setup to use the @remote decorator. You can refer to the remote function documentation for details about how to setup the environment and use a configuration file to set defaults. For more information about the @step decorator, see sagemaker.workflow.function_step.step.

To view to sample notebooks that demonstrate the use of @step decorator, see @step decorator sample notebooks.

The following sections explain how you can annotate your local ML code with a @step decorator to create a step, create and run a pipeline using the step, and customize the experience for your use case.

Topics

- Create a pipeline with @step-decorated functions
- Run a pipeline
- Configure your pipeline
Create a pipeline with @step-decorated functions

You can create a pipeline by converting Python functions into pipeline steps using the @step decorator, creating dependencies between those functions to create a pipeline graph (or directed acyclic graph (DAG)), and passing the leaf nodes of that graph as a list of steps to the pipeline. The following sections explain this procedure in detail with examples.

Topics

- Convert a function to a step
- Create dependencies between the steps
- Use ConditionStep with @step-decorated steps
- Define a pipeline using the DelayedReturn output of steps
- Create a pipeline

Convert a function to a step

To create a step using the @step decorator, annotate the function with @step. The following example shows a @step-decorated function that preprocesses the data.

```python
from sagemaker.workflow.function_step import step

@step
def preprocess(raw_data):
    df = pandas.read_csv(raw_data)
    ...
    return processed_dataframe

step_process_result = preprocess(raw_data)
```

When you invoke a @step-decorated function, SageMaker returns a DelayedReturn instance instead of running the function. A DelayedReturn instance is a proxy for the actual return of that function. The DelayedReturn instance can be passed to another function as an argument or directly to a pipeline instance as a step. For information about the DelayedReturn class, see sagemaker.workflow.function_step.DelayedReturn.
Create dependencies between the steps

When you create a dependency between two steps, you create a connection between the steps in your pipeline graph. The following sections introduce multiple ways you can create a dependency between your pipeline steps.

Data dependencies through input arguments

Passing in the DelayedReturn output of one function as an input to another function automatically creates a data dependency in the pipeline DAG. In the following example, passing in the DelayedReturn output of the preprocess function to the train function creates a dependency between preprocess and train.

```python
from sagemaker.workflow.function_step import step

@step
def preprocess(raw_data):
    df = pandas.read_csv(raw_data)
    ...
    return processed_dataframe

@step
def train(training_data):
    ...
    return trained_model

step_process_result = preprocess(raw_data)
step_train_result = train(step_process_result)
```

The previous example defines a training function which is decorated with @step. When this function is invoked, it receives the DelayedReturn output of the preprocessing pipeline step as input. Invoking the training function returns another DelayedReturn instance. This instance holds the information about all the previous steps defined in that function (i.e, the preprocess step in this example) which form the pipeline DAG.

In the previous example, the preprocess function returns a single value. For more complex return types like lists or tuples, refer to Limitations.

Define custom dependencies

In the previous example, the train function received the DelayedReturn output of preprocess and created a dependency. If you want to define the dependency explicitly without passing
the previous step output, use the `add_depends_on` function with the step. You can use the `get_step()` function to retrieve the underlying step from its `DelayedReturn` instance, and then call `add_depends_on` on with the dependency as input. To view the `get_step()` function definition, see `sagemaker.workflow.step_outputs.get_step`. The following example shows you how to create a dependency between `preprocess` and `train` using `get_step()` and `add_depends_on()`.

```python
from sagemaker.workflow.step_outputs import get_step

@step
def preprocess(raw_data):
    df = pandas.read_csv(raw_data)
    ...
    processed_data = ..
    return s3.upload(processed_data)

@step
def train():
    training_data = s3.download(....)
    ...
    return trained_model

step_process_result = preprocess(raw_data)
step_train_result = train()

get_step(step_train_result).add_depends_on([step_process_result])
```

**Pass data to and from a @step-decorated function to a traditional pipeline step**

You can create a pipeline that includes a @step-decorated step and a traditional pipeline step and passes data between them. For example, you can use `ProcessingStep` to process the data and pass its result to the @step-decorated training function. In the following example, a @step-decorated training step references the output of a processing step.

```python
# Define processing step

from sagemaker.sklearn.processing import SKLearnProcessor
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker.workflow.steps import ProcessingStep

sklearn_processor = SKLearnProcessor(
    framework_version='1.2-1',
    ...)
```
role='arn:aws:iam::123456789012:role/SagemakerExecutionRole',
instance_type='ml.m5.large',
instance_count='1',
)

inputs = [
    ProcessingInput(source=input_data, destination="/opt/ml/processing/input"),
]
outputs = [
    ProcessingOutput(output_name="train", source="/opt/ml/processing/train"),
    ProcessingOutput(output_name="validation", source="/opt/ml/processing/validation"),
    ProcessingOutput(output_name="test", source="/opt/ml/processing/test")
]

process_step = ProcessingStep(
    name="MyProcessStep",
    step_args=skelearn_processor.run(inputs=inputs,
    outputs=outputs,code='preprocessing.py'),
)

# Define a @step-decorated train step which references the 
# output of a processing step

@step
def train(train_data_path, test_data_path):
    ...
    return trained_model

step_train_result = train(
    process_step.properties.ProcessingOutputConfig.Outputs["train"].S3Output.S3Uri,
    process_step.properties.ProcessingOutputConfig.Outputs["test"].S3Output.S3Uri,
)

Use ConditionStep with @step-decorated steps

SageMaker Pipelines supports a ConditionStep class which evaluates the results of preceding steps to decide what action to take in the pipeline. You can use ConditionStep with a @step-decorated step as well. To use the output of any @step-decorated step with ConditionStep, enter the output of that step as an argument to ConditionStep. In the following example, the condition step receives the output of the @step-decorated model evaluation step.

# Define steps
@step(name="evaluate")
def evaluate_model():
    # code to evaluate the model
    return {
        "rmse":rmse_value
    }

@step(name="register")
def register_model():
    # code to register the model
    ...

# Define ConditionStep
from sagemaker.workflow.condition_step import ConditionStep
from sagemaker.workflow.conditions import ConditionGreaterThanOrEqualTo
from sagemaker.workflow.fail_step import FailStep
conditionally_register = ConditionStep(
    name="conditional_register",
    conditions=[
        ConditionGreaterThanOrEqualTo(  # Output of the evaluate step must be json serializable
            left=evaluate_model()["rmse"],  #
            right=5,
        )
    ],
    if_steps=[FailStep(name="Fail", error_message="Model performance is not good enough")],
    else_steps=[register_model()],
)

Define a pipeline using the DelayedReturn output of steps

You define a pipeline the same way whether or not you use a @step decorator. When you pass a DelayedReturn instance to your pipeline, you don't need to pass a full list of steps to build the pipeline. The SDK automatically infers the previous steps based on the dependencies you define. All the previous steps of the Step objects you passed to the pipeline or DelayedReturn objects are included in the pipeline graph. In the following example, the pipeline receives the DelayedReturn object for the train function. SageMaker adds the preprocess step, as a previous step of train, to the pipeline graph.
from sagemaker.workflow.pipeline import Pipeline

pipeline = Pipeline(
    name="<pipeline-name>",
    steps=[step_train_result],
    sagemaker_session=<sagemaker-session>,
)

If there are no data or custom dependencies between the steps and you run multiple steps in parallel, the pipeline graph has more than one leaf node. Pass all of these leaf nodes in a list to the steps argument in your pipeline definition, as shown in the following example:

```python
@step
def process1():
    ...
    return data

@step
def process2():
    ...
    return data

step_process1_result = process1()
step_process2_result = process2()

pipeline = Pipeline(
    name="<pipeline-name>",
    steps=[step_process1_result, step_process2_result],
    sagemaker_session=sagemaker-session,
)
```

When the pipeline runs, both steps run in parallel.

You only pass the leaf nodes of the graph to the pipeline because the leaf nodes contain information about all the previous steps defined through data or custom dependencies. When it compiles the pipeline, SageMaker also infers of all of the subsequent steps that form the pipeline graph and adds each of them as a separate step to the pipeline.

Create a pipeline

Create a pipeline by calling `pipeline.create()`, as shown in the following snippet. For details about `create()`, see `sagemaker.workflow.pipeline.Pipeline.create`. 
role = "pipeline-role"
pipeline.create(role)

When you call pipeline.create(), SageMaker compiles all of the steps defined as part of the pipeline instance. SageMaker uploads the serialized function, arguments, and all the other step-related artifacts to Amazon S3.

Data resides in the S3 bucket according to the following structure:

```
s3_root_uri/
  pipeline_name/
    sm_rf_user_ws/
      workspace.zip  # archive of the current working directory (workdir)
    step_name/
      timestamp/
        arguments/          # serialized function arguments
        function/           # serialized function
        pre_train_dependencies/  # any dependencies and pre_execution scripts
      provided for the step
    execution_id/
    step_name/
      results  # returned output from the serialized function including
                 the model
```

`s3_root_uri` is defined in the SageMaker config file and applies to the entire pipeline. If undefined, the default SageMaker bucket is used.

⚠️ **Note**

Every time SageMaker compiles a pipeline, SageMaker saves the steps' serialized functions, arguments and dependencies in a folder timestamped with the current time. This occurs every time you run `pipeline.create()`, `pipeline.update()`, `pipeline.upsert()` or `pipeline.definition()`.

---

**Run a pipeline**

Start a new pipeline run with the `pipeline.start()` function as you would for a traditional SageMaker pipeline run. For information about the `start()` function, see [sagemaker.workflow.pipeline.Pipeline.start](https://sagemaker.readthedocs.io/en/stable/api/pipeline.html#sagemaker.workflow.pipeline.Pipeline.start).
Note
A step defined using the @step decorator runs as a training job. Therefore, be aware of the following limits:

- Instance limits and training job limits in your accounts. Update your limits accordingly to avoid any throttling or resource limit issues.
- The monetary costs associated with every run of a training step in the pipeline. For more details, refer to Amazon SageMaker Pricing.

Retrieve results from a pipeline run locally

To view the result of any step of a pipeline run, use `execution.result()`, as shown in the following snippet:

```python
execution = pipeline.start()
execution.result(step_name="train")
```

Note
SageMaker Pipelines does not support `execution.result()` in local mode.

You can only retrieve results for one step at a time. If the step name was generated by SageMaker, you can retrieve the step name by calling `list_steps` as follows:

```python
execution.list_step()
```

Run a pipeline locally

You can run a pipeline with @step-decorated steps locally as you would for traditional pipeline steps. For details about local mode pipeline runs, see Local Mode. To use local mode, provide a `LocalPipelineSession` instead of a `SageMakerSession` to your pipeline definition, as shown in the following example:

```python
from sagemaker.workflow.function_step import step
from sagemaker.workflow.pipeline import Pipeline
```
from sagemaker.workflow.pipeline_context import LocalPipelineSession

@step
def train():
    training_data = s3.download(....)
    ...
    return trained_model

step_train_result = train()

local_pipeline_session = LocalPipelineSession()

local_pipeline = Pipeline(
    name="<pipeline-name>",
    steps=[step_train_result],
    sagemaker_session=local_pipeline_session # needed for local mode
)

local_pipeline.create(role_arn="role_arn")

# pipeline runs locally
execution = local_pipeline.start()

Configure your pipeline

You are advised to use the SageMaker config file to set the defaults for the pipeline. For information about the SageMaker configuration file, see Configuring and using defaults with the SageMaker Python SDK. Any configuration added to the config file applies to all steps in the pipeline. If you want to override options for any of the steps, provide new values in the @step decorator arguments.

The @step decorator's configuration in the config file is identical to the @remote decorator's configuration. To set up the pipeline role ARN and pipeline tags in the config file, use the Pipeline section shown in the following snippet:

```
SchemaVersion: '1.0'
SageMaker:
    Pipeline:
        RoleArn: 'arn:aws:iam::555555555555:role/IMRole'
        Tags:
            - Key: 'tag_key'
              Value: 'tag_value'
```
For most of the defaults you can set in the configuration file you can also override by passing new values to the @step decorator. For example, you can override the instance type set in the config file for your preprocessing step, as shown in the following example:

```python
@step(instance_type="ml.m5.large")
def preprocess(raw_data):
    df = pandas.read_csv(raw_data)
    ...
    return processed_dataframe
```

A few arguments are not part of the @step decorator parameters list—these can be configured for the entire pipeline only through the SageMaker configuration file. They are listed as follows:

- **sagemaker_session (sagemaker.session.Session):** The underlying SageMaker session to which SageMaker delegates service calls. If unspecified, a session is created using a default configuration as follows:

  ```
  SageMaker:
  PythonSDK:
  Modules:
  Session:
  DefaultS3Bucket: 'default_s3_bucket'
  DefaultS3ObjectKeyPrefix: 'key_prefix'
  ```

- **custom_file_filter (CustomFileFilter):** A CustomFileFilter object that specifies the local directories and files to include in the pipeline step. If unspecified, this value defaults to None. For custom_file_filter to take effect, you must set IncludeLocalWorkdir to True. The following example shows a configuration that ignores all notebook files, and files and directories named data.

  ```
  SchemaVersion: '1.0'
  SageMaker:
  PythonSDK:
  Modules:
  RemoteFunction:
  IncludeLocalWorkDir: true
  CustomFileFilter:
  IgnoreNamePatterns: # files or directories to ignore
  - "*.ipynb" # all notebook files
  - "data" # folder or file named "data"
  ```
For more details about how to use IncludeLocalWorkdir with CustomFileFilter, see Using modular code with the @remote decorator.

- **s3_root_uri (str)**: The root Amazon S3 folder to which SageMaker uploads the code archives and data. If unspecified, the default SageMaker bucket is used.

- **s3_kms_key (str)**: The key used to encrypt the input and output data. You can only configure this argument in the SageMaker config file and the argument applies to all steps defined in the pipeline. If unspecified, the value defaults to None. See the following snippet for an example S3 KMS key configuration:

```plaintext
SchemaVersion: '1.0'
SageMaker:
  PythonSDK:
    Modules:
      RemoteFunction:
        S3KmsKeyId: 's3kmskeyid'
        S3RootUri: 's3://my-bucket/my-project'
```

**Best Practices**

The following sections suggest best practices to follow when you use the `@step` decorator for your pipeline steps.

**Use warm pools**

For faster pipeline step runs, use the warm pooling functionality provided for training jobs. You can turn on the warm pool functionality by providing the `keep_alive_period_in_seconds` argument to the `@step` decorator as demonstrated in the following snippet:

```python
@step(  
    keep_alive_period_in_seconds=900  
)
```

For more information about warm pools, see Train Using SageMaker Managed Warm Pools.

**Structure your directory**

You are advised to use code modules while using the `@step` decorator. Put the `pipeline.py` module, in which you invoke the step functions and define the pipeline, at the root of the workspace. The recommended structure is shown as follows:
Limitations

Be aware of the following limitations when you use the @step decorator for your pipeline steps.

**Function argument limitations**

When you pass an input argument to the @step-decorated function, the following limitations apply:

- You can pass the DelayedReturn, Properties (of steps of other types), Parameter, and ExecutionVariable objects to @step-decorated functions as arguments. But @step-decorated functions do not support JsonGet and Join objects as arguments.
- You cannot directly access a pipeline variable from a @step function. The following example produces an error:

```python
param = ParameterInteger(name="<parameter-name>", default_value=10)

@step
def func():
    print(param)

func() # this raises a SerializationError
```

- You cannot nest a pipeline variable in another object and pass it to a @step function. The following example produces an error:

```python
param = ParameterInteger(name="<parameter-name>", default_value=10)

@step
def func(arg):
    print(arg)
```
### Function imports

You should import the libraries required by the step inside rather than outside the function. If you import them at global scope, you risk an import collision while serializing the function. For example, `sklearn.pipeline.Pipeline` could be overridden by `sagemaker.workflow.pipeline.Pipeline`.

### Referencing child members of function return value

If you reference child members of a `@step`-decorated function's return value, the following limitations apply:

- You can reference the child members with `[]` if the `DelayedReturn` object represents a tuple, list or dict, as shown in the following example:

  ```python
delayed_return[0]
delayed_return["a_key"]
delayed_return[1]["a_key"]
```

- You cannot unpack a tuple or list output because the exact length of the underlying tuple or list can't be known when you invoke the function. The following example produces an error:

  ```python
  a, b, c = func() # this raises ValueError
  ```

- You cannot iterate over a `DelayedReturn` object. The following example raises an error:

  ```python
  for item in func(): # this raises a NotImplementedError
  ```
• You cannot reference arbitrary child members with '.'. The following example produces an error:

```python
delayed_return.a_child # raises AttributeError
```

### Existing pipeline features that are not supported

You cannot use the `@step` decorator with the following pipeline features:

- **Pipeline step caching**
- **Property files**

### Pass Data Between Steps

When you need to retrieve information from the output of a pipeline step, you can use `JsonGet`. `JsonGet` helps you extract information from Amazon S3 or property files. The following sections explain methods you can use to extract step outputs with `JsonGet`.

#### Pass data between steps with Amazon S3

You can use `JsonGet` in a `ConditionStep` to fetch the JSON output directly from Amazon S3. The Amazon S3 URI can be a `Std:Join` function containing primitive strings, pipeline run variables, or pipeline parameters. The following example shows how you can use `JsonGet` in a `ConditionStep`:

```python
# Example json file in s3 bucket generated by a processing_step
{
    "Output": [5, 10]
}

cond_lte = ConditionLessThanOrEqualTo(
    left=JsonGet(
        step_name="<step-name>",
        s3_uri="<s3-path-to-json>",
        json_path="Output[1]"
    ),
    right=6.0
)
```
If you are using JsonGet with an Amazon S3 path in the condition step, you must explicitly add a dependency between the condition step and the step generating the JSON output. In following example, the condition step is created with a dependency on the processing step:

```python
cond_step = ConditionStep(  
    name="<step-name>",  
    conditions=[cond_lte],  
    if_steps=[fail_step],  
    else_steps=[register_model_step],  
    depends_on=[processing_step],  
)
```

### Pass data between steps with property files

Use property files to store information from the output of a processing step. This is particularly useful when analyzing the results of a processing step to decide how a conditional step should be executed. The JsonGet function processes a property file and enables you to use JsonPath notation to query the property JSON file. For more information on JsonPath notation, see the JsonPath repo.

To store a property file for later use, you must first create a `PropertyFile` instance with the following format. The path parameter is the name of the JSON file to which the property file is saved. Any `output_name` must match the `output_name` of the `ProcessingOutput` that you define in your processing step. This enables the property file to capture the `ProcessingOutput` in the step.

```python
from sagemaker.workflow.properties import PropertyFile

<property_file_instance> = PropertyFile(  
    name="<property_file_name>",  
    output_name="<processingoutput_output_name>",  
    path="<path_to_json_file>"
)
```

When you create your `ProcessingStep` instance, add the `property_files` parameter to list all of the parameter files that the Amazon SageMaker Model Building Pipelines service must index. This saves the property file for later use.

```python
property_files=[<property_file_instance>]
```
To use your property file in a condition step, add the `property_file` to the condition that you pass to your condition step as shown in the following example to query the JSON file for your desired property using the `json_path` parameter.

```python
cond_lte = ConditionLessThanOrEqualTo(
    left=JsonGet(
        step_name=step_eval.name,
        property_file=<property_file_instance>,
        json_path="mse"
    ),
    right=6.0
)
```

For more in-depth examples, see `Property File` in the Amazon SageMaker Python SDK.

**Caching Pipeline Steps**

When you use step signature caching, SageMaker Pipelines tries to find a previous run of your current pipeline step with the same values for certain attributes. If found, SageMaker Pipelines propagates the outputs from the previous run rather than recomputing the step. The attributes checked are specific to the step type, and are listed in Default cache key attributes by pipeline step type.

You must opt in to step caching — it is off by default. When you turn on step caching, you must also define a timeout. This timeout defines how old a previous run can be to remain a candidate for reuse.

Step caching only considers successful runs — it never reuses failed runs. When multiple successful runs exist within the timeout period, SageMaker Pipelines uses the result for the most recent successful run. If no successful runs match in the timeout period, SageMaker Pipelines reruns the step. If the executor finds a previous run that meets the criteria but is still in progress, both steps continue running and update the cache if they're successful.

Step caching is only scoped for individual pipelines, so you can't reuse a step from another pipeline even if there is a step signature match.

Step caching is available for the following step types:

- **Processing**
- **Training**
• Tuning
• AutoML
• Transform
• ClarifyCheck
• QualityCheck
• EMR

Topics
• Turn on step caching
• Turn off step caching
• Default cache key attributes by pipeline step type
• Cached data access control

Turn on step caching

To turn on step caching, you must add a CacheConfig property to the step definition.

CacheConfig properties use the following format in the pipeline definition file:

```json
{
   "CacheConfig": {
      "Enabled": false,
      "ExpireAfter": "<time>"
   }
}
```

The Enabled field indicates whether caching is turned on for the particular step. You can set the field to true, which tells SageMaker to try to find a previous run of the step with the same attributes. Or, you can set the field to false, which tells SageMaker to run the step every time the pipeline runs. ExpireAfter is a string in ISO 8601 duration format that defines the timeout period. The ExpireAfter duration can be a year, month, week, day, hour, or minute value. Each value consists of a number followed by a letter indicating the unit of duration. For example:

• "30d" = 30 days
• "5y" = 5 years
• "T16m" = 16 minutes
• "30dT5h" = 30 days and 5 hours.

The following discussion describes the procedure to turn on caching for new or pre-existing pipelines using the Amazon SageMaker Python SDK.

**Turn on caching for new pipelines**

For new pipelines, initialize a `CacheConfig` instance with `enable_caching=True` and provide it as an input to your pipeline step. The following example turns on caching with a 1-hour timeout period for a training step:

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.workflow.steps import CacheConfig

cache_config = CacheConfig(enable_caching=True, expire_after="PT1H")
estimator = Estimator(..., sagemaker_session=PipelineSession())

step_train = TrainingStep(
    name="TrainAbaloneModel",
    step_args=estimator.fit(inputs=inputs),
    cache_config=cache_config
)
```

**Turn on caching for pre-existing pipelines**

To turn on caching for pre-existing, already-defined pipelines, turn on the `enable_caching` property for the step, and set `expire_after` to a timeout value. Lastly, update the pipeline with `pipeline.upsert()` or `pipeline.update()`. Once you run it again, the following code example turns on caching with a 1-hour timeout period for a training step:

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.workflow.steps import CacheConfig
from sagemaker.workflow.pipeline import Pipeline

cache_config = CacheConfig(enable_caching=True, expire_after="PT1H")
estimator = Estimator(..., sagemaker_session=PipelineSession())

step_train = TrainingStep(
    name="TrainAbaloneModel",
```
Alternatively, update the cache config after you have already defined the (pre-existing) pipeline, allowing one continuous code run. The following code sample demonstrates this method:

```python
# turn on caching with timeout period of one hour
pipeline.steps[0].cache_config.enable_caching = True
pipeline.steps[0].cache_config.expire_after = "PT1H"

# additional step for existing pipelines
pipeline.update()
# or, call upsert() to update the pipeline
# pipeline.upsert()
```

For more detailed code examples and a discussion about how Python SDK parameters affect caching, see [Caching Configuration](#) in the Amazon SageMaker Python SDK documentation.

### Turn off step caching

A pipeline step does not rerun if you change any attributes that are not listed in Default cache key attributes by pipeline step type for its step type. However, you may decide that you want the pipeline step to rerun anyway. In this case, you need to turn off step caching.

To turn off step caching, set the Enabled attribute in the step definition's CacheConfig property in the step definition to `false`, as shown in the following code snippet:

```json
{
    "CacheConfig": {
        "Enabled": false,
        "ExpireAfter": "<time>"
    }
}
Note that the `ExpireAfter` attribute is ignored when `Enabled` is false.

To turn off caching for a pipeline step using the Amazon SageMaker Python SDK, define the pipeline of your pipeline step, turn off the `enable_caching` property, and update the pipeline.

Once you run it again, the following code example turns off caching for a training step:

```python
from sagemaker.workflow.pipeline_context import PipelineSession
from sagemaker.workflow.steps import CacheConfig
from sagemaker.workflow.pipeline import Pipeline

cache_config = CacheConfig(enable_caching=False, expire_after="PT1H")
estimator = Estimator(..., sagemaker_session=PipelineSession())

step_train = TrainingStep(
    name="TrainAbaloneModel",
    step_args=estimator.fit(inputs=inputs),
    cache_config=cache_config
)

# define pipeline
pipeline = Pipeline(
    steps=[step_train]
)

# update the pipeline
pipeline.update()
# or, call upsert() to update the pipeline
# pipeline.upsert()
```

Alternatively, turn off the `enable_caching` property after you have already defined the pipeline, allowing one continuous code run. The following code sample demonstrates this solution:

```python
# turn off caching for the training step
pipeline.steps[0].cache_config.enable_caching = False

# update the pipeline
pipeline.update()
# or, call upsert() to update the pipeline
```
For more detailed code examples and a discussion about how Python SDK parameters affect caching, see [Caching Configuration](#) in the Amazon SageMaker Python SDK documentation.

**Default cache key attributes by pipeline step type**

When deciding whether to reuse a previous pipeline step or rerun the step, SageMaker Pipelines checks to see if certain attributes have changed. If the set of attributes is different from all previous runs within the timeout period, the step runs again. These attributes include input artifacts, app or algorithm specification, and environment variables.

The following list shows each pipeline step type and the attributes that, if changed, initiate a rerun of the step. For more information about which Python SDK parameters are used to create the following attributes, see [Caching Configuration](#) in the Amazon SageMaker Python SDK documentation.

**Processing step**

- AppSpecification
- Environment
- ProcessingInputs. This attribute contains information about the preprocessing script.

**Training step**

- AlgorithmSpecification
- CheckpointConfig
- DebugHookConfig
- DebugRuleConfigurations
- Environment
- HyperParameters
- InputDataConfig. This attribute contains information about the training script.

**Tuning step**

- HyperParameterTuningJobConfig
• TrainingJobDefinition. This attribute is composed of multiple child attributes, not all of which cause the step to rerun. The child attributes that could incur a rerun (if changed) are:
  • AlgorithmSpecification
  • HyperParameterRanges
  • InputDataConfig
  • StaticHyperParameters
  • TuningObjective
  • TrainingJobDefinitions

**AutoML step**

• AutoMLJobConfig. This attribute is composed of multiple child attributes, not all of which cause the step to rerun. The child attributes that could incur a rerun (if changed) are:
  • CompletionCriteria
  • CandidateGenerationConfig
  • DataSplitConfig
  • Mode
  • AutoMLJobObjective
  • InputDataConfig
  • ProblemType

**Transform step**

• DataProcessing
• Environment
• ModelName
• TransformInput

**ClarifyCheck Step**

• ClarifyCheckConfig
• CheckJobConfig
• SkipCheck
• RegisterNewBaseline
• ModelPackageGroupName
• SuppliedBaselineConstraints

**QualityCheck Step**

• QualityCheckConfig
• CheckJobConfig
• SkipCheck
• RegisterNewBaseline
• ModelPackageGroupName
• SuppliedBaselineConstraints
• SuppliedBaselineStatistics

**EMR Step**

• ClusterId
• StepConfig

**Cached data access control**

When a SageMaker pipeline runs, it caches the parameters and metadata associated with the SageMaker jobs launched by the pipeline and saves them for reuse in subsequent runs. This metadata is accessible through a variety of sources in addition to cached pipeline steps, and includes the following types:

• Describe*Job requests
• CloudWatch Logs
• CloudWatch Events
Note that access to each data source in the list is controlled by its own set of IAM permissions. Removing a particular role's access to one data source does not affect the level of access to the others. For example, an account admin might remove IAM permissions for `Describe*Job` requests from a caller's role. While the caller can no longer make `Describe*Job` requests, they can still retrieve the metadata from a pipeline run with cached steps as long as they have permission to run the pipeline. If an account admin wants to remove access to the metadata from a particular SageMaker job completely, they need to remove permissions for each of the relevant services that provide access to the data.

**Retry Policy for Pipeline Steps**

Retry policies help you automatically retry your SageMaker Pipelines steps after an error occurs. Any pipeline step can encounter exceptions, and exceptions happen for various reasons. In some cases, a retry can resolve these issues. With a retry policy for pipeline steps, you can choose whether to retry a particular pipeline step or not.

The retry policy only supports the following pipeline steps:

- Processing Step
- Training Step
- Tuning Step
- AutoML Step
- CreateModel Step
- RegisterModel Step
- Transform Step
- Notebook Job Step

**Note**

Jobs running inside both the tuning and AutoML steps conduct retries internally and will not retry the `SageMaker.JOB_INTERNAL_ERROR` exception type, even if a retry policy is configured. You can program your own [Retry Strategy](#) using the SageMaker API.
Supported exception types for the retry policy

The retry policy for pipeline steps supports the following exception types:

- **Step.SERVICE_FAULT**: These exceptions occur when an internal server error or transient error happens when calling downstream services. SageMaker Pipelines retries on this type of error automatically. With a retry policy, you can override the default retry operation for this exception type.

- **Step.THROTTLING**: Throttling exceptions can occur while calling the downstream services. SageMaker Pipelines retries on this type of error automatically. With a retry policy, you can override the default retry operation for this exception type.

- **SageMaker.JOB_INTERNAL_ERROR**: These exceptions occur when the SageMaker job returns `InternalServerError`. In this case, starting a new job may fix a transient issue.

- **SageMaker.CAPACITY_ERROR**: The SageMaker job may encounter Amazon EC2 `InsufficientCapacityErrors`, which leads to the SageMaker job’s failure. You can retry by starting a new SageMaker job to avoid the issue.

- **SageMaker.RESOURCE_LIMIT**: You can exceed the resource limit quota when running a SageMaker job. You can wait and retry running the SageMaker job after a short period and see if resources are released.

The JSON schema for the retry policy

The retry policy for Pipelines has the following JSON schema:

```json
"RetryPolicy": {
  "ExceptionType": [String]
  "IntervalSeconds": Integer
  "BackoffRate": Double
  "MaxAttempts": Integer
  "ExpireAfterMin": Integer
}
```

- **ExceptionType**: This field requires the following exception types in a string array format.
  - **Step.SERVICE_FAULT**
  - **Step.THROTTLING**
  - **SageMaker.JOB_INTERNAL_ERROR**
• SageMaker.CAPACITY_ERROR
• SageMaker.RESOURCE_LIMIT

• IntervalSeconds (optional): The number of seconds before the first retry attempt (1 by default). IntervalSeconds has a maximum value of 43200 seconds (12 hours).

• BackoffRate (optional): The multiplier by which the retry interval increases during each attempt (2.0 by default).

• MaxAttempts (optional): A positive integer that represents the maximum number of retry attempts (5 by default). If the error recurs more times than MaxAttempts specifies, retries cease and normal error handling resumes. A value of 0 specifies that errors are never retried. MaxAttempts has a maximum value of 20.

• ExpireAfterMin (optional): A positive integer that represents the maximum timespan of retry. If the error recurs after ExpireAfterMin minutes counting from the step gets executed, retries cease and normal error handling resumes. A value of 0 specifies that errors are never retried. ExpireAfterMin has a maximum value of 14,400 minutes (10 days).

**Note**

Only one of MaxAttempts or ExpireAfterMin can be given, but not both; if both are not specified, MaxAttempts becomes the default. If both properties are identified within one policy, then the retry policy generates a validation error.

### Configuring a retry policy

The following is an example of a training step with a retry policy.

```json
{
  "Steps": [
    {
      "Name": "MyTrainingStep",
      "Type": "Training",
      "RetryPolicies": [
        {
          "ExceptionType": [
            "SageMaker.JOB_INTERNAL_ERROR",
            "SageMaker.CAPACITY_ERROR"
          ],
          "IntervalSeconds": 1,
```
The following is an example of how to build a `TrainingStep` in SDK for Python (Boto3) with a retry policy.

```python
from sagemaker.workflow.retry import (StepRetryPolicy,
StepExceptionTypeEnum,
SageMakerJobExceptionTypeEnum,
SageMakerJobStepRetryPolicy
)

step_train = TrainingStep(
    name="MyTrainingStep",
    xxx,
    retry_policies=[
        // override the default
        StepRetryPolicy(
            exception_types=[
                StepExceptionTypeEnum.SERVICE_FAULT,
                StepExceptionTypeEnum.THROTTLING
            ],
            expire_after_mins=5,
            interval_seconds=10,
            backoff_rate=2.0
        ),
        // retry when resource limit quota gets exceeded
        SageMakerJobStepRetryPolicy(
            exception_types=[SageMakerJobExceptionTypeEnum.RESOURCE_LIMIT],
            expire_after_mins=120,
            interval_seconds=60,
            backoff_rate=2.0
        ),
        // retry when job failed due to transient error or EC2 ICE.
        SageMakerJobStepRetryPolicy(
            failure_reason_types=[SageMakerJobExceptionTypeEnum.INTERNAL_ERROR,
```
SageMakerJobExceptionTypeEnum.CAPACITY_ERROR,
    ],
    max_attempts=10,
    interval_seconds=30,
    backoff_rate=2.0
  )
)
)

For more information on configuring retry behavior for certain step types, see Amazon SageMaker Model Building Pipelines - Retry Policy in the Amazon SageMaker Python SDK documentation.

Selective execution of pipeline steps

As you use SageMaker Pipelines to create workflows and orchestrate your ML training steps, you might need to undertake multiple experimentation phases. Instead of running the entire pipeline from start to finish, you might only want to iterate over particular steps. SageMaker Pipelines supports selective execution of pipeline steps to help you optimize your ML training. Selective execution is useful in the following scenarios:

- You want to restart a specific step with updated instance type, hyperparameters, or other variables while keeping the parameters from upstream steps.
- Your pipeline fails an intermediate step. Previous steps in the execution, such as data preparation or feature extraction, are expensive to rerun. You might need to introduce a fix and rerun certain steps manually to complete the pipeline.

Using selective execution, you can choose to run any subset of steps as long as they are connected in the directed acyclic graph (DAG) of your pipeline. The following DAG shows an example pipeline workflow:
You can select steps AbaloneTrain and AbaloneEval in a selective execution, but you cannot select just AbaloneTrain and AbaloneMSECond steps to run a selective execution because these steps are not connected in the DAG. For non-selected steps in the workflow, the selective execution reuses the outputs from a reference pipeline execution rather than recomputing the steps. Also, non-selected steps that are downstream from the selected steps do not run in a selective execution.

If you choose to run a subset of intermediate steps in your pipeline, your steps may have dependencies on upstream steps. SageMaker needs a reference pipeline execution from which to resource these dependencies. For example, if you choose to run the steps AbaloneTrain and AbaloneEval, you need the output collaterals for the AbaloneProcess step from a reference pipeline execution. You can either provide a reference execution ARN or direct SageMaker to use
the latest pipeline execution, which is the default behavior. If you have a reference execution, you can also build the runtime parameters from your reference run and supply them to your selective executive run with any overrides. For details, see Reuse runtime parameter values from a reference execution.

In detail, you specify a configuration for your selective execution pipeline run using SelectiveExecutionConfig. If you specify an ARN for a reference pipeline execution (with the source_pipeline_execution_arn argument), SageMaker uses the upstream step dependencies from the specified pipeline execution. If you do not specify an ARN and a latest pipeline execution exists, SageMaker uses the latest pipeline execution as a reference by default. If you do not specify an ARN and do not want SageMaker to use your latest pipeline execution, set reference_latest_execution to False. The pipeline execution which SageMaker ultimately uses as a reference, whether the latest or user-specified, must be in Success or Failed state.

The following table summarizes how SageMaker chooses a reference execution based on your arguments to SelectiveExecutionConfig.

<table>
<thead>
<tr>
<th>The source_pipeline_execution_arn argument value</th>
<th>The reference_latest_execution argument value</th>
<th>The reference execution used</th>
</tr>
</thead>
<tbody>
<tr>
<td>A pipeline ARN</td>
<td>True or unspecified</td>
<td>The specified pipeline ARN</td>
</tr>
<tr>
<td>A pipeline ARN</td>
<td>False</td>
<td>The specified pipeline ARN</td>
</tr>
<tr>
<td>null or unspecified</td>
<td>True or unspecified</td>
<td>The latest pipeline execution</td>
</tr>
<tr>
<td>null or unspecified</td>
<td>False</td>
<td>None—in this case, select steps without upstream dependencies</td>
</tr>
</tbody>
</table>

For more information about selective execution configuration requirements, see the sagemaker.workflow.selective_execution_config.SelectiveExecutionConfig documentation.
The following discussion includes examples for the cases in which you want to specify a pipeline reference execution, use the latest pipeline execution as a reference, or run selective execution without a reference pipeline execution.

**Selective execution with a user-specified pipeline reference**

The following example demonstrates the use of selective execution to rerun AbaloneTrain and AbaloneEval in the same pipeline rerun using a reference pipeline execution.

```python
from sagemaker.workflow.selective_execution_config import SelectiveExecutionConfig

selective_execution_config = SelectiveExecutionConfig(
    abalone/execution/123ab12cd3ef",
    selected_steps=["AbaloneTrain", "AbaloneEval"]
)

selective_execution = pipeline.start(
    execution_display_name=f"Sample-Selective-Execution-1",
    parameters={"MaxDepth":6, "NumRound":60},
    selective_execution_config=selective_execution_config,
)
```

**Selective execution with the latest pipeline execution as a reference**

The following example demonstrates the use of selective execution to rerun AbaloneTrain and AbaloneEval in the same pipeline rerun using the latest pipeline execution as a reference. Since SageMaker uses the latest pipeline execution by default, you can optionally set the reference_latest_execution argument to True.

```python
# Prepare a new selective execution. Select only the first step in the pipeline without
# providing source_pipeline_execution_arn.
selective_execution_config = SelectiveExecutionConfig(
    selected_steps=["AbaloneTrain", "AbaloneEval"],
    # optional
    reference_latest_execution=True
)

# Start pipeline execution without source_pipeline_execution_arn
pipeline.start(
    execution_display_name=f"Sample-Selective-Execution-1",
)
Selective execution without a reference pipeline

The following example demonstrates the use of selective execution to rerun AbaloneProcess and AbaloneTrain in the same pipeline rerun without providing a reference ARN and disallowing the use of the latest pipeline run as a reference. SageMaker allows this configuration since this subset of steps doesn't have upstream dependencies.

```python
# Prepare a new selective execution. Select only the first step in the pipeline without providing source_pipeline_execution_arn.
selective_execution_config = SelectiveExecutionConfig(
    selected_steps=['AbaloneProcess', 'AbaloneTrain'],
    reference_latest_execution=False
)

# Start pipeline execution without source_pipeline_execution_arn
pipeline.start(
    execution_display_name=f'Sample-Selective-Execution-1',
    parameters={'MaxDepth': 6, 'NumRound': 60},
    selective_execution_config=selective_execution_config,
)
```

Reuse runtime parameter values from a reference execution

You can build the parameters from your reference pipeline execution using `build_parameters_from_execution`, and supply the result to your selective execution pipeline. You can use the original parameters from the reference execution, or apply any overrides using the `parameter_value_overrides` argument.

The following example shows you how to build parameters from a reference execution and apply an override for the MseThreshold parameter.

```python
# Prepare a new selective execution.
selective_execution_config = SelectiveExecutionConfig(
    selected_steps=['AbaloneTrain', 'AbaloneEval', 'AbaloneMSECond'],
)
```
# Define a new parameters list to test.
new_parameters_mse={
    "MseThreshold": 5,
}

# Build parameters from reference execution and override with new parameters to test.
new_parameters = pipeline.build_parameters_from_execution(  
    parameter_value_overrides=new_parameters_mse
)

# Start pipeline execution with new parameters.
execution = pipeline.start(  
    selective_execution_config=selective_execution_config,  
    parameters=new_parameters
)

Baseline calculation, drift detection and lifecycle with ClarifyCheck and QualityCheck steps in Amazon SageMaker Model Building Pipelines

The following topic discusses how baselines and model versions evolve in the Amazon SageMaker Model Building Pipelines when using the ClarifyCheck and QualityCheck steps.

For the ClarifyCheck step, a baseline is a single file that resides in the step properties with the suffix constraints. For the QualityCheck step, a baseline is a combination of two files that resides in the step properties: one with the suffix statistics and the other with the suffix constraints. In the following topics we discuss these properties with a prefix that describes how they are used, impacting baseline behavior and lifecycle in these two pipeline steps. For example, the ClarifyCheck step always calculates and assigns the new baselines in the CalculatedBaselineConstraints property and the QualityCheck step does the same in the CalculatedBaselineConstraints and CalculatedBaselineStatistics properties.

Baseline calculation and registration for ClarifyCheck and QualityCheck steps

Both the ClarifyCheck and QualityCheck steps always calculate new baselines based on step inputs through the underlying processing job run. These newly calculated baselines are accessed through the properties with the prefix CalculatedBaseline. You can record these properties as the ModelMetrics of your model package in the Model Step. This model package can be registered with 5 different baselines. You can register it with one for each check type: data bias, model bias, and model explainability from running the ClarifyCheck step and model
quality, and data quality from running the QualityCheck step. The `register_new_baseline` parameter dictates the value set in the properties with the prefix `BaselineUsedForDriftCheck` after a step runs.

The following table of potential use cases shows different behaviors resulting from the step parameters you can set for the ClarifyCheck and QualityCheck steps:

<table>
<thead>
<tr>
<th>Possible use case that you may consider for selecting this configuration</th>
<th>skip_check/register_new_baseline</th>
<th>Does step do a drift check?</th>
<th>Value of step property <code>CalculateBaseline</code></th>
<th>Value of step property <code>BaselineUsedForDriftCheck</code></th>
</tr>
</thead>
<tbody>
<tr>
<td>You are doing regular retraining with checks enabled to get a new model version, but you want to carry over the previous baselines as the DriftCheck baseline in the model registry for your new model version.</td>
<td>False/False</td>
<td>Drift check runs against existing baselines</td>
<td>New baselines calculated by running the step</td>
<td>Baseline from the latest approved model in Model Registry or the baseline supplied as step parameter</td>
</tr>
<tr>
<td>You are doing regular retraining with checks enabled to get a new model version, but you want to refresh</td>
<td>False/True</td>
<td>Drift check runs against existing baselines</td>
<td>New baselines calculated by running the step</td>
<td>Newly calculated baseline by running the step (value of property <code>CalculateBaseline</code>)</td>
</tr>
</tbody>
</table>
## Possible use case that you may consider for selecting this configuration

<table>
<thead>
<tr>
<th>skip_check/register_new_baseline</th>
<th>Does step do a drift check?</th>
<th>Value of step property CalculatedBaseline</th>
<th>Value of step property BaselineUsedForDriftCheck</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>the DriftCheckBaseline s in the model registry with the newly calculated baselines for your new model version.</code></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Possible use case that you may consider for selecting this configuration</td>
<td>skip_check / register_new_baseline</td>
<td>Does step do a drift check?</td>
<td>Value of step property CalculateBaseline</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>You are initiating the pipeline to retrain a new model version because there is a violation detected by Amazon SageMaker Model Monitor on an endpoint for a particular type of check, and you want to <em>skip this type of check against the previous baseline, but carry over the previous baseline as DriftCheckBaseline</em> in the model registry for your new model version.</td>
<td>True/ False</td>
<td>No drift check</td>
<td>New baselines calculated by running</td>
</tr>
<tr>
<td>Possible use case that you may consider for selecting this configuration</td>
<td>skip_check/check_new_basel ine</td>
<td>Does step do a drift check?</td>
<td>Value of step property CalculateBaseline</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>This happens in the following cases:</td>
<td>True/True</td>
<td>No drift check</td>
<td>New baselines calculated by running the step</td>
</tr>
<tr>
<td>• You are starting the initial run of the pipeline, building your first model version, and generating the initial baselines.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• You are initiating the pipeline to retrain a new model version because there is a violation detected by Model Monitor on the endpoint for a particular type of check. If you want to skip the</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Possible use case that you may consider for selecting this configuration

<table>
<thead>
<tr>
<th>skip_check / register_new_baseline</th>
<th>Does step do a drift check?</th>
<th>Value of step property CalculateBaseline</th>
<th>Value of step property BaselineUsedForDriftCheck</th>
</tr>
</thead>
<tbody>
<tr>
<td>check against the previous baseline and refresh the DriftCheckBaselines with the newly calculated baseline in the model registry directly.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Note

If you use scientific notation in your constraint, you need to convert to float. For a preprocessing script example of how to do this, see [Create a Model Quality Baseline](#).

When you register a model with **Model Step**, you can register the BaselineUsedForDriftCheck property as DriftCheckBaselines. These baseline files can then be used by Model Monitor for model and data quality checks. In addition, these baselines can also be used in the ClarifyCheckStep and QualityCheck step to compare newly trained models against the existing models that are registered in the model registry for future pipeline runs.

## Drift Detection against Previous Baselines in SageMaker Pipelines

In the case of the QualityCheck step, when you initiate the pipeline for regular retraining to get a new model version, you may not want to run the training step if the data quality and the data bias has [Schema for Violations](constraintViolations.json file) on the baselines of your previous
approved model version. You also may not want to register the newly trained model version if the model quality, model bias, or model explainability violates the registered baseline of your previous approved model version when running the ClarifyCheck step. In these cases, you can enable the checks you want by setting the skip_check property of the corresponding check step set to False, resulting in the ClarifyCheck and QualityCheck step failing if violation is detected against previous baselines. The pipeline process then does not proceed so that the model drifted from the baseline isn't registered. ClarifyCheck and QualityCheck steps are able to get DriftCheckBaselines of the latest approved model version of a given model package group against which to compare. Previous baselines can also be supplied directly through supplied_baseline_constraints (in addition to supplied_baseline_statistics if it is a QualityCheck step) and are always prioritized over any baselines pulled from the model package group.

Baseline and model version lifecycle and evolution with SageMaker Pipelines

By setting register_new_baseline of your ClarifyCheck and QualityCheck step to False, your previous baseline is accessible through the step property prefix BaselineUsedForDriftCheck. You can then register these baselines as the DriftCheckBaselines in the new model version when you register a model with Model Step. Once you approve this new model version in the model registry, the DriftCheckBaseline in this model version becomes available for the ClarifyCheck and QualityCheck steps in the next pipeline process. If you want to refresh the baseline of a certain check type for future model versions, you can set register_new_baseline to True so that the properties with prefix BaselineUsedForDriftCheck become the newly calculated baseline. In these ways, you can preserve your preferred baselines for a model trained in the future, or refresh the baselines for drift checks when needed, managing your baseline evolution and lifecycle throughout your model training iterations.

The following diagram illustrates a model-version-centric view of the baseline evolution and lifecycle.
You can schedule your Amazon SageMaker Model Building Pipelines executions using Amazon EventBridge. Amazon SageMaker Model Building Pipelines is supported as a target in Amazon EventBridge. This allows you to initiate the execution of your model building pipeline based on any event in your event bus. With EventBridge, you can automate your pipeline executions and respond automatically to events such as training job or endpoint status changes. Events include a new file being uploaded to your Amazon S3 bucket, a change in status of your Amazon SageMaker endpoint due to drift, and Amazon Simple Notification Service (SNS) topics.

The following SageMaker Pipelines actions can be automatically initiated:

- **StartPipelineExecution**
For more information on scheduling SageMaker jobs, see Automating SageMaker with Amazon EventBridge.

Topics
- Schedule a Pipeline with Amazon EventBridge
- Schedule a pipeline with the SageMaker Python SDK

Schedule a Pipeline with Amazon EventBridge

To start a pipeline execution with Amazon CloudWatch Events, you must create an EventBridge rule. When you create a rule for events, you specify a target action to take when EventBridge receives an event that matches the rule. When an event matches the rule, EventBridge sends the event to the specified target and initiates the action defined in the rule.

The following tutorials show how to schedule a pipeline execution with EventBridge using the EventBridge console or the AWS CLI.

Prerequisites

- A role that EventBridge can assume with the SageMaker::StartPipelineExecution permission. This role can be created automatically if you create a rule from the EventBridge console; otherwise, you need to create this role yourself. For information on creating a SageMaker role, see SageMaker Roles.
- An Amazon SageMaker Pipeline to schedule. To create an Amazon SageMaker Pipeline, see Define a Pipeline.

Create an EventBridge rule using the EventBridge console

The following procedure shows how to create an EventBridge rule using the EventBridge console.

1. Navigate to the EventBridge console.
2. Select Rules on the left hand side.
3. Select Create Rule.
4. Enter a name and description for your rule.
5. Select how you want to initiate this rule. You have the following choices for your rule:
   - Event pattern: Your rule is initiated when an event matching the pattern occurs. You can choose a predefined pattern that matches a certain type of event, or you can create a
custom pattern. If you select a predefined pattern, you can edit the pattern to customize it. For more information on Event patterns, see Event Patterns in CloudWatch Events.

- **Schedule**: Your rule is initiated regularly on a specified schedule. You can use a fixed-rate schedule that initiates regularly for a specified number of minutes, hour, or weeks. You can also use a cron expression to create a more fine-grained schedule, such as “the first Monday of each month at 8am.” Schedule is not supported on a custom or partner event bus.

6. Select your desired Event bus.

7. Select the target(s) to invoke when an event matches your event pattern or when the schedule is initiated. You can add up to 5 targets per rule. Select SageMaker Pipeline in the target dropdown list.

8. Select the pipeline you want to initiate from the pipeline dropdown list.

9. Add parameters to pass to your pipeline execution using a name and value pair. Parameter values can be static or dynamic. For more information on Amazon SageMaker Pipeline parameters, see AWS::Events::Rule SagemakerPipelineParameters.

   - Static values are passed to the pipeline execution every time the pipeline is initiated. For example, if{"Name": "Instance_type", "Value": "ml.4xlarge"} is specified in the parameter list, then it is passed as a parameter in StartPipelineExecutionRequest every time EventBridge initiates the pipeline.
   - Dynamic values are specified using a JSON path. EventBridge parses the value from an event payload, then passes it to the pipeline execution. For example: $.detail.param.value

10. Select the role to use for this rule. You can either use an existing role or create a new one.

11. (Optional) Add tags.

12. Select Create to finalize your rule.

Your rule is now in effect and ready to initiate your pipeline executions.

**Create an EventBridge rule using the AWS CLI**

The following procedure shows how to create an EventBridge rule using the AWS CLI.

1. Create a rule to be initiated. When creating an EventBridge rule using the AWS CLI, you have two options for how your rule is initiated, event pattern and schedule.

   - **Event pattern**: Your rule is initiated when an event matching the pattern occurs. You can choose a predefined pattern that matches a certain type of event, or you can create a
custom pattern. If you select a predefined pattern, you can edit the pattern to customize it. You can create a rule with event pattern using the following command:

```
aws events put-rule --name <RULE_NAME> --event-pattern <YOUR_EVENT_PATTERN> --description <RULE_DESCRIPTION> --role-arn <ROLE_TO_EXECUTE_PIPELINE> --tags <TAGS>
```

- **Schedule**: Your rule is initiated regularly on a specified schedule. You can use a fixed-rate schedule that initiates regularly for a specified number of minutes, hour, or weeks. You can also use a cron expression to create a more fine-grained schedule, such as “the first Monday of each month at 8am.” Schedule is not supported on a custom or partner event bus. You can create a rule with schedule using the following command:

```
aws events put-rule --name <RULE_NAME> --schedule-expression <YOUR_CRON_EXPRESSION> --description <RULE_DESCRIPTION> --role-arn <ROLE_TO_EXECUTE_PIPELINE> --tags <TAGS>
```

2. Add target(s) to invoke when an event matches your event pattern or when the schedule is initiated. You can add up to 5 targets per rule. For each target, you must specify:

- ARN: The resource ARN of your pipeline.
- Role ARN: The ARN of the role EventBridge should assume to execute the pipeline.
- Parameters: Amazon SageMaker pipeline parameters to pass.

3. Run the following command to pass a Amazon SageMaker pipeline as a target to your rule using `put-targets`:

```
aws events put-targets --rule <RULE_NAME> --event-bus-name <EVENT_BUS_NAME> --targets "{\"Id\": <ID>, \"Arn\": <RESOURCE_ARN>, \"RoleArn\": <ROLE_ARN>, \"SageMakerPipelineParameter\": { \"Name\": <NAME>, \"Value\": <VALUE>}}
```

**Schedule a pipeline with the SageMaker Python SDK**

The following sections show you how to set up permissions to access EventBridge resources and create your pipeline schedule using the SageMaker Python SDK.
Required permissions

You need to have necessary permissions to use the pipeline scheduler. Complete the following steps to set up your permissions:

1. Attach the following minimum privilege policy to the IAM role used to create the pipeline triggers, or use the AWS managed policy AmazonEventBridgeSchedulerFullAccess.

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Action": ["scheduler:ListSchedules",
                  "scheduler:GetSchedule",
                  "scheduler:CreateSchedule",
                  "scheduler:UpdateSchedule",
                  "scheduler:DeleteSchedule"],
      "Effect": "Allow",
      "Resource": ["*"]
    },
    {
      "Effect": "Allow",
      "Action": "iam:PassRole",
      "Resource": "arn:aws:iam::*:role/*",
      "Condition": {
        "StringLike": {
          "iam:PassedToService": "scheduler.amazonaws.com"
        }
      }
    }
  ]
}
```

2. Establish a trust relationship with EventBridge by adding the service principal scheduler.amazonaws.com to this role’s trust policy. Make sure you attach the following trust policy to the execution role if you launch the notebook in SageMaker Studio.
Create a pipeline schedule

Using the PipelineSchedule constructor, you can schedule a pipeline to run once or at a predetermined interval. A pipeline schedule must be of the type at, rate, or cron. This set of scheduling types is an extension of the EventBridge scheduling options. For more information about how to use the PipelineSchedule class, see sagemaker.workflow.triggers.PipelineSchedule. The following example demonstrates how to create each scheduling type with PipelineSchedule.

```python
from sagemaker.workflow.triggers import PipelineSchedule

# schedules a pipeline run for 12/13/2023 at time 10:15:20 UTC
my_datetime_schedule = PipelineSchedule(
    name="<schedule-name>",
    at=datetime(2023, 12, 13, 10, 15, 20)
)

# schedules a pipeline run every 5 minutes
my_rate_schedule = PipelineSchedule(
    name="<schedule-name>",
    rate=(5, "minutes")
)

# schedules a pipeline run at 10:15am UTC on the last Friday of each month during the years 2022 to 2023
my_cron_schedule = PipelineSchedule(
    name="<schedule-name>",
    rate="rate="cron",
    schedule=("day_of_month"="L", "day_of_week"="F")
)
```
name="<schedule-name>",
cron="15 10 ? 6L 2022-2023"
)

Note

If you create a one-time schedule and need to access the current time, use
datetime.utcnow() instead of datetime.now(). The latter does not store the current
zone context and results in an incorrect time passed to EventBridge.

Attach the trigger to your pipeline

To attach your PipelineSchedule to your pipeline, invoke the put_triggers call on your
created pipeline object with a list of triggers. If you get a response ARN, you successfully created
the schedule in your account and EventBridge begins to invoke the target pipeline at the time
or rate specified. You must specify a role with correct permissions to attach triggers to a parent
pipeline. If you don’t provide one, SageMaker Pipelines fetches the default role used to create the
pipeline from the configuration file.

The following example demonstrates how to attach a schedule to a pipeline.

```python
scheduled_pipeline = Pipeline(
    name="<pipeline-name>",
    steps=[...],
    sagemaker_session=<sagemaker-session>,
)
custom_schedule = PipelineSchedule(
    name="<schedule-name>",
    at=datetime(year=2023, month=12, date=25, hour=10, minute=30, second=30)
)
scheduled_pipeline.put_triggers(triggers=[custom_schedule], role_arn=<role>)
```

Describe current triggers

To retrieve information about your created pipeline triggers, you can invoke the
describe_trigger() API with the trigger name. This command returns details about the
created schedule expression such as its start time, enabled state, and other useful information. The
following snippet shows a sample invocation:
Cleanup trigger resources

Before you delete your pipeline, clean up existing triggers to avoid a resource leak in your account. You should delete the triggers before destroying the parent pipeline. You can delete your triggers by passing a list of trigger names to the `delete_triggers` API. The following snippet demonstrates how to delete triggers.

```
pipeline.delete_triggers(trigger_names=['<schedule-name>'])
```

**Note**

Be aware of the following limitations when you delete your triggers:

- The option to delete the triggers by specifying trigger names is only available in the SageMaker Python SDK. Deleting the pipeline in the CLI or a `DeletePipeline` API call does not delete your triggers. As a result, the triggers become orphaned and SageMaker attempts to start a run for a non-existent pipeline.

- Also, if you are using another notebook session or already deleted the pipeline target, clean up orphaned schedules through the scheduler CLI or EventBridge console.

Amazon SageMaker Experiments Integration

Amazon SageMaker Model Building Pipelines is closely integrated with Amazon SageMaker Experiments. By default, when SageMaker Pipelines creates and executes a pipeline, the following SageMaker Experiments entities are created if they don't exist:

- An experiment for the pipeline
- A run group for every execution of the pipeline
- A run that's added to the run group for each SageMaker job created in a pipeline execution step

You can compare metrics such as model training accuracy across multiple pipeline executions just as you can compare such metrics across multiple run groups of a SageMaker model training experiment.
The following sample shows the relevant parameters of the `Pipeline` class in the Amazon SageMaker Python SDK.

```python
Pipeline(
    name="MyPipeline",
    parameters=[...],
    pipeline_experiment_config=PipelineExperimentConfig(
        ExecutionVariables.PIPELINE_NAME,
        ExecutionVariables.PIPELINE_EXECUTION_ID
    ),
    steps=[...]
)
```

If you don't want an experiment and run group created for the pipeline, set `pipeline_experiment_config` to `None`.

**Note**

Experiments integration was introduced in the Amazon SageMaker Python SDK v2.41.0.

The following naming rules apply based on what you specify for the `ExperimentName` and `TrialName` parameters of `pipeline_experiment_config`:

- If you don't specify `ExperimentName`, the pipeline name is used for the experiment name.

  If you do specify `ExperimentName`, it's used for the experiment name. If an experiment with that name exists, the pipeline-created run groups are added to the existing experiment. If an experiment with that name doesn't exist, a new experiment is created.

- If you don't specify `TrialName`, the pipeline execution ID is used for the run group name.

  If you do specify `TrialName`, it's used for the run group name. If a run group with that name exists, the pipeline-created runs are added to the existing run group. If a run group with that name doesn't exist, a new run group is created.
Note
The experiment entities aren't deleted when the pipeline that created the entities is deleted. You can use the SageMaker Experiments API to delete the entities. For more information, see Clean Up Amazon SageMaker Experiment Resources.

For information about how to view the SageMaker Experiment entities associated with a pipeline, see View Experiment Entities Created by SageMaker Pipelines. For more information on SageMaker Experiments, see Manage Machine Learning with Amazon SageMaker Experiments.

The following sections show examples of the previous rules and how they are represented in the pipeline definition file. For more information on pipeline definition files, see SageMaker Pipelines Overview.

Topics
• Default Behavior
• Disable Experiments Integration
• Specify a Custom Experiment Name
• Specify a Custom Run Group Name

Default Behavior

Create a pipeline

The pipeline_experiment_config is omitted. ExperimentName defaults to the pipeline name. TrialName defaults to the execution ID.

```python
pipeline_name = f"MyPipeline"
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[...],
    steps=[step_train]
)
```

Pipeline definition file

```json
{
    "Version": "2020-12-01",
```
"Parameters": [
    {
        "Name": "InputDataSource"
    },
    {
        "Name": "InstanceCount",
        "Type": "Integer",
        "DefaultValue": 1
    }
],
"PipelineExperimentConfig": {
    "ExperimentName": {"Get": "Execution.PipelineName"},
    "TrialName": {"Get": "Execution.PipelineExecutionId"}
},
"Steps": [...]  
}

Disable Experiments Integration

Create a pipeline

The pipeline_experiment_config is set to None.

```python
pipeline_name = f"MyPipeline"
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[...],
    pipeline_experiment_config=None,
    steps=[step_train]
)
```

Pipeline definition file

This is the same as the preceding default example, without the PipelineExperimentConfig.

Specify a Custom Experiment Name

A custom experiment name is used. The run group name is set to the execution ID, as with the default behavior.

Create a pipeline

```python
pipeline_name = f"MyPipeline"
pipeline = Pipeline(
```
Pipeline definition file

```json
{
  ...
  "PipelineExperimentConfig": {
    "ExperimentName": "CustomExperimentName",
    "TrialName": {"Get": "Execution.PipelineExecutionId"}
  },
  "Steps": [...]
}
```

Specify a Custom Run Group Name

A custom run group name is used and appended with the execution ID. The experiment name is set to the pipeline name, as with the default behavior.

Create a pipeline

```python
pipeline_name = f"MyPipeline"
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[...],
    pipeline_experiment_config=PipelineExperimentConfig(
        ExecutionVariables.PIPELINE_NAME,
        Join(on="-", values=["CustomTrialName",
            ExecutionVariables.PIPELINE_EXECUTION_ID])
    ),
    steps=[step_train]
)
```

Pipeline definition file

```json
{
  ...
  "PipelineExperimentConfig": {
    "ExperimentName": "CustomExperimentName",
    "TrialName": {"Get": "Execution.PipelineExecutionId"}
  },
  "Steps": [...]
}
```
Local Mode

SageMaker Pipelines local mode is an easy way to test your training, processing and inference scripts, as well as the runtime compatibility of pipeline parameters before you execute your pipeline on the managed SageMaker service. By using local mode, you can test your SageMaker pipeline locally using a smaller dataset. This allows quick and easy debugging of errors in user scripts and the pipeline definition itself without incurring the costs of using the managed service.

Pipelines local mode leverages SageMaker jobs local mode under the hood. This is a feature in the SageMaker Python SDK that allows you to run SageMaker built-in or custom images locally using Docker containers. Pipelines local mode is built on top of SageMaker jobs local mode. Therefore, you can expect to see the same results as if you were running those jobs separately. For example, local mode still uses Amazon S3 to upload model artifacts and processing outputs. If you want data generated by local jobs to reside on local disk, you can use the setup mentioned in Local Mode.

Pipeline local mode currently supports the following step types:

- Training Step
- Processing Step
- Transform Step
- Model Step (with Create Model arguments only)
- Condition Step
- Fail Step

As opposed to the managed Pipelines service which allows multiple steps to execute in parallel using Parallelism Configuration, the local pipeline executor runs the steps sequentially. Therefore,
overall execution performance of a local pipeline may be poorer than one that runs on the cloud - this mostly depends on the size of the dataset, algorithm, as well as the power of your local computer. Also note that Pipelines runs in local mode are not recorded in SageMaker Experiments.

**Note**

Pipelines local mode is not compatible with SageMaker algorithms such as XGBoost. If you want to use these algorithms, you must use them in script mode.

In order to execute a pipeline locally, the sagemaker_session fields associated with the pipeline steps and the pipeline itself need to be of type LocalPipelineSession. The following example shows how you can define a SageMaker pipeline to execute locally.

```python
from sagemaker.workflow.pipeline_context import LocalPipelineSession
from sagemaker.pytorch import PyTorch
from sagemaker.workflow.steps import TrainingStep
from sagemaker.workflow.pipeline import Pipeline

local_pipeline_session = LocalPipelineSession()

pytorch_estimator = PyTorch(
    sagemaker_session=local_pipeline_session,
    role=sagemaker.get_execution_role(),
    instance_type="ml.c5.xlarge",
    instance_count=1,
    framework_version="1.8.0",
    py_version="py36",
    entry_point="./entry_point.py",
)

step = TrainingStep(
    name="MyTrainingStep",
    step_args=pytorch_estimator.fit(
        inputs=TrainingInput(s3_data="s3://my-bucket/my-data/train"),
    ),
)

pipeline = Pipeline(
    name="MyPipeline",
    steps=[step],
)```
Once you are ready to execute the pipeline on the managed SageMaker Pipelines service, you can do so by replacing `LocalPipelineSession` in the previous code snippet with `PipelineSession` (as shown in the following code sample) and rerunning the code.

```python
from sagemaker.workflow.pipeline_context import PipelineSession
pipeline_session = PipelineSession()
```

**Troubleshooting Amazon SageMaker Model Building Pipelines**

When using Amazon SageMaker Model Building Pipelines, you might run into issues for various reasons. This topic provides information about common errors and how to resolve them.

**Pipeline Definition Issues**

Your pipeline definition might not be formatted correctly. This can result in your execution failing or your job being inaccurate. These errors can be caught when the pipeline is created or when an execution occurs. If your definition doesn't validate, SageMaker Pipelines returns an error message identifying the character where the JSON file is malformed. To fix this problem, review the steps created using the SageMaker Python SDK for accuracy.
You can only include steps in a pipeline definition once. Because of this, steps cannot exist as part of a condition step and a pipeline in the same pipeline.

Examing Pipeline Logs

You can view the status of your steps using the following command:

```python
execution.list_steps()
```

Each step includes the following information:

- The ARN of the entity launched by the pipeline, such as SageMaker job ARN, model ARN, or model package ARN.
- The failure reason includes a brief explanation of the step failure.
- If the step is a condition step, it includes whether the condition is evaluated to true or false.
- If the execution reuses a previous job execution, the `CacheHit` lists the source execution.

You can also view the error messages and logs in the Amazon SageMaker Studio interface. For information about how to see the logs in Studio, see [View a Pipeline Execution](#).

Missing Permissions

Correct permissions are required for the role that creates the pipeline execution, and the steps that create each of the jobs in your pipeline execution. Without these permissions, you may not be able to submit your pipeline execution or run your SageMaker jobs as expected. To ensure that your permissions are properly set up, see [IAM Access Management](#).

Job Execution Errors

You may run into issues when executing your steps because of issues in the scripts that define the functionality of your SageMaker jobs. Each job has a set of CloudWatch logs. To view these logs from Studio, see [View a Pipeline Execution](#). For information about using CloudWatch logs with SageMaker, see [Log Amazon SageMaker Events with Amazon CloudWatch](#).

Property File Errors

You may have issues when incorrectly implementing property files with your pipeline. To ensure that your implementation of property files works as expected, see [Pass Data Between Steps](#).
Create and Manage SageMaker Pipelines

You can use Amazon SageMaker Model Building Pipelines to create end-to-end workflows that manage and deploy SageMaker jobs. SageMaker Pipelines comes with SageMaker Python SDK integration, so you can build each step of your pipeline using a Python-based interface.

After your pipeline is deployed, you can view the directed acyclic graph (DAG) for your pipeline and manage your executions using Amazon SageMaker Studio. Using SageMaker Studio, you can get information about your current and historical pipelines, compare executions, see the DAG for your executions, get metadata information, and more. To learn how to view pipelines from SageMaker Studio, see View, Track, and Execute SageMaker Pipelines in SageMaker Studio.

Topics

- Define a Pipeline
- Run a pipeline
- View, Track, and Execute SageMaker Pipelines in SageMaker Studio

Define a Pipeline

To orchestrate your workflows with Amazon SageMaker Model Building Pipelines, you need to generate a directed acyclic graph (DAG) in the form of a JSON pipeline definition. The following image is a representation of the pipeline DAG that you create in this tutorial:
You can generate your JSON pipeline definition using the SageMaker Python SDK. The following tutorial shows how to generate a pipeline definition for a pipeline that solves a regression problem to determine the age of an abalone based on its physical measurements. For a Jupyter notebook that includes the content in this tutorial that you can run, see Orchestrating Jobs with Amazon SageMaker Model Building Pipelines.

Topics

- Prerequisites
- Create a Pipeline
Prerequisites

To run the following tutorial you must do the following:

- Set up your notebook instance as outlined in [Create a notebook instance](#). This gives your role permissions to read and write to Amazon S3, and create training, batch transform, and processing jobs in SageMaker.
- Grant your notebook permissions to get and pass its own role as shown in [Modifying a role permissions policy](#). Add the following JSON snippet to attach this policy to your role. Replace `<your-role-arn>` with the ARN used to create your notebook instance.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "iam:GetRole",
            "iam:PassRole"
         ],
         "Resource": "<your-role-arn>"
      }
   }
}
```

- Trust the SageMaker service principal by following the steps in [Modifying a role trust policy](#). Add the following statement fragment to the trust relationship of your role:

```json
{
   "Sid": "",
   "Effect": "Allow",
   "Principal": {
      "Service": "sagemaker.amazonaws.com"
   },
   "Action": "sts:AssumeRole"
}
```

Set Up Your Environment

Create a new SageMaker session using the following code block. This returns the role ARN for the session. This role ARN should be the execution role ARN that you set up as a prerequisite.
Create a Pipeline

Run the following steps from your SageMaker notebook instance to create a pipeline including steps for preprocessing, training, evaluation, conditional evaluation, and model registration.

Step 1: Download the Dataset

This notebook uses the UCI Machine Learning Abalone Dataset. The dataset contains the following features:

- **length** – The longest shell measurement of the abalone.
- **diameter** – The diameter of the abalone perpendicular to its length.
- **height** – The height of the abalone with meat in the shell.
- **whole_weight** – The weight of the whole abalone.
- **shucked_weight** – The weight of the meat removed from the abalone.
- **viscera_weight** – The weight of the abalone viscera after bleeding.
- **shell_weight** – The weight of the abalone shell after meat removal and drying.
- **sex** – The sex of the abalone. One of 'M', 'F', or 'I', where 'I' is an infant abalone.
- **rings** – The number of rings in the abalone shell.

The number of rings in the abalone shell is a good approximation for its age using the formula \( \text{age} = \text{rings} + 1.5 \). However, obtaining this number is a time-consuming task. You must cut the shell through the cone, stain the section, and count the number of rings through a microscope.
However, the other physical measurements are easier to determine. This notebook uses the dataset to build a predictive model of the variable rings using the other physical measurements.

**To download the dataset**

1. Download the dataset into your account's default Amazon S3 bucket.

```bash
!mkdir -p data
local_path = "data/abalone-dataset.csv"

s3 = boto3.resource("s3")
s3.Bucket(f"sagemaker-servicecatalog-seedcode-{region}").download_file(
    "dataset/abalone-dataset.csv",
    local_path
)

base_uri = f"s3://{default_bucket}/abalone"
input_data_uri = sagemaker.s3.S3Uploader.upload(
    local_path=local_path,
    desired_s3_uri=base_uri,
)
print(input_data_uri)
```

2. Download a second dataset for batch transformation after your model is created.

```bash
local_path = "data/abalone-dataset-batch.csv"

s3 = boto3.resource("s3")
s3.Bucket(f"sagemaker-servicecatalog-seedcode-{region}").download_file(
    "dataset/abalone-dataset-batch",
    local_path
)

base_uri = f"s3://{default_bucket}/abalone"
batch_data_uri = sagemaker.s3.S3Uploader.upload(
    local_path=local_path,
    desired_s3_uri=base_uri,
)
print(batch_data_uri)
```
Step 2: Define Pipeline Parameters

This code block defines the following parameters for your pipeline:

- `processing_instance_count` – The instance count of the processing job.
- `input_data` – The Amazon S3 location of the input data.
- `batch_data` – The Amazon S3 location of the input data for batch transformation.
- `model_approval_status` – The approval status to register the trained model with for CI/CD.

For more information, see Automate MLOps with SageMaker Projects.

```python
from sagemaker.workflow.parameters import (  
    ParameterInteger,  
    ParameterString,  
)

processing_instance_count = ParameterInteger(  
    name="ProcessingInstanceCount",  
    default_value=1  
)
model_approval_status = ParameterString(  
    name="ModelApprovalStatus",  
    default_value="PendingManualApproval"  
)
input_data = ParameterString(  
    name="InputData",  
    default_value=input_data_uri,  
)
batch_data = ParameterString(  
    name="BatchData",  
    default_value=batch_data_uri,  
)
```

Step 3: Define a Processing Step for Feature Engineering

This section shows how to create a processing step to prepare the data from the dataset for training.

To create a processing step

1. Create a directory for the processing script.
2. Create a file in the `/abalone` directory named `preprocessing.py` with the following content. This preprocessing script is passed in to the processing step for execution on the input data. The training step then uses the preprocessed training features and labels to train a model, and the evaluation step uses the trained model and preprocessed test features and labels to evaluate the model. The script uses scikit-learn to do the following:

- Fill in missing sex categorical data and encode it so it's suitable for training.
- Scale and normalize all numerical fields except for `rings` and `sex`.
- Split the data into training, test, and validation datasets.

```python
# Because this is a headerless CSV file, specify the column names here.
feature_columns_names = [
    "sex",
    "length",
    "diameter",
    "height",
    "whole_weight",
    "shucked_weight",
    "viscera_weight",
    "shell_weight",
]
label_column = "rings"
```
feature_columns_dtype = {
    "sex": str,
    "length": np.float64,
    "diameter": np.float64,
    "height": np.float64,
    "whole_weight": np.float64,
    "shucked_weight": np.float64,
    "viscera_weight": np.float64,
    "shell_weight": np.float64
}
label_column_dtype = {"rings": np.float64}

def merge_two_dicts(x, y):
    z = x.copy()
    z.update(y)
    return z

if __name__ == "__main__":
    base_dir = "/opt/ml/processing"

    df = pd.read_csv(
        f"{base_dir}/input/abalone-dataset.csv",
        header=None,
        names=feature_columns_names + [label_column],
        dtype=merge_two_dicts(feature_columns_dtype, label_column_dtype)
    )
    numeric_features = list(feature_columns_names)
    numeric_features.remove("sex")
    numeric_transformer = Pipeline(
        steps=[
            ("imputer", SimpleImputer(strategy="median")),
            ("scaler", StandardScaler())
        ]
    )

    categorical_features = ["sex"]
    categorical_transformer = Pipeline(
        steps=[
            ("imputer", SimpleImputer(strategy="constant", fill_value="missing")),
            ("onehot", OneHotEncoder(handle_unknown="ignore"))
        ]
    )
preprocess = ColumnTransformer(
    transformers=[
        ("num", numeric_transformer, numeric_features),
        ("cat", categorical_transformer, categorical_features)
    ]
)

y = df.pop("rings")
X_pre = preprocess.fit_transform(df)
y_pre = y.to_numpy().reshape(len(y), 1)

X = np.concatenate((y_pre, X_pre), axis=1)

np.random.shuffle(X)
train, validation, test = np.split(X, [int(.7*len(X)), int(.85*len(X))])

pd.DataFrame(train).to_csv(f"{base_dir}/train/train.csv", header=False, index=False)
pd.DataFrame(validation).to_csv(f"{base_dir}/validation/validation.csv", header=False, index=False)
pd.DataFrame(test).to_csv(f"{base_dir}/test/test.csv", header=False, index=False)

3. Create an instance of an SKLearnProcessor to pass in to the processing step.

```python
from sagemaker.sklearn.processing import SKLearnProcessor

framework_version = "0.23-1"

sklearn_processor = SKLearnProcessor(  
    framework_version=framework_version,  
    instance_type="ml.m5.xlarge",  
    instance_count=processing_instance_count,  
    base_job_name="sklearn-abalone-process",  
    sagemaker_session=pipeline_session,  
    role=role,
)
```

4. Create a processing step. This step takes in the SKLearnProcessor, the input and output channels, and the preprocessing.py script that you created. This is very similar to a
processor instance's run method in the SageMaker Python SDK. The input_data parameter passed into ProcessingStep is the input data of the step itself. This input data is used by the processor instance when it runs.

Note the "train", "validation", and "test" named channels specified in the output configuration for the processing job. Step Properties such as these can be used in subsequent steps and resolve to their runtime values at execution.

```python
from sagemaker.processing import ProcessingInput, ProcessingOutput
from sagemaker.workflow.steps import ProcessingStep

processor_args = sklearn_processor.run(
    inputs=[
        ProcessingInput(source=input_data, destination="/opt/ml/processing/input"),
    ],
    outputs=[
        ProcessingOutput(output_name="train", source="/opt/ml/processing/train"),
        ProcessingOutput(output_name="validation", source="/opt/ml/processing/validation"),
        ProcessingOutput(output_name="test", source="/opt/ml/processing/test")
    ],
    code="abalone/preprocessing.py",
)

step_process = ProcessingStep(
    name="AbaloneProcess",
    step_args=processor_args
)
```

**Step 4: Define a Training step**

This section shows how to use the SageMaker [XGBoost Algorithm](https://en.wikipedia.org/wiki/XGBoost) to train a model on the training data output from the processing steps.

**To define a training step**

1. Specify the model path where you want to save the models from training.

```python
model_path = f"s3://{default_bucket}/AbaloneTrain"
```
2. Configure an estimator for the XGBoost algorithm and the input dataset. The training instance
type is passed into the estimator. A typical training script loads data from the input channels,
configures training with hyperparameters, trains a model, and saves a model to `model_dir`
so that it can be hosted later. SageMaker uploads the model to Amazon S3 in the form of
a `model.tar.gz` at the end of the training job.

```python
from sagemaker.estimator import Estimator

image_uri = sagemaker.image_uris.retrieve(
    framework="xgboost",
    region=region,
    version="1.0-1",
    py_version="py3",
    instance_type="ml.m5.xlarge"
)

xgb_train = Estimator(
    image_uri=image_uri,
    instance_type="ml.m5.xlarge",
    instance_count=1,
    output_path=model_path,
    sagemaker_session=pipeline_session,
    role=role,
)

xgb_train.set_hyperparameters(
    objective="reg:linear",
    num_round=50,
    max_depth=5,
    eta=0.2,
    gamma=4,
    min_child_weight=6,
    subsample=0.7,
    silent=0
)
```

3. Create a `TrainingStep` using the estimator instance and properties of
the `ProcessingStep`. In particular, pass in the S3Uri of the "train" and "validation"
output channel to the `TrainingStep`.

```python
from sagemaker.inputs import TrainingInput
from sagemaker.workflow.steps import TrainingStep
```
Step 5: Define a Processing Step for Model Evaluation

This section shows how to create a processing step to evaluate the accuracy of the model. The result of this model evaluation is used in the condition step to determine which execute path to take.

To define a processing step for model evaluation

1. Create a file in the /abalone directory named evaluation.py. This script is used in a processing step to perform model evaluation. It takes a trained model and the test dataset as input, then produces a JSON file containing classification evaluation metrics.

```python
%%writefile abalone/evaluation.py
import json
import pathlib
import pickle
import tarfile
import joblib
import numpy as np
```
import pandas as pd
import xgboost

from sklearn.metrics import mean_squared_error

if __name__ == '__main__':
    model_path = f'/opt/ml/processing/model/model.tar.gz'
    with tarfile.open(model_path) as tar:
        tar.extractall(path='.')

    model = pickle.load(open("xgboost-model", "rb"))

    test_path = '/opt/ml/processing/test/test.csv'
    df = pd.read_csv(test_path, header=None)

    y_test = df.iloc[:, 0].to_numpy()
    df.drop(df.columns[0], axis=1, inplace=True)

    X_test = xgboost.DMatrix(df.values)

    predictions = model.predict(X_test)

    mse = mean_squared_error(y_test, predictions)
    std = np.std(y_test - predictions)

    report_dict = {
        "regression_metrics": {
            "mse": {
                "value": mse,
                "standard_deviation": std
            },
        },
    }

    output_dir = '/opt/ml/processing/evaluation'
    pathlib.Path(output_dir).mkdir(parents=True, exist_ok=True)

    evaluation_path = f'{output_dir}/evaluation.json'
    with open(evaluation_path, "w") as f:
        f.write(json.dumps(report_dict))

2. Create an instance of a ScriptProcessor that is used to create a ProcessingStep.
from sagemaker.processing import ScriptProcessor

script_eval = ScriptProcessor(
    image_uri=image_uri,
    command=['python3'],
    instance_type='ml.m5.xlarge',
    instance_count=1,
    base_job_name='script-abalone-eval',
    sagemaker_session=pipeline_session,
    role=role,
)

3. Create a `ProcessingStep` using the processor instance, the input and output channels, and the `evaluation.py` script. In particular, pass in the `S3ModelArtifacts` property from the `step_train` training step, as well as the `S3Uri` of the "test" output channel of the `step_process` processing step. This is very similar to a processor instance's `run` method in the SageMaker Python SDK.

from sagemaker.workflow.properties import PropertyFile

evaluation_report = PropertyFile(
    name='EvaluationReport',
    output_name='evaluation',
    path='evaluation.json'
)

eval_args = script_eval.run(
    inputs=[
        ProcessingInput(
            source=step_train.properties.ModelArtifacts.S3ModelArtifacts,
            destination='/opt/ml/processing/model'
        ),
        ProcessingInput(
            source=step_process.properties.ProcessingOutputConfig.Outputs['test'].S3Output.S3Uri,
            destination='/opt/ml/processing/test'
        )
    ],
    outputs=[
    ]
)
Step 6: Define a CreateModelStep for Batch Transformation

⚠️ Important

We recommend using `Model Step` to create models as of v2.90.0 of the SageMaker Python SDK. CreateModelStep will continue to work in previous versions of the SageMaker Python SDK, but is no longer actively supported.

This section shows how to create a SageMaker model from the output of the training step. This model is used for batch transformation on a new dataset. This step is passed into the condition step and only executes if the condition step evaluates to `true`.

To define a CreateModelStep for batch transformation


   ```python
   from sagemaker.model import Model

   model = Model(
       image_uri=image_uri,
       model_data=step_train.properties.ModelArtifacts.S3ModelArtifacts,
       sagemaker_session=pipeline_session,
       role=role,
   )
   ```

2. Define the model input for your SageMaker model.
from sagemaker.inputs import CreateModelInput

inputs = CreateModelInput(
    instance_type="ml.m5.large",
    accelerator_type="ml.eia1.medium",
)

3. Create your CreateModelStep using the CreateModelInput and SageMaker model instance you defined.

from sagemaker.workflow.steps import CreateModelStep

step_create_model = CreateModelStep(
    name="AbaloneCreateModel",
    model=model,
    inputs=inputs,
)

**Step 7: Define a TransformStep to Perform Batch Transformation**

This section shows how to create a TransformStep to perform batch transformation on a dataset after the model is trained. This step is passed into the condition step and only executes if the condition step evaluates to true.

**To define a TransformStep to perform batch transformation**

1. Create a transformer instance with the appropriate compute instance type, instance count, and desired output Amazon S3 bucket URI. Pass in the ModelName property from the step_create_model CreateModel step.

from sagemaker.transformer import Transformer

transformer = Transformer(
    model_name=step_create_model.properties.ModelName,
    instance_type="ml.m5.xlarge",
    instance_count=1,
    output_path=f"s3://{default_bucket}/AbaloneTransform"
2. Create a `TransformStep` using the transformer instance you defined and the `batch_data` pipeline parameter.

```python
from sagemaker.inputs import TransformInput
from sagemaker.workflow.steps import TransformStep

step_transform = TransformStep(
    name="AbaloneTransform",
    transformer=transformer,
    inputs=TransformInput(data=batch_data)
)
```

---

**Step 8: Define a RegisterModel Step to Create a Model Package**

⚠️ **Important**

We recommend using [Model Step](#) to register models as of v2.90.0 of the SageMaker Python SDK. `RegisterModel` will continue to work in previous versions of the SageMaker Python SDK, but is no longer actively supported.

This section shows how to construct an instance of `RegisterModel`. The result of executing `RegisterModel` in a pipeline is a model package. A model package is a reusable model artifacts abstraction that packages all ingredients necessary for inference. It consists of an inference specification that defines the inference image to use along with an optional model weights location. A model package group is a collection of model packages. You can use a [ModelPackageGroup](#) for SageMaker Pipelines to add a new version and model package to the group for every pipeline execution. For more information about model registry, see [Register and Deploy Models with Model Registry](#).

This step is passed into the condition step and only executes if the condition step evaluates to `true`. 
To define a RegisterModel step to create a model package

- Construct a RegisterModel step using the estimator instance you used for the training step. Pass in the S3ModelArtifacts property from the step_train training step and specify a ModelPackageGroup. SageMaker Pipelines creates this ModelPackageGroup for you.

```python
from sagemaker.model_metrics import MetricsSource, ModelMetrics
from sagemaker.workflow.step_collections import RegisterModel

model_metrics = ModelMetrics(
    model_statistics=MetricsSource(
        s3_uri="{}/evaluation.json".format(
            step_eval.arguments["ProcessingOutputConfig"]["Outputs"][0]["S3Output"]["S3Uri"]
        ),
        content_type="application/json"
    )
)

step_register = RegisterModel(
    name="AbaloneRegisterModel",
    estimator=xgb_train,
    model_data=step_train.properties.ModelArtifacts.S3ModelArtifacts,
    content_types=["text/csv"],
    response_types=["text/csv"],
    inference_instances=["ml.t2.medium", "ml.m5.xlarge"],
    transform_instances=["ml.m5.xlarge"],
    model_package_group_name=model_package_group_name,
    approval_status=model_approval_status,
    model_metrics=model_metrics
)
```

Step 9: Define a Condition Step to Verify Model Accuracy

A ConditionStep allows SageMaker Pipelines to support conditional execution in your pipeline DAG based on the condition of step properties. In this case, you only want to register a model package if the accuracy of that model, as determined by the model evaluation step, exceeds the required value. If the accuracy exceeds the required value, the pipeline also creates a SageMaker Model and runs batch transformation on a dataset. This section shows how to define the Condition step.
To define a condition step to verify model accuracy

1. Define a `ConditionLessThanOrEqualTo` condition using the accuracy value found in the output of the model evaluation processing step, `step_eval`. Get this output using the property file you indexed in the processing step and the respective JSONPath of the mean squared error value, "mse".

```python
from sagemaker.workflow.conditions import ConditionLessThanOrEqualTo
from sagemaker.workflow.condition_step import ConditionStep
from sagemaker.workflow.functions import JsonGet

cond_lte = ConditionLessThanOrEqualTo(
    left=JsonGet(
        step_name=step_eval.name,
        property_file=evaluation_report,
        json_path="regression_metrics.mse.value"),
    right=6.0)
```

2. Construct a `ConditionStep`. Pass the `ConditionEquals` condition in, then set the model package registration and batch transformation steps as the next steps if the condition passes.

```python
step_cond = ConditionStep(
    name="AbaloneMSECond",
    conditions=[cond_lte],
    if_steps=[step_register, step_create_model, step_transform],
    else_steps=[],
)
```

Step 10: Create a pipeline

Now that you’ve created all of the steps, combine them into a pipeline.

To create a pipeline

1. Define the following for your pipeline: name, parameters, and steps. Names must be unique within an (account, region) pair.
A step can only appear once in either the pipeline's step list or the if/else step lists of the condition step. It cannot appear in both.

```python
from sagemaker.workflow.pipeline import Pipeline

pipeline_name = f"AbalonePipeline"
pipeline = Pipeline(
    name=pipeline_name,
    parameters=[
        processing_instance_count,
        model_approval_status,
        input_data,
        batch_data,
    ],
    steps=[step_process, step_train, step_eval, step_cond],
)
```

2. (Optional) Examine the JSON pipeline definition to ensure that it's well-formed.

```python
import json

json.loads(pipeline.definition())
```

This pipeline definition is ready to submit to SageMaker. In the next tutorial, you submit this pipeline to SageMaker and start an execution.

**Next step:** [Run a pipeline](#)

**Run a pipeline**

After you've created a pipeline definition using the SageMaker Python SDK, you can submit it to SageMaker to start your execution. The following tutorial shows how to submit a pipeline, start an execution, examine the results of that execution, and delete your pipeline.

**Topics**

Amazon SageMaker Model Building Pipelines
Prerequisites

This tutorial requires the following:

- A SageMaker notebook instance.
- A SageMaker Pipelines pipeline definition. This tutorial assumes you're using the pipeline definition created by completing the Define a Pipeline tutorial.

Step 1: Start the Pipeline

First, you need to start the pipeline.

To start the pipeline

1. Examine the JSON pipeline definition to ensure that it's well-formed.

   ```python
   import json
   json.loads(pipeline.definition())
   ```

2. Submit the pipeline definition to the SageMaker Pipelines service to create a pipeline if it doesn't exist, or update the pipeline if it does. The role passed in is used by SageMaker Pipelines to create all of the jobs defined in the steps.

   ```python
   pipeline.upsert(role_arn=role)
   ```

3. Start a pipeline execution.

   ```python
   execution = pipeline.start()
   ```
Step 2: Examine a Pipeline Execution

Next, you need to examine the pipeline execution.

To examine a pipeline execution

1. Describe the pipeline execution status to ensure that it has been created and started successfully.

   ```python
   execution.describe()
   ```

2. Wait for the execution to finish.

   ```python
   execution.wait()
   ```

3. List the execution steps and their status.

   ```python
   execution.list_steps()
   ```

Your output should look like the following:

```json
[{'StepName': 'AbaloneTransform',
  'StartTime': datetime.datetime(2020, 11, 21, 2, 41, 27, 870000, tzinfo=tzlocal()),
  'EndTime': datetime.datetime(2020, 11, 21, 2, 45, 50, 492000, tzinfo=tzlocal()),
  'StepStatus': 'Succeeded',
  'CacheHitResult': {'SourcePipelineExecutionArn': ''},
{'StepName': 'AbaloneRegisterModel',
  'StartTime': datetime.datetime(2020, 11, 21, 2, 41, 26, 929000, tzinfo=tzlocal()),
  'EndTime': datetime.datetime(2020, 11, 21, 2, 41, 28, 15000, tzinfo=tzlocal()),
  'StepStatus': 'Succeeded',
  'CacheHitResult': {'SourcePipelineExecutionArn': ''},
  'Metadata': {'RegisterModel': {'Arn': 'arn:aws:sagemaker:us-east-2:11112223333:model-package/abalonemodelpackagegroupname/1'}}},
{'StepName': 'AbaloneCreateModel',
  'StartTime': datetime.datetime(2020, 11, 21, 2, 41, 26, 895000, tzinfo=tzlocal()),
  'EndTime': datetime.datetime(2020, 11, 21, 2, 41, 27, 708000, tzinfo=tzlocal()),
```
4. After your pipeline execution is complete, download the resulting `evaluation.json` file from Amazon S3 to examine the report.

```python
evaluation_json = sagemaker.s3.S3Downloader.read_file("/{}/evaluation.json".format(
)"
**Step 3: Override Default Parameters for a Pipeline Execution**

You can run additional executions of the pipeline by specifying different pipeline parameters to override the defaults.

### To override default parameters

1. Create the pipeline execution. This starts another pipeline execution with the model approval status override set to "Approved". This means that the model package version generated by the `RegisterModel` step is automatically ready for deployment through CI/CD pipelines, such as with SageMaker Projects. For more information, see Automate MLOps with SageMaker Projects.

   ```python
   execution = pipeline.start(
       parameters=dict(
           ModelApprovalStatus="Approved",
       )
   )
   ```

2. Wait for the execution to finish.

   ```python
   execution.wait()
   ```

3. List the execution steps and their status.

   ```python
   execution.list_steps()
   ```

4. After your pipeline execution is complete, download the resulting `evaluation.json` file from Amazon S3 to examine the report.

   ```python
   evaluation_json = sagemaker.s3.S3Downloader.read_file("{}/evaluation.json".format(
       step_eval.arguments["ProcessingOutputConfig"]["Outputs"][0]["S3Output"]
   ["S3Uri"]
   ))
   json.loads(evaluation_json)
   ```
Step 4: Stop and Delete a Pipeline Execution

When you're finished with your pipeline, you can stop any ongoing executions and delete the pipeline.

To stop and delete a pipeline execution

1. Stop the pipeline execution.

   ```python
   execution.stop()
   ``

2. Delete the pipeline.

   ```python
   pipeline.delete()
   ``

View, Track, and Execute SageMaker Pipelines in SageMaker Studio

To view, track, and execute Amazon SageMaker Pipelines in Amazon SageMaker Studio, you must sign in to Studio. For more information, see Launch Amazon SageMaker Studio.

Topics

- View a Pipeline
- View a Pipeline Execution
- Download a Pipeline Definition
- View Experiment Entities Created by SageMaker Pipelines
- Start (and Stop) a Pipeline Execution
- Track the Lineage of a SageMaker ML Pipeline

View a Pipeline

This procedure shows you how to find a pipeline directly and view its details page. You can also find pipelines that are part of a project listed in the project's details page. For information about finding a pipeline that is part of a project, see Automate MLOps with SageMaker Projects.

To view a list of pipelines in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.
Studio

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, select Pipelines.
3. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
4. Select a pipeline name to view details about the pipeline. The pipeline's Executions page opens and displays a list of pipeline executions. Use the Column icon ( ) to choose which columns to display.
5. From the pipeline's Executions page, choose one of the following pages in the Overview, Settings, or Details dropdown menus (to the left of the pipeline executions table) to view pipeline details:
   - Executions – Details about the executions.
   - Graph – The DAG for the pipeline.
   - Parameters – Includes the model approval status.
   - Information – The metadata associated with the pipeline, such as the pipeline Amazon Resource Name (ARN) and role ARN. You can also edit the pipeline description from this page.

Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the Studio Classic sidebar, choose the Home icon ( )
3. Select Pipelines from the menu.
4. To narrow the list of pipelines by name, enter a full or partial pipeline name in the search field.
5. Select a pipeline name to view details about the pipeline. The pipeline details tab opens and displays a list of pipeline executions. You can start an execution or choose one of the
other tabs for more information about the pipeline. Use the Property Inspector icon ( ⚙️ ) to choose which columns to display.

6. From the pipeline details page, choose one of the following tabs to view details about the pipeline:

   - **Executions** – Details about the executions. You can create an execution from this tab or the Graph tab.
   - **Graph** – The DAG for the pipeline.
   - **Parameters** – Includes the model approval status.
   - **Settings** – The metadata associated with the pipeline. You can download the pipeline definition file and edit the pipeline name and description from this tab.

**View a Pipeline Execution**

This procedure shows you how to view a pipeline execution. For information about how to view a list of pipeline executions, and how to use SageMaker search to narrow the executions in the list, see [View a Pipeline](#).

To view a pipeline execution in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](#).
2. In the left navigation pane, select **Pipelines**.
3. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
4. Select a pipeline name to view details about the pipeline. The pipeline's **Executions** page opens and displays a list of pipeline executions.
5. Select the name of a pipeline execution to view. The pipeline graph of the execution appears.
6. (Optional) Select a step in the **Select step** dropdown menu to the right of the graph to center the graph on your chosen step. Use the resizing icons on the lower-right side of the graph to zoom in and out of the graph, fit the graph to screen, and expand the graph to
full screen. To focus on a specific part of the graph, you can select a blank area of the graph and drag the graph to center on that area.

7. Choose one of the pipeline steps in the graph to see details about the step. You can view step execution details in the following tabs:

   - **Overview** — Details related to the step execution, including status and runtime, related metrics and charts, and file locations of output collaterals.
   - **Settings** — Parameters and values related to your pipeline step, as defined by the JSON definition for the step. Includes input scripts and datasets.
   - **Details** — General information about the step, including step type (such as processing or training), and log file locations.

**Studio Classic**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
2. In the Studio Classic sidebar, choose the **Home** icon (  

3. Select **Pipelines** from the menu.

4. To narrow the list of pipelines by name, enter a full or partial pipeline name in the search field.

5. Select a pipeline name. The pipeline's **Executions** page opens.

6. In the **Executions** page, select an execution name to view details about the execution. The execution details tab opens and displays a graph of the steps in the pipeline.

7. To search for a step by name, type characters that match a step name in the search field. Use the resizing icons on the lower-right side of the graph to zoom in and out of the graph, fit the graph to screen, and expand the graph to full screen. To focus on a specific part of the graph, you can select a blank area of the graph and drag the graph to center on that area.

8. Choose one of the pipeline steps in the graph to see details about the step. In the preceding screenshot, a training step is chosen and displays the following tabs:
• **Input** – The training inputs. If an input source is from Amazon Simple Storage Service (Amazon S3), choose the link to view the file in the Amazon S3 console.

• **Output** – The training outputs, such as metrics, charts, files, and evaluation outcome. The graphs are produced using the Tracker APIs.

• **Logs** – The Amazon CloudWatch logs produced by the step.

• **Info** – The parameters and metadata associated with the step.

![Image of output, logs, and info tables]

### Download a Pipeline Definition

You can download a pipeline definition in the Amazon SageMaker Studio console. To download a pipeline definition, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](#).
2. In the left navigation pane, select **Pipelines**.
3. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
4. Select a pipeline name. The **Executions** page opens and displays a list of pipeline executions.
5. Stay on the **Executions** page or choose the **Graph**, **Information**, or **Parameters** page to the left of the pipeline executions table. You can download the pipeline definition from any of these pages.

6. At the top right of the page, choose the vertical ellipsis and choose **Download pipeline definition (JSON)**.

### Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).

2. In the Studio Classic sidebar, choose the **Home** icon ( ).

3. Select **Pipelines** from the menu.

4. To narrow the list of pipelines by name, enter a full or partial pipeline name in the search field.

5. Select a pipeline name.

6. Choose the **Settings** tab.

7. Choose **Download pipeline definition file**.

### View Experiment Entities Created by SageMaker Pipelines

**Note**

SageMaker Experiments is a feature provided in Studio Classic only.

When you create a pipeline and specify `pipeline_experiment_config`, SageMaker Pipelines creates the following SageMaker Experiments entities by default if they don't exist:

- An experiment for the pipeline
- A run group for every execution of the pipeline
- A run for each SageMaker job created in a pipeline step
For information about how experiments are integrated with pipelines, see Amazon SageMaker Experiments Integration. For more information about SageMaker Experiments, see Manage Machine Learning with Amazon SageMaker Experiments.

You can get to the list of runs associated with a pipeline from either the pipeline executions list or the experiments list.

**To view the runs list from the pipeline executions list**

1. To view the pipeline executions list, follow the first five steps in the Studio Classic tab of View a Pipeline.
2. On the top right of the screen, choose the Filter icon ( ![Filter](filter_icon.png)).
3. Choose Experiment. If experiment integration wasn't deactivated when the pipeline was created, the experiment name is displayed in the executions list.

   **Note**
   
   Experiments integration was introduced in v2.41.0 of the Amazon SageMaker Python SDK. Pipelines created with an earlier version of the SDK aren't integrated with experiments by default.

4. Select the experiment of your choice to view run groups and runs related to that experiment.

**To view the runs list from the experiments list**

1. In the left sidebar of Studio Classic, choose the Home icon ( ![Home](home_icon.png)).
2. Select Experiments from the menu.
3. Use search bar or Filter icon ( ![Filter](filter_icon.png)) to filter the list to experiments created by a pipeline.
4. Open an experiment name and view a list of runs created by the pipeline.
Start (and Stop) a Pipeline Execution

You can start and stop a pipeline execution in the Amazon SageMaker Studio console. For information about how to view a list of pipeline executions, see View a Pipeline.

To start and stop a pipeline execution in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

Studio

To start a pipeline execution

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, select Pipelines.
3. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
4. Select a pipeline name. The Executions page opens and displays a list of pipeline executions.
5. You can create an execution from either the Executions or Graph pages. To create an execution from the Executions page, choose Create. To create an execution from the Graph page, choose Graph to the left of the executions table and then Create execution in the top right of the DAG.
6. Enter or update the following required information:
   - **Name** – A name unique to your account in the AWS Region.
   - **Description** – An optional description for your execution.
   - **ProcessingInstanceType** – The Amazon EC2 instance type to use for the processing job.
   - **TrainingInstanceType** – The Amazon EC2 instance type to use for the training job
   - **InputData** – The Amazon S3 URI to the input data.
   - **PreprocessScript** – The Amazon S3 URI to the preprocessing script.
   - **EvaluateScript** – The Amazon S3 URI to the model evaluation script.
   - **AccuracyConditionThreshold** – The threshold of model accuracy to achieve to register the model into the registry.
   - **ModelGroup** – The registry into which to register the model.
- **MaximumParallelTrainingJobs** – The maximum number of training jobs to run in parallel.
- **MaximumTrainingJobs** – The maximum number of training jobs to run.

7. Choose Create.

**To stop a pipeline execution**

1. In the left navigation pane, select Pipelines.
2. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
3. Select a pipeline name. The Executions page opens and displays a list of pipeline executions.
4. Select the execution to stop.
5. Choose Stop.

**To resume a stopped pipeline execution**

1. In the left navigation pane, select Pipelines.
2. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
3. Select a pipeline name. The Executions page opens and displays a list of pipeline executions.
4. Select the execution to resume.
5. Choose Resume.

**Studio Classic**

**To start, stop, or resume a pipeline execution**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the Studio Classic sidebar, choose the Home icon ().
3. Select Pipelines from the menu.
4. To narrow the list of pipelines by name, enter a full or partial pipeline name in the search field.

5. Select a pipeline name.

6. From the **Executions** or **Graph** tab in the execution list, choose **Create execution**.

7. Enter or update the following required information:
   - **Name** – Must be unique to your account in the AWS Region.
   - **ProcessingInstanceCount** – The number of instances to use for processing.
   - **ModelApprovalStatus** – For your convenience.
   - **InputDataUrl** – The Amazon S3 URI of the input data.

8. Choose **Start**.

   - To see details of the execution or to stop the execution, choose **View details** on the status banner.
   - To stop the execution, choose **Stop** on the status banner.
   - To resume the execution from where it was stopped, choose **Resume** on the status banner.

**Note**

If your pipeline fails, the status banner will show **Failed** status. After troubleshooting the failed step, choose **Retry** on the status banner to resume running the pipeline from that step.

For a list of registered models, see [Automate MLOps with SageMaker Projects](#).

**Track the Lineage of a SageMaker ML Pipeline**

In this tutorial, you use Amazon SageMaker Studio to track the lineage of an Amazon SageMaker ML Pipeline.

The pipeline was created by the [Orchestrating Jobs with Amazon SageMaker Model Building Pipelines](#) notebook in the [Amazon SageMaker example GitHub repository](#). For detailed information on how the pipeline was created, see [Define a Pipeline](#).
Lineage tracking in Studio is centered around a directed acyclic graph (DAG). The DAG represents the steps in a pipeline. From the DAG you can track the lineage from any step to any other step. The following diagram displays the steps in the pipeline. These steps appear as a DAG in Studio.

To track the lineage of a pipeline in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

**To track the lineage of a pipeline**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](#).
2. In the left navigation pane, select **Pipelines**.
3. (Optional) To filter the list of pipelines by name, enter a full or partial pipeline name in the search field.
4. In the **Name** column, select a pipeline name to view details about the pipeline. The pipeline's **Executions** page opens and displays a list of pipeline executions.
5. In the **Name** column of the **Executions** table, select the name of a pipeline execution to view.
6. At the top right of the **Executions** page, choose the vertical ellipsis and choose **Download pipeline definition (JSON)**. You can view the file to see how the pipeline graph was defined.
7. Use the resizing icons on the lower-right side of the graph to zoom in and out of the graph, fit the graph to screen, or expand the graph to full screen. To focus on a specific part of the graph, you can select a blank area of the graph and drag the graph to center on that area. The inset on the lower-right side of the graph displays your location in the graph.

The following image shows an example pipeline graph with inset and resizing icons. Also, the tabs to the right of the graph contain detailed information about your pipeline run.
8. To view your training, validation, and test datasets, complete the following steps:
   
a. Choose the Processing step in your pipeline graph.

b. In the **Overview** tab, in the **Files** section, find the Amazon S3 paths to the training, validation, and test datasets.

9. To view your model artifacts, complete the following steps:
   
a. Choose the Training step in your pipeline graph.

b. In the **Overview** tab, in the **Files** section, find the Amazon S3 paths to the model artifact.

10. To find the model package ARN, complete the following steps:
   
a. Choose the model register (**RegisterModel**) step.

b. In the **Overview** tab, in the **Files** section, find the ARN of the model package.
To track the lineage of a pipeline

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left sidebar of Studio, choose the Home icon ( )
3. In the menu, select Pipelines.
4. Use the Search box to filter the pipelines list.
5. Choose the AbalonePipeline pipeline to view the execution list and other details about the pipeline.
6. Choose the Property Inspector icon ( ) in the right sidebar to open the TABLE PROPERTIES pane, where you can choose which properties to view.
7. Choose the Settings tab and then choose Download pipeline definition file. You can view the file to see how the pipeline graph was defined.
8. On the Execution tab, select the first row in the execution list to view its execution graph and other details about the execution. Note that the graph matches the diagram displayed at the beginning of the tutorial.

Use the resizing icons on the lower-right side of the graph to zoom in and out of the graph, fit the graph to screen, or expand the graph to full screen. To focus on a specific part of the graph, you can select a blank area of the graph and drag the graph to center on that area. The inset on the lower-right side of the graph displays your location in the graph.
9. On the **Graph** tab, choose the AbaloneProcess step to view details about the step.

10. Find the Amazon S3 paths to the training, validation, and test datasets in the **Output** tab, under **Files**.

    ❗ **Note**
    
    To get the full paths, right-click the path and then choose **Copy cell contents**.

    `s3://sagemaker-eu-west-1-acct-id/sklearn-abalone-process-2020-12-05-17-28-28-509/output/train`
    `s3://sagemaker-eu-west-1-acct-id/sklearn-abalone-process-2020-12-05-17-28-28-509/output/validation`
    `s3://sagemaker-eu-west-1-acct-id/sklearn-abalone-process-2020-12-05-17-28-28-509/output/test`

11. Choose the AbaloneTrain step.

12. Find the Amazon S3 path to the model artifact in the **Output** tab, under **Files**:
13. Choose the AbaloneRegisterModel step.

14. Find the ARN of the model package in the Output tab, under Files:

```
arn:aws:sagemaker:eu-west-1:acct-id:model-package/abalonemodelpackagegroupname/2
```

## Kubernetes Orchestration

You can orchestrate your SageMaker training and inference jobs with SageMaker Operators for Kubernetes and SageMaker Components for Kubeflow Pipelines. SageMaker Operators for Kubernetes make it easier for developers and data scientists using Kubernetes to train, tune, and deploy machine learning (ML) models in SageMaker. SageMaker Components for Kubeflow Pipelines allow you to move your data processing and training jobs from the Kubernetes cluster to SageMaker's machine learning-optimized managed service.

### Contents
- [SageMaker Operators for Kubernetes](#)
- [SageMaker Components for Kubeflow Pipelines](#)

## SageMaker Operators for Kubernetes

SageMaker Operators for Kubernetes make it easier for developers and data scientists using Kubernetes to train, tune, and deploy machine learning (ML) models in SageMaker. You can install these SageMaker Operators on your Kubernetes cluster in Amazon Elastic Kubernetes Service (Amazon EKS) to create SageMaker jobs natively using the Kubernetes API and command-line Kubernetes tools such as kubectl. This guide shows how to set up and use the operators to run model training, hyperparameter tuning, or inference (real-time and batch) on SageMaker from a Kubernetes cluster. The procedures and guidelines in this chapter assume that you are familiar with Kubernetes and its basic commands.

⚠️ **Important**

We are stopping the development and technical support of the original version of [SageMaker Operators for Kubernetes](#).
If you are currently using version v1.2.2 or below of SageMaker Operators for Kubernetes, we recommend migrating your resources to the ACK service controller for Amazon SageMaker. The ACK service controller is a new generation of SageMaker Operators for Kubernetes based on AWS Controllers for Kubernetes (ACK). For information on the migration steps, see Migrate resources to the latest Operators. For answers to frequently asked questions on the end of support of the original version of SageMaker Operators for Kubernetes, see Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes.

Note

There is no additional charge to use these operators. You do incur charges for any SageMaker resources that you use through these operators.

What is an operator?

A Kubernetes operator is an application controller managing applications on behalf of a Kubernetes user. Controllers of the control plane encompass various control loops listening to a central state manager (ETCD) to regulate the state of the application they control. Examples of such applications include the Cloud-controller-manager and kube-controller-manager. Operators typically provide a higher-level abstraction than raw Kubernetes API, making it easier for users to deploy and manage applications. To add new capabilities to Kubernetes, developers can extend the Kubernetes API by creating a custom resource that contains their application-specific or domain-specific logic and components. Operators in Kubernetes allow users to natively invoke these custom resources and automate associated workflows.

How does AWS Controllers for Kubernetes (ACK) work?

The SageMaker Operators for Kubernetes allow you to manage jobs in SageMaker from your Kubernetes cluster. The latest version of SageMaker Operators for Kubernetes is based on AWS Controllers for Kubernetes (ACK). ACK includes a common controller runtime, a code generator, and a set of AWS service-specific controllers, one of which is the SageMaker controller.

The following diagram illustrates how ACK works.
In this diagram, a Kubernetes user wants to run model training on SageMaker from within the Kubernetes cluster using the Kubernetes API. The user issues a call to `kubectl apply`, passing in a file that describes a Kubernetes custom resource describing the SageMaker training job. `kubectl apply` passes this file, called a manifest, to the Kubernetes API server running in the Kubernetes controller node (Step 1 in the workflow diagram). The Kubernetes API server receives the manifest with the SageMaker training job specification and determines whether the user has permissions to create a custom resource of kind `sagemaker.services.k8s.aws/TrainingJob`, and whether the custom resource is properly formatted (Step 2). If the user is authorized and the custom resource is valid, the Kubernetes API server writes (Step 3) the custom resource to its etcd data store and then responds back (Step 4) to the user that the custom resource has been created. The SageMaker controller, which is running on a Kubernetes worker node within the context of a normal Kubernetes Pod, is notified (Step 5) that a new custom resource of kind `sagemaker.services.k8s.aws/TrainingJob` has been created. The SageMaker controller then communicates (Step 6) with the SageMaker API, calling the SageMaker `CreateTrainingJob` API to create the training job in AWS. After communicating with the SageMaker API, the SageMaker controller calls the Kubernetes API server to update (Step 7) the custom resource's status with information it received from SageMaker. The SageMaker controller therefore provides the same information to the developers that they would have received using the AWS SDK.
Permissions overview

The operators access SageMaker resources on your behalf. The IAM role that the operator assumes to interact with AWS resources differs from the credentials you use to access the Kubernetes cluster. The role also differs from the role that AWS assumes when running your machine learning jobs.

The following image explains the various authentication layers.
Latest SageMaker Operators for Kubernetes

This section is based on the latest version of SageMaker Operators for Kubernetes using AWS Controllers for Kubernetes (ACK).

⚠️ Important

If you are currently using version v1.2.2 or below of SageMaker Operators for Kubernetes, we recommend migrating your resources to the ACK service controller for Amazon SageMaker. The ACK service controller is a new generation of SageMaker Operators for Kubernetes based on AWS Controllers for Kubernetes (ACK). For information on the migration steps, see Migrate resources to the latest Operators. For answers to frequently asked questions on the end of support of the original version of SageMaker Operators for Kubernetes, see Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes.

The latest version of SageMaker Operators for Kubernetes is based on AWS Controllers for Kubernetes (ACK), a framework for building Kubernetes custom controllers where each controller communicates with an AWS service API. These controllers allow Kubernetes users to provision AWS resources like databases or message queues using the Kubernetes API.

Use the following steps to install and use ACK to train, tune, and deploy machine learning models with Amazon SageMaker.

Contents

- Install SageMaker Operators for Kubernetes
- Use SageMaker Operators for Kubernetes
- Reference

Install SageMaker Operators for Kubernetes

To set up the latest available version of SageMaker Operators for Kubernetes, see the Setup section in Machine Learning with the ACK SageMaker Controller.

Use SageMaker Operators for Kubernetes

For a tutorial on how to train a machine learning model with the ACK service controller for Amazon SageMaker using Amazon EKS, see Machine Learning with the ACK SageMaker Controller.
For an autoscaling example, see Scale SageMaker Workloads with Application Auto Scaling

Reference

See also the ACK service controller for Amazon SageMaker GitHub repository or read AWS Controllers for Kubernetes Documentation.

Old SageMaker Operators for Kubernetes

This section is based on the original version of SageMaker Operators for Kubernetes.

⚠️ Important

We are stopping the development and technical support of the original version of SageMaker Operators for Kubernetes.

If you are currently using version v1.2.2 or below of SageMaker Operators for Kubernetes, we recommend migrating your resources to the ACK service controller for Amazon SageMaker. The ACK service controller is a new generation of SageMaker Operators for Kubernetes based on AWS Controllers for Kubernetes (ACK).

For information on the migration steps, see Migrate resources to the latest Operators.

For answers to frequently asked questions on the end of support of the original version of SageMaker Operators for Kubernetes, see Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes.

Contents

- Install SageMaker Operators for Kubernetes
- Use Amazon SageMaker Jobs
- Migrate resources to the latest Operators
- Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes

Install SageMaker Operators for Kubernetes

Use the following steps to install and use SageMaker Operators for Kubernetes to train, tune, and deploy machine learning models with Amazon SageMaker.

Contents

- IAM role-based setup and operator deployment
IAM role-based setup and operator deployment

The following sections describe the steps to set up and deploy the original version of the operator.

⚠️ Warning

**Reminder:** The following steps do not install the latest version of SageMaker Operators for Kubernetes. To install the new ACK-based SageMaker Operators for Kubernetes, see [Latest SageMaker Operators for Kubernetes](#).

## Prerequisites

This guide assumes that you have completed the following prerequisites:

- Install the following tools on the client machine used to access your Kubernetes cluster:
  - **kubectl** Version 1.13 or later. Use a kubectl version that is within one minor version of your Amazon EKS cluster control plane. For example, a 1.13 kubectl client works with Kubernetes 1.13 and 1.14 clusters. OpenID Connect (OIDC) is not supported in versions earlier than 1.13.
  - **eksctl** Version 0.7.0 or later
  - **AWS CLI** Version 1.16.232 or later
  - (optional) **Helm** Version 3.0 or later
  - **aws-iam-authenticator**
- Have IAM permissions to create roles and attach policies to roles.
- Created a Kubernetes cluster on which to run the operators. It should either be Kubernetes version 1.13 or 1.14. For automated cluster creation using eksctl, see [Getting Started with eksctl]. It takes 20–30 minutes to provision a cluster.

## Cluster-scoped deployment

Before you can deploy your operator using an IAM role, associate an OpenID Connect (OIDC) Identity Provider (IdP) with your role to authenticate with the IAM service.
Create an OIDC provider for your cluster

The following instructions show how to create and associate an OIDC provider with your Amazon EKS cluster.

1. Set the local CLUSTER_NAME and AWS_REGION environment variables as follows:

   ```shell
   # Set the Region and cluster
   export CLUSTER_NAME="<your cluster name>"
   export AWS_REGION="<your region>"
   ```

2. Use the following command to associate the OIDC provider with your cluster. For more information, see [Enabling IAM Roles for Service Accounts on your Cluster](#).

   ```shell
   eksctl utils associate-iam-oidc-provider --cluster ${CLUSTER_NAME} \
   --region ${AWS_REGION} --approve
   ```

   Your output should look like the following:

   ```text
   [✓] eksctl version 0.10.1
   [✓] using region us-east-1
   [✓] IAM OpenID Connect provider is associated with cluster "my-cluster" in "us-east-1"
   ```

Now that the cluster has an OIDC identity provider, you can create a role and give a Kubernetes ServiceAccount permission to assume the role.

Get the OIDC ID

To set up the ServiceAccount, obtain the OIDC issuer URL using the following command:

```shell
aws eks describe-cluster --name ${CLUSTER_NAME} --region ${AWS_REGION} \
--query cluster.identity.oidc.issuer --output text
```

The command returns a URL like the following:

```text
https://oidc.eks.${AWS_REGION}.amazonaws.com/id/D48675832CA65BD10A532F5970IDCID
```

In this URL, the value D48675832CA65BD10A532F5970IDCID is the OIDC ID. The OIDC ID for your cluster is different. You need this OIDC ID value to create a role.
If your output is None, it means that your client version is old. To work around this, run the following command:

```
aws eks describe-cluster --region ${AWS_REGION} --query cluster --name ${CLUSTER_NAME} --output text | grep OIDC
```

The OIDC URL is returned as follows:

```
OIDC https://oidc.eks.us-east-1.amazonaws.com/id/D48675832CA65BD10A532F59701D01D
```

**Create an IAM role**

1. Create a file named `trust.json` and insert the following trust relationship code block into it. Be sure to replace all `<OIDC ID>`, `<AWS account number>`, and `<EKS Cluster region>` placeholders with values corresponding to your cluster.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Federated": "arn:aws:iam::<AWS account number>:oidc-provider/oidc.eks.<EKS Cluster region>.amazonaws.com/id/<OIDC ID>"
            },
            "Action": "sts:AssumeRoleWithWebIdentity",
            "Condition": {
                "StringEquals": {
                    "oidc.eks.<EKS Cluster region>.amazonaws.com/id/<OIDC ID>:aud": "sts.amazonaws.com",
                }
            }
        }
    ]
}
```

2. Run the following command to create a role with the trust relationship defined in `trust.json`. This role allows the Amazon EKS cluster to get and refresh credentials from IAM.
aws iam create-role --region ${AWS_REGION} --role-name <role name> --assume-role-policy-document file://trust.json --output=text

Your output should look like the following:

ABCDDEFSDN7EXAMPLE    my-role
ASSUMEROLEPOLICYDOCUMENT    2012-10-17
STATEMENT    sts:AssumeRoleWithWebIdentity    Allow
PRINCIPAL    arn:aws:iam::123456789012:oidc-provider/oidc.eks.us-east-1.amazonaws.com/id/

Take note of ROLE ARN; you pass this value to your operator.

Attach the AmazonSageMakerFullAccess policy to the role

To give the role access to SageMaker, attach the AmazonSageMakerFullAccess policy. If you want to limit permissions to the operator, you can create your own custom policy and attach it.

To attach AmazonSageMakerFullAccess, run the following command:

aws iam attach-role-policy --role-name <role name> --policy-arn arn:aws:iam::aws:policy/AmazonSageMakerFullAccess

The Kubernetes ServiceAccount sagemaker-k8s-operator-default should have AmazonSageMakerFullAccess permissions. Confirm this when you install the operator.

Deploy the operator

When deploying your operator, you can use either a YAML file or Helm charts.

Deploy the operator using YAML

This is the simplest way to deploy your operators. The process is as follows:

1. Download the installer script using the following command:

2. Edit the installer.yaml file to replace eks.amazonaws.com/role-arn. Replace the ARN here with the Amazon Resource Name (ARN) for the OIDC-based role you’ve created.

3. Use the following command to deploy the cluster:

   kubectl apply -f installer.yaml

**Deploy the operator using Helm Charts**

Use the provided Helm Chart to install the operator.

1. Clone the Helm installer directory using the following command:

   git clone https://github.com/aws/amazon-sagemaker-operator-for-k8s.git

2. Navigate to the amazon-sagemaker-operator-for-k8s/hack/charts/installer folder. Edit the rolebased/values.yaml file, which includes high-level parameters for the chart. Replace the role ARN here with the Amazon Resource Name (ARN) for the OIDC-based role you’ve created.

3. Install the Helm Chart using the following command:

   kubectl create namespace sagemaker-k8s-operator-system
   helm install --namespace sagemaker-k8s-operator-system sagemaker-operator rolebased/

   If you decide to install the operator into a namespace other than the one specified, you need to adjust the namespace defined in the IAM role trust.json file to match.

4. After a moment, the chart is installed with a randomly generated name. Verify that the installation succeeded by running the following command:

   helm ls

   Your output should look like the following:
<table>
<thead>
<tr>
<th>NAME</th>
<th>NAMESPACE</th>
<th>REVISION</th>
<th>UPDATED</th>
<th>STATUS</th>
<th>CHART</th>
<th>APP</th>
</tr>
</thead>
<tbody>
<tr>
<td>sagemaker-operator</td>
<td>sagemaker-k8s-operator-system</td>
<td>1</td>
<td>2019-11-20 23:14:59.6777082 +0000 UTC</td>
<td>deployed</td>
<td>sagemaker-k8s-operator-0.1.0</td>
<td></td>
</tr>
</tbody>
</table>

Verify the operator deployment

1. You should be able to see the SageMaker Custom Resource Definitions (CRDs) for each operator deployed to your cluster by running the following command:

   kubectl get crd | grep sagemaker

Your output should look like the following:

```
batchtransformjobs.sagemaker.aws.amazon.com   2019-11-20T17:12:34Z
endpointconfigs.sagemaker.aws.amazon.com      2019-11-20T17:12:34Z
hostingdeployments.sagemaker.aws.amazon.com   2019-11-20T17:12:34Z
hyperparametertuningjobs.sagemaker.aws.amazon.com 2019-11-20T17:12:34Z
models.sagemaker.aws.amazon.com               2019-11-20T17:12:34Z
trainingjobs.sagemaker.aws.amazon.com         2019-11-20T17:12:34Z
```

2. Ensure that the operator pod is running successfully. Use the following command to list all pods:

   kubectl -n sagemaker-k8s-operator-system get pods

You should see a pod named sagemaker-k8s-operator-controller-manager-***** in the namespace sagemaker-k8s-operator-system as follows:

<table>
<thead>
<tr>
<th>NAME</th>
<th>AGE</th>
<th>READY</th>
<th>STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>sagemaker-k8s-operator-controller-manager-12345678-r8abc</td>
<td>23s</td>
<td>2/2</td>
<td>Running 0</td>
</tr>
</tbody>
</table>
Namespace-scoped deployment

You have the option to install your operator within the scope of an individual Kubernetes namespace. In this mode, the controller only monitors and reconciles resources with SageMaker if the resources are created within that namespace. This allows for finer-grained control over which controller is managing which resources. This is useful for deploying to multiple AWS accounts or controlling which users have access to particular jobs.

This guide outlines how to install an operator into a particular, predefined namespace. To deploy a controller into a second namespace, follow the guide from beginning to end and change out the namespace in each step.

Create an OIDC provider for your Amazon EKS cluster

The following instructions show how to create and associate an OIDC provider with your Amazon EKS cluster.

1. Set the local CLUSTER_NAME and AWS_REGION environment variables as follows:

```
# Set the Region and cluster
export CLUSTER_NAME="<your cluster name>"
export AWS_REGION="<your region>"
```

2. Use the following command to associate the OIDC provider with your cluster. For more information, see Enabling IAM Roles for Service Accounts on your Cluster.

```
eksctl utils associate-iam-oidc-provider --cluster ${CLUSTER_NAME} --region ${AWS_REGION} --approve
```

Your output should look like the following:

```
[] eksctl version 0.10.1
[.] using region us-east-1
[.] IAM OpenID Connect provider is associated with cluster "my-cluster" in "us-east-1"
```

Now that the cluster has an OIDC identity provider, create a role and give a Kubernetes ServiceAccount permission to assume the role.
Get your OIDC ID

To set up the ServiceAccount, first obtain the OpenID Connect issuer URL using the following command:

```bash
aws eks describe-cluster --name ${CLUSTER_NAME} --region ${AWS_REGION} \
--query cluster.identity.oidc.issuer --output text
```

The command returns a URL like the following:

```bash
https://oidc.eks.${AWS_REGION}.amazonaws.com/id/D48675832CA65BD10A532F597OIDCID
```

In this URL, the value D48675832CA65BD10A532F597OIDCID is the OIDC ID. The OIDC ID for your cluster is different. You need this OIDC ID value to create a role.

If your output is None, it means that your client version is old. To work around this, run the following command:

```bash
aws eks describe-cluster --region ${AWS_REGION} --query cluster --name ${CLUSTER_NAME} \
--output text | grep OIDC
```

The OIDC URL is returned as follows:

```bash
OIDC https://oidc.eks.us-east-1.amazonaws.com/id/D48675832CA65BD10A532F597OIDCID
```

Create your IAM role

1. Create a file named `trust.json` and insert the following trust relationship code block into it. Be sure to replace all `<OIDC ID>`, `<AWS account number>`, `<EKS Cluster region>`, and `<Namespace>` placeholders with values corresponding to your cluster. For the purposes of this guide, `my-namespace` is used for the `<Namespace>` value.

   ```json
   {
      "Version": "2012-10-17",
      "Statement": [
      {
          "Effect": "Allow",
          "Principal": {
              "Federated": "arn:aws:iam::<AWS account number>:oidc-provider/oidc.eks.<EKS Cluster region>.amazonaws.com/id/<OIDC ID>"
          },
      }
   }
   ```
2. Run the following command to create a role with the trust relationship defined in trust.json. This role allows the Amazon EKS cluster to get and refresh credentials from IAM.

```
aws iam create-role --region ${AWS_REGION} --role-name <role name> --assume-role-policy-document file://trust.json --output=text
```

Your output should look like the following:

```
ASSUMEROLEPOLICYDOCUMENT        2012-10-17
STATEMENT   sts:AssumeRoleWithWebIdentity   Allow
STRINGEQUALS    sts.amazonaws.com       system:serviceaccount:my-namespace:sagemaker-k8s-operator-default
PRINCIPAL    arn:aws:iam::123456789012:oidc-provider/oidc.eks.us-east-1.amazonaws.com/id/
```

Take note of ROLE ARN. You pass this value to your operator.

**Attach the AmazonSageMakerFullAccess policy to your role**

To give the role access to SageMaker, attach the [AmazonSageMakerFullAccess](#) policy. If you want to limit permissions to the operator, you can create your own custom policy and attach it.

To attach AmazonSageMakerFullAccess, run the following command:

```
aws iam attach-role-policy --role-name <role name> --policy-arn arn:aws:iam::aws:policy/AmazonSageMakerFullAccess
```
The Kubernetes ServiceAccount sagemaker-k8s-operator-default should have AmazonSageMakerFullAccess permissions. Confirm this when you install the operator.

**Deploy the operator to your namespace**

When deploying your operator, you can use either a YAML file or Helm charts.

**Deploy the operator to your namespace using YAML**

There are two parts to deploying an operator within the scope of a namespace. The first is the set of CRDs that are installed at a cluster level. These resource definitions only need to be installed once per Kubernetes cluster. The second part is the operator permissions and deployment itself.

If you have not already installed the CRDs into the cluster, apply the CRD installer YAML using the following command:

```
kubectl apply -f https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/release/rolebased/namespaced/crd.yaml
```

To install the operator onto the cluster:

1. **Download the operator installer YAML using the following command:**

   ```
   ```

2. **Update the installer YAML to place the resources into your specified namespace using the following command:**

   ```
   sed -i -e 's/PLACEHOLDER-NAMESPACE/<YOUR NAMESPACE>/g' operator.yaml
   ```

3. **Edit the operator.yaml file to place resources into your eks.amazonaws.com/role-arn. Replace the ARN here with the Amazon Resource Name (ARN) for the OIDC-based role you've created.**

4. **Use the following command to deploy the cluster:**

   ```
   kubectl apply -f operator.yaml
   ```
Deploy the operator to your namespace using Helm Charts

There are two parts needed to deploy an operator within the scope of a namespace. The first is the set of CRDs that are installed at a cluster level. These resource definitions only need to be installed once per Kubernetes cluster. The second part is the operator permissions and deployment itself. When using Helm Charts you have to first create the namespace using `kubectl`.

1. Clone the Helm installer directory using the following command:

```bash
git clone https://github.com/aws/amazon-sagemaker-operator-for-k8s.git
```

2. Navigate to the `amazon-sagemaker-operator-for-k8s/hack/charts/installer/namespaced` folder. Edit the `rolebased/values.yaml` file, which includes high-level parameters for the chart. Replace the role ARN here with the Amazon Resource Name (ARN) for the OIDC-based role you've created.

3. Install the Helm Chart using the following command:

```bash
helm install crds crd_chart/
```

4. Create the required namespace and install the operator using the following command:

```bash
kubectl create namespace <namespace>
helm install --n <namespace> op operator_chart/
```

5. After a moment, the chart is installed with the name `sagemaker-operator`. Verify that the installation succeeded by running the following command:

```bash
helm ls
```

Your output should look like the following:

<table>
<thead>
<tr>
<th>NAME</th>
<th>NAMESPACE</th>
<th>REVISION</th>
<th>UPDATED</th>
<th>CHART</th>
</tr>
</thead>
<tbody>
<tr>
<td>sagemaker-operator</td>
<td>my-namespace</td>
<td>1</td>
<td>2019-11-20</td>
<td>sagemaker-k8s-operator-0.1.0</td>
</tr>
</tbody>
</table>
**Verify the operator deployment to your namespace**

1. You should be able to see the SageMaker Custom Resource Definitions (CRDs) for each operator deployed to your cluster by running the following command:

   ```bash
ekubectl get crd | grep sagemaker
   ```

   Your output should look like the following:

   ```
   batchtransformjobs.sagemaker.aws.amazon.com         2019-11-20T17:12:34Z
   endpointconfigs.sagemaker.aws.amazon.com            2019-11-20T17:12:34Z
   hostingdeployments.sagemaker.aws.amazon.com         2019-11-20T17:12:34Z
   hyperparametertuningjobs.sagemaker.aws.amazon.com   2019-11-20T17:12:34Z
   models.sagemaker.aws.amazon.com                     2019-11-20T17:12:34Z
   trainingjobs.sagemaker.aws.amazon.com               2019-11-20T17:12:34Z
   ```

2. Ensure that the operator pod is running successfully. Use the following command to list all pods:

   ```bash
   kubectl -n my-namespace get pods
   ```

   You should see a pod named `sagemaker-k8s-operator-controller-manager-*****` in the namespace `my-namespace` as follows:

   ```
   NAME                                                         READY   STATUS      AGE
   sagemaker-k8s-operator-controller-manager-12345678-r8abc     2/2     Running   23s
   ```

**Install the SageMaker logs kubectl plugin**

As part of the SageMaker Operators for Kubernetes, you can use the `smlogs` plugin for `kubectl`. This allows SageMaker CloudWatch logs to be streamed with `kubectl`. `kubectl` must be installed onto your PATH. The following commands place the binary in the `sagemaker-k8s-bin` directory in your home directory, and add that directory to your PATH.

```bash
export os="linux"
```
```
wget https://amazon-sagemaker-operator-for-k8s-us-east-1.s3.amazonaws.com/kubectl-smlogs-plugin/v1/${os}.amd64.tar.gz
  tar xvzf ${os}.amd64.tar.gz

# Move binaries to a directory in your homedir.
  mkdir ~/sagemaker-k8s-bin
  cp ./kubectl-smlogs.${os}.amd64/kubectl-smlogs ~/sagemaker-k8s-bin/.

# This line adds the binaries to your PATH in your .bashrc.
  echo 'export PATH=$PATH:~/sagemaker-k8s-bin' >> ~/.bashrc

# Source your .bashrc to update environment variables:
  source ~/.bashrc
```

Use the following command to verify that the kubectl plugin is installed correctly:

```
kubectl smlogs
```

If the kubectl plugin is installed correctly, your output should look like the following:

---

**View SageMaker logs via Kubernetes**

**Usage:**

```
smlogs [command]
```

**Aliases:**

```
smlogs, SMLogs, Smlogs
```

**Available Commands:**

<table>
<thead>
<tr>
<th>Command</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BatchTransformJob</td>
<td>View BatchTransformJob logs via Kubernetes</td>
</tr>
<tr>
<td>TrainingJob</td>
<td>View TrainingJob logs via Kubernetes</td>
</tr>
<tr>
<td>help</td>
<td>Help about any command</td>
</tr>
</tbody>
</table>

**Flags:**

```
-h, --help  help for smlogs
```

Use "smlogs [command] --help" for more information about a command.
Clean up resources

To uninstall the operator from your cluster, you must first make sure to delete all SageMaker resources from the cluster. Failure to do so causes the operator delete operation to hang. Run the following commands to stop all jobs:

```
# Delete all SageMaker jobs from Kubernetes
kubectl delete --all --all-namespaces hyperparametertuningjob.sagemaker.aws.amazon.com
kubectl delete --all --all-namespaces trainingjobs.sagemaker.aws.amazon.com
kubectl delete --all --all-namespaces batchtransformjob.sagemaker.aws.amazon.com
kubectl delete --all --all-namespaces hostingdeployment.sagemaker.aws.amazon.com
```

You should see output similar to the following:

```
$ kubectl delete --all --all-namespaces trainingjobs.sagemaker.aws.amazon.com
trainingjobs.sagemaker.aws.amazon.com "xgboost-mnist-from-for-s3" deleted

$ kubectl delete --all --all-namespaces hyperparametertuningjob.sagemaker.aws.amazon.com
hyperparametertuningjob.sagemaker.aws.amazon.com "xgboost-mnist-hpo" deleted

$ kubectl delete --all --all-namespaces batchtransformjob.sagemaker.aws.amazon.com
batchtransformjob.sagemaker.aws.amazon.com "xgboost-mnist" deleted

$ kubectl delete --all --all-namespaces hostingdeployment.sagemaker.aws.amazon.com
hostingdeployment.sagemaker.aws.amazon.com "host-xgboost" deleted
```

After you delete all SageMaker jobs, see Delete operators to delete the operator from your cluster.

Delete operators

Delete cluster-based operators

Operators installed using YAML

To uninstall the operator from your cluster, make sure that all SageMaker resources have been deleted from the cluster. Failure to do so causes the operator delete operation to hang.

⚠️ Note
Before deleting your cluster, be sure to delete all SageMaker resources from the cluster. See Clean up resources for more information.
After you delete all SageMaker jobs, use `kubectl` to delete the operator from the cluster:

```bash
# Delete the operator and its resources
kubectl delete -f /installer.yaml
```

You should see output similar to the following:

```bash
$ kubectl delete -f raw-yaml/installer.yaml
namespace "sagemaker-k8s-operator-system" deleted
customresourcedefinition.apiextensions.k8s.io "batchtransformjobs.sagemaker.aws.amazon.com" deleted
customresourcedefinition.apiextensions.k8s.io "endpointconfigs.sagemaker.aws.amazon.com" deleted
customresourcedefinition.apiextensions.k8s.io "hostingdeployments.sagemaker.aws.amazon.com" deleted
customresourcedefinition.apiextensions.k8s.io "hyperparametertuningjobs.sagemaker.aws.amazon.com" deleted
customresourcedefinition.apiextensions.k8s.io "models.sagemaker.aws.amazon.com" deleted
customresourcedefinition.apiextensions.k8s.io "trainingjobs.sagemaker.aws.amazon.com" deleted
role.rbac.authorization.k8s.io "sagemaker-k8s-operator-leader-election-role" deleted
clusterrole.rbac.authorization.k8s.io "sagemaker-k8s-operator-manager-role" deleted
clusterrole.rbac.authorization.k8s.io "sagemaker-k8s-operator-proxy-role" deleted
rolebinding.rbac.authorization.k8s.io "sagemaker-k8s-operator-leader-election-rolebinding" deleted
clusterrolebinding.rbac.authorization.k8s.io "sagemaker-k8s-operator-manager-rolebinding" deleted
clusterrolebinding.rbac.authorization.k8s.io "sagemaker-k8s-operator-proxy-rolebinding" deleted
service "sagemaker-k8s-operator-controller-manager-metrics-service" deleted
deployment.apps "sagemaker-k8s-operator-controller-manager" deleted
secrets "sagemaker-k8s-operator-abcde" deleted
```

**Operators installed using Helm Charts**

To delete the operator CRDs, first delete all the running jobs. Then delete the Helm Chart that was used to deploy the operators using the following commands:

```bash
# get the helm charts
helm ls

# delete the charts
```
Delete namespace-based operators

Operators installed with YAML

To uninstall the operator from your cluster, first make sure that all SageMaker resources have been deleted from the cluster. Failure to do so causes the operator delete operation to hang.

⚠️ Note
Before deleting your cluster, be sure to delete all SageMaker resources from the cluster. See Clean up resources for more information.

After you delete all SageMaker jobs, use kubectl to first delete the operator from the namespace and then the CRDs from the cluster. Run the following commands to delete the operator from the cluster:

```bash
# Delete the operator using the same yaml file that was used to install the operator
kubectl delete -f operator.yaml

# Now delete the CRDs using the CRD installer yaml

# Now you can delete the namespace if you want
kubectl delete namespace <namespace>
```

Operators installed with Helm Charts

To delete the operator CRDs, first delete all the running jobs. Then delete the Helm Chart that was used to deploy the operators using the following commands:

```bash
# Delete the operator
helm delete <chart_name>

# delete the crds
helm delete crds
```
# optionally delete the namespace
kubect1 delete namespace <namespace>

## Troubleshooting

### Debugging a failed job

Use these steps to debug a failed job.

- Check the job status by running the following:

  ```bash
kubectl get <CRD Type> <job name>
  ```

- If the job was created in SageMaker, you can use the following command to see the STATUS and the SageMaker Job Name:

  ```bash
kubectl get <crd type> <job name>
  ```

- You can use `smlogs` to find the cause of the issue using the following command:

  ```bash
kubect1 smlogs <crd type> <job name>
  ```

- You can also use the `describe` command to get more details about the job using the following command. The output has an additional field that has more information about the status of the job.

  ```bash
kubect1 describe <crd type> <job name>
  ```

- If the job was not created in SageMaker, then use the logs of the operator's pod to find the cause of the issue as follows:

  ```bash
$ kubectl get pods -A | grep sagemaker
  
  # Output:
  sagemaker-k8s-operator-system   sagemaker-k8s-operator-controller-manager-5cd7df4d74-wh22z   2/2     Running   0          3h33m

$ kubectl logs -p <pod name> -c manager -n sagemaker-k8s-operator-system
  ```
Deleting an operator CRD

If deleting a job is not working, check if the operator is running. If the operator is not running, then you have to delete the finalizer using the following steps:

1. In a new terminal, open the job in an editor using `kubectl edit` as follows:

```
kubectl edit <crd type> <job name>
```

2. Edit the job to delete the finalizer by removing the following two lines from the file. Save the file and the job is be deleted.

```
finalizers:
- sagemaker-operator-finalizer
```

Images and SMlogs in each Region

The following table lists the available operator images and SMLogs in each Region.

<table>
<thead>
<tr>
<th>Region</th>
<th>Controller Image</th>
<th>Linux SMLogs</th>
</tr>
</thead>
</table>
Use Amazon SageMaker Jobs

This section is based on the original version of SageMaker Operators for Kubernetes.

⚠️ Important

We are stopping the development and technical support of the original version of SageMaker Operators for Kubernetes.

If you are currently using version v1.2.2 or below of SageMaker Operators for Kubernetes, we recommend migrating your resources to the ACK service controller for Amazon SageMaker. The ACK service controller is a new generation of SageMaker Operators for Kubernetes based on AWS Controllers for Kubernetes (ACK).

For information on the migration steps, see Migrate resources to the latest Operators.

For answers to frequently asked questions on the end of support of the original version of SageMaker Operators for Kubernetes, see Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes.

To run an Amazon SageMaker job using the Operators for Kubernetes, you can either apply a YAML file or use the supplied Helm Charts.

All sample operator jobs in the following tutorials use sample data taken from a public MNIST dataset. In order to run these samples, download the dataset into your Amazon S3 bucket. You can find the dataset in Download the MNIST Dataset.

Contents

- The TrainingJob operator
- The HyperParameterTuningJob operator
- The BatchTransformJob operator
- The HostingDeployment operator
- The ProcessingJob operator
- HostingAutoscalingPolicy (HAP) Operator
The TrainingJob operator

Training job operators reconcile your specified training job spec to SageMaker by launching it for you in SageMaker. You can learn more about SageMaker training jobs in the SageMaker CreateTrainingJob API documentation.

Topics

- Create a TrainingJob using a YAML file
- Create a TrainingJob Using a Helm Chart
- List TrainingJobs
- Describe a TrainingJob
- View logs from TrainingJobs
- Delete TrainingJobs

Create a TrainingJob using a YAML file

1. Download the sample YAML file for training using the following command:

   ```
   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/xgboost-mnist-trainingjob.yaml
   ```

2. Edit the xgboost-mnist-trainingjob.yaml file to replace the roleArn parameter with your `<sagemaker-execution-role>`, and outputPath with your Amazon S3 bucket to which the SageMaker execution role has write access. The roleArn must have permissions so that SageMaker can access Amazon S3, Amazon CloudWatch, and other services on your behalf. For more information on creating an SageMaker ExecutionRole, see SageMaker Roles. Apply the YAML file using the following command:

   ```
   kubectl apply -f xgboost-mnist-trainingjob.yaml
   ```

Create a TrainingJob Using a Helm Chart

You can use Helm Charts to run TrainingJobs.

1. Clone the GitHub repository to get the source using the following command:

   ```
   git clone https://github.com/aws/amazon-sagemaker-operator-for-k8s.git
   ```
2. Navigate to the `amazon-sagemaker-operator-for-k8s/hack/charts/training-jobs/` folder and edit the `values.yaml` file to replace values like `rolearn` and `outputpath` with values that correspond to your account. The RoleARN must have permissions so that SageMaker can access Amazon S3, Amazon CloudWatch, and other services on your behalf. For more information on creating an SageMaker ExecutionRole, see [SageMaker Roles](#).

Create the TrainingJob

With the roles and Amazon S3 buckets replaced with appropriate values in `values.yaml`, you can create a training job using the following command:

```
helm install . --generate-name
```

Your output should look like the following:

```
NAME: chart-12345678
LAST DEPLOYED: Wed Nov 20 23:35:49 2019
NAMESPACE: default
STATUS: deployed
REVISION: 1
TEST SUITE: None
NOTES:
Thanks for installing the sagemaker-k8s-trainingjob.
```

Verify your training Helm Chart

To verify that the Helm Chart was created successfully, run:

```
helm ls
```

Your output should look like the following:

```
NAME                    NAMESPACE       REVISION        UPDATED
STATUS          CHART                           APP VERSION
chart-12345678        default         1               2019-11-20 23:35:49.9136092 +0000
                     chart-12345678 UTC deployed   sagemaker-k8s-trainingjob-0.1.0
rolebased-12345678    default         1               2019-11-20 23:14:59.6777082 +0000
                     rolebased-12345678 UTC deployed    sagemaker-k8s-operator-0.1.0
```

Kubernetes Orchestration
Helm install creates a TrainingJob Kubernetes resource. The operator launches the actual training job in SageMaker and updates the TrainingJob Kubernetes resource to reflect the status of the job in SageMaker. You incur charges for SageMaker resources used during the duration of your job. You do not incur any charges once your job completes or stops.

**Note:** SageMaker does not allow you to update a running training job. You cannot edit any parameter and re-apply the config file. Either change the metadata name or delete the existing job and create a new one. Similar to existing training job operators like TFJob in Kubeflow, update is not supported.

**List TrainingJobs**

Use the following command to list all jobs created using the Kubernetes operator:

```
kubectl get TrainingJob
```

The output listing all jobs should look like the following:

```
kubectl get trainingjobs
NAME                        STATUS       SECONDARY-STATUS   CREATION-TIME
SAGEMAKER-JOB-NAME
xgboost-mnist-from-for-s3   InProgress   Starting           2019-11-20T23:42:35Z
xgboost-mnist-from-for-s3-examplef11eab94e0ed4671d5a8f
```

A training job continues to be listed after the job has completed or failed. You can remove a TrainingJob job from the list by following the [Delete TrainingJobs](#) steps. Jobs that have completed or stopped do not incur any charges for SageMaker resources.

**TrainingJob status values**

The STATUS field can be one of the following values:

- Completed
- InProgress
- Failed
- Stopped
- Stopping

These statuses come directly from the SageMaker official [API documentation](#).
In addition to the official SageMaker status, it is possible for STATUS to be `SynchronizingK8sJobWithSageMaker`. This means that the operator has not yet processed the job.

**Secondary status values**

The secondary statuses come directly from the SageMaker official [API documentation](#). They contain more granular information about the status of the job.

**Describe a TrainingJob**

You can get more details about the training job by using the `describe kubectl` command. This is typically used for debugging a problem or checking the parameters of a training job. To get information about your training job, use the following command:

```
kubectl describe trainingjob xgboost-mnist-from-for-s3
```

The output for your training job should look like the following:

<table>
<thead>
<tr>
<th>Name:</th>
<th>xgboost-mnist-from-for-s3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Namespace:</td>
<td>default</td>
</tr>
<tr>
<td>Labels:</td>
<td>&lt;none&gt;</td>
</tr>
<tr>
<td>Annotations:</td>
<td>&lt;none&gt;</td>
</tr>
<tr>
<td>API Version:</td>
<td>sagemaker.aws.amazon.com/v1</td>
</tr>
<tr>
<td>Kind:</td>
<td>TrainingJob</td>
</tr>
<tr>
<td>Metadata:</td>
<td></td>
</tr>
<tr>
<td>Finalizers:</td>
<td>sagemaker-operator-finalizer</td>
</tr>
<tr>
<td>Generation:</td>
<td>2</td>
</tr>
<tr>
<td>Resource Version:</td>
<td>23119</td>
</tr>
<tr>
<td>Self Link:</td>
<td>/apis/sagemaker.aws.amazon.com/v1/namespaces/default/trainingjobs/xgboost-mnist-from-for-s3</td>
</tr>
<tr>
<td>UID:</td>
<td>6d7uiui-0bef-11ea-b94e-0ed467example</td>
</tr>
<tr>
<td>Spec:</td>
<td></td>
</tr>
<tr>
<td>Algorithm Specification:</td>
<td></td>
</tr>
<tr>
<td>Training Image:</td>
<td>8256416981234.dkr.ecr.us-east-2.amazonaws.com/xgboost:1</td>
</tr>
<tr>
<td>Training Input Mode:</td>
<td>File</td>
</tr>
<tr>
<td>Hyper Parameters:</td>
<td></td>
</tr>
<tr>
<td>Name:</td>
<td>eta</td>
</tr>
<tr>
<td>Value:</td>
<td>0.2</td>
</tr>
<tr>
<td>Name:</td>
<td>gamma</td>
</tr>
<tr>
<td>Value:</td>
<td>4</td>
</tr>
</tbody>
</table>
Name:  max_depth
Value:  5
Name:  min_child_weight
Value:  6
Name:  num_class
Value:  10
Name:  num_round
Value:  10
Name:  objective
Value:  multi:softmax
Name:  silent
Value:  0

Input Data Config:
Channel Name:  train
Compression Type:  None
Content Type:  text/csv
Data Source:
  S3 Data Source:
    S3 Data Distribution Type:  FullyReplicated
    S3 Data Type:  S3Prefix
    S3 Uri:  https://s3-us-east-2.amazonaws.com/my-bucket/sagemaker/xgboost-mnist/train/

Channel Name:  validation
Compression Type:  None
Content Type:  text/csv
Data Source:
  S3 Data Source:
    S3 Data Distribution Type:  FullyReplicated
    S3 Data Type:  S3Prefix
    S3 Uri:  https://s3-us-east-2.amazonaws.com/my-bucket/sagemaker/xgboost-mnist/validation/

Output Data Config:
S3 Output Path:  s3://my-bucket/sagemaker/xgboost-mnist/xgboost/
Region:  us-east-2

Resource Config:
Instance Count:  1
Instance Type:  ml.m4.xlarge
Volume Size In GB:  5
Role Arn:  arn:aws:iam::12345678910:role/service-role/AmazonSageMaker-ExecutionRole

Stopping Condition:
Max Runtime In Seconds:  86400
Training Job Name:  xgboost-mnist-from-for-s3-6d7fa0af0bef11eab94e0example

Status:
View logs from TrainingJobs

Use the following command to see the logs from the kmeans-mnist training job:

```
kubectl smlogs trainingjob xgboost-mnist-from-for-s3
```

Your output should look similar to the following. The logs from instances are ordered chronologically.

"xgboost-mnist-from-for-s3" has SageMaker TrainingJobName "xgboost-mnist-from-for-s3-123456789" in region "us-east-2", status "InProgress" and secondary status "Starting"
xgboost-mnist-from-for-s3-6d7fa0af0bef11eab94e0ed46example/algo-1-1574293123 2019-11-20 23:45:24.7 +0000 UTC Arguments: train
xgboost-mnist-from-for-s3-6d7fa0af0bef11eab94e0ed46example/algo-1-1574293123 2019-11-20 23:45:24.7 +0000 UTC [2019-11-20:23:45:22:INFO] File size need to be processed in the node: 1122.95mb. Available memory size in the node: 8586.0mb
xgboost-mnist-from-for-s3-6d7fa0af0bef11eab94e0ed46example/algo-1-1574293123 2019-11-20 23:45:24.7 +0000 UTC [2019-11-20:23:45:22:INFO] Determined delimiter of CSV input is ','
xgboost-mnist-from-for-s3-6d7fa0af0bef11eab94e0ed46example/algo-1-1574293123 2019-11-20 23:45:24.7 +0000 UTC [23:45:22] S3DistributionType set as FullyReplicated

Delete TrainingJobs

Use the following command to stop a training job on Amazon SageMaker:

```
kubectl delete trainingjob xgboost-mnist-from-for-s3
```

This command removes the SageMaker training job from Kubernetes. This command returns the following output:
If the job is still in progress on SageMaker, the job stops. You do not incur any charges for SageMaker resources after your job stops or completes.

**Note**: SageMaker does not delete training jobs. Stopped jobs continue to show on the SageMaker console. The delete command takes about 2 minutes to clean up the resources from SageMaker.

**The HyperParameterTuningJob operator**

Hyperparameter tuning job operators reconcile your specified hyperparameter tuning job spec to SageMaker by launching it in SageMaker. You can learn more about SageMaker hyperparameter tuning jobs in the SageMaker [CreateHyperParameterTuningJob API documentation](#).

**Topics**

- [Create a HyperparameterTuningJob using a YAML file](#)
- [Create a HyperparameterTuningJob using a Helm Chart](#)
- [List HyperparameterTuningJobs](#)
- [Describe a HyperparameterTuningJob](#)
- [View logs from HyperparameterTuningJobs](#)
- [Delete a HyperparameterTuningJob](#)

**Create a HyperparameterTuningJob using a YAML file**

1. Download the sample YAML file for the hyperparameter tuning job using the following command:

   ```bash
   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/xgboost-mnist-hpo.yaml
   ```

2. Edit the `xgboost-mnist-hpo.yaml` file to replace the `roleArn` parameter with your `sagemaker-execution-role`. For the hyperparameter tuning job to succeed, you must also change the `s3InputPath` and `s3OutputPath` to values that correspond to your account. Apply the updates YAML file using the following command:

   ```bash
   kubectl apply -f xgboost-mnist-hpo.yaml
   ```
Create a HyperparameterTuningJob using a Helm Chart

You can use Helm Charts to run hyperparameter tuning jobs.

1. Clone the GitHub repository to get the source using the following command:

   ```
   git clone https://github.com/aws/amazon-sagemaker-operator-for-k8s.git
   ```

2. Navigate to the `amazon-sagemaker-operator-for-k8s/hack/charts/hyperparameter-tuning-jobs/` folder.

3. Edit the `values.yaml` file to replace the `roleArn` parameter with your `sagemaker-execution-role`. For the hyperparameter tuning job to succeed, you must also change the `s3InputPath` and `s3OutputPath` to values that correspond to your account.

Create the HyperparameterTuningJob

With the roles and Amazon S3 paths replaced with appropriate values in `values.yaml`, you can create a hyperparameter tuning job using the following command:

```
helm install . --generate-name
```

Your output should look similar to the following:

```
NAME: chart-1574292948
LAST DEPLOYED: Wed Nov 20 23:35:49 2019
NAMESPACE: default
STATUS: deployed
REVISION: 1
TEST SUITE: None
NOTES:
Thanks for installing the sagemaker-k8s-hyperparametertuningjob.
```

Verify chart installation

To verify that the Helm Chart was created successfully, run the following command:

```
helm ls
```

Your output should look like the following:
**NAME**  | **NAMESPACE**  | **REVISION**  | **UPDATED**  
--- | --- | --- | ---  
chart-1474292948 | default | 1 | 2019-11-20 23:35:49.913609+0000 UTC  
| | | | deployed  
| | | | sagemaker-k8s-hyperparametertuningjob-0.1.0  
| chart-1574292948 | default | 1 | 2019-11-20 23:35:49.913609+0000 UTC  
| | | | deployed  
| | | | sagemaker-k8s-trainingjob-0.1.0  
| rolebased-1574291698 | default | 1 | 2019-11-20 23:14:59.677708+0000 UTC  
| | | | deployed  
| | | | sagemaker-k8s-operator-0.1.0

**helm install** creates a **HyperParameterTuningJob** Kubernetes resource. The operator launches the actual hyperparameter optimization job in SageMaker and updates the **HyperParameterTuningJob** Kubernetes resource to reflect the status of the job in SageMaker. You incur charges for SageMaker resources used during the duration of your job. You do not incur any charges once your job completes or stops.

**Note**: SageMaker does not allow you to update a running hyperparameter tuning job. You cannot edit any parameter and re-apply the config file. You must either change the metadata name or delete the existing job and create a new one. Similar to existing training job operators like **TFJob** in Kubeflow, update is not supported.

**List HyperparameterTuningJobs**

Use the following command to list all jobs created using the Kubernetes operator:

```bash
test -e helm-install  && helm install
```

Your output should look like the following:

<table>
<thead>
<tr>
<th>NAME</th>
<th>STATUS</th>
<th>CREATION-TIME</th>
<th>COMPLETED</th>
<th>INPROGRESS</th>
<th>ERRORS</th>
<th>SAGEMAKER-JOB-NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>xgboost-mnist-hpo</td>
<td>Completed</td>
<td>2019-10-17T01:15:52Z</td>
<td>10</td>
<td>0</td>
<td>0</td>
<td>xgboostha92f5e3cf07b11e9bf6c06d6-009-4c7a123</td>
</tr>
<tr>
<td>xgboostha92f5e3cf07b11e9bf6c123</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

A hyperparameter tuning job continues to be listed after the job has completed or failed. You can remove a hyperparametertuningjob from the list by following the steps in **Delete a HyperparameterTuningJob**. Jobs that have completed or stopped do not incur any charges for SageMaker resources.
Hyperparameter tuning job status values

The STATUS field can be one of the following values:

- Completed
- InProgress
- Failed
- Stopped
- Stopping

These statuses come directly from the SageMaker official [API documentation](https://docs.aws.amazon.com/sagemaker/latest/dg/). In addition to the official SageMaker status, it is possible for STATUS to be **SynchronizingK8sJobWithSageMaker**. This means that the operator has not yet processed the job.

Status counters

The output has several counters, like **COMPLETED** and **INPROGRESS**. These represent how many training jobs have completed and are in progress, respectively. For more information about how these are determined, see [TrainingJobStatusCounters](https://docs.aws.amazon.com/sagemaker/latest/dg/) in the SageMaker API documentation.

Best TrainingJob

This column contains the name of the TrainingJob that best optimized the selected metric.

To see a summary of the tuned hyperparameters, run:

```
kubectl describe hyperparametertuningjob xgboost-mnist-hpo
```

To see detailed information about the TrainingJob, run:

```
kubectl describe trainingjobs <job name>
```

Spawned TrainingJobs

You can also track all 10 training jobs in Kubernetes launched by HyperparameterTuningJob by running the following command:

```
kubectl get trainingjobs
```
Describe a HyperparameterTuningJob

You can obtain debugging details using the describe kubectl command.

kubectl describe hyperparametertuningjob xgboost-mnist-hpo

In addition to information about the tuning job, the SageMaker Operator for Kubernetes also exposes the best training job found by the hyperparameter tuning job in the describe output as follows:

<table>
<thead>
<tr>
<th>Name:</th>
<th>xgboost-mnist-hpo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Namespace:</td>
<td>default</td>
</tr>
<tr>
<td>Labels:</td>
<td>&lt;none&gt;</td>
</tr>
<tr>
<td>Annotations:</td>
<td>kubectl.kubernetes.io/last-applied-configuration:</td>
</tr>
<tr>
<td></td>
<td>{&quot;apiVersion&quot;:&quot;sagemaker.aws.amazon.com/v1&quot;,&quot;kind&quot;:&quot;HyperparameterTuningJob&quot;,&quot;metadata&quot;:{&quot;annotations&quot;:{},&quot;name&quot;:&quot;xgboost-mnist-hpo&quot;,&quot;namespace&quot;:...}</td>
</tr>
<tr>
<td>API Version:</td>
<td>sagemaker.aws.amazon.com/v1</td>
</tr>
<tr>
<td>Kind:</td>
<td>HyperparameterTuningJob</td>
</tr>
<tr>
<td>Metadata:</td>
<td></td>
</tr>
<tr>
<td>Creation Timestamp:</td>
<td>2019-10-17T01:15:52Z</td>
</tr>
<tr>
<td>Finalizers:</td>
<td>sagemaker-operator-finalizer</td>
</tr>
<tr>
<td>Generation:</td>
<td>2</td>
</tr>
<tr>
<td>Resource Version:</td>
<td>8167</td>
</tr>
<tr>
<td>Self Link:</td>
<td>/apis/sagemaker.aws.amazon.com/v1/namespaces/default/hyperparametertuningjobs/xgboost-mnist-hpo</td>
</tr>
<tr>
<td>UID:</td>
<td>a92f5e3c-f07b-11e9-bf6c-06d6f303uidu</td>
</tr>
<tr>
<td>Spec:</td>
<td></td>
</tr>
<tr>
<td>Hyper Parameter Tuning Job Config:</td>
<td></td>
</tr>
<tr>
<td>Hyper Parameter Tuning Job Objective:</td>
<td></td>
</tr>
<tr>
<td>Metric Name:</td>
<td>validation:error</td>
</tr>
<tr>
<td>Type:</td>
<td>Minimize</td>
</tr>
<tr>
<td>Parameter Ranges:</td>
<td></td>
</tr>
<tr>
<td>Integer Parameter Ranges:</td>
<td></td>
</tr>
<tr>
<td>Max Value:</td>
<td>20</td>
</tr>
<tr>
<td>Min Value:</td>
<td>10</td>
</tr>
<tr>
<td>Name:</td>
<td>num_round</td>
</tr>
<tr>
<td>Scaling Type:</td>
<td>Linear</td>
</tr>
<tr>
<td>Resource Limits:</td>
<td></td>
</tr>
<tr>
<td>Max Number Of Training Jobs:</td>
<td>10</td>
</tr>
<tr>
<td>Max Parallel Training Jobs:</td>
<td>10</td>
</tr>
<tr>
<td>Strategy:</td>
<td>Bayesian</td>
</tr>
</tbody>
</table>
Training Job Early Stopping Type: Off
Hyper Parameter Tuning Job Name: xgboosta92f5e3cf07b11e9bf6c06d6
Region: us-east-2
Training Job Definition:
  Algorithm Specification:
    Training Image: 12345678910.dkr.ecr.us-east-2.amazonaws.com/xgboost:1
    Training Input Mode: File
  Input Data Config:
    Channel Name: train
    Content Type: text/csv
    Data Source:
      s3DataSource:
        s3DataDistributionType: FullyReplicated
        s3DataType: S3Prefix
        s3Uri: https://s3-us-east-2.amazonaws.com/my-bucket/sagemaker/xgboost-mnist/train/
    Channel Name: validation
    Content Type: text/csv
    Data Source:
      s3DataSource:
        s3DataDistributionType: FullyReplicated
        s3DataType: S3Prefix
        s3Uri: https://s3-us-east-2.amazonaws.com/my-bucket/sagemaker/xgboost-mnist/validation/
  Output Data Config:
    s3OutputPath: https://s3-us-east-2.amazonaws.com/my-bucket/sagemaker/xgboost-mnist/xgboost
  Resource Config:
    Instance Count: 1
    Instance Type: ml.m4.xlarge
    Volume Size In GB: 5
    Role Arn: arn:aws:iam::123456789012:role/service-role/AmazonSageMaker-ExecutionRole
  Static Hyper Parameters:
    Name: base_score
    Value: 0.5
    Name: booster
    Value: gbtree
    Name: csv_weights
    Value: 0
    Name: dsplit
    Value: row
    Name: grow_policy
    Value: depthwise
Name: lambda_bias
Value: 0.0
Name: max_bin
Value: 256
Name: max_leaves
Value: 0
Name: normalize_type
Value: tree
Name: objective
Value: reg:linear
Name: one_drop
Value: 0
Name: prob_buffer_row
Value: 1.0
Name: process_type
Value: default
Name: rate_drop
Value: 0.0
Name: refresh_leaf
Value: 1
Name: sample_type
Value: uniform
Name: scale_pos_weight
Value: 1.0
Name: silent
Value: 0
Name: sketch_eps
Value: 0.03
Name: skip_drop
Value: 0.0
Name: tree_method
Value: auto
Name: tweedie_variance_power
Value: 1.5

Stopping Condition:
Max Runtime In Seconds: 86400

Status:
Best Training Job:
Creation Time: 2019-10-17T01:16:14Z
Final Hyper Parameter Tuning Job Objective Metric:
Metric Name: validation:error
Value:
Objective Status: Succeeded
Training End Time: 2019-10-17T01:20:24Z
Training Job Arn: arn:aws:sagemaker:us-east-2:123456789012:training-job/xgboostha92f5e3cf07b11e9bf6c06d6-009-4sample
Training Job Name: xgboostha92f5e3cf07b11e9bf6c06d6-009-4c7a3059
Training Job Status: Completed
Training Start Time: 2019-10-17T01:18:35Z
Tuned Hyper Parameters:
  Name: num_round
  Value: 18
Hyper Parameter Tuning Job Status: Completed
Last Check Time: 2019-10-17T01:21:01Z
Sage Maker Hyper Parameter Tuning Job Name: xgboostha92f5e3cf07b11e9bf6c06d6
Training Job Status Counters:
  Completed: 10
  In Progress: 0
  Non Retryable Error: 0
  Retryable Error: 0
  Stopped: 0
  Total Error: 0
Events: <none>

View logs from HyperparameterTuningJobs

Hyperparameter tuning jobs do not have logs, but all training jobs launched by them do have logs. These logs can be accessed as if they were a normal training job. For more information, see View logs from TrainingJobs.

Delete a HyperparameterTuningJob

Use the following command to stop a hyperparameter job in SageMaker.

```bash
kubectl delete hyperparametertuningjob xgboost-mnist-hpo
```

This command removes the hyperparameter tuning job and associated training jobs from your Kubernetes cluster and stops them in SageMaker. Jobs that have stopped or completed do not incur any charges for SageMaker resources. SageMaker does not delete hyperparameter tuning jobs. Stopped jobs continue to show on the SageMaker console.

Your output should look like the following:

```
hyperparametertuningjob.sagemaker.aws.amazon.com "xgboost-mnist-hpo" deleted
```

Note: The delete command takes about 2 minutes to clean up the resources from SageMaker.
The BatchTransformJob operator

Batch transform job operators reconcile your specified batch transform job spec to SageMaker by launching it in SageMaker. You can learn more about SageMaker batch transform job in the SageMaker CreateTransformJob API documentation.

Topics

- Create a BatchTransformJob using a YAML File
- Create a BatchTransformJob using a Helm Chart
- List BatchTransformJobs
- Describe a BatchTransformJob
- View logs from BatchTransformJobs
- Delete a BatchTransformJob

Create a BatchTransformJob using a YAML File

1. Download the sample YAML file for the batch transform job using the following command:

   ```bash
   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/xgboost-mnist-batchtransform.yaml
   ```

2. Edit the file `xgboost-mnist-batchtransform.yaml` to change necessary parameters to replace the inputdataconfig with your input data and s3outputPath with your Amazon S3 buckets that the SageMaker execution role has write access to.

3. Apply the YAML file using the following command:

   ```bash
   kubectl apply -f xgboost-mnist-batchtransform.yaml
   ```

Create a BatchTransformJob using a Helm Chart

You can use Helm Charts to run batch transform jobs.

Get the Helm installer directory

Clone the GitHub repository to get the source using the following command:

```bash
git clone https://github.com/aws/amazon-sagemaker-operator-for-k8s.git
```
Configure the Helm Chart

Navigate to the amazon-sagemaker-operator-for-k8s/hack/charts/batch-transform-jobs/ folder.

Edit the values.yaml file to replace the inputdataconfig with your input data and outputPath with your S3 buckets to which the SageMaker execution role has write access.

Create a BatchTransformJob

1. Use the following command to create a batch transform job:

   ```
   helm install . --generate-name
   ```

   Your output should look like the following:

   ```
   NAME: chart-1574292948
   LAST DEPLOYED: Wed Nov 20 23:35:49 2019
   NAMESPACE: default
   STATUS: deployed
   REVISION: 1
   TEST SUITE: None
   NOTES:
   Thanks for installing the sagemaker-k8s-batch-transform-job.
   ```

2. To verify that the Helm Chart was created successfully, run the following command:

   ```
   helm ls
   ```

   This command creates a BatchTransformJob Kubernetes resource. The operator launches the actual transform job in SageMaker and updates the BatchTransformJob Kubernetes resource to reflect the status of the job in SageMaker. You incur charges for SageMaker
resources used during the duration of your job. You do not incur any charges once your job completes or stops.

**Note**: SageMaker does not allow you to update a running batch transform job. You cannot edit any parameter and re-apply the config file. You must either change the metadata name or delete the existing job and create a new one. Similar to existing training job operators like TFJob in Kubeflow, update is not supported.

**List BatchTransformJobs**

Use the following command to list all jobs created using the Kubernetes operator:

```
kubectl get batchtransformjob
```

Your output should look like the following:

<table>
<thead>
<tr>
<th>NAME</th>
<th>STATUS</th>
<th>CREATION-TIME</th>
<th>SAGEMAKER-JOB-NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>xgboost-mnist-batch-transform</td>
<td>Completed</td>
<td>2019-11-18T03:44:00Z</td>
<td>xgboost-mnist-</td>
</tr>
<tr>
<td>a88fb19809b511eaac440aa8axgboost</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

A batch transform job continues to be listed after the job has completed or failed. You can remove a hyperparametertuningjob from the list by following the [Delete a BatchTransformJob](#) steps. Jobs that have completed or stopped do not incur any charges for SageMaker resources.

**Batch transform status values**

The STATUS field can be one of the following values:

- Completed
- InProgress
- Failed
- Stopped
- Stopping

These statuses come directly from the SageMaker official [API documentation](#).
In addition to the official SageMaker status, it is possible for STATUS to be `SynchronizingK8sJobWithSageMaker`. This means that the operator has not yet processed the job.

**Describe a BatchTransformJob**

You can obtain debugging details using the describe kubectl command.

```
kubectl describe batchtransformjob xgboost-mnist-batch-transform
```

Your output should look like the following:

```
Name:         xgboost-mnist-batch-transform
Namespace:    default
Labels:       <none>
Annotations:  kubectl.kubernetes.io/last-applied-configuration:
               {"apiVersion":"sagemaker.aws.amazon.com/v1","kind":"BatchTransformJob","metadata":{"annotations":{},"name":"xgboost-mnist","namespace"...}
API Version:  sagemaker.aws.amazon.com/v1
Kind:         BatchTransformJob
Metadata:
   Creation Timestamp:  2019-11-18T03:44:00Z
   Finalizers:
      sagemaker-operator-finalizer
   Generation:        2
   Resource Version:  21990924
   Self Link:         /apis/sagemaker.aws.amazon.com/v1/namespaces/default/batchtransformjobs/xgboost-mnist
   UID:               a88fb198-09b5-11ea-ac44-0aa8a9UIDNUM
Spec:
   Model Name:  TrainingJob-20190814SMJob-IKEB
   Region:      us-east-1
   Transform Input:
      Content Type:  text/csv
      Data Source:
         S3 Data Source:
            S3 Data Type:  S3Prefix
            S3 Uri:        s3://my-bucket/mnist_kmeans_example/input
   Transform Job Name:   xgboost-mnist-a88fb19809b511eaac440aa8a9SMJOB
   Transform Output:
      S3 Output Path:  s3://my-bucket/mnist_kmeans_example/output
   Transform Resources:
```
Instance Count: 1
Instance Type: ml.m4.xlarge
Status:
Last Check Time: 2019-11-19T22:50:40Z
SageMaker Transform Job Name: xgboost-mnist-a88fb19809b511eaac440aaSMJOB
Transform Job Status: Completed
Events: <none>

View logs from BatchTransformJobs

Use the following command to see the logs from the xgboost-mnist batch transform job:

```bash
ekubectl smlogs batchtransformjob xgboost-mnist-batch-transform
```

Delete a BatchTransformJob

Use the following command to stop a batch transform job in SageMaker.

```bash
ekubectl delete batchTransformJob xgboost-mnist-batch-transform
```

Your output should look like the following:

```bash
batchtransformjob.sagemaker.aws.amazon.com "xgboost-mnist" deleted
```

This command removes the batch transform job from your Kubernetes cluster, as well as stops them in SageMaker. Jobs that have stopped or completed do not incur any charges for SageMaker resources. Delete takes about 2 minutes to clean up the resources from SageMaker.

**Note:** SageMaker does not delete batch transform jobs. Stopped jobs continue to show on the SageMaker console.

The HostingDeployment operator

HostingDeployment operators support creating and deleting an endpoint, as well as updating an existing endpoint, for real-time inference. The hosting deployment operator reconciles your specified hosting deployment job spec to SageMaker by creating models, endpoint-configs and endpoints in SageMaker. You can learn more about SageMaker inference in the SageMaker CreateEndpoint API documentation.

Topics

- [Configure a HostingDeployment resource](#)
Configure a HostingDeployment resource

Download the sample YAML file for the hosting deployment job using the following command:

```bash
wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/xgboost-mnist-hostingdeployment.yaml
```

The `xgboost-mnist-hostingdeployment.yaml` file has the following components that can be edited as required:

- **ProductionVariants**. A production variant is a set of instances serving a single model. SageMaker load-balances between all production variants according to set weights.
- **Models**. A model is the containers and execution role ARN necessary to serve a model. It requires at least a single container.
- **Containers**. A container specifies the dataset and serving image. If you are using your own custom algorithm instead of an algorithm provided by SageMaker, the inference code must meet SageMaker requirements. For more information, see [Using Your Own Algorithms with SageMaker](#).

Create a HostingDeployment

To create a HostingDeployment, use `kubectl` to apply the file `hosting.yaml` with the following command:

```bash
kubectl apply -f hosting.yaml
```

SageMaker creates an endpoint with the specified configuration. You incur charges for SageMaker resources used during the lifetime of your endpoint. You do not incur any charges once your endpoint is deleted.
The creation process takes approximately 10 minutes.

**List HostingDeployments**

To verify that the HostingDeployment was created, use the following command:

```
kubectl get hostingdeployments
```

Your output should look like the following:

<table>
<thead>
<tr>
<th>NAME</th>
<th>STATUS</th>
<th>SAGEMAKER-ENDPOINT-NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>host-xgboost</td>
<td>Creating</td>
<td>host-xgboost-def0e83e0d5f11eaa450aSMLOGS</td>
</tr>
</tbody>
</table>

**HostingDeployment status values**

The status field can be one of several values:

- **SynchronizingK8sJobWithSageMaker**: The operator is preparing to create the endpoint.
- **ReconcilingEndpoint**: The operator is creating, updating, or deleting endpoint resources. If the HostingDeployment remains in this state, use `kubectl describe` to see the reason in the Additional field.
- **OutOfService**: The endpoint is not available to take incoming requests.
- **Creating**: `CreateEndpoint` is running.
- **Updating**: `UpdateEndpoint` or `UpdateEndpointWeightsAndCapacities` is running.
- **SystemUpdating**: The endpoint is undergoing maintenance and cannot be updated or deleted or re-scaled until it has completed. This maintenance operation does not change any customer-specified values such as VPC config, AWS KMS encryption, model, instance type, or instance count.
- **RollingBack**: The endpoint fails to scale up or down or change its variant weight and is in the process of rolling back to its previous configuration. Once the rollback completes, the endpoint returns to an InService status. This transitional status only applies to an endpoint that has autoscaling turned on and is undergoing variant weight or capacity changes as part of an `UpdateEndpointWeightsAndCapacities` call or when the `UpdateEndpointWeightsAndCapacities` operation is called explicitly.
- **InService**: The endpoint is available to process incoming requests.
- **Deleting**: `DeleteEndpoint` is running.
- **Failed**: The endpoint could not be created, updated, or re-scaled. Use `DescribeEndpoint:FailureReason` for information about the failure. `DeleteEndpoint` is the only operation that can be performed on a failed endpoint.
Amazon SageMaker

Developer Guide

Describe a HostingDeployment
You can obtain debugging details using the describe kubectl command.
kubectl describe hostingdeployment

Your output should look like the following:
Name:
Namespace:
Labels:
Annotations:

host-xgboost
default
<none>
kubectl.kubernetes.io/last-applied-configuration:
{"apiVersion":"sagemaker.aws.amazon.com/
v1","kind":"HostingDeployment","metadata":{"annotations":{},"name":"hostxgboost","namespace":"def..."
API Version: sagemaker.aws.amazon.com/v1
Kind:
HostingDeployment
Metadata:
Creation Timestamp: 2019-11-22T19:40:00Z
Finalizers:
sagemaker-operator-finalizer
Generation:
1
Resource Version: 4258134
Self Link:
/apis/sagemaker.aws.amazon.com/v1/namespaces/default/
hostingdeployments/host-xgboost
UID:
def0e83e-0d5f-11ea-aa45-0a3507uiduid
Spec:
Containers:
Container Hostname: xgboost
Image:
123456789012.dkr.ecr.us-east-2.amazonaws.com/xgboost:latest
Model Data URL:
s3://my-bucket/inference/xgboost-mnist/model.tar.gz
Models:
Containers:
xgboost
Execution Role Arn: arn:aws:iam::123456789012:role/service-role/AmazonSageMakerExecutionRole
Name:
xgboost-model
Primary Container:
xgboost
Production Variants:
Initial Instance Count: 1
Instance Type:
ml.c5.large
Model Name:
xgboost-model
Variant Name:
all-traffic
Kubernetes Orchestration

4910


The status field provides more information using the following fields:

- **Additional**: Additional information about the status of the hosting deployment. This field is optional and only gets populated in case of error.
- **Creation Time**: When the endpoint was created in SageMaker.
- **Endpoint ARN**: The SageMaker endpoint ARN.
- **Endpoint Config Name**: The SageMaker name of the endpoint configuration.
- **Endpoint Name**: The SageMaker name of the endpoint.
- **Endpoint Status**: The status of the endpoint.
- **Endpoint URL**: The HTTPS URL that can be used to access the endpoint. For more information, see [Deploy a Model on SageMaker Hosting Services](#).
- **FailureReason**: If a create, update, or delete command fails, the cause is shown here.
- **Last Check Time**: The last time the operator checked the status of the endpoint.
- **Last Modified Time**: The last time the endpoint was modified.
- **Model Names**: A key-value pair of HostingDeployment model names to SageMaker model names.

**Invoking the endpoint**

Once the endpoint status is InService, you can invoke the endpoint in two ways: using the AWS CLI, which does authentication and URL request signing, or using an HTTP client like cURL. If you use your own client, you need to do AWS v4 URL signing and authentication on your own.
To invoke the endpoint using the AWS CLI, run the following command. Make sure to replace the Region and endpoint name with your endpoint's Region and SageMaker endpoint name. This information can be obtained from the output of kubectl describe.

```bash
# Invoke the endpoint with mock input data.
aws sagemaker-runtime invoke-endpoint \
  --region us-east-2 \
  --endpoint-name <endpoint name> \
  --body $(seq 784 | xargs echo | sed 's/ /,/g') \
  > (cat) \
  --content-type text/csv > /dev/null
```

For example, if your Region is us-east-2 and your endpoint config name is host-xgboost-f56b6b280d7511ea824b129926example, then the following command would invoke the endpoint:

```bash
aws sagemaker-runtime invoke-endpoint \
  --region us-east-2 \
  --endpoint-name host-xgboost-f56b6b280d7511ea824b1299example \
  --body $(seq 784 | xargs echo | sed 's/ /,/g') \
  > (cat) \
  --content-type text/csv > /dev/null
```

Here, 4.95847082138 is the prediction from the model for the mock data.

**Update HostingDeployment**

1. Once a HostingDeployment has a status of InService, it can be updated. It might take about 10 minutes for HostingDeployment to be in service. To verify that the status is InService, use the following command:

   ```bash
   kubectl get hostingdeployments
   ```

2. The HostingDeployment can be updated before the status is InService. The operator waits until the SageMaker endpoint is InService before applying the update.

   To apply an update, modify the hosting.yaml file. For example, change the initialInstanceCount field from 1 to 2 as follows:

   ```yaml
   apiVersion: sagemaker.aws.amazon.com/v1
   ```
3. Save the file, then use `kubectl` to apply your update as follows. You should see the status change from InService to ReconcilingEndpoint, then Updating.

```bash
$ kubectl apply -f hosting.yaml
hostingdeployment.sagemaker.aws.amazon.com/host-xgboost configured

$ kubectl get hostingdeployments
NAME           STATUS                SAGEMAKER-ENDPOINT-NAME
host-xgboost   ReconcilingEndpoint   host-xgboost-def0e83e0d5f11eaaa450a350abcdef

$ kubectl get hostingdeployments
NAME           STATUS     SAGEMAKER-ENDPOINT-NAME
host-xgboost   Updating   host-xgboost-def0e83e0d5f11eaaa450a3507abcdef
```

SageMaker deploys a new set of instances with your models, switches traffic to use the new instances, and drains the old instances. As soon as this process begins, the status becomes Updating. After the update is complete, your endpoint becomes InService. This process takes approximately 10 minutes.
Delete the HostingDeployment

1. Use kubectl to delete a HostingDeployment with the following command:

   kubectl delete hostingdeployments host-xgboost

   Your output should look like the following:

   hostingdeployment.sagemaker.aws.amazon.com "host-xgboost" deleted

2. To verify that the hosting deployment has been deleted, use the following command:

   kubectl get hostingdeployments
   No resources found.

Endpoints that have been deleted do not incur any charges for SageMaker resources.

The ProcessingJob operator

ProcessingJob operators are used to launch Amazon SageMaker processing jobs. For more information on SageMaker processing jobs, see CreateProcessingJob.

Topics

- Create a ProcessingJob using a YAML file
- List ProcessingJobs
- Describe a ProcessingJob
- Delete a ProcessingJob

Create a ProcessingJob using a YAML file

Follow these steps to create an Amazon SageMaker processing job by using a YAML file:

1. Download the kmeans_preprocessing.py pre-processing script.

   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/kmeans_preprocessing.py
2. In one of your Amazon Simple Storage Service (Amazon S3) buckets, create a `mnist_kmeans_example/processing_code` folder and upload the script to the folder.

3. Download the `kmeans-mnist-processingjob.yaml` file.

   ```bash
   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/kmeans-mnist-processingjob.yaml
   ```

4. Edit the YAML file to specify your `sagemaker-execution-role` and replace all instances of `my-bucket` with your S3 bucket.

   ```yaml
   ...  
   metadata:
     name: kmeans-mnist-processing
   ...
   roleArn: arn:aws:iam::<acct-id>:role/service-role/<sagemaker-execution-role>
   ...
   processingOutputConfig:
     outputs:
       ...
       s3Output:
         s3Uri: s3://<my-bucket>/mnist_kmeans_example/output/
   ...
   processingInputs:
     ...
     s3Input:
       s3Uri: s3://<my-bucket>/mnist_kmeans_example/processing_code/
   kmeans_preprocessing.py
   ```

   The `sagemaker-execution-role` must have permissions so that SageMaker can access your S3 bucket, Amazon CloudWatch, and other services on your behalf. For more information on creating an execution role, see [SageMaker Roles](#).

5. Apply the YAML file using one of the following commands.

   For cluster-scoped installation:

   ```bash
   kubectl apply -f kmeans-mnist-processingjob.yaml
   ```

   For namespace-scoped installation:
List ProcessingJobs

Use one of the following commands to list all the jobs created using the ProcessingJob operator. `SAGEMAKER-JOB-NAME` comes from the metadata section of the YAML file.

For cluster-scoped installation:

```
kubectl get ProcessingJob kmeans-mnist-processing
```

For namespace-scoped installation:

```
kubectl get ProcessingJob -n <NAMESPACE> kmeans-mnist-processing
```

Your output should look similar to the following:

<table>
<thead>
<tr>
<th>NAME</th>
<th>STATUS</th>
<th>CREATION-TIME</th>
<th>SAGEMAKER-JOB-NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>kmeans-mnist-processing</td>
<td>InProgress</td>
<td>2020-09-22T21:13:25Z</td>
<td>kmeans-mnist-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>processing-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>7410ed52fd1811eab19a165ae9f9e385</td>
</tr>
</tbody>
</table>

The output lists all jobs regardless of their status. To remove a job from the list, see Delete a Processing Job.

ProcessingJob Status

- **SynchronizingK8sJobWithSageMaker** – The job is first submitted to the cluster. The operator has received the request and is preparing to create the processing job.
- **Reconciling** – The operator is initializing or recovering from transient errors, along with others. If the processing job remains in this state, use the `kubectl describe` command to see the reason in the Additional field.
- **InProgress | Completed | Failed | Stopping | Stopped** – Status of the SageMaker processing job. For more information, see DescribeProcessingJob.
- **Error** – The operator cannot recover by reconciling.

Jobs that have completed, stopped, or failed do not incur further charges for SageMaker resources.
Describe a ProcessingJob

Use one of the following commands to get more details about a processing job. These commands are typically used for debugging a problem or checking the parameters of a processing job.

For cluster-scoped installation:

```
kubectl describe processingjob kmeans-mnist-processing
```

For namespace-scoped installation:

```
kubectl describe processingjob kmeans-mnist-processing -n <NAMESPACE>
```

The output for your processing job should look similar to the following.

```
$ kubectl describe ProcessingJob kmeans-mnist-processing
Name:         kmeans-mnist-processing
Namespace:    default
Labels:       <none>
Annotations:  kubectl.kubernetes.io/last-applied-configuration:
               {
                   "apiVersion": "sagemaker.aws.amazon.com/v1",
                   "kind": "ProcessingJob",
                   "metadata": {
                       "annotations": {},
                       "name": "kmeans-mnist-processing",
                   },
               ...
API Version:  sagemaker.aws.amazon.com/v1
Kind:         ProcessingJob
Metadata:
   Finalizers:
      sagemaker-operator-finalizer
   Generation:        2
   Resource Version:  21746658
   Self Link:         /apis/sagemaker.aws.amazon.com/v1/namespaces/default/processingjobs/kmeans-mnist-processing
   UID:               7410ed52-fd18-11ea-b19a-165ae9f9e385
Spec:
   App Specification:
      Container Entrypoint:
         python
         /opt/ml/processing/code/kmeans_preprocessing.py
   Image Uri:  763104351884.dkr.ecr.us-west-2.amazonaws.com/pytorch-training:1.5.0-cpu-py36-ubuntu16.04
   Environment:
      Name:   MYVAR
```
Value: my_value
Name: MYVAR2
Value: my_value2

Network Config:

Processing Inputs:
- Input Name: mnist_tar
  s3Input:
    Local Path: /opt/ml/processing/input
    s3DataType: S3Prefix
    s3InputMode: File
    s3Uri: s3://<s3bucket>-us-west-2/algorithms/kmeans/mnist/mnist.pkl.gz

- Input Name: source_code
  s3Input:
    Local Path: /opt/ml/processing/code
    s3DataType: S3Prefix
    s3InputMode: File
    s3Uri: s3://<s3bucket>/mnist_kmeans_example/processing_code/kmeans_preprocessing.py

Processing Output Config:
- Output Name: train_data
  s3Output:
    Local Path: /opt/ml/processing/output_train/
    s3UploadMode: EndOfJob
    s3Uri: s3://<s3bucket>/mnist_kmeans_example/output/

- Output Name: test_data
  s3Output:
    Local Path: /opt/ml/processing/output_test/
    s3UploadMode: EndOfJob
    s3Uri: s3://<s3bucket>/mnist_kmeans_example/output/

- Output Name: valid_data
  s3Output:
    Local Path: /opt/ml/processing/output_valid/
    s3UploadMode: EndOfJob
    s3Uri: s3://<s3bucket>/mnist_kmeans_example/output/

Processing Resources:
- Cluster Config:
  Instance Count: 1
  Instance Type: ml.m5.xlarge
  Volume Size In GB: 20
  Region: us-west-2
  Role Arn: arn:aws:iam::<acct-id>:role/m-sagemaker-role
- Stopping Condition:
  Max Runtime In Seconds: 1800
Delete a ProcessingJob

When you delete a processing job, the SageMaker processing job is removed from Kubernetes but the job isn't deleted from SageMaker. If the job status in SageMaker is InProgress the job is stopped. Processing jobs that are stopped do not incur any charges for SageMaker resources. Use one of the following commands to delete a processing job.

For cluster-scoped installation:

```
kubectl delete processingjob kmeans-mnist-processing
```

For namespace-scoped installation:

```
kubectl delete processingjob kmeans-mnist-processing -n <NAMESPACE>
```

The output for your processing job should look similar to the following.

```
processingjob.sagemaker.aws.amazon.com "kmeans-mnist-processing" deleted
```

**Note**

SageMaker does not delete the processing job. Stopped jobs continue to show in the SageMaker console. The delete command takes a few minutes to clean up the resources from SageMaker.
HostingAutoscalingPolicy (HAP) Operator

The HostingAutoscalingPolicy (HAP) operator takes a list of resource IDs as input and applies the same policy to each of them. Each resource ID is a combination of an endpoint name and a variant name. The HAP operator performs two steps: it registers the resource IDs and then applies the scaling policy to each resource ID. Delete undoes both actions. You can apply the HAP to an existing SageMaker endpoint or you can create a new SageMaker endpoint using the HostingDeployment operator. You can read more about SageMaker autoscaling in the Application Autoscaling Policy documentation.

Note

In your kubectl commands, you can use the short form, hap, in place of hostingautoscalingpolicy.

Topics

- Create a HostingAutoscalingPolicy using a YAML file
- List HostingAutoscalingPolicies
- Describe a HostingAutoscalingPolicy
- Update a HostingAutoscalingPolicy
- Delete a HostingAutoscalingPolicy
- Update or delete an endpoint with a HostingAutoscalingPolicy

Create a HostingAutoscalingPolicy using a YAML file

Use a YAML file to create a HostingAutoscalingPolicy (HAP) that applies a predefined or custom metric to one or multiple SageMaker endpoints.

Amazon SageMaker requires specific values in order to apply autoscaling to your variant. If these values are not specified in the YAML spec, the HAP operator applies the following default values.

```yaml
# Do not change
Namespace = "sagemaker"
# Do not change
ScalableDimension = "sagemaker:variant:DesiredInstanceCount"
```
Use the following samples to create a HAP that applies a predefined or custom metric to one or multiple endpoints.

**Sample 1: Apply a predefined metric to a single endpoint variant**

1. Download the sample YAML file for a predefined metric using the following command:

   ```bash
   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/hap-predefined-metric.yaml
   ```

2. Edit the YAML file to specify your `endpointName`, `variantName`, and `Region`.

3. Use one of the following commands to apply a predefined metric to a single resource ID (endpoint name and variant name combination).

   For cluster-scoped installation:
   ```bash
   kubectl apply -f hap-predefined-metric.yaml
   ```

   For namespace-scoped installation:
   ```bash
   kubectl apply -f hap-predefined-metric.yaml -n <NAMESPACE>
   ```

**Sample 2: Apply a custom metric to a single endpoint variant**

1. Download the sample YAML file for a custom metric using the following command:

   ```bash
   wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/hap-custom-metric.yaml
   ```

2. Edit the YAML file to specify your `endpointName`, `variantName`, and `Region`.

3. Use one of the following commands to apply a custom metric to a single resource ID (endpoint name and variant name combination) in place of the recommended `SageMakerVariantInvocationsPerInstance`.

   ```bash
   kubectl apply -f hap-custom-metric.yaml -n <NAMESPACE>
   ```
Note
Amazon SageMaker does not check the validity of your YAML spec.

For cluster-scoped installation:

```
kubectl apply -f hap-custom-metric.yaml
```

For namespace-scoped installation:

```
kubectl apply -f hap-custom-metric.yaml -n <NAMESPACE>
```

**Sample 3: Apply a scaling policy to multiple endpoints and variants**

You can use the HAP operator to apply the same scaling policy to multiple resource IDs. A separate `scaling_policy` request is created for each resource ID (endpoint name and variant name combination).

1. Download the sample YAML file for a predefined metric using the following command:

   ```
wget https://raw.githubusercontent.com/aws/amazon-sagemaker-operator-for-k8s/master/samples/hap-predefined-metric.yaml
```

2. Edit the YAML file to specify your Region and multiple `endpointName` and `variantName` values.

3. Use one of the following commands to apply a predefined metric to multiple resource IDs (endpoint name and variant name combinations).

   For cluster-scoped installation:
   ```
kubectl apply -f hap-predefined-metric.yaml
```

   For namespace-scoped installation:
   ```
kubectl apply -f hap-predefined-metric.yaml -n <NAMESPACE>
```
Considerations for HostingAutoscalingPolicies for multiple endpoints and variants

The following considerations apply when you use multiple resource IDs:

- If you apply a single policy across multiple resource IDs, one PolicyARN is created per resource ID. Five endpoints have five PolicyARNs. When you run the describe command on the policy, the responses show up as one job and include a single job status.

- If you apply a custom metric to multiple resource IDs, the same dimension or value is used for all the resource ID (variant) values. For example, if you apply a customer metric for instances 1-5, and the endpoint variant dimension is mapped to variant 1, when variant 1 exceeds the metrics, all endpoints are scaled up or down.

- The HAP operator supports updating the list of resource IDs. If you modify, add, or delete resource IDs to the spec, the autoscaling policy is removed from the previous list of variants and applied to the newly specified resource ID combinations. Use the describe command to list the resource IDs to which the policy is currently applied.

List HostingAutoscalingPolicies

Use one of the following commands to list all HostingAutoscalingPolicies (HAPs) created using the HAP operator.

For cluster-scoped installation:

```sh
cubectl get hap
```

For namespace-scoped installation:

```sh
cubectl get hap -n <NAMESPACE>
```

Your output should look similar to the following:

<table>
<thead>
<tr>
<th>NAME</th>
<th>STATUS</th>
<th>CREATION-TIME</th>
</tr>
</thead>
<tbody>
<tr>
<td>hap-predefined</td>
<td>Created</td>
<td>2021-07-13T21:32:21Z</td>
</tr>
</tbody>
</table>

Use the following command to check the status of your HostingAutoscalingPolicy (HAP).

```sh
cubectl get hap <job-name>
```

One of the following values is returned:
• Reconciling – Certain types of errors show the status as Reconciling instead of Error. Some examples are server-side errors and endpoints in the Creating or Updating state. Check the Additional field in status or operator logs for more details.

• Created

• Error

To view the autoscaling endpoint to which you applied the policy

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the left side panel, expand Inference.
3. Choose Endpoints.
4. Select the name of the endpoint of interest.
5. Scroll to the Endpoint runtime settings section.

Describe a HostingAutoscalingPolicy

Use the following command to get more details about a HostingAutoscalingPolicy (HAP). These commands are typically used for debugging a problem or checking the resource IDs (endpoint name and variant name combinations) of a HAP.

```
kubectl describe hap <job-name>
```

Update a HostingAutoscalingPolicy

The HostingAutoscalingPolicy (HAP) operator supports updates. You can edit your YAML spec to change the values and then reapply the policy. The HAP operator deletes the existing policy and applies the new policy.

Delete a HostingAutoscalingPolicy

Use one of the following commands to delete a HostingAutoscalingPolicy (HAP) policy.

For cluster-scoped installation:

```
kubectl delete hap hap-predefined
```

For namespace-scoped installation:
kubectl delete hap hap-predefined -n <NAMESPACE>

This command deletes the scaling policy and deregisters the scaling target from Kubernetes. This command returns the following output:

hostingautoscalingpolicies.sagemaker.aws.amazon.com "hap-predefined" deleted

Update or delete an endpoint with a HostingAutoscalingPolicy

To update an endpoint that has a HostingAutoscalingPolicy (HAP), use the `kubectl delete` command to remove the HAP, update the endpoint, and then reapply the HAP.

To delete an endpoint that has a HAP, use the `kubectl delete` command to remove the HAP before you delete the endpoint.

Migrate resources to the latest Operators

We are stopping the development and technical support of the original version of SageMaker Operators for Kubernetes.

If you are currently using version v1.2.2 or below of SageMaker Operators for Kubernetes, we recommend migrating your resources to the ACK service controller for Amazon SageMaker. The ACK service controller is a new generation of SageMaker Operators for Kubernetes based on AWS Controllers for Kubernetes (ACK).

For answers to frequently asked questions on the end of support of the original version of SageMaker Operators for Kubernetes, see Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes

Use the following steps to migrate your resources and use ACK to train, tune, and deploy machine learning models with Amazon SageMaker.

Note
The latest SageMaker Operators for Kubernetes are not backwards compatible.

Contents
- Prerequisites
- Adopt resources
• **Clean up old resources**

• **Use the new SageMaker Operators for Kubernetes**

**Prerequisites**

To successfully migrate resources to the latest SageMaker Operators for Kubernetes, you must do the following:


2. If you are using **HostingAutoscalingPolicy resources**, install the new Application Auto Scaling Operators. See [Setup](https://docs.aws.amazon.com/sagemaker/latest/dg/ack-controller.html) in *Scale SageMaker Workloads with Application Auto Scaling* for step-by-step instructions. This step is optional if you are not using HostingAutoScalingPolicy resources.

If permissions are configured correctly, then the ACK SageMaker service controller can determine the specification and status of the AWS resource and reconcile the resource as if the ACK controller originally created it.

**Adopt resources**

The new SageMaker Operators for Kubernetes provide the ability to adopt resources that were not originally created by the ACK service controller. For more information, see [Adopt Existing AWS Resources](https://docs.aws.amazon.com/sagemaker/latest/dg/ack-controller.html) in the ACK documentation.

The following steps show how the new SageMaker Operators for Kubernetes can adopt an existing SageMaker endpoint. Save the following sample to a file named `adopt-endpoint-sample.yaml`.

```yaml
apiVersion: services.k8s.aws/v1alpha1
kind: AdoptedResource
metadata:
  name: adopt-endpoint-sample
spec:
  aws:
    # resource to adopt, not created by ACK
    nameOrID: xgboost-endpoint
  kubernetes:
    group: sagemaker.services.k8s.aws
    kind: Endpoint
```
metadata:
  # target K8s CR name
  name: xgboost-endpoint

Submit the custom resource (CR) using `kubectl apply`:

```
kubectl apply -f adopt-endpoint-sample.yaml
```

Use `kubectl describe` to check the status conditions of your adopted resource.

```
kubectl describe adoptedresource adopt-endpoint-sample
```

Verify that the `ACK.Adopted` condition is `True`. The output should look similar to the following example:

```yaml
---
kind: AdoptedResource
metadata:
  annotations:
    kubectl.kubernetes.io/last-applied-configuration: |
      {"apiVersion": "services.k8s.aws/v1alpha1", "kind": "AdoptedResource", "metadata": {"annotations": [], "name": "xgboost-endpoint", "namespace": "default"}, "spec": {"aws": {"nameOrID": "xgboost-endpoint"}, "kubernetes": {"group": "sagemaker.services.k8s.aws", "kind": "Endpoint", "metadata": {"name": "xgboost-endpoint"}}}}'
  creationTimestamp: '2021-04-27T02:49:14Z'
  finalizers:
    - finalizers.services.k8s.aws/AdoptedResource
  generation: 1
  name: adopt-endpoint-sample
  namespace: default
  resourceVersion: '12669876'
  selfLink: /apis/services.k8s.aws/v1alpha1/namespaces/default/adoptedresources/adopt-endpoint-sample
  uid: 35f8fa92-29dd-4040-9d0d-0b07b7ca0b
spec:
  aws:
    nameOrID: xgboost-endpoint
  kubernetes:
    group: sagemaker.services.k8s.aws
    kind: Endpoint
    metadata:
```
name: xgboost-endpoint
status:
  conditions:
  - status: 'True'
  type: ACK.Adopted

Check that your resource exists in your cluster:

```
kubectl describe endpoints.sagemaker xgboost-endpoint
```

**HostingAutoscalingPolicy resources**

The HostingAutoscalingPolicy (HAP) resource consists of multiple Application Auto Scaling resources: ScalableTarget and ScalingPolicy. When adopting a HAP resource with ACK, first install the Application Auto Scaling controller. To adopt HAP resources, you need to adopt both ScalableTarget and ScalingPolicy resources. You can find the resource identifier for these resources in the status of the HostingAutoscalingPolicy resource (status.ResourceIDList).

**HostingDeployment resources**

The HostingDeployment resource consists of multiple SageMaker resources: Endpoint, EndpointConfig, and each Model. If you adopt a SageMaker endpoint in ACK, you need to adopt the Endpoint, EndpointConfig, and each Model separately. The Endpoint, EndpointConfig, and Model names can be found in status of the HostingDeployment resource (status.endpointName, status.endpointConfigName, and status.modelNames).

For a list of all supported SageMaker resources, refer to the [ACK API Reference](#).

**Clean up old resources**

After the new SageMaker Operators for Kubernetes adopt your resources, you can uninstall old operators and clean up old resources.

**Step 1: Uninstall the old operator**

To uninstall the old operator, see [Delete operators](#).

⚠️ **Warning**

Uninstall the old operator before deleting any old resources.
Step 2: Remove finalizers and delete old resources

⚠️ Warning
Before deleting old resources, be sure that you have uninstalled the old operator.

After uninstalling the old operator, you must explicitly remove the finalizers to delete old operator resources. The following sample script shows how to delete all training jobs managed by the old operator in a given namespace. You can use a similar pattern to delete additional resources once they are adopted by the new operator.

💡 Note
You must use full resource names to get resources. For example, use kubectl get trainingjobs.sagemaker.aws.amazon.com instead of kubectl get trainingjob.

```
namespace=sagemaker_namespace
training_jobs=$(kubectl get trainingjobs.sagemaker.aws.amazon.com -n $namespace -ojson | jq -r '.items | .[] | .metadata.name')

for job in $training_jobs
do
echo "Deleting $job resource in $namespace namespace"
kubectl patch trainingjobs.sagemaker.aws.amazon.com $job -n $namespace -p '{"metadata":{"finalizers":null}}' --type=merge
kubectl delete trainingjobs.sagemaker.aws.amazon.com $job -n $namespace
done
```

Use the new SageMaker Operators for Kubernetes

For in-depth guides on using the new SageMaker Operators for Kubernetes, see Use SageMaker Operators for Kubernetes

Announcing the End of Support of the Original Version of SageMaker Operators for Kubernetes

This page announces the end of support for the original version of SageMaker Operators for Kubernetes and provides answers to frequently asked questions as well as migration information.
about the ACK service controller for Amazon SageMaker, a new generation of fully supported SageMaker Operators for Kubernetes. For general information about the new SageMaker Operators for Kubernetes, see Latest SageMaker Operators for Kubernetes.

End of Support Frequently Asked Questions

Contents

- Why are we ending support for the original version of SageMaker Operators for Kubernetes?
- Where can I find more information about the new SageMaker Operators for Kubernetes and ACK?
- What does end of support (EOS) mean?
- How can I migrate my workload to the new SageMaker Operators for Kubernetes for training and inference?
- Which version of ACK should I migrate to?
- Are the initial SageMaker Operators for Kubernetes and the new Operators (ACK service controller for Amazon SageMaker) functionally equivalent?

Why are we ending support for the original version of SageMaker Operators for Kubernetes?

Users can now take advantage of the ACK service controller for Amazon SageMaker. The ACK service controller is a new generation of SageMaker Operators for Kubernetes based on AWS Controllers for Kubernetes (ACK), a community-driven project optimized for production, standardizing the way to expose AWS services via a Kubernetes operator. We are therefore announcing the end of support (EOS) for the original version (not ACK-based) of SageMaker Operators for Kubernetes. The support ends on Feb 15, 2023 along with Amazon Elastic Kubernetes Service Kubernetes 1.21.

For more information on ACK, see ACK history and tenets.

Where can I find more information about the new SageMaker Operators for Kubernetes and ACK?

- For more information about the new SageMaker Operators for Kubernetes, see the ACK service controller for Amazon SageMaker GitHub repository or read AWS Controllers for Kubernetes Documentation.
- For a tutorial on how to train a machine learning model with the ACK service controller for Amazon SageMaker using Amazon EKS, see this SageMaker example.
For an autoscaling example, see Scale SageMaker Workloads with Application Auto Scaling.

- For information on AWS Controller for Kubernetes (ACK), see the AWS Controllers for Kubernetes (ACK) documentation.
- For a list of supported SageMaker resources, see ACK API Reference.

What does end of support (EOS) mean?

While users can continue to use their current operators, we are no longer developing new features for the operators, nor will we release any patches or security updates for any issues found. v1.2.2 is the last release of SageMaker Operators for Kubernetes. Users should migrate their workloads to use the ACK service controller for Amazon SageMaker.

How can I migrate my workload to the new SageMaker Operators for Kubernetes for training and inference?

For information about migrating resources from the old to the new SageMaker Operators for Kubernetes, follow Migrate resources to the latest Operators.

Which version of ACK should I migrate to?

Users should migrate to the most recent released version of the ACK service controller for Amazon SageMaker.

Are the initial SageMaker Operators for Kubernetes and the new Operators (ACK service controller for Amazon SageMaker) functionally equivalent?

Yes, they are at feature parity.

A few highlights of the main notable differences between the two versions include:

- The Custom Resources Definitions (CRD) used by the ACK-based SageMaker Operators for Kubernetes follow the AWS API definition making it incompatible with the custom resources specifications from the SageMaker Operators for Kubernetes in its original version. Refer to the CRDs in the new controller or use the migration guide to adopt the resources and use the new controller.
- The Hosting Autoscaling policy is no longer part of the new SageMaker Operators for Kubernetes and has been migrated to the Application autoscaling ACK controller. To learn how
to use the application autoscaling controller to configure autoscaling on SageMaker Endpoints, follow this [autoscaling example](#).

- The HostingDeployment resource was used to create Models, Endpoint Configurations, and Endpoints in one CRD. The new SageMaker Operators for Kubernetes has a separate CRD for each of these resources.

## SageMaker Components for Kubeflow Pipelines

This document outlines how to use SageMaker Components for Kubeflow Pipelines. With these pipeline components, you can create and monitor native SageMaker training, tuning, endpoint deployment, and batch transform jobs from your Kubeflow Pipelines. By running Kubeflow Pipeline jobs on SageMaker, you move data processing and training jobs from the Kubernetes cluster to SageMaker's machine learning-optimized managed service. This document assumes prior knowledge of Kubernetes and Kubeflow.

### Contents

- [What are Kubeflow Pipelines?](#)
- [What are Kubeflow Pipeline components?](#)
- [Why use SageMaker Components for Kubeflow Pipelines?](#)
- [SageMaker Components for Kubeflow Pipelines versions](#)
- [List of SageMaker Components for Kubeflow Pipelines](#)
- [IAM permissions](#)
- [Converting pipelines to use SageMaker](#)
- [Install Kubeflow Pipelines](#)
- [Use SageMaker components](#)

### What are Kubeflow Pipelines?

Kubeflow Pipelines (KFP) is a platform for building and deploying portable, scalable machine learning (ML) workflows based on Docker containers. The Kubeflow Pipelines platform consists of the following:

- A user interface (UI) for managing and tracking experiments, jobs, and runs.
- An engine (Argo) for scheduling multi-step ML workflows.
• An SDK for defining and manipulating pipelines and components.
• Notebooks for interacting with the system using the SDK.

A pipeline is a description of an ML workflow expressed as a directed acyclic graph. Every step in the workflow is expressed as a Kubeflow Pipeline component, which is a AWS SDK for Python (Boto3) module.

For more information on Kubeflow Pipelines, see the Kubeflow Pipelines documentation.

What are Kubeflow Pipeline components?

A Kubeflow Pipeline component is a set of code used to execute one step of a Kubeflow pipeline. Components are represented by a Python module built into a Docker image. When the pipeline runs, the component's container is instantiated on one of the worker nodes on the Kubernetes cluster running Kubeflow, and your logic is executed. Pipeline components can read outputs from the previous components and create outputs that the next component in the pipeline can consume. These components make it fast and easy to write pipelines for experimentation and production environments without having to interact with the underlying Kubernetes infrastructure.

You can use SageMaker Components in your Kubeflow pipeline. Rather than encapsulating your logic in a custom container, you simply load the components and describe your pipeline using the Kubeflow Pipelines SDK. When the pipeline runs, your instructions are translated into a SageMaker job or deployment. The workload then runs on the fully managed infrastructure of SageMaker.

Why use SageMaker Components for Kubeflow Pipelines?

SageMaker Components for Kubeflow Pipelines offer an alternative to launching your compute-intensive jobs from SageMaker. The components integrate SageMaker with the portability and orchestration of Kubeflow Pipelines. Using the SageMaker Components for Kubeflow Pipelines, you can create and monitor your SageMaker resources as part of a Kubeflow Pipelines workflow. Each of the jobs in your pipelines runs on SageMaker instead of the local Kubernetes cluster allowing you to take advantage of key SageMaker features such as data labeling, large-scale hyperparameter tuning and distributed training jobs, or one-click secure and scalable model deployment. The job parameters, status, logs, and outputs from SageMaker are still accessible from the Kubeflow Pipelines UI.

The SageMaker components integrate key SageMaker features into your ML workflows from preparing data, to building, training, and deploying ML models. You can create a Kubeflow Pipeline built entirely using these components, or integrate individual components into your...
workflow as needed. The components are available in one or two versions. Each version of a component leverages a different backend. For more information on those versions, see [SageMaker Components for Kubeflow Pipelines versions](#).

There is no additional charge for using SageMaker Components for Kubeflow Pipelines. You incur charges for any SageMaker resources you use through these components.

**SageMaker Components for Kubeflow Pipelines versions**

SageMaker Components for Kubeflow Pipelines come in two versions. Each version leverages a different backend to create and manage resources on SageMaker.

- The SageMaker Components for Kubeflow Pipelines version 1 (v1.x or below) use [Boto3](https://aws.amazon.com/boto3/) as backend.

AWS introduced [ACK](https://github.com/aws-sagemaker-extensions/sagemaker-ack) to facilitate a Kubernetes-native way of managing AWS Cloud resources. ACK includes a set of AWS service-specific controllers, one of which is the SageMaker controller. The SageMaker controller makes it easier for machine learning developers and data scientists using Kubernetes as their control plane to train, tune, and deploy machine learning (ML) models in SageMaker. For more information, see [SageMaker Operators for Kubernetes](https://github.com/aws-sagemaker-extensions/sagemaker-ack).

Both versions of the SageMaker Components for Kubeflow Pipelines are supported. However, the version 2 provides some additional advantages. In particular, it offers:

1. A consistent experience to manage your SageMaker resources from any application; whether you are using Kubeflow pipelines, or Kubernetes CLI (`kubectl`) or other Kubeflow applications such as Notebooks.
2. The flexibility to manage and monitor your SageMaker resources outside of the Kubeflow pipeline workflow.
3. Zero setup time to use the SageMaker components if you deployed the full [Kubeflow on AWS](https://github.com/aws-sagemaker-extensions/sagemaker-ack) release since the SageMaker Operator is part of its deployment.

**List of SageMaker Components for Kubeflow Pipelines**

The following is a list of all SageMaker Components for Kubeflow Pipelines and their available versions. Alternatively, you can find all [SageMaker Components for Kubeflow Pipelines in GitHub](https://github.com/aws-sagemaker-extensions/sagemaker-ack).
We encourage users to utilize Version 2 of a SageMaker component wherever it is available.

Ground Truth components

- Ground Truth

The Ground Truth component enables you to submit SageMaker Ground Truth labeling jobs directly from a Kubeflow Pipelines workflow.

<table>
<thead>
<tr>
<th>Version 1 of the component</th>
<th>Version 2 of the component</th>
</tr>
</thead>
<tbody>
<tr>
<td>SageMaker Ground Truth Kubeflow Pipelines component version 1</td>
<td>X</td>
</tr>
</tbody>
</table>

- Workteam

The Workteam component enables you to create SageMaker private workteam jobs directly from a Kubeflow Pipelines workflow.

<table>
<thead>
<tr>
<th>Version 1 of the component</th>
<th>Version 2 of the component</th>
</tr>
</thead>
<tbody>
<tr>
<td>SageMaker create private workteam Kubeflow Pipelines component version 1</td>
<td>X</td>
</tr>
</tbody>
</table>

Data processing components

- Processing

The Processing component enables you to submit processing jobs to SageMaker directly from a Kubeflow Pipelines workflow.
### Training components

**Training**

The Training component allows you to submit SageMaker Training jobs directly from a Kubeflow Pipelines workflow.

<table>
<thead>
<tr>
<th>Version 1 of the component</th>
<th>Version 2 of the component</th>
</tr>
</thead>
<tbody>
<tr>
<td>SageMaker Processing Kubeflow Pipeline component version 1</td>
<td>X</td>
</tr>
</tbody>
</table>

**Hyperparameter Optimization**

The Hyperparameter Optimization component enables you to submit hyperparameter tuning jobs to SageMaker directly from a Kubeflow Pipelines workflow.

<table>
<thead>
<tr>
<th>Version 1 of the component</th>
<th>Version 2 of the component</th>
</tr>
</thead>
<tbody>
<tr>
<td>SageMaker Training Kubeflow Pipelines component version 1</td>
<td>SageMaker Training Kubeflow Pipelines component version 2</td>
</tr>
</tbody>
</table>

### Inference components

**Hosting Deploy**

The Hosting components allow you to deploy a model using SageMaker hosting services from a Kubeflow Pipelines workflow.
## SageMaker Hosting Services

Version 2 of the Hosting components consists of the three sub-components needed to create a hosting deployment on SageMaker.

- A **SageMaker Model** [Kubeflow Pipelines component version 2](#) responsible for the model artifacts and the model image registry path that contains the inference code.
- A **SageMaker Endpoint Configuration** [Kubeflow Pipelines component version 2](#) responsible for defining the configuration of the endpoint such as the instance type, models, number of instances, and serverless inference option.
- A **SageMaker Endpoint** [Kubeflow Pipelines component version 2](#) responsible for creating or updating the endpoint on SageMaker as specified in the endpoint configuration.

### Batch Transform

The Batch Transform component allows you to run inference jobs for an entire dataset in SageMaker from a Kubeflow Pipelines workflow.

## Batch Transform

The Batch Transform component allows you to run inference jobs for an entire dataset in SageMaker from a Kubeflow Pipelines workflow.

### Version 1 of the component | Version 2 of the component
---|---
**SageMaker Hosting Services - Create Endpoint** [Kubeflow Pipeline component version 1](#) | Version 2 of the Hosting components consists of the three sub-components needed to create a hosting deployment on SageMaker.
- A **SageMaker Model** [Kubeflow Pipelines component version 2](#) responsible for the model artifacts and the model image registry path that contains the inference code.
- A **SageMaker Endpoint Configuration** [Kubeflow Pipelines component version 2](#) responsible for defining the configuration of the endpoint such as the instance type, models, number of instances, and serverless inference option.
- A **SageMaker Endpoint** [Kubeflow Pipelines component version 2](#) responsible for creating or updating the endpoint on SageMaker as specified in the endpoint configuration.

### Model Monitor

- **Model Monitor**

---

**Kubernetes Orchestration**

4937
The Model Monitor components allow you to monitor the quality of SageMaker machine learning models in production from a Kubeflow Pipelines workflow.

<table>
<thead>
<tr>
<th>Version 1 of the component</th>
<th>Version 2 of the component</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>The Model Monitor components consist of four sub-components for monitoring drift in a model.</td>
</tr>
<tr>
<td></td>
<td>• A <a href="#">SageMaker Data Quality Job Definition Kubeflow Pipelines component version 2</a> responsible for monitoring drift in data quality.</td>
</tr>
<tr>
<td></td>
<td>• A <a href="#">SageMaker Model Quality Job Definition Kubeflow Pipelines component version 2</a> responsible for monitoring drift in model quality metrics.</td>
</tr>
<tr>
<td></td>
<td>• A <a href="#">SageMaker Model Bias Job Definition Kubeflow Pipelines component version 2</a> responsible for monitoring bias in a model's predictions.</td>
</tr>
<tr>
<td></td>
<td>• A <a href="#">SageMaker Model Explainability Job Definition Kubeflow Pipelines component version 2</a> responsible for monitoring drift in feature attribution.</td>
</tr>
</tbody>
</table>

Additionally, for on-schedule monitoring at a specified frequency, a fifth component, [SageMaker Monitoring Schedule Kubeflow Pipelines component version 2](#), is responsible for monitoring the data collected from a real-time endpoint on a schedule.

For more information on Amazon SageMaker Model Monitor, see [Monitor data and model quality](#).
Deploying Kubeflow Pipelines with SageMaker components requires the following three layers of authentication:

- An IAM role granting your gateway node (which can be your local machine or a remote instance) access to the Amazon Elastic Kubernetes Service (Amazon EKS) cluster.

The user accessing the gateway node assumes this role to:
- Create an Amazon EKS cluster and install KFP
- Create IAM roles
- Create Amazon S3 buckets for your sample input data

The role requires the following permissions:
- CloudWatchLogsFullAccess
- AWSCloudFormationFullAccess
- IAMFullAccess
- AmazonS3FullAccess
- AmazonEC2FullAccess
- AmazonEKSAdminPolicy (Create this policy using the schema from Amazon EKS Identity-Based Policy Examples)

- A Kubernetes IAM execution role assumed by Kubernetes pipeline pods (kfp-example-pod-role) or the SageMaker Operator for Kubernetes controller pod to access SageMaker. This role is used to create and monitor SageMaker jobs from Kubernetes.

The role requires the following permission:
- AmazonSageMakerFullAccess

You can limit permissions to the KFP and controller pods by creating and attaching your own custom policy.

- A SageMaker IAM execution role assumed by SageMaker jobs to access AWS resources such as Amazon S3 or Amazon ECR (kfp-example-sagemaker-execution-role).

SageMaker jobs use this role to:
- Access SageMaker resources
- Input Data from Amazon S3
- Store your output model to Amazon S3

The role requires the following permissions:
Converting pipelines to use SageMaker

You can convert an existing pipeline to use SageMaker by porting your generic Python processing containers and training containers. If you are using SageMaker for inference, you also need to attach IAM permissions to your cluster and convert an artifact to a model.

Install Kubeflow Pipelines

Kubeflow Pipelines (KFP) is the pipeline orchestration component of Kubeflow.

You can deploy Kubeflow Pipelines (KFP) on an existing Amazon Elastic Kubernetes Service (Amazon EKS) or create a new Amazon EKS cluster. Use a gateway node to interact with your cluster. The gateway node can be your local machine or an Amazon EC2 instance.

The following section guides you through the steps to set up and configure these resources.

Topics

- Choose an installation option
- Configure your pipeline permissions to access SageMaker
- Access the KFP UI (Kubeflow Dashboard)

Choose an installation option

Kubeflow Pipelines is available as a core component of the full distribution of Kubeflow on AWS or as a standalone installation.

Select the option that applies to your use case:

1. Full Kubeflow on AWS Deployment

   To use other Kubeflow components in addition to Kubeflow Pipelines, choose the full AWS distribution of Kubeflow deployment.

2. Standalone Kubeflow Pipelines Deployment

   To use the Kubeflow Pipelines without the other components of Kubeflow, install Kubeflow pipelines standalone.
Full Kubeflow on AWS Deployment

To install the full release of Kubeflow on AWS, choose the vanilla deployment option from Kubeflow on AWS deployment guide or any other deployment option supporting integrations with various AWS services (Amazon S3, Amazon RDS, Amazon Cognito).

Standalone Kubeflow Pipelines Deployment

This section assumes that your user has permissions to create roles and define policies for the role.

Set up a gateway node

You can use your local machine or an Amazon EC2 instance as your gateway node. A gateway node is used to create an Amazon EKS cluster and access the Kubeflow Pipelines UI.

Complete the following steps to set up your node.

1. **Create a gateway node.**

   You can use an existing Amazon EC2 instance or create a new instance with the latest Ubuntu 18.04 DLAMI version using the steps in Launching and Configuring a DLAMI.

2. **Create an IAM role to grant your gateway node access to AWS resources.**

   Create an IAM role with permissions to the following resources: CloudWatch, AWS CloudFormation, IAM, Amazon EC2, Amazon S3, Amazon EKS.

   Attach the following policies to the IAM role:

   - CloudWatchLogsFullAccess
   - AWSCloudFormationFullAccess
   - IAMFullAccess
   - AmazonS3FullAccess
   - AmazonEC2FullAccess
   - AmazonEKSAutoPolicy (Create this policy using the schema from Amazon EKS Identity-Based Policy Examples)

   For information on adding IAM permissions to an IAM role, see Adding and removing IAM identity permissions.
3. **Install the following tools and clients**

Install and configure the following tools and resources on your gateway node to access the Amazon EKS cluster and KFP User Interface (UI).

- **AWS CLI**: The command line tool for working with AWS services. For AWS CLI configuration information, see [Configuring the AWS CLI](#).
- **aws-iam-authenticator** version 0.1.31 and above: A tool to use AWS IAM credentials to authenticate to a Kubernetes cluster.
- **eksctl** version above 0.15: The command line tool for working with Amazon EKS clusters.
- **kubectl**: The command line tool for working with Kubernetes clusters. The version needs to match your Kubernetes version within one minor version.
- **AWS SDK for Python (Boto3)**.

```
pip install boto3
```

**Set up an Amazon EKS cluster**

1. If you do not have an existing Amazon EKS cluster, run the following steps from the command line of your gateway node, skip this step otherwise.

   a. Run the following command to create an Amazon EKS cluster with version 1.17 or above. Replace `<clustername>` with any name for your cluster.

   ```bash
   eksctl create cluster --name <clustername> --region us-east-1 --auto-kubeconfig --timeout=50m --managed --nodes=1
   ```

   b. When the cluster creation is complete, ensure that you have access to your cluster by listing the cluster's nodes.

   ```bash
   kubectl get nodes
   ```

2. Ensure that the current `kubectl` context points to your cluster with the following command. The current context is marked with an asterisk (*) in the output.

   ```bash
   kubectl config get-contexts
   ```
3. If the desired cluster is not configured as your current default, update the default with the following command.

```
aws eks update-kubeconfig --name <clusternam e> --region us-east-1
```

### Install Kubeflow Pipelines

Run the following steps from the terminal of your gateway node to install Kubeflow Pipelines on your cluster.

1. **Install all cert-manager components.**

   ```
kubectl apply -f https://github.com/cert-manager/cert-manager/releases/download/v1.9.1/cert-manager.yaml
   ```

2. **Install the Kubeflow Pipelines.**

   ```
   export PIPELINE_VERSION=2.0.0-alpha.5
ekubectl apply -k "github.com/kubeflow/pipelines/manifests/kustomize/env/cert-manager/cluster-scoped-resources?ref=$KFP_VERSION"
kubectl wait --for condition=established --timeout=60s crd/applications.app.k8s.io
kubectl apply -k "github.com/kubeflow/pipelines/manifests/kustomize/env/cert-manager/dev?ref=$KFP_VERSION"
   ```

3. **Ensure that the Kubeflow Pipelines service and other related resources are running.**

   ```
kubectl -n kubeflow get all | grep pipeline
   ```

   Your output should look like the following.

<table>
<thead>
<tr>
<th>Pod Name</th>
<th>Phase</th>
<th>IP</th>
</tr>
</thead>
<tbody>
<tr>
<td>pod/ml-pipeline-6b88c67994-kdtjv</td>
<td>1/1</td>
<td>Running</td>
</tr>
<tr>
<td>pod/ml-pipeline-persistenceagent-64d74dfdbf-66stk</td>
<td>1/1</td>
<td>Running</td>
</tr>
<tr>
<td>pod/ml-pipeline-scheduledworkflow-65bdf46db7-5x9qj</td>
<td>1/1</td>
<td>Running</td>
</tr>
</tbody>
</table>
Configure your pipeline permissions to access SageMaker

In this section, you create an IAM execution role granting Kubeflow Pipeline pods access to SageMaker services.
Configuration for SageMaker components version 2

To run SageMaker Components version 2 for Kubeflow Pipelines, you need to install SageMaker Operator for Kubernetes and configure Role-Based Access Control (RBAC) allowing the Kubeflow Pipelines pods to create SageMaker custom resources in your Kubernetes cluster.

⚠️ Important

Follow this section if you are using Kubeflow pipelines standalone deployment. If you are using AWS distribution of Kubeflow version 1.6.0-aws-b1.0.0 or above, SageMaker components version 2 are already set up.


   Follow the Setup section of Machine Learning with ACK SageMaker Controller tutorial.

2. Configure RBAC permissions for the execution role (service account) used by Kubeflow Pipelines pods. In Kubeflow Pipelines standalone deployment, pipeline runs are executed in the namespace kubeflow using the pipeline-runner service account.

   a. Create a RoleBinding that gives the service account permission to manage SageMaker custom resources.

```
cat > manage_sagemaker_cr.yaml <<EOF
apiVersion: rbac.authorization.k8s.io/v1
kind: RoleBinding
metadata:
  name: manage-sagemaker-cr
  namespace: kubeflow
subjects:
  - kind: ServiceAccount
    name: pipeline-runner
    namespace: kubeflow
roleRef:
  kind: ClusterRole
  name: ack-sagemaker-controller
  apiGroup: rbac.authorization.k8s.io
EOF

kubectl apply -f manage_sagemaker_cr.yaml
```
b. Ensure that the rolebinding was created by running:

```
kubectl get rolebinding manage-sagemaker-cr -n kubeflow -o yaml
```

**Configuration for SageMaker components version 1**

To run SageMaker Components version 1 for Kubeflow Pipelines, the Kubeflow Pipeline pods need access to SageMaker.

⚠️ **Important**

Follow this section whether you are using the full Kubeflow on AWS deployment or Kubeflow Pipelines standalone.

To create an IAM execution role granting Kubeflow pipeline pods access to SageMaker, follow those steps:

1. Export your cluster name (e.g., `my-cluster-name`) and cluster region (e.g., `us-east-1`).

   ```
   export CLUSTER_NAME=my-cluster-name
   export CLUSTER_REGION=us-east-1
   ```

2. Export the namespace and service account name according to your installation.

   - For the full Kubeflow on AWS installation, export your profile namespace (e.g., `kubeflow-user-example-com`) and `default-editor` as the service account.

     ```
     export NAMESPACE=kubeflow-user-example-com
     export KUBEFLOW_PIPELINE_POD_SERVICE_ACCOUNT=default-editor
     ```

   - For the standalone Pipelines deployment, export `kubeflow` as the namespace and `pipeline-runner` as the service account.

     ```
     export NAMESPACE=kubeflow
     export KUBEFLOW_PIPELINE_POD_SERVICE_ACCOUNT=pipeline-runner
     ```

3. Create an [IAM OIDC provider for the Amazon EKS cluster](#) with the following command.
4. Create an IAM execution role for the KFP pods to access AWS services (SageMaker, CloudWatch).

```bash
eksctl create iamserviceaccount \
  --name ${KUBEFLOW_PIPELINE_POD_SERVICE_ACCOUNT} \ 
  --namespace ${NAMESPACE} --cluster ${CLUSTER_NAME} \ 
  --region ${CLUSTER_REGION} \
  --attach-policy-arn arn:aws:iam::aws:policy/AmazonSageMakerFullAccess \ 
  --attach-policy-arn arn:aws:iam::aws:policy/CloudWatchLogsFullAccess \ 
  --override-existing-serviceaccounts \ 
  --approve
```

Once your pipeline permissions are configured to access SageMaker Components version 1, follow the SageMaker components for Kubeflow pipelines guide on the [Kubeflow on AWS documentation](https://kubeflow.org/docs/guide/)

**Access the KFP UI (Kubeflow Dashboard)**

The Kubeflow Pipelines UI is used for managing and tracking experiments, jobs, and runs on your cluster. For instructions on how to access the Kubeflow Pipelines UI from your gateway node, follow the steps that apply to your deployment option in this section.

**Full Kubeflow on AWS Deployment**

Follow the instructions on the [Kubeflow on AWS website](https://kubeflow.org/docs/guide/) to connect to the Kubeflow dashboard and navigate to the pipelines tab.

**Standalone Kubeflow Pipelines Deployment**

Use port forwarding to access the Kubeflow Pipelines UI from your gateway node by following those steps.

**Set up port forwarding to the KFP UI service**

Run the following command from the command line of your gateway node.

1. Verify that the KFP UI service is running using the following command.

```bash
kubect1 -n kubeflow get service ml-pipeline-ui
```
2. Run the following command to set up port forwarding to the KFP UI service. This forwards the KFP UI to port 8080 on your gateway node and allows you to access the KFP UI from your browser.

```
kubectl port-forward -n kubeflow service/ml-pipeline-ui 8080:80
```

The port forward from your remote machine drops if there is no activity. Run this command again if your dashboard is unable to get logs or updates. If the commands return an error, ensure that there is no process already running on the port you are trying to use.

**Access the KFP UI service**

Your method of accessing the KFP UI depends on your gateway node type.

- **Local machine as the gateway node:**
  1. Access the dashboard in your browser as follows:

      `http://localhost:8080`

  2. Choose **Pipelines** to access the pipelines UI.

- **Amazon EC2 instance as the gateway node:**
  1. You need to set up an SSH tunnel on your Amazon EC2 instance to access the Kubeflow dashboard from your local machine's browser.

      From a new terminal session in your local machine, run the following. Replace `<public-DNS-of-gateway-node>` with the IP address of your instance found on the Amazon EC2 console. You can also use the public DNS. Replace `<path_to_key>` with the path to the pem key used to access the gateway node.

      ```
      public_DNS_address=<public-DNS-of-gateway-node>
      key=<path_to_key>
      
      on Ubuntu:
      ssh -i ${key} -L 9000:localhost:8080 ubuntu@$public_DNS_address
      ```
or on Amazon Linux:
```
ssh -i ${key} -L 9000:localhost:8080 ec2-user@${public_DNS_address}
```

2. Access the dashboard in your browser.

```
http://localhost:9000
```

3. Choose **Pipelines** to access the KFP UI.

(Optional) Grant SageMaker notebook instances access to Amazon EKS, and run KFP pipelines from your notebook.

A SageMaker notebook instance is a fully managed Amazon EC2 compute instance that runs the Jupyter Notebook App. You can use a notebook instance to create and manage Jupyter notebooks then define, compile, deploy, and run your KFP pipelines using AWS SDK for Python (Boto3) or the KFP CLI.

1. Follow the steps in [Create a SageMaker Notebook Instance](#) to create your notebook instance, then attach the S3FullAccess policy to its IAM execution role.

2. From the command line of your gateway node, run the following command to retrieve the IAM role ARN of the notebook instance you created. Replace `<instance-name>` with the name of your instance.

```
aws sagemaker describe-notebook-instance --notebook-instance-name <instance-name> --region <region> --output text --query 'RoleArn'
```

This command outputs the IAM role ARN in the `arn:aws:iam::<account-id>:role/<role-name>` format. Take note of this ARN.

3. Run this command to attach the following policies (AmazonSageMakerFullAccess, AmazonEKSWorkerNodePolicy, AmazonS3FullAccess) to this IAM role. Replace `<role-name>` with the `<role-name>` in your ARN.

```
aws iam attach-role-policy --role-name <role-name> --policy-arn arn:aws:iam::aws:policy/AmazonSageMakerFullAccess
aws iam attach-role-policy --role-name <role-name> --policy-arn arn:aws:iam::aws:policy/AmazonEKSWorkerNodePolicy
aws iam attach-role-policy --role-name <role-name> --policy-arn arn:aws:iam::aws:policy/AmazonS3FullAccess
```
4. Amazon EKS clusters use IAM roles to control access to the cluster. The rules are implemented in a config map named `aws-auth`. `eksctl` provides commands to read and edit the `aws-auth` config map. Only the users that have access to the cluster can edit this config map.

    system:masters is one of the default user groups with super user permissions to the cluster. Add your user to this group or create a group with more restrictive permissions.

5. Bind the role to your cluster by running the following command. Replace `<IAM-Role-arn>` with the ARN of the IAM role. `<your_username>` can be any unique username.

   ```bash
   eksctl create iamidentitymapping \
   --cluster <cluster-name> \
   --arn <IAM-Role-arn> \
   --group system:masters \
   --username <your-username> \
   --region <region>
   ```

6. Open a Jupyter notebook on your SageMaker instance and run the following command to ensure that it has access to the cluster.

   ```bash
   aws eks --region <region> update-kubeconfig --name <cluster-name>
   kubectl -n kubeflow get all | grep pipeline
   ```

**Use SageMaker components**

In this tutorial, you run a pipeline using SageMaker Components for Kubeflow Pipelines to train a classification model using Kmeans with the MNIST dataset on SageMaker. The workflow uses Kubeflow Pipelines as the orchestrator and SageMaker to execute each step of the workflow. The example was taken from an existing SageMaker example and modified to work with SageMaker Components for Kubeflow Pipelines.

You can define your pipeline in Python using AWS SDK for Python (Boto3) then use the KFP dashboard, KFP CLI, or Boto3 to compile, deploy, and run your workflows. The full code for the MNIST classification pipeline example is available in the Kubeflow Github repository. To use it, clone the Python files to your gateway node.

You can find additional SageMaker Kubeflow Pipelines examples on GitHub. For information on the components used, see the KubeFlow Pipelines GitHub repository.
To run the classification pipeline example, create a SageMaker IAM execution role granting your training job the permission to access AWS resources, then continue with the steps that correspond to your deployment option.

**Create a SageMaker execution role**

The `kfp-example-sagemaker-execution-role` IAM role is a runtime role assumed by SageMaker jobs to access AWS resources. In the following command, you create an IAM execution role named `kfp-example-sagemaker-execution-role`, attach two managed policies (`AmazonSageMakerFullAccess`, `AmazonS3FullAccess`), and create a trust relationship with SageMaker to grant SageMaker jobs access to those AWS resources.

You provide this role as an input parameter when running the pipeline.

Run the following command to create the role. Note the ARN that is returned in your output.

```
SAGEMAKER_EXECUTION_ROLE_NAME=kfp-example-sagemaker-execution-role

aws iam create-role --role-name ${SAGEMAKER_EXECUTION_ROLE_NAME} --assume-role-policy-document "$TRUST"
aws iam attach-role-policy --role-name ${SAGEMAKER_EXECUTION_ROLE_NAME} --policy-arn arn:aws:iam::aws:policy/AmazonSageMakerFullAccess
aws iam attach-role-policy --role-name ${SAGEMAKER_EXECUTION_ROLE_NAME} --policy-arn arn:aws:iam::aws:policy/AmazonS3FullAccess

aws iam get-role --role-name ${SAGEMAKER_EXECUTION_ROLE_NAME} --output text --query 'Role.Arn'
```

**Full Kubeflow on AWS Deployment**

Follow the instructions of the [SageMaker Training Pipeline tutorial for MNIST Classification with K-Means](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-training-pipeline.html).

**Standalone Kubeflow Pipelines Deployment**

**Prepare datasets**

To run the pipelines, you need to upload the data extraction pre-processing script to an Amazon S3 bucket. This bucket and all resources for this example must be located in the `us-east-1` region. For information on creating a bucket, see [Creating a bucket](https://docs.aws.amazon.com/AmazonS3/latest/userguide/creating-a-bucket.html).
From the `mnist-kmeans-sagemaker` folder of the Kubeflow repository you cloned on your gateway node, run the following command to upload the `kmeans_preprocessing.py` file to your Amazon S3 bucket. Change `<bucket-name>` to the name of your Amazon S3 bucket.

```bash
aws s3 cp mnist-kmeans-sagemaker/kmeans_preprocessing.py s3://<bucket-name>/mnist_kmeans_example/processing_code/kmeans_preprocessing.py
```

**Compile and deploy your pipeline**

After defining the pipeline, you must compile it to an intermediate representation before you submit it to the Kubeflow Pipelines service on your cluster. The intermediate representation is a workflow specification in the form of a YAML file compressed into a tar.gz file. You need the KFP SDK to compile your pipeline.

**Install KFP SDK**

Run the following from the command line of your gateway node:

1. Install the KFP SDK following the instructions in the [Kubeflow pipelines documentation](https://www.kubeflow.org/docs/user-guide/)
2. Verify that the KFP SDK is installed with the following command:
   ```bash
   pip show kfp
   ```
3. Verify that `dsl-compile` has been installed correctly as follows:
   ```bash
   which dsl-compile
   ```

**Compile your pipeline**

You have three options to interact with Kubeflow Pipelines: KFP UI, KFP CLI, or the KFP SDK. The following sections illustrate the workflow using the KFP UI and CLI.

Complete the following steps from your gateway node.

1. Modify your Python file with your Amazon S3 bucket name and IAM role ARN.
2. Use the `dsl-compile` command from the command line to compile your pipeline as follows. Replace `<path-to-python-file>` with the path to your pipeline and `<path-to-output>` with the location where you want your tar.gz file to be.
Upload and run the pipeline using the KFP CLI

Complete the following steps from the command line of your gateway node. KFP organizes runs of your pipeline as experiments. You have the option to specify an experiment name. If you do not specify one, the run will be listed under Default experiment.

1. Upload your pipeline as follows:

```bash
dsl-compile --py <path-to-python-file> --output <path-to-output>
```

Your output should look like the following. Take note of the pipeline ID.

```
Pipeline 29c3ff21-49f5-4dfe-94f6-618c0e2420fe has been submitted

Pipeline Details
------------------
ID 29c3ff21-49f5-4dfe-94f6-618c0e2420fe
Name sm-pipeline
Description
 Uploaded at 2020-04-30T20:22:39+00:00
...
...
```

2. Create a run using the following command. The KFP CLI run command currently does not support specifying input parameters while creating the run. You need to update your parameters in the AWS SDK for Python (Boto3) pipeline file before compiling. Replace <experiment-name> and <job-name> with any names. Replace <pipeline-id> with the ID of your submitted pipeline. Replace <your-role-arn> with the ARN of kfp-example-pod-role. Replace <your-bucket-name> with the name of the Amazon S3 bucket you created.

```bash
kfp run submit --experiment-name <experiment-name> --run-name <job-name> --pipeline-id <pipeline-id> role_arn="<your-role-arn>" bucket_name="<your-bucket-name>"
```
You can also directly submit a run using the compiled pipeline package created as the output of the `dsl-compile` command.

```
kfp run submit --experiment-name <experiment-name> --run-name <job-name> --package-file <path-to-output> role_arn="<your-role-arn>" bucket_name="<your-bucket-name>"
```

Your output should look like the following:

```
Creating experiment aws.
Run 95084a2c-f18d-4b77-a9da-eba00bf01e63 is submitted
+--------------------------------------+--------+----------+
| run id )| name )| status )| created at |
+--------------------------------------+--------+----------+
| 95084a2c-f18d-4b77-a9da-eba00bf01e63 | sm-job |         | 2020-04-30T20:36:41+00:00 |
+--------------------------------------+--------+----------+
```

3. Navigate to the UI to check the progress of the job.

**Upload and run the pipeline using the KFP UI**

1. On the left panel, choose the **Pipelines** tab.
2. In the upper-right corner, choose **+UploadPipeline**.
3. Enter the pipeline name and description.
4. Choose **Upload a file** and enter the path to the tar.gz file you created using the CLI or with AWS SDK for Python (Boto3).
5. On the left panel, choose the **Pipelines** tab.
6. Find the pipeline you created.
7. Choose **+CreateRun**.
8. Enter your input parameters.
9. Choose **Run**.
Run predictions

Once your classification pipeline is deployed, you can run classification predictions against the endpoint that was created by the Deploy component. Use the KFP UI to check the output artifacts for `sagemaker-deploy-model-endpoint_name`. Download the .tgz file to extract the endpoint name or check the SageMaker console in the region you used.

Configure permissions to run predictions

If you want to run predictions from your gateway node, skip this section.

To use any other machine to run predictions, assign the `sagemaker:InvokeEndpoint` permission to the IAM role used by the client machine.

1. On your gateway node, run the following to create an IAM policy file:

   ```
cat <<EoF > ./sagemaker-invoke.json
{
   "Version": "2012-10-17",
   "Statement": [
     {
       "Effect": "Allow",
       "Action": [
         "sagemaker:InvokeEndpoint"
       ],
       "Resource": "*
     }
   ]
}
EoF
```

2. Attach the policy to the IAM role of the client node.

   Run the following command. Replace `<your-instance-IAM-role>` with the name of the IAM role. Replace `<path-to-sagemaker-invoke-json>` with the path to the policy file you created.

   ```
   aws iam put-role-policy --role-name <your-instance-IAM-role> --policy-name sagemaker-invoke-for-worker --policy-document file://<path-to-sagemaker-invoke-json>
   ```
Run predictions

1. Create a AWS SDK for Python (Boto3) file from your client machine named `mnist-predictions.py` with the following content. Replace the `ENDPOINT_NAME` variable. The script loads the MNIST dataset, creates a CSV from those digits, then sends the CSV to the endpoint for prediction and prints the results.

```python
import boto3
import gzip
import io
import json
import numpy
import pickle

ENDPOINT_NAME='<endpoint-name>'
region = boto3.Session().region_name

# S3 bucket where the original mnist data is downloaded and stored
downloaded_data_bucket = f"jumpstart-cache-prod-{region}"
download_data_prefix = "lp-notebooks-datasets/mnist"

# Download the dataset
s3 = boto3.client("s3")
s3.download_file(downloaded_data_bucket, f"{download_data_prefix}/mnist.pkl.gz", "mnist.pkl.gz")

# Load the dataset
with gzip.open('mnist.pkl.gz', 'rb') as f:
    train_set, valid_set, test_set = pickle.load(f, encoding='latin1')

# Simple function to create a csv from our numpy array
def np2csv(arr):
    csv = io.BytesIO()
    numpy.savetxt(csv, arr, delimiter=',', fmt='%g')
    return csv.getvalue().decode().rstrip()

runtime = boto3.Session(region).client('sagemaker-runtime')

payload = np2csv(train_set[0][30:31])

response = runtime.invoke_endpoint(EndpointName=ENDPOINT_NAME,
                                   ContentType='text/csv',
                                   Body=payload)
```
result = json.loads(response['Body'].read().decode())
print(result)

2. Run the AWS SDK for Python (Boto3) file as follows:

    python mnist-predictions.py

View results and logs

When the pipeline is running, you can choose any component to check execution details, such as inputs and outputs. This lists the names of created resources.

If the KFP request is successfully processed and an SageMaker job is created, the component logs in the KFP UI provide a link to the job created in SageMaker. The CloudWatch logs are also provided if the job is successfully created.

If you run too many pipeline jobs on the same cluster, you may see an error message that indicates that you do not have enough pods available. To fix this, log in to your gateway node and delete the pods created by the pipelines you are not using:

    kubectl get pods -n kubeflow
    kubectl delete pods -n kubeflow <name-of-pipeline-pod>

Cleanup

When you're finished with your pipeline, you need to clean up your resources.

1. From the KFP dashboard, terminate your pipeline runs if they do not exit properly by choosing **Terminate**.

2. If the **Terminate** option doesn't work, log in to your gateway node and manually terminate all the pods created by your pipeline run as follows:

    kubectl get pods -n kubeflow
    kubectl delete pods -n kubeflow <name-of-pipeline-pod>

3. Using your AWS account, log in to the SageMaker service. Manually stop all training, batch transform, and HPO jobs. Delete models, data buckets, and endpoints to avoid incurring any additional costs. Terminating the pipeline runs does not stop the jobs in SageMaker.
SageMaker Notebook Jobs

You can use Amazon SageMaker to interactively build, train, and deploy machine learning models from your Jupyter notebook in any JupyterLab environment. However, there are various scenarios in which you might want to run your notebook as a noninteractive, scheduled job. For example, you might want to create regular audit reports that analyze all training jobs run over a certain time frame and analyze the business value of deploying those models into production. Or you might want to scale up a feature engineering job after testing the data transformation logic on a small subset of data. Other common use cases include:

- Scheduling jobs for model drift monitoring
- Exploring the parameter space for better models

In these scenarios, you can use SageMaker Notebook Jobs to create a noninteractive job (which SageMaker runs as an underlying training job) to either run on demand or on a schedule. SageMaker Notebook Jobs provides an intuitive user interface so you can schedule your jobs right from JupyterLab by choosing the Notebook Jobs widget in your notebook. You can also schedule your jobs using the SageMaker Python SDK, which offers the flexibility of scheduling multiple notebook jobs in a pipeline workflow. You can run multiple notebooks in parallel, and parameterize cells in your notebooks to customize the input parameters.

This feature leverages the Amazon EventBridge, SageMaker Training and SageMaker Pipelines services and is available for use in your Jupyter notebook in any of the following environments:

- Studio, Studio Lab, Studio Classic, or Notebook Instances
- Local setup, such as your local machine, where you run JupyterLab

Prerequisites

To schedule a notebook job, make sure you meet the following criteria:

- Ensure your Jupyter notebook and any initialization or startup scripts are self-contained with respect to code and software packages. Otherwise, your noninteractive job may incur errors.
- Review Constraints and considerations to make sure you properly configured your Jupyter notebook, network settings, and container settings.
- Ensure your notebook can access needed external resources, such as Amazon EMR clusters.
• If you are setting up Notebook Jobs in a local Jupyter notebook, complete the installation. For instructions, see [Installation Guide](#).

• If you connect to an Amazon EMR cluster in your notebook and want to parameterize your Amazon EMR connection command, you must apply a workaround using environment variables to pass parameters. For details, see [Connect to an Amazon EMR cluster from your notebook](#).

• If you connect to an Amazon EMR cluster using Kerberos, LDAP, or HTTP Basic Auth authentication, you must use the AWS Secrets Manager to pass your security credentials to your Amazon EMR connection command. For details, see [Connect to an Amazon EMR cluster from your notebook](#).

• (optional) If you want the UI to preload a script to run upon notebook startup, your admin must install it with a Lifecycle Configuration (LCC). For information about how to use a LCC script, see [Customize a Notebook Instance Using a Lifecycle Configuration Script](#).

---

**Installation Guide**

The following discussion includes detailed instructions about additional installation you need to perform so you can use Notebook Jobs in your JupyterLab environment.

**For Amazon SageMaker Studio and Amazon SageMaker Studio Lab**

If your notebook is in Amazon SageMaker Studio or Amazon SageMaker Studio Lab, you don't need to perform additional installation—SageMaker Notebook Jobs is built into the platform. To set up required permissions for Studio, see [Install policies and permissions for Studio](#).

**For local Jupyter notebooks**

If you want to use SageMaker Notebook Jobs for your local JupyterLab environment, you need to perform additional installation.

To install SageMaker Notebook Jobs, complete the following steps:

1. Install Python 3. For details, see [Installing Python 3 and Python Packages](#).
2. Install JupyterLab version 3 or higher. For details, see [JupyterLab SDK documentation](#).
3. Install the AWS CLI. For details, see [Installing or updating the latest version of the AWS CLI](#).
4. Install two sets of permissions. The IAM user needs permissions to submit jobs to SageMaker, and once submitted, the notebook job itself assumes an IAM role that needs permissions to access resources depending on the job tasks.
a. If you haven’t yet created an IAM user, see Creating an IAM user in your AWS account.

b. If you haven’t yet created your notebook job role, see Creating a role to delegate permissions to an IAM user.

c. Attach the necessary permissions and trust policy to attach to your user and role. For step-by-step instructions and permission details, see Install policies and permissions for local Jupyter environments.

5. Generate AWS credentials for your newly-created IAM user and save them in the credentials file (~/.aws/credentials) of your JupyterLab environment. You can do this with the CLI command `aws configure`. For instructions, see section Set and view configuration settings using commands in Configuration and credential file settings.

6. (optional) By default, the scheduler extension uses a pre-built SageMaker Docker image with Python 2.0. Any non-default kernel used in the notebook should be installed in the container. If you want to run your notebook in a container or Docker image, you need to create an Amazon Elastic Container Registry (Amazon ECR) image. For information about how to push a Docker image to an Amazon ECR, see Pushing a Docker Image.

7. Add the JupyterLab extension for SageMaker Notebook Jobs. You can add it to your JupyterLab environment with the command: `pip install amazon_sagemaker_jupyter_scheduler`. You may need to restart your Jupyter server with the command: `sudo systemctl restart jupyter-server`.

8. Start JupyterLab with the command: `jupyter lab`.

9. Verify that the Notebook Jobs widget appears in your Jupyter notebook taskbar.

Install policies and permissions for Studio

Before you schedule your first notebook run, make sure that you install the proper policies and permissions. The following instructions show you how to configure the following permissions:

- Job execution role trust relationships
- Additional IAM permissions attached to the job execution role
- (optional) The AWS KMS permission policy to use a custom KMS key
Important

If your AWS account belongs to an organization with service control policies (SCP) in place, your effective permissions are the logical intersection between what is allowed by the SCPs and what is allowed by your IAM role and user policies. For example, if your organization's SCP specifies that you can only access resources in us-east-1 and us-west-1, and your policies only allow you to access resources in us-west-1 and us-west-2, then ultimately you can only access resources in us-west-1. If you want to exercise all the permissions allowed in your role and user policies, your organization's SCPs should grant the same set of permissions as your own IAM user and role policies. For details about how to determine your allowed requests, see Determining whether a request is allowed or denied within an account.

Trust relationships

To modify the trust relationships, complete the following steps:

1. Open the IAM console.
2. Select Roles in the left panel.
3. Find the job execution role for your notebook job and choose the role name.
4. Choose the Trust relationships tab.
5. Choose Edit trust policy.
6. Copy and paste the following policy:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Service": "sagemaker.amazonaws.com"
            },
            "Action": "sts:AssumeRole"
        },
        {
            "Effect": "Allow",
            "Principal": {
                "Service": "events.amazonaws.com"
            }
        }
    ]
}
```
7. Choose **Update Policy**.

### Additional IAM permissions

You might need to include additional IAM permissions in the following situations:

- Your Studio execution and notebook job roles differ
- You need to access Amazon S3 resources through a S3 VPC endpoint
- You want to use a custom KMS key to encrypt your input and output Amazon S3 buckets

The following discussion provides the policies you need for each case.

#### Permissions needed if your Studio execution and notebook job roles differ

The following JSON snippet is an example policy that you should add to the Studio execution and notebook job roles if you don’t use the Studio execution role as the notebook job role. Review and modify this policy if you need to further restrict privileges.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": "iam:PassRole",
      "Resource": "arn:aws:iam::*:role/*",
      "Condition": {
        "StringLike": {
          "iam:PassedToService": [
            "sagemaker.amazonaws.com",
            "events.amazonaws.com"
          ]
        }
      }
    }
  ]
}
```
"Effect": "Allow",
"Action": [
    "events:TagResource",
    "events:DeleteRule",
    "events:PutTargets",
    "events:DescribeRule",
    "events:PutRule",
    "events:RemoveTargets",
    "events:DisableRule",
    "events:EnableRule"
],
"Resource": "*",
"Condition": {
    "StringEquals": {
        "aws:ResourceTag/sagemaker:is-scheduling-notebook-job": "true"
    }
}
],
{
    "Effect": "Allow",
    "Action": [
        "s3:CreateBucket",
        "s3:PutBucketVersioning",
        "s3:PutEncryptionConfiguration"
    ],
    "Resource": "arn:aws:s3:::sagemaker-automated-execution-*"
},
{
    "Sid": "S3DriverAccess",
    "Effect": "Allow",
    "Action": [
        "s3:ListBucket",
        "s3:GetObject",
        "s3:GetBucketLocation"
    ],
    "Resource": [
        "arn:aws:s3:::sagemaker-headless-execution-*"
    ]
}
]
"Resource": [
    "arn:aws:sagemaker:*:*:user-profile/**",
    "arn:aws:sagemaker:*:*:space/**",
    "arn:aws:sagemaker:*:*:training-job/**",
    "arn:aws:sagemaker:*:*:pipeline/**"
  ],
  {
    "Effect":"Allow",
    "Action": [
      "sagemaker:AddTags"
    ],
    "Resource": [
      "arn:aws:sagemaker:*:*:training-job/**",
      "arn:aws:sagemaker:*:*:pipeline/**"
    ]
  },
  {
    "Effect":"Allow",
    "Action": [
      "ec2:CreateNetworkInterface",
      "ec2:CreateNetworkInterfacePermission",
      "ec2:CreateVpcEndpoint",
      "ec2:DeleteNetworkInterface",
      "ec2:DeleteNetworkInterfacePermission",
      "ec2:DescribeDhcpOptions",
      "ec2:DescribeNetworkInterfaces",
      "ec2:DescribeRouteTables",
      "ec2:DescribeSecurityGroups",
      "ec2:DescribeSubnets",
      "ec2:DescribeVpcEndpoints",
      "ec2:DescribeVpcs",
      "ecr:BatchCheckLayerAvailability",
      "ecr:BatchGetImage",
      "ecr:GetDownloadUrlForLayer",
      "ecr:GetAuthorizationToken",
      "s3:ListBucket",
      "s3:GetBucketLocation",
      "s3:GetEncryptionConfiguration",
      "s3:PutObject",
      "s3:DeleteObject",
      "s3:GetObject",
      "sagemaker:DescribeApp",
      "sagemaker:DescribeDomain",
      "sagemaker:DescribeModel`
    ]
  },
Permissions needed to access Amazon S3 resources through a S3 VPC endpoint

If you run SageMaker Studio in private VPC mode and access S3 through the S3 VPC endpoint, you can add permissions to the VPC endpoint policy to control which S3 resources are accessible through the VPC endpoint. Add the following permissions to your VPC endpoint policy. You can modify the policy if you need to further restrict permissions—for example, you can provide a more narrow specification for the Principal field.

```json
{
    "Sid": "S3DriverAccess",
    "Effect": "Allow",
    "Principal": "*",
    "Action": [
        "s3:GetBucketLocation",
        "s3:GetObject",
        "s3:ListBucket"
    ],
    "Resource": "arn:aws:s3:::sagemakerheadlessexecution-*"
}
```

For details about how to set up a S3 VPC endpoint policy, see [Edit the VPC endpoint policy](#).
Permissions needed to use a custom KMS key (optional)

By default, the input and output Amazon S3 buckets are encrypted using server side encryption, but you can specify a custom KMS key to encrypt your data in the output Amazon S3 bucket and the storage volume attached to the notebook job.

If you want to use a custom KMS key, attach the following policy and supply your own KMS key ARN.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "kms:Encrypt",
            "kms:Decrypt",
            "kms:ReEncrypt*",
            "kms:GenerateDataKey*",
            "kms:DescribeKey",
            "kms:CreateGrant"
         ],
         "Resource": "your_KMS_key_ARN"
      }
   ]
}
```

Install policies and permissions for local Jupyter environments

As previously stated, you install two sets of permissions—permissions for the IAM user and for the IAM role that the notebook job assumes. As shown in the following diagram, the IAM user needs to set up IAM permissions in order to submits jobs to SageMaker. Once the user submits the notebook job, the job itself assumes an IAM role that has permissions to access resources depending on the job tasks.
The following sections help you install necessary policies and permissions for both the IAM user and the job execution role.

**IAM user permissions**

**Permissions to submit jobs to SageMaker**

To add permissions to submit jobs, complete the following steps:

1. Open the [IAM console](#).
2. Select **Users** in the left panel.
3. Find the IAM user for your notebook job and choose the user name.
4. Choose **Add Permissions**, and choose **Create inline policy** from the dropdown menu.
5. Choose the **JSON** tab.
6. Copy and paste the following policy:
Amazon SageMaker

Developer Guide

{
"Version": "2012-10-17",
"Statement": [
{
"Sid": "EventBridgeSchedule",
"Effect": "Allow",
"Action": [
"events:TagResource",
"events:DeleteRule",
"events:PutTargets",
"events:DescribeRule",
"events:EnableRule",
"events:PutRule",
"events:RemoveTargets",
"events:DisableRule"
],
"Resource": "*",
"Condition": {
"StringEquals": {
"aws:ResourceTag/sagemaker:is-scheduling-notebook-job": "true"
}
}
},
{
"Sid": "IAMPassrole",
"Effect": "Allow",
"Action": "iam:PassRole",
"Resource": "arn:aws:iam::*:role/*",
"Condition": {
"StringLike": {
"iam:PassedToService": [
"sagemaker.amazonaws.com",
"events.amazonaws.com"
]
}
}
},
{
"Sid": "IAMListRoles",
"Effect": "Allow",
"Action": "iam:ListRoles",
"Resource": "*"
},

Notebook Jobs

4968


```json
{
    "Sid": "S3ArtifactsAccess",
    "Effect": "Allow",
    "Action": [
        "s3:PutEncryptionConfiguration",
        "s3:CreateBucket",
        "s3:PutBucketVersioning",
        "s3:ListBucket",
        "s3:PutObject",
        "s3:GetObject",
        "s3:GetEncryptionConfiguration",
        "s3:DeleteObject",
        "s3:GetBucketLocation"
    ],
    "Resource": [
        "arn:aws:s3:::sagemaker-automated-execution-*"
    ]
},
{
    "Sid": "S3DriverAccess",
    "Effect": "Allow",
    "Action": [
        "s3:ListBucket",
        "s3:GetObject",
        "s3:GetBucketLocation"
    ],
    "Resource": [
        "arn:aws:s3:::sagemakerheadlessexecution-*"
    ]
},
{
    "Sid": "SagemakerJobs",
    "Effect": "Allow",
    "Action": [
        "sagemaker:DescribeTrainingJob",
        "sagemaker:StopTrainingJob",
        "sagemaker:DescribePipeline",
        "sagemaker:CreateTrainingJob",
        "sagemaker:DeletePipeline",
        "sagemaker:CreatePipeline"
    ],
    "Resource": "*",
    "Condition": {
        "StringEquals": {
```
"aws:ResourceTag/sagemaker:is-scheduling-notebook-job": "true"

},

},

{
    "Sid": "AllowSearch",
    "Effect": "Allow",
    "Action": "sagemaker:Search",
    "Resource": "*"
},

{
    "Sid": "SagemakerTags",
    "Effect": "Allow",
    "Action": [
        "sagemaker:ListTags",
        "sagemaker:AddTags"
    ],
    "Resource": [
        "arn:aws:sagemaker:*:*:pipeline/*",
        "arn:aws:sagemaker:*:*:space/*",
        "arn:aws:sagemaker:*:*:training-job/*",
        "arn:aws:sagemaker:*:*:user-profile/*"
    ]
},

{
    "Sid": "ECRImage",
    "Effect": "Allow",
    "Action": [
        "ecr:GetAuthorizationToken",
        "ecr:BatchGetImage"
    ],
    "Resource": "*"
}
}

AWS KMS permission policy (optional)

By default, the input and output Amazon S3 buckets are encrypted using server side encryption, but you can specify a custom KMS key to encrypt your data in the output Amazon S3 bucket and the storage volume attached to the notebook job.
If you want to use a custom KMS key, repeat the previous instructions, attaching the following policy, and supply your own KMS key ARN.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "kms:Encrypt",
                "kms:Decrypt",
                "kms:ReEncrypt*",
                "kms:GenerateDataKey*",
                "kms:DescribeKey",
                "kms:CreateGrant"
                ],
            "Resource": "your_KMS_key_ARN"
        }
    ]
}
```

Job execution role permissions

Trust relationships

To modify the job execution role trust relationships, complete the following steps:

1. Open the IAM console.
2. Select Roles in the left panel.
3. Find the job execution role for your notebook job and choose the role name.
4. Choose the Trust relationships tab.
5. Choose Edit trust policy.
6. Copy and paste the following policy:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Service": [Notebook Jobs 4971]
Additional permissions

Once submitted, the notebook job needs permissions to access resources. The following instructions show you how to add a minimal set of permissions. If needed, add more permissions based on your notebook job needs. To add permissions to your job execution role, complete the following steps:

1. Open the IAM console.
2. Select Roles in the left panel.
3. Find the job execution role for your notebook job and choose the role name.
4. Choose Add Permissions, and choose Create inline policy from the dropdown menu.
5. Choose the JSON tab.
6. Copy and paste the following policy:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "PassroleForJobCreation",
         "Effect": "Allow",
         "Action": "iam:PassRole",
         "Resource": "arn:aws:iam::*:role/*",
         "Condition": {
            "StringLike": {
               "iam:PassedToService": "sagemaker.amazonaws.com"
            }
         }
      },
      {
         "Sid": "S3ForStoringArtifacts",
         "Effect": "Allow",
         "Action": "s3:*",
         "Resource": "arn:aws:s3:::bucketname/*"
      }
   ]
}
```
"Action": [
  "s3:PutObject",
  "s3:GetObject",
  "s3:ListBucket",
  "s3:GetBucketLocation"
],
"Resource": "arn:aws:s3:::sagemaker-automated-execution-*"
},
{
  "Sid": "S3DriverAccess",
  "Effect": "Allow",
  "Action": [
    "s3:ListBucket",
    "s3:GetObject",
    "s3:GetBucketLocation"
  ],
  "Resource": [
    "arn:aws:s3:::sagemakerheadlessexecution-*"
  ]
},
{
  "Sid": "SagemakerJobs",
  "Effect": "Allow",
  "Action": [
    "sagemaker:StartPipelineExecution",
    "sagemaker:CreateTrainingJob"
  ],
  "Resource": "*"
},
{
  "Sid": "ECRImage",
  "Effect": "Allow",
  "Action": [
    "ecr:GetDownloadUrlForLayer",
    "ecr:BatchGetImage",
    "ecr:GetAuthorizationToken",
    "ecr:BatchCheckLayerAvailability"
  ],
  "Resource": "*"
}
]

7. Add permissions to other resources your notebook job accesses.
8. Choose **Review policy**.

9. Enter a name for your policy.

10. Choose **Create policy**.

### Create a notebook job

If you want to create a notebook job, you have multiple options. You can create a job in your JupyterLab notebook in the Studio UI, or you can programmatically create a job with the SageMaker Python SDK.

If you create your notebook job in the Studio UI, you supply details about the image and kernel, security configurations, and any custom variables or scripts, and your job is scheduled. For details about how to schedule your job using SageMaker Notebook Jobs, see [Create a notebook job in Studio](#).

To create a notebook job with the SageMaker Python SDK, you create a pipeline with a Notebook Job step and initiate an on-demand run or optionally use the pipeline scheduling feature to schedule future runs. The SageMaker SDK gives you the flexibility to customize your pipeline—you can expand your pipeline to a workflow with multiple notebook job steps. Since you create both a SageMaker Notebook Job step and a pipeline, you can track your pipeline execution status in the SageMaker Notebook Jobs job dashboard and also view your pipeline graph in Studio. For details about how to schedule your job with the SageMaker Python SDK and links to example notebooks, see [Create a notebook job with SageMaker Python SDK](#).

### Create a notebook job with SageMaker Python SDK

To run a standalone notebook using the SageMaker Python SDK, you need to create a Notebook Job step, attach it into a pipeline, and use the utilities provided by SageMaker Pipelines to run your job on demand or optionally schedule one or more future jobs.

The following sections describe the basic steps to create an on-demand or scheduled notebook job and track the run. In addition, refer to the following discussion if you need to pass parameters to your notebook job or connect to Amazon EMR in your notebook—additional preparation of your Jupyter notebook is required in these cases. You can also apply defaults for a subset of the arguments of `NotebookJobStep` so you don’t have to specify them every time you create a Notebook Job step.

To view sample notebooks that demonstrate how to schedule notebook jobs with the SageMaker Python SDK, see [notebook job sample notebooks](#).
Topics

- Steps to create a notebook job
- View your notebook jobs in the Studio UI dashboard
- View your pipeline graph in Studio
- Passing parameters to your notebook
- Connecting to an Amazon EMR cluster in your input notebook
- Set up default options

Steps to create a notebook job

You can either create a notebook job that runs immediately or on a schedule. The following instructions describe both methods.

To schedule a notebook job, complete the following basic steps:

1. Create a NotebookJobStep instance. For details about NotebookJobStep parameters, see `sagemaker.workflow.steps.NotebookJobStep`. At minimum, you can provide the following arguments as shown in the following code snippet:

   ```python
   Important
   If you schedule your notebook job using the SageMaker Python SDK, you can only specify certain images to run your notebook job. For more information, see Image constraints for SageMaker Python SDK notebook jobs.
   ```

   ```python
   notebook_job_step = NotebookJobStep(
       input_notebook='input-notebook',
       image_uri='image-uri',
       kernel_name='kernel-name'
   )
   ```

2. Create a pipeline with your NotebookJobStep as a single step, as shown in the following snippet:

   ```python
   pipeline = Pipeline(
       name='pipeline-name',
   )
   ```
3. Run the pipeline on demand or optionally schedule future pipeline runs. To initiate an immediate run, use the following command:

```python
evaluation = pipeline.start(
    parameters={...}
)
```

Optionally, you can schedule a single future pipeline run or multiple runs at a predetermined interval. You specify your schedule in `PipelineSchedule` and then pass the schedule object to your pipeline with `put_triggers`. For more information about pipeline scheduling, see [Schedule a pipeline with the SageMaker Python SDK](#).

The following example schedules your pipeline to run once at December 12, 2023 at 10:31:32 UTC.

```python
my_schedule = PipelineSchedule(
    name="my-schedule",
    at=datetime(year=2023, month=12, date=25, hour=10, minute=31, second=32)
)
pipeline.put_triggers(triggers=[my_schedule])
```

The following example schedules your pipeline to run at 10:15am UTC on the last Friday of each month during the years 2022 to 2023. For details about cron-based scheduling, see [Cron-based schedules](#).

```python
my_schedule = PipelineSchedule(
    name="my-schedule",
    cron="15 10 ? * 6L 2022-2023"
)
pipeline.put_triggers(triggers=[my_schedule])
```

4. (Optional) View your notebook jobs in the SageMaker Notebook Jobs dashboard. The values you supply for the `tags` argument of your Notebook Job step control how the Studio UI captures and displays the job. For more information, see [View your notebook jobs in the Studio UI dashboard](#).
View your notebook jobs in the Studio UI dashboard

The notebook jobs you create as pipeline steps appear in the Studio Notebook Job dashboard if you specify certain tags.

Note

Only notebook jobs created in Studio or local JupyterLab environments create job definitions. Therefore, if you create your notebook job with the SageMaker Python SDK, you don’t see job definitions in the Notebook Jobs dashboard. You can, however, view your notebook jobs as described in View notebook jobs.

You can control which team members can view your notebook jobs with the following tags:

- To display the notebook to all user profiles or spaces in a domain, add the domain tag with your domain name. An example is shown as follows:
  - key: sagemaker:domain-name, value: d-abcdefghij5k

- To display the notebook job to a certain user profile in a domain, add both the user profile and the domain tags. An example of a user profile tag is shown as follows:
  - key: sagemaker:user-profile-name, value: studio-user

- To display the notebook job to a space, add both the space and the domain tags. An example of a space tag is shown as follows:
  - key: sagemaker:shared-space-name, value: my-space-name

- If you do not attach any domain or user profile or space tags, then the Studio UI does not show the notebook job created by pipeline step. In this case, you can view the underlying training job in the training job console or you can view the status in the list of pipeline executions.

Once you set up the necessary tags to view your jobs in the dashboard, see View notebook jobs for instructions about how to view your jobs and download outputs.

View your pipeline graph in Studio

Since your notebook job step is part of a pipeline, you can view the pipeline graph (DAG) in Studio. In the pipeline graph, you can view the status of the pipeline run and track lineage. For details, see View a Pipeline Execution.
Passing parameters to your notebook

If you want to pass parameters to your notebook job (using the parameters argument of NotebookJobStep), you need to prepare your input notebook to receive the parameters.

The Papermill-based notebook job executor searches for a Jupyter cell tagged with the parameters tag and applies the new parameters or parameter overrides immediately after this cell. For details, see Parameterize your notebook.

Once you have performed this step, pass your parameters to your NotebookJobStep, as shown in the following example:

```python
notebook_job_parameters = {
    "company": "Amazon"
}

notebook_job_step = NotebookJobStep(
    image_uri=image-uri,
    kernel_name=kernel-name,
    role=role-name,
    input_notebook=input-notebook,
    parameters=notebook_job_parameters,
    ...
)
```

Connecting to an Amazon EMR cluster in your input notebook

If you connect to an Amazon EMR cluster from your Jupyter notebook in Studio, you might need to further modify your Jupyter notebook. See Connect to an Amazon EMR cluster from your notebook if you need to perform any of the following tasks in your notebook:

- **Pass parameters into your Amazon EMR connection command.** Studio uses Papermill to run notebooks. In SparkMagic kernels, parameters you pass to your Amazon EMR connection command may not work as expected due to how Papermill passes information to SparkMagic.

- **Passing user credentials to Kerberos, LDAP, or HTTP Basic Auth-authenticated Amazon EMR clusters.** You have to pass user credentials through the AWS Secrets Manager.

Set up default options

The SageMaker SDK gives you the option to set defaults for a subset of parameters so you don’t have to specify these parameters every time you create a NotebookJobStep instance.
These parameters are `role`, `s3_root_uri`, `s3_kms_key`, `volume_kms_key`, `subnets`, and `security_group_ids`. Use the SageMaker config file to set the defaults for the step. For information about the SageMaker configuration file, see [Configuring and using defaults with the SageMaker Python SDK](#).

To set up the notebook job defaults, apply your new defaults to the notebook job section of the config file as shown in the following snippet:

```yaml
SageMaker:
  PythonSDK:
    Modules:
      NotebookJob:
        RoleArn: 'arn:aws:iam::555555555555:role/IMRole'
        S3RootUri: 's3://my-bucket/my-project'
        S3KmsKeyId: 's3kmskeyid'
        VolumeKmsKeyId: 'volumekmskeyid1'
        VpcConfig:
          SecurityGroupIds:
            - 'sg123'
          Subnets:
            - 'subnet-1234'
```

Create a notebook job in Studio

**Note**

The notebook scheduler is built from the Amazon EventBridge, SageMaker Training, and SageMaker Pipelines services. If your notebook jobs fail, you might see errors related to these services.

SageMaker Notebook Jobs gives you the tools to create and manage your noninteractive notebook jobs using the Notebook Jobs widget. You can create jobs, view the jobs you created, and pause, stop, or resume existing jobs. You can also modify notebook schedules.

When you create your scheduled notebook job with the widget, the scheduler tries to infer a selection of default options and automatically populates the form to help you get started quickly. If you are using Studio, at minimum you can submit an on-demand job without setting any options. You can also submit a (scheduled) notebook job definition supplying just the time-specific schedule information. However, you can customize other fields if your scheduled job requires specialized
settings. If you are running a local Jupyter notebook, the scheduler extension provides a feature for you to specify your own defaults (for a subset of options) so you don't have to manually insert the same values every time.

To schedule a notebook job, complete the following steps.

1. Open the **Create Job** form.

   In local JupyterLab environments, choose the **Create a notebook job** icon in the taskbar. If you don't see the icon, follow the instructions in **Installation Guide** to install it.

   In Studio, open the form in one of two ways:
   - **Using the File Browser**
     1. In the **File Browser** in the left panel, right-click on the notebook you want to run as a scheduled job.
     2. Choose **Create Notebook Job**.
   - **Within the Studio notebook**
     - Inside the Studio notebook you want to run as a scheduled job, choose the **Create a notebook job** icon in the Studio toolbar.

2. Complete the popup form. The form displays the following fields:
   - **Job name**: A descriptive name you specify for your job.
   - **Input file**: The name of the notebook which you are scheduling to run in noninteractive mode.
   - **Compute type**: The type of Amazon EC2 instance in which you want to run your notebook.
   - **Parameters**: Custom parameters you can optionally specify as inputs to your notebook. To use this feature, you might optionally want to tag a specific cell in your Jupyter notebook with the **parameters** tag to control where your parameters are applied. For more details, see **Parameterize your notebook**.
   - **Additional Options**: You can specify additional customizations for your job. For example, you can specify an image or kernel, input and output folders, job retry and timeout options, encryption details, and custom initialization scripts. For the complete listing of customizations you can apply, see **Available options**.

3. Schedule your job. You can run your notebook on demand or on a fixed schedule.
To run the notebook on demand, complete the following steps:

- Select **Run Now**.
- Choose **Create**.
- The **Notebook Jobs** tab appears. Choose **Reload** to load your job into the dashboard.

To run the notebook on a fixed schedule, complete the following steps:

- Choose **Run on a schedule**.
- Choose the **Interval** dropdown list and select an interval. The intervals range from every minute to monthly. You can also select **Custom schedule**.
- Based on the interval you choose, additional fields appear to help you further specify your desired run day and time. For example, if you select **Day** for a daily run, an additional field appears for you to specify the desired time. Note that any time you specify is in UTC format. Note also that if you choose a small interval, such as one minute, your jobs overlap if the previous job is not complete when the next job starts.

If you select a custom schedule, you use cron syntax in the expression box to specify your exact run date and time. The cron syntax is a space-separated list of digits, each of which represent a unit of time from seconds to years. For help with cron syntax, you can choose **Get help with cron syntax** under the expression box.

- Choose **Create**.
- The **Notebook Job Definitions** tab appears. Choose **Reload** to load your job definition into the dashboard.

### Set up default options for local notebooks

⚠️ **Important**

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see [Amazon SageMaker Studio](mailto:SageMaker Studio Link).

If you have to manually type (or paste in) custom values in the **Create Job** form, you can store new default values and the scheduler extension inserts your new values every time you create a new job definition. This feature is available for the following options:
• Role ARN
• S3 Input Folder
• S3 Output Folder
• Output encryption KMS key (if you turn on Configure Job Encryption)
• Job instance volume encryption KMS key (if you turn on Configure Job Encryption)

This feature saves you time if you insert different values than the provided defaults and continue to use those values for future job runs. Your chosen user settings are stored on the machine that runs your JupyterLab server and are retrieved with the help of native API. If you provide new default values for one or more but not all five options, the previous defaults are taken for the ones you don’t customize.

The following instructions show you how to preview the existing default values, set new default values, and reset your default values for your notebook jobs.

To preview existing default values for your notebook jobs, complete the following steps:

1. Open the Amazon SageMaker Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.
2. In the File Browser in the left panel, right-click on the notebook you want to run as a scheduled job.
3. Choose Create Notebook Job.
4. Choose Additional options to expand the tab of notebook job settings. You can view the default settings here.

To set new default values for your future notebook jobs, complete the following steps:

1. Open the Amazon SageMaker Studio Classic console by following the instructions in Launch Amazon SageMaker Studio Classic.
2. From the top menu in Studio Classic, choose Settings, then choose Advanced Settings Editor.
3. Choose Amazon SageMaker Scheduler from the list below Settings. This may already be open by default.
4. You can update the default settings directly in this UI page or by using the JSON editor.

   • In the UI you can insert new values for Role ARN, S3 Input Folder, S3 Output Folder, Output encryption KMS key, or Job instance volume encryption KMS key. If you change
these values, you will see the new defaults for these fields while you create your next notebook job under Additional options.

- (Optional) To update the user defaults using the JSON Settings Editor, complete the following steps:

  1. In the top right corner, choose JSON Settings Editor.
  2. In the Settings left sidebar, choose Amazon SageMaker Scheduler. This may already be open by default.

     You can see your current default values in the User Preferences panel.

     You can see the system default values in the System Defaults panel.

  3. To update your default values, copy and paste the JSON snippet from the System Defaults panel to the User Preferences panel, and update the fields.

  4. If you updated the default values, choose the Save User Settings icon in the top right corner. Closing the editor does not save the changes.

If you previously changed and now want to reset the user-defined default values, complete following steps:

  1. From the top menu in Studio Classic, choose Settings, then choose Advanced Settings Editor.
  2. Choose Amazon SageMaker Scheduler from the list below Settings. This may already be open by default.
  3. You can restore the defaults by directly using this UI page or using the JSON editor.

     - In the UI you can choose Restore to Defaults in the top right corner. Your defaults are restored to empty strings. You only see this option if you previously changed your default values.

     - (Optional) To restart the default settings using the JSON Settings Editor, complete the following steps:

        1. In the top right corner, choose JSON Settings Editor.
        2. In the Settings left sidebar, choose Amazon SageMaker Scheduler. This may already be open by default.

           You can see your current default values in the User Preferences panel.
You can see the system default values in the **System Defaults** panel.

3. To restore your current default settings copy the content from the **System Defaults** panel to the **User Preferences** panel.

4. Choose the **Save User Settings** icon in the top right corner. Closing the editor does not save the changes.

### Create a workflow of notebook jobs

Since a notebook job runs your custom code, you can create a pipeline that includes one or more notebook job steps. ML workflows often contain multiple steps, such as a processing step to preprocess data, a training step to build your model, and a model evaluation step, among others. One possible use of notebook jobs is to handle preprocessing—you might have a notebook that performs data transformation or ingestion, an EMR step that performs data cleaning, and another notebook job that performs featurization of your inputs before initiating a training step. A notebook job may require information from previous steps in the pipeline or from user-specified customization as parameters in the input notebook. For examples that show how to pass environment variables and parameters to your notebook and retrieve information from prior steps, see [Pass information to and from your notebook step](#).

In another use case, one of your notebook jobs might call another notebook to perform some tasks during your notebook run—in this scenario you need to specify these sourced notebooks as dependencies with your notebook job step. For information about how to call another notebook, see [Invoke another notebook in your notebook job](#).

To view sample notebooks that demonstrate how to schedule notebook jobs with the SageMaker Python SDK, see [notebook job sample notebooks](#).

### Pass information to and from your notebook step

The following sections describe ways to pass information to your notebook as environment variables and parameters.

#### Pass environment variables

Pass environment variables as a dictionary to the `environment_variable` argument of your `NotebookJobStep`, as shown in the following example:
environment_variables = {"RATE": 0.0001, "BATCH_SIZE": 1000}

notebook_job_step = NotebookJobStep(
    ...
    environment_variables=environment_variables,
    ...
)

You can use the environment variables in the notebook using `os.getenv()`, as shown in the following example:

```python
# inside your notebook
import os
print(f"ParentNotebook: env_key={os.getenv('env_key')}")
```

**Pass parameters**

When you pass parameters to the first Notebook Job step in your `NotebookJobStep` instance, you might optionally want to tag a cell in your Jupyter notebook to indicate where to apply new parameters or parameter overrides. For instructions about how to tag a cell in your Jupyter notebook, see [Parameterize your notebook](#).

You pass parameters through the Notebook Job step's `parameters` parameter, as shown in the following snippet:

```python
notebook_job_parameters = {
    "company": "Amazon",
}

notebook_job_step = NotebookJobStep(
    ...
    parameters=notebook_job_parameters,
    ...
)
```

Inside your input notebook, your parameters are applied after the cell tagged with `parameters` or at the beginning of the notebook if you don't have a tagged cell.

```python
# this cell is in your input notebook and is tagged with 'parameters'
# your parameters and parameter overrides are applied after this cell
```
company='default'

# in this cell, your parameters are applied
# prints "company is Amazon"
print(f'company is {company}')

Retrieve information from a previous step

The following discussion explains how you can extract data from a previous step to pass to your Notebook Job step.

Use properties attribute

You can use the following properties with the previous step's properties attribute:

- **ComputingJobName**—The training job name
- **ComputingJobStatus**—The training job status
- **NotebookJobInputLocation**—The input Amazon S3 location
- **NotebookJobOutputLocationPrefix**—The path to your training job outputs, more specifically \{NotebookJobOutputLocationPrefix\}/\{training-job-name\}/output/\{output.tar.gz\} containing outputs
- **InputNotebookName**—The input notebook file name
- **OutputNotebookName**—The output notebook file name (which may not exist in the training job output folder if the job fails)

The following code snippet shows how to extract parameters from the properties attribute.

```python
notebook_job_step2 = NotebookJobStep(
    ....
    parameters={
        "step1_JobName": notebook_job_step1.properties.ComputingJobName,
        "step1_JobStatus": notebook_job_step1.properties.ComputingJobStatus,
        "step1_NotebookJobInput":
            notebook_job_step1.properties.NotebookJobInputLocation,
        "step1_NotebookJobOutput":
            notebook_job_step1.properties.NotebookJobOutputLocationPrefix,
    }
)
```

Use JsonGet
If you want to pass parameters other than the ones previously mentioned and the JSON outputs of your previous step reside in Amazon S3, use `JsonGet`. `JsonGet` is a general mechanism that can directly extract data from JSON files in Amazon S3.

To extract JSON files in Amazon S3 with `JsonGet`, complete the following steps:

1. Upload your JSON file to Amazon S3. If your data is already uploaded to Amazon S3, skip this step. The following example demonstrates uploading a JSON file to Amazon S3.

```python
import json
from sagemaker.s3 import S3Uploader

output = {
    "key1": "value1",
    "key2": [0,5,10]
}

json_output = json.dumps(output)

with open("notebook_job_params.json", "w") as file:
    file.write(json_output)

S3Uploader.upload(
    local_path="notebook_job_params.json",
    desired_s3_uri="s3://path/to/bucket"
)
```

2. Provide your S3 URI and the JSON path to the value you want to extract. In the following example, `JsonGet` returns an object representing index 2 of the value associated with key `key2` (10).

```python
NotebookJobStep(
    ....
    parameters={
        # the key job_key1 returns an object representing the value 10
        "job_key1": JsonGet(
            s3_uri=Join(on="/", values=["s3:/", ..]),
            json_path="key2[2]" # value to reference in that json file
        ),
        "job_key2": "Amazon"
    }
)
```
Invoke another notebook in your notebook job

The following discussion sets up an example of a pipeline with a Notebook Job step in which the notebook calls two other notebooks. The input notebook contains the following lines:

```python
%run 'subfolder/notebook_to_call_in_subfolder.ipynb'
%run 'notebook_to_call.ipynb'
```

Pass these notebooks into your NotebookJobStep instances with `additional_dependencies`, as shown in the following snippet. Note that the paths provided for the notebooks in `additional_dependencies` are provided from the root location. For information about how SageMaker uploads your dependent files and folders to Amazon S3 so you can correctly provide paths to your dependencies, see the description for `additional_dependencies` in `NotebookJobStep`.

```python
input_notebook = "inputs/input_notebook.ipynb"
simple_notebook_path = "inputs/notebook_to_call.ipynb"
folder_with_sub_notebook = "inputs/subfolder"

notebook_job_step = NotebookJobStep(
    image_uri=image-uri,
    kernel_name=kernel-name,
    role=role-name,
    input_notebook=input_notebook,
    additional_dependencies=[simple_notebook_path, folder_with_sub_notebook],
    tags=tags,
)
```

Available options

The following table displays all available options you can use to customize your notebook job, whether you run your Notebook Job in Studio, a local Jupyter environment, or using the SageMaker Python SDK. The table includes the type of custom option, a description, additional guidelines about how to use the option, a field name for the option in Studio (if available) and the parameter name for the notebook job step in the SageMaker Python SDK (if available).

For some options, you can also preset custom default values so you don’t have to specify them every time you set up a notebook job. For Studio, these options are **Role**, **Input folder**, **Output folder**, and **KMS Key ID**, and are specified in the following table. If you preset custom defaults for these options, these fields are prepopulated in the **Create Job** form when you create your
notebook job. For details about how to create custom defaults in Studio and local Jupyter environments, see [Set up default options for local notebooks](#).

The SageMaker SDK also gives you the option to set intelligent defaults so that you don't have to specify these parameters when you create a NotebookJobStep. These parameters are `role`, `s3_root_uri`, `s3_kms_key`, `volume_kms_key`, `subnets`, `security_group_ids`, and are specified in the following table. For information about how to set intelligent defaults, see [Set up default options](#).

<table>
<thead>
<tr>
<th>Custom option</th>
<th>Description</th>
<th>Studio-specific guideline</th>
<th>Local Jupyter environment guideline</th>
<th>SageMaker Python SDK guideline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Job name</td>
<td>Your job name as it should appear in the Notebook Jobs dashboard.</td>
<td>Field <strong>Job name</strong>.</td>
<td>Same as Studio.</td>
<td>Parameter <code>notebook_job_name</code>. Defaults to None.</td>
</tr>
<tr>
<td>Image</td>
<td>The container image used to run the notebook noninteractively on the chosen compute type.</td>
<td>Field <strong>Image</strong>. This field defaults to your notebook's current image. Change this field from the default to a custom value if needed. If Studio cannot infer this value, the form displays a validation error requiring you to specify it. This image can be a custom, <a href="#">bring-your-own image</a> or an available Amazon SageMaker image. For a list of available SageMaker images supported by</td>
<td>Field <strong>Image</strong>. This field requires an ECR URI of a Docker image that can run the provided notebook on the selected compute type. By default, the scheduler extension uses a pre-built SageMaker Docker image—base Python 2.0. This is the official Python 3.8 image from DockerHub with boto3, AWS CLI, and the Python 3 kernel. You can also provide any ECR URI that meets the</td>
<td><strong>Required.</strong> Parameter <code>image_uri</code>. URI location of a Docker image on ECR. You can use specific SageMaker</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>the notebook scheduler, see <a href="#">Available Amazon SageMaker Images</a>.</td>
<td>notebook custom image specification. For details, see <a href="#">Custom SageMaker image specifications</a>. This image should have all the kernels and libraries needed for the notebook run.</td>
<td>Distribution Images or custom image based on those images, or your own image pre-installed with notebook job dependencies that meets additional requirements. For details, see Image Notebook Jobs.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>Instance type</td>
<td>The EC2 instance type to use to run the notebook job. The notebook job uses a SageMaker Training Job as a computing layer, so the specified instance type should be a SageMaker Training supported instance type.</td>
<td>Field <strong>Compute type</strong>. Defaults to <code>ml.m5.large</code>.</td>
<td>Same as Studio.</td>
<td>Parameter <code>instance_type</code>. Defaults to <code>ml.m5.large</code>.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>Kernel</td>
<td>The Jupyter kernel used to run the notebook job.</td>
<td>Field <strong>Kernel</strong>. This field defaults to your notebook’s current kernel. Change this field from the default to a custom value if needed. If Studio cannot infer this value, the form displays a validation error requiring you to specify it.</td>
<td>Field <strong>Kernel</strong>. This kernel should be present in the image and follow the Jupyter kernel specs. This field defaults to the Python3 kernel found in the base Python 2.0 SageMaker image. Change this field to a custom value if needed.</td>
<td>Required. Parameter <strong>kernel_name</strong>. This kernel should be present in the image and follow the Jupyter kernel specs. To see the kernel identifiers for your image, see <a href="#">LINK</a>.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>SageMaker session</td>
<td>The underlyin g SageMaker session to which SageMaker service calls are delegated.</td>
<td>N/A</td>
<td>N/A</td>
<td>Parameter sagemaker_session. If unspecified, one is created using a default configuration chain.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>Role ARN</td>
<td>The role’s Amazon Resource Name (ARN) used with the notebook job.</td>
<td>Field <strong>Role ARN</strong>. This field defaults to the Studio execution role. Change this field to a custom value if needed.</td>
<td>Field <strong>Role ARN</strong>. This field defaults to any role prefixed with Sagemaker JupyterScheduler. If you have multiple roles with the prefix, the extension chooses one. Change this field to a custom value if needed. For this field, you can set your own user default that pre-populates whenever you create a new job definition. For details, see Set up default options for local notebooks.</td>
<td>Parameter role. Defaults to the SageMaker default IAM role if the SDK is running in SageMaker Notebooks or SageMaker Studio Notebooks. Otherwise, it throws a ValueError. Allows intelligent Notebook Jobs</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline defaults</td>
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<td>---------------</td>
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</tr>
<tr>
<td><strong>Input notebook</strong></td>
<td>The name of the notebook which you are scheduling to run.</td>
<td><strong>Required.</strong> Field <strong>Input file.</strong></td>
<td>Same as Studio.</td>
<td><strong>Required.</strong> Parameter input_notebook.</td>
</tr>
<tr>
<td><strong>Input folder</strong></td>
<td>The folder containing your inputs. The job inputs, including the input notebook and any optional start-up or initialization scripts, are put in this folder.</td>
<td>Field <strong>Input folder.</strong> If you don’t provide a folder, the scheduler creates a default Amazon S3 bucket for your inputs.</td>
<td>Same as Studio. For this field, you can set your own user default that pre-populates whenever you create a new job definition. For details, see <a href="#">Set up default options for local notebooks</a>.</td>
<td>N/A. The input folder is placed inside the location specified by parameter s3_root_uri.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
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</tr>
<tr>
<td>Output folder</td>
<td>The folder containing your outputs. The job outputs, including the output notebook and logs, are put in this folder.</td>
<td>Field <strong>Output folder</strong>. If you don’t specify a folder, the scheduler creates a default Amazon S3 bucket for your outputs.</td>
<td>Same as Studio. For this field, you can set your own user default that pre-populates whenever you create a new job definition. For details, see <a href="#">Set up default options for local notebooks</a>.</td>
<td>N/A. The output folder is placed inside the location specified by parameter <code>s3_root_uri</code>.</td>
</tr>
<tr>
<td>Parameters</td>
<td>A dictionary of variables and values to pass to your notebook job.</td>
<td>Field <strong>Parameters</strong>. You need to <a href="#">parameterize your notebook</a> to accept parameters.</td>
<td>Same as Studio.</td>
<td>Parameter parameters. You need to <a href="#">parameterize your notebook</a> to accept parameters.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>Additiona l (file or folder) dependencies</td>
<td>The list of file or folder dependencies which the notebook job uploads to S3 staged folder.</td>
<td>Not supported.</td>
<td>Not supported.</td>
<td>Parameter additiona l_dependencies. The notebook job uploads these dependencies to an S3 staged folder so they can be consumed during execution.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
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</tr>
<tr>
<td>S3 root URI</td>
<td>The folder containing your inputs. The job inputs, including the input notebook and any optional start-up or initialization scripts, are put in this folder.</td>
<td>N/A. Use <strong>Input Folder</strong> and <strong>Output folder</strong>.</td>
<td>Same as Studio.</td>
<td>Parameter <code>s3_root_uri</code>. Defaults to a default S3 bucket. Allows intelligent defaults.</td>
</tr>
<tr>
<td>Environment variables</td>
<td>Any existing environment variables that you want to override, or new environment variables that you want to introduce and use in your notebook.</td>
<td>Field <strong>Environment variables</strong>.</td>
<td>Same as Studio.</td>
<td>Parameter <code>environment_variables</code>. Defaults to None.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>Tags</td>
<td>A list of tags attached to the job.</td>
<td>N/A</td>
<td>N/A</td>
<td>Parameter tags. Defaults to None. Your tags control how the Studio UI captures and displays the job created by the pipeline. For details, see <a href="#">View your notebook jobs in the Studio</a>.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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</tr>
<tr>
<td>Start-up script</td>
<td>A script preloaded in the notebook startup menu that you can choose to run before you run the notebook.</td>
<td>Field <strong>Start-up script.</strong> Select a Lifecycle Configuration (LCC) script that runs on the image at start-up.</td>
<td>Not supported.</td>
<td>Not supported.</td>
</tr>
</tbody>
</table>

**Note**

A start-up script runs in a shell outside of the Studio environment. Therefore, this script cannot depend on the Studio local storage, environment variables, or app metadata (in `/opt/ml/metadata`). Also, if you use a start-up script and an initialization script, the start-up script runs first.
<table>
<thead>
<tr>
<th>Custom option</th>
<th>Description</th>
<th>Studio-specific guideline</th>
<th>Local Jupyter environment guideline</th>
<th>SageMaker Python SDK guideline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initialization script</td>
<td>A path to a local script you can run when your notebook starts up.</td>
<td>Field <strong>Initialization script</strong>. Enter the EFS file path where a local script or a Lifecycle Configuration (LCC) script is located. If you use a start-up script and an initialization script, the start-up script runs first.</td>
<td>Field <strong>Initialization script</strong>. Enter the local file path where a local script or a Lifecycle Configuration (LCC) script is located.</td>
<td>Parameter initialization_script. Defaults to None.</td>
</tr>
</tbody>
</table>

**Note**

An initialization script is sourced from the same shell as the notebook job. This is not the case for a start-up script described previously. Also, if you use a start-up script and an initialization script, the start-up script runs first.
<table>
<thead>
<tr>
<th>Custom option</th>
<th>Description</th>
<th>Studio-specific guideline</th>
<th>Local Jupyter environment guideline</th>
<th>SageMaker Python SDK guideline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max retry attempts</td>
<td>The number of times Studio tries to rerun a failed job run.</td>
<td>Field <strong>Max retry attempts</strong>.</td>
<td>Same as Studio.</td>
<td>Parameter <code>max_retry_attempts</code>. Defaults to 1.</td>
</tr>
<tr>
<td>Max run time (in seconds)</td>
<td>The maximum length of time, in seconds, that a notebook job can run before it is stopped. If you configure both <strong>Max run time</strong> and <strong>Max retry attempts</strong>, the run time applies to each retry. If a job does not complete in this time, its status is set to Failed.</td>
<td>Field <strong>Max run time (in seconds)</strong>. Defaults to 172800 seconds (2 days).</td>
<td>Same as Studio.</td>
<td>Parameter <code>max_run_time_in_seconds</code>. Defaults to 172800 seconds (2 days).</td>
</tr>
<tr>
<td>Retry policies</td>
<td>A list of retry policies, which govern actions to take in case of failure.</td>
<td>Not supported.</td>
<td>Not supported.</td>
<td>Parameter <code>retry_policies</code>. Defaults to None.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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<td>---------------------------------</td>
</tr>
<tr>
<td>Add Step or StepCollection dependencies</td>
<td>A list of Step or StepCollection names or instances on which the job depends.</td>
<td>Not supported.</td>
<td>Not supported.</td>
<td>Parameter depends_on. Defaults to None. Use this to define explicit dependencies between steps in your pipeline graph.</td>
</tr>
<tr>
<td>Volume size</td>
<td>The size in GB of the storage volume for storing input and output data during training.</td>
<td>Not supported.</td>
<td>Not supported.</td>
<td>Parameter volume_size. Defaults to 30GB.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
<td>Studio-specific guideline</td>
<td>Local Jupyter environment guideline</td>
<td>SageMaker Python SDK guideline</td>
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<td>---------------------------------</td>
</tr>
<tr>
<td>Encrypt traffic between containers</td>
<td>A flag that specifies whether traffic between training containers is encrypted for the training job.</td>
<td>N/A. Enabled by default.</td>
<td>N/A. Enabled by default.</td>
<td>Parameter encrypt_inter_container_traffic. Defaults to True.</td>
</tr>
<tr>
<td>Configure job encryption</td>
<td>An indicator that you want to encrypt your notebook job outputs, job instance volume, or both.</td>
<td>Field <strong>Configure job encryption</strong>. Check this box to choose encryption. If left unchecked, the job outputs are encrypted with the account's default KMS key and the job instance volume is not encrypted.</td>
<td>Same as Studio.</td>
<td>Not supported.</td>
</tr>
<tr>
<td>Output encryption KMS key</td>
<td>A KMS key to use if you want to customize the encryption key used for your notebook job outputs. This field is only applicable if you checked <strong>Configure job encryption</strong>.</td>
<td>Field <strong>Output encryption KMS key</strong>. If you do not specify this field, your notebook job outputs are encrypted with SSE-KMS using the default Amazon S3 KMS key. Also, if you create the Amazon S3 bucket yourself and use encryption, your encryption method is preserved.</td>
<td>Same as Studio. For this field, you can set your own user default that pre-populates whenever you create a new job definition. For details, see <a href="#">Set up default options for local notebooks</a>.</td>
<td>Parameter s3_kms_key. Defaults to None. Allows intelligent defaults.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
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<td>------------------------------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>Job instance volume encryption KMS key</td>
<td>A KMS key to use if you want to encrypt your job instance volume. This field is only applicable if you checked Configure job encryption.</td>
<td>Field <strong>Job instance volume encryption KMS key</strong>. For this field, you can set your own user default that pre-populates whenever you create a new job definition. For details, see <a href="#">Set up default options for local notebooks</a>.</td>
<td>Field <strong>Job instance volume encryption KMS key</strong>. For this field, you can set your own user default that pre-populates whenever you create a new job definition. For details, see <a href="#">Set up default options for local notebooks</a>.</td>
<td>Parameter volume_kms_key. Defaults to None. Allows intelligent defaults.</td>
</tr>
<tr>
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</tr>
<tr>
<td>Use a Virtual Private Cloud to run this job (for VPC users)</td>
<td>An indicator that you want to run this job in a Virtual Private Cloud (VPC). For better security, it is recommend that you use a private VPC.</td>
<td>Field <strong>Use a Virtual Private Cloud to run this job</strong>. Check this box if you want to use a VPC. At minimum, create the following VPC endpoints to enable your notebook job to privately connect to those AWS resources:</td>
<td>Same as Studio.</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• <strong>SageMaker</strong>: For information on how to connect to SageMaker through a VPC interface endpoint, see <a href="#">Connect to SageMaker Within your VPC</a>.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• <strong>Amazon S3</strong>: For information on how to connect to Amazon S3 through a VPC interface endpoint, see <a href="#">Gateway endpoints for Amazon S3</a>.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• <strong>Amazon EC2</strong>: For information on how to connect to Amazon EC2 through a VPC interface endpoint, see <a href="#">Access Amazon EC2 using an interface VPC endpoint</a>.</td>
<td></td>
<td></td>
</tr>
<tr>
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</tr>
<tr>
<td></td>
<td></td>
<td>Amazon EventBridge: This endpoint is only needed when setting up a scheduled notebook. It is not needed when launching a job on demand. For information on how to connect to EventBridge through a VPC interface endpoint, see <a href="#">Using Amazon EventBridge with interface VPC Endpoints</a>.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

If you choose to use a VPC, you need to specify at least one private subnet and at least one security group in the following options. If you don’t use any private subnets, you need to consider other configuration options. For details, see [Public VPC subnets not supported](#) in Constraints and considerations.
<table>
<thead>
<tr>
<th>Custom option (for VPC users)</th>
<th>Description</th>
<th>Studio-specific guideline</th>
<th>Local Jupyter environment guideline</th>
<th>SageMaker Python SDK guideline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subnet(s)</td>
<td>Your subnets. This field must contain at least one and at most five, and all the subnets you provide should be private. For details, see Public VPC subnets not supported in Constraints and considerations.</td>
<td>Field <strong>Subnet(s)</strong>. This field defaults to the subnets associated with the Studio domain, but you can change this field if needed.</td>
<td>Field <strong>Subnet(s)</strong>. The scheduler cannot detect your subnets, so you need to enter any subnets you configured for your VPC.</td>
<td>Parameter subnets. Defaults to None. Allows intelligent defaults.</td>
</tr>
<tr>
<td>Security group(s) (for VPC users)</td>
<td>Your security groups. This field must contain at least one and at most 15. For details, see Public VPC subnets not supported in Constraints and considerations.</td>
<td>Field <strong>Security groups</strong>. This field defaults to the security groups associated with the domain VPC, but you can change this field if needed.</td>
<td>Field <strong>Security groups</strong>. The scheduler cannot detect your security groups, so you need to enter any security groups you configured for your VPC.</td>
<td>Parameter security_group_ids. Defaults to None. Allows intelligent defaults.</td>
</tr>
<tr>
<td>Custom option</td>
<td>Description</td>
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<td>-------------------------------</td>
</tr>
<tr>
<td>Name</td>
<td>The name of the notebook job step.</td>
<td>N/A</td>
<td>N/A</td>
<td>Parameter name. If unspecified, it is derived from the notebook file name.</td>
</tr>
<tr>
<td>Display name</td>
<td>Your job name as it should appear in your list of pipeline executions.</td>
<td>N/A</td>
<td>N/A</td>
<td>Parameter display_name. Defaults to None.</td>
</tr>
<tr>
<td>Description</td>
<td>A description of your job.</td>
<td>N/A</td>
<td>N/A</td>
<td>Parameter description.</td>
</tr>
</tbody>
</table>

**Parameterize your notebook**

To pass new parameters or parameter overrides to your scheduled notebook job, you might optionally want to modify your Jupyter notebook if you want your new parameters values to be applied after a cell. When you pass a parameter, the notebook job executor uses the methodology enforced by Papermill. The notebook job executor searches for a Jupyter cell tagged with the `parameters` tag and applies the new parameters or parameter overrides immediately after this cell. If you don’t have any cells tagged with `parameters`, the parameters are applied at
the beginning of the notebook. If you have more than one cell tagged with `parameters`, the parameters are applied after the first cell tagged with `parameters`.

To tag a cell in your notebook with the `parameters` tag, complete the following steps:

1. Select the cell to parameterize.
2. Choose the Property Inspector icon ( ) in the right sidebar.
3. Type `parameters` in the Add Tag box.
4. Choose the + sign.
5. The `parameters` tag appears under Cell Tags with a check mark, which means the tag is applied to the cell.

Connect to an Amazon EMR cluster from your notebook

If you connect to an Amazon EMR cluster from your Jupyter notebook in Studio, you might need to perform additional setup. In particular, the following discussion addresses two issues:

- **Passing parameters into your Amazon EMR connection command.** In SparkMagic kernels, parameters you pass to your Amazon EMR connection command may not work as expected due to differences in how Papermill passes parameters and how SparkMagic receives parameters. The workaround to address this limitation is to pass parameters as environment variables. For more details about the issue and workaround, see [Pass parameters to your EMR connection command](#).

- **Passing user credentials to Kerberos, LDAP, or HTTP Basic Auth-authenticated Amazon EMR clusters.** In interactive mode, Studio asks for credentials in a popup form where you can enter your sign-in credentials. In your noninteractive scheduled notebook, you have to pass them through the AWS Secrets Manager. For more details about how to use the AWS Secrets Manager in your scheduled notebook jobs, see [Pass user credentials to your Kerberos, LDAP, or HTTP Basic Auth-authenticated Amazon EMR cluster](#).

Pass parameters to your EMR connection command

If you are using images with the SparkMagic PySpark and Spark kernels and want to parameterize your EMR connection command, provide your parameters in the **Environment variables** field instead of the Parameters field in the Create Job form (in the **Additional Options** dropdown.
Make sure your EMR connection command in the Jupyter notebook passes these parameters as environment variables. For example, suppose you pass `cluster-id` as an environment variable when you create your job. Your EMR connection command should look like the following:

```python
%%local
import os

%s_m Analytics emr connect --cluster-id {os.getenv('cluster_id')} --auth-type None
```

You need this workaround to meet requirements by SparkMagic and Papermill. For background context, the SparkMagic kernel expects that the `%%local` magic command accompany any local variables you define. However, Papermill does not pass the `%%local` magic command with your overrides. In order to work around this Papermill limitation, you must supply your parameters as environment variables in the **Environment variables** field.

**Pass user credentials to your Kerberos, LDAP, or HTTP Basic Auth-authenticated Amazon EMR cluster**

To establish a secure connection to an Amazon EMR cluster that uses Kerberos, LDAP, or HTTP Basic Auth authentication, you use the AWS Secrets Manager to pass user credentials to your connection command. For information about how to create a Secrets Manager secret, see [Create an AWS Secrets Manager secret](#). Your secret must contain your username and password. You pass the secret with the `--secrets` argument, as shown in the following example:

```bash
%s_m Analytics emr connect --cluster-id j_abcde12345
--auth Kerberos
--secret aws_secret_id_123
```

Your administrator can set up a flexible access policy using an attribute-based-access-control (ABAC) method, which assigns access based on special tags. You can set up flexible access to create a single secret for all users in the account or a secret for each user. The following code samples demonstrate these scenarios:

**Create a single secret for all users in the account**

```json
{
    "Version" : "2012-10-17",
    "Statement" : [
```
Create a different secret for each user

You can create a different secret for each user using the `PrincipalTag` tag, as shown in the following example:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::AWS_ACCOUNT_ID:role/service-role/AmazonSageMaker-ExecutionRole-20190101T012345"},
            "Condition": {
                "StringEquals": {
                    "aws:ResourceTag/user-identity": "${aws:PrincipalTag/user-identity}"
                }
            },
            "Action": "secretsmanager:GetSecretValue",
            "Resource": [
                "arn:aws:secretsmanager:us-west-2:AWS_ACCOUNT_ID:secret:aes789-7g8h9i"
            ]
        }
    ]
}
```
Track notebook jobs and job definitions

SageMaker Notebook Jobs dashboards help organize the job definitions that you schedule, and also keep track of the actual jobs that run from your job definitions. There are two important concepts to understand when scheduling notebook jobs: job definitions and job runs. Job definitions are schedules you set to run specific notebooks. For example, you can create a job definition that runs notebook XYZ.ipynb every Wednesday. This job definition launches the actual job runs which occur this coming Wednesday, next Wednesday, the Wednesday after that, and so on.

**Note**

The SageMaker Python SDK notebook job step does not create job definitions. However, you can view your jobs in the Notebook Jobs dashboard. Both jobs and job definitions are available if you schedule your job in a JupyterLab environment.

The interface provides two main tabs that help you track your existing job definitions and job runs:

- **Notebook Jobs** tab: This tab displays a list of all your job runs from your on-demand jobs and job definitions. From this tab, you can directly access the details for a single job run. For example, you can view a single job run that occurred two Wednesdays ago.

- **Notebook Job Definitions** tab: This tab displays a list of all your job definitions. From this tab, you can directly access the details for a single job definition. For example, you can view the schedule you created to run XYZ.ipynb every Wednesday.

For details about the **Notebook Jobs** tab, see [View notebook jobs](#).

For details about the **Notebook Job Definitions** tab, see [View notebook job definitions](#).

View notebook jobs

**Note**

You can automatically view your notebook jobs if you scheduled your notebook job from the Studio UI. If you used the SageMaker Python SDK to schedule your notebook job, you
need to supply additional tags when you create the notebook job step. For details, see View your notebook jobs in the Studio UI dashboard.

The Notebook Jobs tab (which you access by choosing the Create a notebook job icon in the Studio toolbar) shows a history of your on-demand jobs and all the jobs that run from the job definitions you created. This tab opens after you create an on-demand job, or you can just view this tab yourself to see a history of past and current jobs. If you select the Job name for any job, you can view details for a single job in its Job Detail page. For more information about the Job Detail page, see the following section View a single job.

The Notebook Jobs tab includes the following information for each job:

- **Output files**: Displays the availability of output files. This column can contain one of the following:
  - A download icon: The output notebook and log are available for download; choose this button to download them. Note that a failed job can still generate output files if the failure occurred after the files were created. In this case, it is helpful to view the output notebook to identify the failure point.
  - Links to the Notebook and Output log: The notebook and output log are downloaded. Choose the links to view their contents.
  - (blank): The job was stopped by the user, or a failure occurred in the job run before it could generate output files. For example, network failures could prevent the job from starting.

  The output notebook is the result of running all cells in the notebook, and also incorporates any new or overriding parameters or environment variables you included. The output log captures the details of the job run to help you troubleshoot failed jobs.

- **Created at**: The time the on-demand job or scheduled job was created.

- **Status**: The current status of the job, which is one of the following values:
  - In progress: The job is running
  - Failed: The job failed from configuration or notebook logic errors
  - Stopped: The job was stopped by the user
  - Completed: The job completed
• **Actions**: This column provides shortcuts to help you stop or remove any job directly in the interface.

**View a single job**

From the **Notebook Jobs** tab, you can select a job name to view the **Job Detail** page for a specific job. The **Job Detail** page includes all the details you provided in the **Create Job** form. Use this page to confirm the settings you specified when you created the job definition.

In addition, you can access shortcuts to help you perform the following actions in the page itself:

• **Delete Job**: Remove the job from the **Notebook Jobs** tab.
• **Stop Job**: Stop your running job.

**View notebook job definitions**

**Note**

If you scheduled your notebook job with the SageMaker Python SDK, skip this section. Only notebook jobs created in Studio or local JupyterLab environments create job definitions. Therefore, if you created your notebook job with the SageMaker Python SDK, you won’t see job definitions in the Notebook Jobs dashboard. You can, however, view your notebook jobs as described in **View notebook jobs**.

When you create a job definition, you create a schedule for a job. The **Notebook Job Definitions** tab lists these schedules. For example, you might create a job definition that runs a specific notebook every minute. Once this job definition is active, you see a new job every minute in the **Notebook Jobs** tab.

The **Notebook Job Definitions** tab displays a dashboard with all your job definitions and includes the input notebook, the creation time, the schedule, and the status for each job definition. The value in the **Status** column is one of the following values:

• **Paused**: You paused the job definition. Studio does not initiate any jobs until you resume the definition.
• **Active**: The schedule is on and Studio can run the notebook according to the schedule you specified.
In addition, the **Actions** column provides shortcuts to help you perform the following tasks directly in the interface:

- **Pause**: Pauses the job definition. Studio won’t create any jobs until you resume the definition.
- **Delete**: Removes the job definition from the **Notebook Job Definitions** tab.
- **Resume**: Continues a paused job definition so that it can start jobs.

If you created a job definition but it doesn’t initiate jobs, see [Job definition doesn’t create jobs](#) in the **Troubleshooting guide**.

**View a single job definition**

If you select a job definition name in the **Notebook Job Definitions** tab, you see the **Job Definition** page where you can view specific details for a job definition. Use this page to confirm the settings you specified when you created the job definition. If you don’t see any jobs created from your job definition, see [Job definition doesn’t create jobs](#) in the **Troubleshooting guide**.

This page also contains a section listing the jobs that run from this job definition. Viewing your jobs in the **Job Definition** page may be a more productive way to help you organize your jobs instead of viewing jobs in the **Notebook Jobs** tab, which combines all jobs from all your job definitions.

In addition, this page provides shortcuts for the following actions:

- **Pause/Resume**: Pause your job definition, or resume a paused definition. Note that if a job is currently running for this definition, Studio does not stop it.
- **Run**: Run a single on-demand job from this job definition. This option also lets you specify different input parameters to your notebook before starting the job.
- **Edit Job Definition**: Change the schedule of your job definition. You can select a different time interval, or you can opt for a custom schedule using cron syntax.
- **Delete Job Definition**: Remove the job definition from the **Notebook Job Definitions** tab. Note that if a job is currently running for this definition, Studio does not stop it.

**Troubleshooting guide**

Refer to this troubleshooting guide to help you debug failures you might experience when your scheduled notebook job runs.
Job definition doesn't create jobs

If your job definition doesn’t initiate any jobs, see the following possible causes:

Missing permissions

- The role assigned to the job definition does not have a trust relationship with Amazon EventBridge. That is, EventBridge cannot assume the role.
- The role assigned to the job definition does not have permission to call SageMaker:StartPipelineExecution.
- The role assigned to the job definition does not have permission to call SageMaker:CreateTrainingJob.

EventBridge quota exceeded

If you see a Put* error such as the following example, you exceeded an EventBridge quota. To resolve this, you can clean up unused EventBridge runs, or ask AWS Support to increase your quota.

```
LimitExceeded: The account-level service limit 'Maximum number of pipelines allowed per account' is XXX Pipelines, with current utilization of XXX Pipelines and a request delta of 1 Pipelines.
```

For more information about EventBridge quotas, see Amazon EventBridge quotas.

Pipeline quota limit exceeded

If you see an error such as the following example, you exceeded the number of pipelines that you can run. To resolve this, you can clean up unused pipelines in your account, or ask AWS Support to increase your quota.

```
ResourceLimitExceeded: The account-level service limit
'Maximum number of pipelines allowed per account' is XXX Pipelines,
with current utilization of XXX Pipelines and a request delta of 1 Pipelines.
```

For more information about pipeline quotas, see Amazon SageMaker endpoints and quotas.

Training job limit exceeded

If you see an error such as the following example, you exceeded the number of training jobs that you can run. To resolve this, reduce the number of training jobs in your account, or ask AWS Support to increase your quota.
ResourceLimitExceeded: The account-level service limit 'ml.m5.2xlarge for training job usage' is 0 Instances, with current utilization of 0 Instances and a request delta of 1 Instances. Please contact AWS support to request an increase for this limit.

For more information about training job quotas, see Amazon SageMaker endpoints and quotas.

Auto visualizations disabled in SparkMagic notebooks

If your notebook uses the SparkMagic PySpark kernel and you run the notebook as a Notebook Job, you may see that your auto visualizations are disabled in the output. Turning on auto visualization causes the kernel to hang, so the notebook job executor currently disables auto visualizations as a workaround.

Constraints and considerations

Review the following constraints to ensure your notebook jobs complete successfully. Studio uses Papermill to run notebooks. You might need to update Jupyter notebooks to align to Papermill's requirements. There are also restrictions on the content of LCC scripts and important details to understand regarding VPC configuration.

JupyterLab version

JupyterLab versions 3.0 and above are supported.

Installation of packages that require kernel restart

Papermill does not support calling `pip install` to install packages that require a kernel restart. In this situation, use `pip install` in an initialization script. For a package installation that does not require kernel restart, you can still include `pip install` in the notebook.

Kernel and language names registered with Jupyter

Papermill registers a translator for specific kernels and languages. If you bring your own instance (BYOI), use a standard kernel name as shown in the following snippet:

```python
papermill_translators.register("python", PythonTranslator)
papermill_translators.register("R", RTranslator)
papermill_translators.register("scala", ScalaTranslator)
papermill_translators.register("julia", JuliaTranslator)
papermill_translators.register("matlab", MatlabTranslator)
papermill_translators.register(".net-csharp", CSharpTranslator)
```
Parameters and environment variable limits

Parameters and environment variable limits. When you create your notebook job, it receives the parameters and environment variables you specify. You can pass up to 100 parameters. Each parameter name can be up to 256 characters long, and the associated value can be up to 2500 characters long. If you pass environment variables, you can pass up to 28 variables. The variable name and associated value can be up to 512 characters long. If you need more than 28 environment variables, use additional environment variables in an initialization script which has no limit on the number of environment variables you can use.

Viewing jobs and job definitions

Viewing jobs and job definitions. If you schedule your notebook job in the Studio UI in the JupyterLab notebook, you can view your notebook jobs and your notebook job definitions in the Studio UI. If you scheduled your notebook job with the SageMaker Python SDK, you can view your jobs only—the SageMaker Python SDK notebook job step does not create job definitions. To view your jobs, you also need to supply additional tags to your notebook job step instance. For details, see View your notebook jobs in the Studio UI dashboard.

Image

You need to manage image constraints depending on whether you run notebook jobs in Studio or the SageMaker Python SDK notebook job step in a pipeline.

Image constraints for SageMaker Notebook Jobs (Studio)

Image and kernel support. The driver that launches your notebook job assumes the following:

- A base Python runtime environment is installed in the Studio or bring-your-own (BYO) images and is the default in the shell.
- The base Python runtime environment includes the Jupyter client with kernelspecs properly configured.
- The base Python runtime environment includes the pip function so the notebook job can install system dependencies.
• For images with multiple environments, your initialization script should switch to the proper kernel-specific environment before installing notebook-specific packages. You should switch back to the default Python runtime environment, if different from the kernel runtime environment, after configuring the kernel Python runtime environment.

The driver that launches your notebook job is a bash script, and Bash v4 must be available at /bin/bash.

**Root privileges on bring-your-own-images (BYOI).** You must have root privileges on your own Studio images, either as the root user or through sudo access. If you are not a root user but accessing root privileges through sudo, use 1000/100 as the UID/GID.

**Image constraints for SageMaker Python SDK notebook jobs**

The notebook job step supports the following images:

• SageMaker Distribution Images listed in [Available Amazon SageMaker Images](#).

• A custom image based on the SageMaker Distribution images in the previous list. Use a [SageMaker Distribution image](#) as a base.

• A custom image (BYOI) pre-installed with notebook job dependencies (i.e., sagemaker-headless-execution-driver). Your image must meet the following requirements:
  • The image is pre-installed with notebook job dependencies.
  • A base Python runtime environment is installed and is default in the shell environment.
  • The base Python runtime environment includes the Jupyter client with kernelspecs properly configured.
  • You have root privileges, either as the root user or through sudo access. If you are not a root user but accessing root privileges through sudo, use 1000/100 as the UID/GID.

**VPC subnets used during job creation**

If you use a VPC, Studio uses your private subnets to create your job. Specify one to five private subnets (and 1–15 security groups).

If you use a VPC with private subnets, you must choose one of the following options to ensure the notebook job can connect to dependent services or resources:

• If the job needs access to an AWS service that supports interface VPC endpoints, create an endpoint to connect to the service. For a list of services that support interface endpoints, see
AWS services that integrate with AWS PrivateLink. For information about creating an interface VPC endpoint, see Access an AWS service using an interface VPC endpoint. At minimum, an Amazon S3 VPC endpoint gateway must be provided.

- If a notebook job needs access to an AWS service that doesn't support interface VPC endpoints or to a resource outside of AWS, create a NAT gateway and configure your security groups to allow outbound connections. For information about setting up a NAT gateway for your VPC, see VPC with public and private Subnets (NAT) in the Amazon Virtual Private Cloud User Guide.

Service limits

Since the notebook job scheduler is built from SageMaker Pipelines, SageMaker Training, and Amazon EventBridge services, your notebook jobs are subject to their service-specific quotas. If you exceed these quotas, you may see error messages related to these services. For example, there are limits for how many pipelines you can run at one time, and how many rules you can set up for a single event bus. For more information about SageMaker quotas, see Amazon SageMaker Endpoints and Quotas. For more information about EventBridge quotas, see Amazon EventBridge Quotas.

Pricing for SageMaker Notebook Jobs

When you schedule notebook jobs, your Jupyter notebooks run on SageMaker training instances. After you select an Image and Kernel in your Create Job form, the form provides a list of available compute types. You are charged for the compute type you choose, based on the combined duration of use for all notebook jobs that run from the job definition. If you don't specify a compute type, SageMaker assigns you a default Amazon EC2 instance type of ml.m5.large. For a breakdown of SageMaker pricing by compute type, see Amazon SageMaker Pricing.

Amazon SageMaker ML Lineage Tracking

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.
Amazon SageMaker ML Lineage Tracking creates and stores information about the steps of a machine learning (ML) workflow from data preparation to model deployment. With the tracking information, you can reproduce the workflow steps, track model and dataset lineage, and establish model governance and audit standards.

With SageMaker Lineage Tracking data scientists and model builders can do the following:

- Keep a running history of model discovery experiments.
- Establish model governance by tracking model lineage artifacts for auditing and compliance verification.

The following diagram shows an example lineage graph that Amazon SageMaker automatically creates in an end-to-end model training and deployment ML workflow.

**Lineage Metadata**

[SageMaker automatically creates a connected graph of lineage entity metadata tracking your workflow.]

### Topics

- [Lineage Tracking Entities](#)
- [Amazon SageMaker–Created Tracking Entities](#)
- [Manually Create Tracking Entities](#)
- [Querying Lineage Entities](#)
- [Cross-Account Lineage Tracking](#)
Lineage Tracking Entities

Tracking entities maintain a representation of all the elements of your end-to-end machine learning workflow. You can use this representation to establish model governance, reproduce your workflow, and maintain a record of your work history.

Amazon SageMaker automatically creates tracking entities for trial components and their associated trials and experiments when you create SageMaker jobs such as processing jobs, training jobs, and batch transform jobs. In addition to auto tracking, you can also manually create tracking entities to model custom steps in your workflow. For more information, see Manage Machine Learning with Amazon SageMaker Experiments.

SageMaker also automatically creates tracking entities for the other steps in a workflow so you can track the workflow from end to end. For more information, see Amazon SageMaker–Created Tracking Entities.

You can create additional entities to supplement those created by SageMaker. For more information, see Manually Create Tracking Entities.

SageMaker reuses any existing entities rather than creating new ones. For example, there can be only one artifact with a unique SourceUri.

Key concepts for querying lineage

- **Lineage** – Metadata that tracks the relationships between various entities in your ML workflows.
- **QueryLineage** – The action to inspect your lineage and discover relationships between entities.
- **Lineage entities** – The metadata elements of which your lineage is composed.
- **Cross-account lineage** – Your ML workflow may span more than one account. With cross-account lineage, you can configure multiple accounts to automatically create lineage associations between shared entity resources. QueryLineage then can return entities even from these shared accounts.

The following tracking entities are defined:

**Experiment entities**

- **Trial component** – A stage of a machine learning trial. Includes processing jobs, training jobs, and batch transform jobs.
- **Trial** – A combination of trial components that generally produces a model.
• **Experiment** – A grouping of trials generally focused on solving a specific use case.

**Lineage entities**

• **Trial Component** – Represents processing, training, and transform jobs in the lineage. Also part of experiment management.

• **Context** – Provides a logical grouping of other tracking or experiment entities. Conceptually, experiments and trials are contexts. Some examples are an endpoint and a model package.

• **Action** – Represents an action or activity. Generally, an action involves at least one input artifact or output artifact. Some examples are a workflow step and a model deployment.

• **Artifact** – Represents a URI addressable object or data. An artifact is generally either an input or an output to a trial component or action. Some examples include a dataset (S3 bucket URI), or an image (Amazon ECR registry path).

• **Association** – Links other tracking or experiment entities, such as an association between the location of training data and a training job.

An association has an optional AssociationType property. The following values are available along with the suggested use for each type. SageMaker places no restrictions on their use:

• **ContributedTo** – The source contributed to the destination or had a part in enabling the destination. For example, the training data contributed to the training job.

• **AssociatedWith** – The source is connected to the destination. For example, an approval workflow is associated with a model deployment.

• **DerivedFrom** - The destination is a modification of the source. For example, a digest output of a channel input for a processing job is derived from the original inputs.

• **Produced** – The source generated the destination. For example, a training job produced a model artifact.

• **SameAs** – When the same lineage entity used in different accounts.

**Common properties**

• **Type property**

The action, artifact, and context entities have a `type` property, `ActionType`, `ArtifactType`, and `ContextType`, respectively. This property is a custom string which can associate meaningful information with the entity and be used as a filter in the List APIs.
• **Source property**

The action, artifact, and context entities have a Source property. This property provides the underlying URI that the entity represents. Some examples are:

- An UpdateEndpoint action where the source is the EndpointArn.
- An image artifact for a processing job where the source is the ImageUri.
- An Endpoint context where the source is the EndpointArn.

• **Metadata property**

The action and artifact entities have an optional Metadata property which can provide the following information:

- ProjectId – For example, the ID of the SageMaker MLOps project to which a model belongs.
- GeneratedBy – For example, the SageMaker pipeline execution that registered a model package version.
- Repository – For example, the repository that contains an algorithm.
- CommitId – For example, the commit ID of an algorithm version.

**Amazon SageMaker–Created Tracking Entities**

Amazon SageMaker automatically creates tracking entities for SageMaker jobs, models, model packages, and endpoints if the data is available. There is no limit to the number of lineage entities created by SageMaker.

For information on how you can manually create tracking entities, see [Manually Create Tracking Entities](#).

**Topics**

- [Tracking Entities for SageMaker Jobs](#)
- [Tracking Entities for Model Packages](#)
- [Tracking Entities for Endpoints](#)

**Tracking Entities for SageMaker Jobs**

SageMaker creates a trial component for and associated with each SageMaker job. SageMaker creates artifacts to track the job metadata and associations between each artifact and the job.
Artifacts are created for the following job properties and associated with the Amazon Resource Name (ARN) of the SageMaker job. The artifact SourceUri is listed in parentheses.

**Training Job**

- The image that contains the training algorithm (TrainingImage).
- The data source of each input channel (S3Uri).
- The location for the model (S3OutputPath).
- The location for the managed spot checkpoint data (S3Uri).

**Processing Job**

- The container to be run by the processing job (ImageUri).
- The data location for each processing input and processing output (S3Uri).

**Transform Job**

- The input data source to be transformed (S3Uri).
- The results of the transform (S3OutputPath).

**Note**

Amazon Simple Storage Service (Amazon S3) artifacts are tracked based on the Amazon S3 URI values provided to the Create API, for example CreateTrainingJob, and not on the Amazon S3 key and hash or etag values from each file.

**Tracking Entities for Model Packages**

The following entities are created:

**Model Packages**

- A context for each model package group.
- An artifact for each model package.
- An association between each model package artifact and the context for each model package group to which the package belongs to.
• An action for the creation of a model package version.

• An association between the model package artifact and the creation action.

• An association between the model package artifact and each model package group context to which the package belongs to.

Inference containers

• An artifact for the image used in each container defined in the model package.

• An artifact for the model used in each container.

• An association between each artifact and the model package artifact.

Algorithms

• An artifact for each algorithm defined in the model package.

• An artifact for the model created by each algorithm.

• An association between each artifact and the model package artifact.

Tracking Entities for Endpoints

The following entities are created by Amazon SageMaker:

Endpoints

• A context for each endpoint

• An action for the model deployment that created each endpoint

• An artifact for each model deployed to the endpoint

• An artifact for the image used in the model

• An artifact for the model package for the model

• An artifact for each image deployed to the endpoint

• An association between each artifact and the model deployment action

Manually Create Tracking Entities

You can manually create tracking entities for any property. For information on the tracking entities that Amazon SageMaker automatically creates, see Amazon SageMaker–Created Tracking Entities.
You can add tags to all entities except associations. Tags are arbitrary key-value pairs that provide custom information. You can filter or sort a list or search query by tags. For more information, see Tagging AWS resources in the AWS General Reference.

For a sample notebook that demonstrates how to create lineage entities, see the Amazon SageMaker Lineage notebook in the Amazon SageMaker example GitHub repository.

Topics

- Manually Create Entities
- Manually Track a Workflow
- Limits

Manually Create Entities

The following procedure shows you how to create and associate artifacts between a SageMaker training job and endpoint. You perform the following steps:

Import tracking entities and associations

1. Import the lineage tracking entities.

   ```python
   import sys
   !{sys.executable} -m pip install -q sagemaker
   from sagemaker import get_execution_role
   from sagemaker.session import Session
   from sagemaker.lineage import context, artifact, association, action
   import boto3
   boto_session = boto3.Session(region_name=region)
   sagemaker_client = boto_session.client("sagemaker")
   ```

2. Create the input and output artifacts.

   ```python
   code_location_arn = artifact.Artifact.create(
       artifact_name='source-code-location',
       source_uri='s3://...',
       artifact_type='code-location')
   .artifact_arn
   ```
# Similar constructs for train_data_location_arn and test_data_location_arn

```python
model_location_arn = artifact.Artifact.create(
    artifact_name='model-location',
    source_uri='s3://...',
    artifact_type='model-location')
).artifact_arn
```

3. Train the model and get the `trial_component_arn` that represents the training job.

4. Associate the input artifacts and output artifacts with the training job (trial component).

```python
input_artifacts = [code_location_arn, train_data_location_arn, test_data_location_arn]
for artifact_arn in input_artifacts:
    try:
        association.Association.create(
            source_arn=artifact_arn,
            destination_arn=trial_component_arn,
            association_type='ContributedTo'
        )
    except:
        logging.info('association between {} and {} already exists', artifact_arn, trial_component_arn)

output_artifacts = [model_location_arn]
for artifact_arn in output_artifacts:
    try:
        association.Association.create(
            source_arn=trial_component_arn,
            destination_arn=artifact_arn,
            association_type='Produced'
        )
    except:
        logging.info('association between {} and {} already exists', artifact_arn, trial_component_arn)
```

5. Create the inference endpoint.

```python
predictor = mnist_estimator.deploy(initial_instance_count=1, instance_type='ml.m4.xlarge')
```

6. Create the endpoint context.
from sagemaker.lineage import context

describe_endpoint = sagemaker_client.describe_endpoint(EndpointName=predictor.endpoint_name)

describe_endpoint_arn = endpoint['EndpointArn']

describe_endpoint_context_arn = context.Context.create(
  context_name=predictor.endpoint_name,
  context_type='Endpoint',
  source_uri=describe_endpoint_arn
).context_arn

7. Associate the training job (trial component) and endpoint context.

association.Association.create(
    source_arn=trial_component_arn,
    destination_arn=endpoint_context_arn
)

**Manually Track a Workflow**

You can manually track the workflow created in the previous section.

Given the endpoint Amazon Resource Name (ARN) from the previous example, the following procedure shows you how to track the workflow back to the datasets used to train the model that was deployed to the endpoint. You perform the following steps:

**To track a workflow from endpoint to training data source**

1. Import the tracking entities.

import sys
!{sys.executable} -m pip install -q sagemaker

from sagemaker import get_execution_role
from sagemaker.session import Session
from sagemaker.lineage import context, artifact, association, action
import boto3
boto_session = boto3.Session(region_name=region)
sagemaker_client = boto_session.client("sagemaker")
2. Get the endpoint context from the endpoint ARN.

```python
endpoint_context_arn = sagemaker_client.list_contexts(
    SourceUri=endpoint_arn)["ContextSummaries"]
[0]["ContextArn"]
```

3. Get the trial component from the association between the trial component and the endpoint context.

```python
trial_component_arn = sagemaker_client.list_associations(
    DestinationArn=endpoint_context_arn)["AssociationSummaries"]
[0]["SourceArn"]
```

4. Get the training data location artifact from the association between the trial component and the endpoint context.

```python
train_data_location_artifact_arn = sagemaker_client.list_associations(
    DestinationArn=trial_component_arn, SourceType='Model')["AssociationSummaries"]
[0]["SourceArn"]
```

5. Get the training data location from the training data location artifact.

```python
train_data_location = sagemaker_client.describe_artifact(
    ArtifactArn=train_data_location_artifact_arn)["Source"]['SourceUri']
print(train_data_location)
```

Response:

```
s3://sagemaker-sample-data-us-east-2/mxnet/mnist/train
```

**Limits**

You can create an association between any entities, experiment and lineage, except the following:

- You cannot create an association between two experiment entities. Experiment entities consist of experiments, trials, and trial components.
- You can create an association with another association.

An error occurs if you try to create an entity that already exists.
Maximum number of manually created lineage entities

- Actions: 3000
- Artifacts: 6000
- Associations: 6000
- Contexts: 500

There is no limit to the number of lineage entities automatically created by Amazon SageMaker.

Querying Lineage Entities

Amazon SageMaker automatically generates graphs of lineage entities as you use them. You can query this data to answer a variety of questions. You can query your lineage entities to:

- Retrieve all data sets that went into the creation of a model.
- Retrieve all jobs that went into the creation of an endpoint.
- Retrieve all models that use a data set.
- Retrieve all endpoints that use a model.
- Retrieve which endpoints are derived from a certain data set.
- Retrieve the pipeline execution that created a training job.
- Retrieve the relationships between entities for investigation, governance, and reproducibility.
- Retrieve all downstream trials that use the artifact.
- Retrieve all upstream trials that use the artifact.
- Retrieve a list of artifacts that use the provided S3 uri.
- Retrieve upstream artifacts that use the dataset artifact.
- Retrieve downstream artifacts that use the dataset artifact.
- Retrieve datasets that use the image artifact.
- Retrieve actions that use the context.
- Retrieve processing jobs that use the endpoint.
- Retrieve transform jobs that use the endpoint.
- Retrieve trial components that use the endpoint.
- Retrieve the ARN for the pipeline execution associated with the model package group.
• Retrieve all artifacts that use the action.
• Retrieve all upstream datasets that use the model package approval action.
• Retrieve model package from model package approval action.
• Retrieve downstream endpoint contexts that use the endpoint.
• Retrieve the ARN for the pipeline execution associated with the trial component.
• Retrieve datasets that use the trial component.
• Retrieve models that use the trial component.
• Explore your lineage for visualization.

Limitations

• Lineage querying is not available in the following Regions:
  • Africa (Cape Town) – af-south
  • Asia Pacific (Jakarta) – ap-southeast-3
  • Asia Pacific (Osaka) – ap-northeast-3
  • Europe (Milan) – eu-south-1
  • Europe (Spain) – eu-south-2
  • Israel (Tel Aviv) – il-central-1
• The maximum depth of relationships to discover is currently limited to 10.
• Filtering is limited to the following properties: last modified date, created date, type, and lineage entity type.

Topics

• Getting Started with Querying Lineage Entities

Getting Started with Querying Lineage Entities

The easiest way to get started is either via the:

• Amazon SageMaker SDK for Python which has defined many common use cases.
• For a notebook that demonstrates how to use SageMaker Lineage APIs to query relationships across the lineage graph, see sagemaker-lineage-multihop-queries.ipynb.
The following examples show how to use the LineageQuery and LineageFilter APIs to construct queries to answer questions about the Lineage Graph and extract entity relationships for a few use cases.

**Example Using the LineageQuery API to find entity associations**

```python
from sagemaker.lineage.context import Context, EndpointContext
from sagemaker.lineage.action import Action
from sagemaker.lineage.association import Association
from sagemaker.lineage.artifact import Artifact, ModelArtifact, DatasetArtifact

from sagemaker.lineage.query import (LineageQuery,
                                     LineageFilter,
                                     LineageSourceEnum,
                                     LineageEntityEnum,
                                     LineageQueryDirectionEnum,
                                     )

# Find the endpoint context and model artifact that should be used for the lineage queries.
contexts = Context.list(source_uri=endpoint_arn)
context_name = list(contexts)[0].context_name
endpoint_context = EndpointContext.load(context_name=context_name)
```

**Example Find all the datasets associated with an endpoint**

```python
# Define the LineageFilter to look for entities of type `ARTIFACT` and the source of type `DATASET`.
query_filter = LineageFilter(
    entities=[LineageEntityEnum.ARTIFACT], sources=[LineageSourceEnum.DATASET]
)

# Providing this `LineageFilter` to the `LineageQuery` constructs a query that traverses through the given context `endpoint_context`
# and find all datasets.
query_result = LineageQuery(sagemaker_session).query(
    start_arns=[endpoint_context.context_arn],
    query_filter=query_filter,
    direction=LineageQueryDirectionEnum.ASCENDANTS,
)```
include_edges=False,
)

# Parse through the query results to get the lineage objects corresponding to the datasets
dataset_artifacts = []
for vertex in query_result.vertices:
    dataset_artifacts.append(vertex.to_lineage_object().source.source_uri)

pp.pprint(dataset_artifacts)

Example Find the models associated with an endpoint

# Define the LineageFilter to look for entities of type 'ARTIFACT' and the source of type 'MODEL'.
query_filter = LineageFilter(
    entities=[LineageEntityEnum.ARTIFACT], sources=[LineageSourceEnum.MODEL]
)

# Providing this 'LineageFilter' to the 'LineageQuery' constructs a query that traverses through the given context 'endpoint_context'
# and find all datasets.
query_result = LineageQuery(sagemaker_session).query(
    start_arns=[endpoint_context.context_arn],
    query_filter=query_filter,
    direction=LineageQueryDirectionEnum.ASCENDANTS,
    include_edges=False,
)

# Parse through the query results to get the lineage objects corresponding to the model
model_artifacts = []
for vertex in query_result.vertices:
    model_artifacts.append(vertex.to_lineage_object().source.source_uri)

# The results of the 'LineageQuery' API call return the ARN of the model deployed to the endpoint along with
# the S3 URI to the model.tar.gz file associated with the model
pp.pprint(model_artifacts)
Example Find the trial components associated with the endpoint

```python
# Define the LineageFilter to look for entities of type 'TRIAL_COMPONENT' and the source of type 'TRAINING_JOB'.

query_filter = LineageFilter(
    entities=[LineageEntityEnum.TRIAL_COMPONENT],
    sources=[LineageSourceEnum.TRAINING_JOB],
)

# Providing this `LineageFilter` to the `LineageQuery` constructs a query that traverses through the given context `endpoint_context` and find all datasets.

query_result = LineageQuery(sagemaker_session).query(
    start_arns=[endpoint_context.context_arn],
    query_filter=query_filter,
    direction=LineageQueryDirectionEnum.ASCENDANTS,
    include_edges=False,
)

# Parse through the query results to get the ARNs of the training jobs associated with this Endpoint

trial_components = []
for vertex in query_result.vertices:
    trial_components.append(vertex.arn)

pp.pprint(trial_components)
```

Example Changing the focal point of lineage

The LineageQuery can be modified to have different start_arns which changes the focal point of lineage. In addition, the LineageFilter can take multiple sources and entities to expand the scope of the query.

In the following we use the model as the lineage focal point and find the endpoints and datasets associated with it.

```python
# Get the ModelArtifact

model_artifact_summary = list(Artifact.list(source_uri=model_package_arn))[0]
model_artifact = ModelArtifact.load(artifact_arn=model_artifact_summary.artifact_arn)
query_filter = LineageFilter(
```
Example Using `LineageQueryDirectionEnum.BOTH` to find ascendent and descendent relationships

When the direction is set to BOTH, the query traverses the graph to find ascendant and descendant relationships. This traversal takes place not only from the starting node, but from each node that is visited. For example; if a training job is run twice and both models generated by the training job are deployed to endpoints, the result of the query with direction set to BOTH shows both endpoints. This is because the same image is used for training and deploying the model. Since the image is common to the model, the start_arn and both the endpoints, appear in the query result.

```python
query_filter = LineageFilter(
    entities=[LineageEntityEnum.ARTIFACT],
    sources=[LineageSourceEnum.ENDPOINT, LineageSourceEnum.DATASET],
)
```
query_result = LineageQuery(sagemaker_session).query(
    start_arns=[model_artifact.artifact_arn],  # Model is the starting artifact
    query_filter=query_filter,
    # This specifies that the query should look for associations both ascending and
descending for the start
direction=LineageQueryDirectionEnum.BOTH,
    include_edges=False,
)

associations = []
for vertex in query_result.vertices:
    associations.append(vertex.to_lineage_object().source.source_uri)

pp.pprint(associations)

Example Directions in LineageQuery - ASCENDANTS vs. DESCENDANTS

To understand the direction in the Lineage Graph, take the following entity relationship graph -
Dataset -> Training Job -> Model -> Endpoint

The endpoint is a descendant of the model, and the model is a descendant of the dataset. Similarly,
the model is an ascendant of the endpoint. The direction parameter can be used to specify
whether the query should return entities that are descendants or ascendants of the entity in
start_arns. If the start_arns contains a model and the direction is DESCENDANTS, the query
returns the endpoint. If the direction is ASCENDANTS, the query returns the dataset.

# In this example, we'll look at the impact of specifying the direction as ASCENDANT or
DESCENDANT in a `LineageQuery`.

query_filter = LineageFilter(
    entities=[LineageEntityEnum.ARTIFACT],
    sources=[
        LineageSourceEnum.ENDPOINT,
        LineageSourceEnum.MODEL,
        LineageSourceEnum.DATASET,
        LineageSourceEnum.TRAINING_JOB,
    ],
)

query_result = LineageQuery(sagemaker_session).query(
    start_arns=[model_artifact.artifact_arn],
)
query_result = LineageQuery(sagemaker_session).query(
    start_arns=[model_artifact.artifact_arn],
    query_filter=query_filter,
    direction=LineageQueryDirectionEnum.DESCENDANTS,
    include_edges=False,
)

# The lineage entity returned for the Training Job is a TrialComponent which can't be converted to a lineage object using the method `to_lineage_object()` so we extract the TrialComponent ARN.
for vertex in query_result.vertices:
    try:
        descendant_artifacts.append(vertex.to_lineage_object().source.source_uri)
    except:
        # Handling TrialComponents.
        descendant_artifacts.append(vertex.arn)

print("Descendant artifacts : ")
pp.pprint(descendant_artifacts)

Example SDK helper functions to make lineage queries easier

The classes EndpointContext, ModelArtifact, and DatasetArtifact have helper functions that are wrappers over the LineageQuery API to make certain lineage queries easier to leverage. The following example shows how to use these helper function.
# Find all the datasets associated with this endpoint

datasets = []
dataset_artifacts = endpoint_context.dataset_artifacts()
for dataset in dataset_artifacts:
    datasets.append(dataset.source.source_uri)
print("Datasets : ", datasets)

# Find the training jobs associated with the endpoint
training_job_artifacts = endpoint_context.training_job_arns()
training_jobs = []
for training_job in training_job_artifacts:
    training_jobs.append(training_job)
print("Training Jobs : ", training_jobs)

# Get the ARN for the pipeline execution associated with this endpoint (if any)
pipeline_executions = endpoint_context.pipeline_execution_arn()
if pipeline_executions:
    for pipeline in pipelines_executions:
        print(pipeline)

# Here we use the `ModelArtifact` class to find all the datasets and endpoints associated with the model

dataset_artifacts = model_artifact.dataset_artifacts()
endpoint_contexts = model_artifact.endpoint_contexts()

datasets = [dataset.source.source_uri for dataset in dataset_artifacts]
endpoints = [endpoint.source.source_uri for endpoint in endpoint_contexts]

print("Datasets associated with this model : ")
pp.pprint(datasets)

print("Endpoints associated with this model : ")
pp.pprint(endpoints)

# Here we use the `DatasetArtifact` class to find all the endpoints hosting models that were trained with a particular dataset
# Find the artifact associated with the dataset

dataset_artifact_arn = list(Artifact.list(source_uri=training_data))[0].artifact_arn
dataset_artifact = DatasetArtifact.load(artifact_arn=dataset_artifact_arn)
# Find the endpoints that used this training dataset

```python
endpoint_contexts = dataset_artifact.endpoint_contexts()
endpoints = [endpoint.source.source_uri for endpoint in endpoint_contexts]

print("Endpoints associated with the training dataset {}".format(training_data))
pp.pprint(endpoints)
```

### Example Getting a Lineage graph visualization

A helper class `Visualizer` is provided in the sample notebook `visualizer.py` to help plot the lineage graph. When the query response is rendered, a graph with the lineage relationships from the `StartArns` is displayed. From the `StartArns` the visualization shows the relationships with the other lineage entities returned in the `query_lineage` API action.

```python
# Graph APIs
# Here we use the boto3 `query_lineage` API to generate the query response to plot.

from visualizer import Visualizer

query_response = sm_client.query_lineage(
    StartArns=[endpoint_context.context_arn], Direction="Ascendants", IncludeEdges=True
)

t = Visualizer()
t.render(query_response, "Endpoint")

query_response = sm_client.query_lineage(
    StartArns=[model_artifact.artifact_arn], Direction="Ascendants", IncludeEdges=True
)
t.render(query_response, "Model")
```

### Cross-Account Lineage Tracking

Amazon SageMaker supports tracking lineage entities from a different AWS account. Other AWS accounts can share their lineage entities with you and you can access these lineage entities through direct API calls or SageMaker lineage queries.

SageMaker uses [AWS Resource Access Manager](https://docs.aws.amazon.com/ram/latest/userguide/what-is-ram.html) to help you securely share your lineage resources. You can share your resources through the [AWS RAM console](https://console.aws.amazon.com/ram/).
Set Up Cross-Account Lineage Tracking

You can group and share your Lineage Tracking Entities through a lineage group in Amazon SageMaker. SageMaker supports only one default lineage group per account. SageMaker creates the default lineage group whenever a lineage entity is created in your account. Every lineage entity owned by your account is assigned to this default lineage group. To share lineage entities with another account, you share this default lineage group with that account.

Note
You can share all lineage tracking entities in a lineage group or none.

Create a resource share for your lineage entities using AWS Resource Access Manager console. For more information, see Sharing your AWS resources in the AWS Resource Access Manager User Guide.

Note
After the resource share is created, it can take a few minutes for the resource and principal associations to complete. Once the association is set, the shared account receives an invitation to join the resource share. The shared account must accept the invite to gain access to shared resources. For more information on accepting a resource share invite in AWS RAM, see Using shared AWS resources in the AWS Resource Access Manager User Guide.

Your cross-account lineage tracking resource policy

Amazon SageMaker supports only one type of resource policy. The SageMaker resource policy must allow all of the following operations:

"sagemaker:DescribeAction"
"sagemaker:DescribeArtifact"
"sagemaker:DescribeContext"
"sagemaker:DescribeTrialComponent"
"sagemaker:AddAssociation"
"sagemaker:DeleteAssociation"
"sagemaker:QueryLineage"
Example The following is a SageMaker resource policy created using AWS Resource Access Manager for creating a resource share for an accounts lineage group.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "FullLineageAccess",
            "Effect": "Allow",
            "Principal": {
                "AWS": "123456789012" #account-id
            },
            "Action": [
                "sagemaker:DescribeAction",
                "sagemaker:DescribeArtifact",
                "sagemaker:DescribeContext",
                "sagemaker:DescribeTrialComponent",
                "sagemaker:AddAssociation",
                "sagemaker:DeleteAssociation",
                "sagemaker:QueryLineage"
            ],
        }
    ]
}
```

**Tracking Cross-Account Lineage Entities**

With cross-account lineage tracking you can associate lineage entities in different accounts using the same AddAssociation API action. When you associate two lineage entities, SageMaker validates if you have permissions to perform the AddAssociation API action on both lineage entities. SageMaker then establishes the association. If you don’t have the permissions, SageMaker does not create the association. Once the cross-account association is established, you can access either lineage entity from the other through the QueryLineage API action. For more information, see [Querying Lineage Entities](#).

In addition to SageMaker automatically creating lineage entities, if you have cross-account access, SageMaker connects artifacts that reference the same object or data. If the data from one account is used in lineage tracking by different accounts, SageMaker creates an artifact in each account to track that data. With cross-account lineage, whenever SageMaker creates new artifacts,
SageMaker checks if there are other artifacts created for the same data that are also shared with you. SageMaker then establishes associations between the newly created artifact and each of the artifacts shared with you with the AssociationType set to SameAs. You can then use the QueryLineage API action to traverse the lineage entities in your own account to lineage entities shared with you but owned by a different AWS account. For more information, see Querying Lineage Entities.

Topics

- Accessing lineage resources from a different accounts
- Authorization for querying cross-account lineage entities

Accessing lineage resources from a different accounts

Once the cross-account access for sharing lineage has been set up, you can call the following SageMaker API actions directly with the ARN to describe the shared lineage entities from another account:

- DescribeAction
- DescribeArtifact
- DescribeContext
- DescribeTrialComponent

You can also manage Associations for lineage entities owned by different accounts that are shared with you, using the following SageMaker API actions:

- AddAssociation
- DeleteAssociation

For a notebook that demonstrates how to use SageMaker Lineage APIs to query lineage across accounts, see sagemaker-lineage-cross-account-with-ram.ipynb.

Authorization for querying cross-account lineage entities

Amazon SageMaker must validate that you have permissions to perform the QueryLineage API action on the StartArns. This is enforced through the resource policy attached to the LineageGroup. The result from this action includes all the lineage entities to which you
have access, whether they are owned by your account or shared by another account. For more information, see Querying Lineage Entities.

Register and Deploy Models with Model Registry

With the SageMaker Model Registry you can do the following:

- Catalog models for production.
- Manage model versions.
- Associate metadata, such as training metrics, with a model.
- Manage the approval status of a model.
- Deploy models to production.
- Automate model deployment with CI/CD.

Catalog models by creating SageMaker Model Registry Model (Package) Groups that contain different versions of a model. You can create a Model Group that tracks all of the models that you train to solve a particular problem. You can then register each model you train and the Model Registry adds it to the Model Group as a new model version. Lastly, you can create categories of Model Groups by further organizing them into SageMaker Model Registry Collections. A typical workflow might look like the following:

- Create a Model Group.
- Create an ML pipeline that trains a model. For information about SageMaker pipelines, see Create and Manage SageMaker Pipelines.
- For each run of the ML pipeline, create a model version that you register in the Model Group you created in the first step.
- Add your Model Group into one or more Model Registry Collections.

For details about how to create and work with models, model versions, and Model Groups, see Model Registry Models, Model Versions, and Model Groups. Optionally, if you want to further group your Model Groups into Collections, see Model Registry Collections.

Model Registry Models, Model Versions, and Model Groups

The SageMaker Model Registry is structured as several Model (Package) Groups with model packages in each group. These Model Groups can optionally be added to one or more Collections.
Each model package in a Model Group corresponds to a trained model. The version of each model package is a numerical value that starts at 1 and is incremented with each new model package added to a Model Group. For example, if 5 model packages are added to a Model Group, the model package versions will be 1, 2, 3, 4, and 5.

There are two types of model packages in SageMaker. One type is used in the AWS Marketplace, and the other is used in the Model Registry. Model packages used in the AWS Marketplace are not versionable entities and are not associated with Model Groups in the Model Registry. For more information about model packages used in the AWS Marketplace, see Sell algorithms and packages in the AWS Marketplace.

The model packages used in the Model Registry are versioned, and must be associated with a Model Group. The ARN of this model package type has the structure: 'arn:aws:sagemaker:region:account:model-package-group/version'

The following topics show you how to create and work with models, model versions, and Model Groups in the Model Registry.

Topics

- Create a Model Group
- Delete a Model Group
- Register a Model Version
- View Model Groups and Versions
- View the Details of a Model Version
- Compare Model Versions
- View and Manage Model Group and Model Version Tags
- Share Models with SageMaker Canvas Users
- Delete a Model Version
- Update the Approval Status of a Model
- Deploy a Model from the Registry
- View the Deployment History of a Model

Create a Model Group

A Model Group contains a group of versioned models. Create a Model Group by using either the AWS SDK for Python (Boto3) or the Amazon SageMaker Studio console.
Create a Model Group (Boto3)

To create a Model Group by using Boto3, call the `create_model_package_group` API operation and specify a name and description as parameters. The following example shows how to create a Model Group. The response from the `create_model_package_group` call is the Amazon Resource Name (ARN) of the new Model Group.

First, import the required packages and set up the SageMaker Boto3 client.

```python
import time
import os
from sagemaker import get_execution_role, session
import boto3

region = boto3.Session().region_name
role = get_execution_role()

sm_client = boto3.client('sagemaker', region_name=region)

Now create the Model Group.

```python
model_package_group_name = "scikit-iris-detector-" + str(round(time.time()))
model_package_group_input_dict = {
    "ModelPackageGroupName" : model_package_group_name,
    "ModelPackageGroupDescription" : "Sample model package group"
}

create_model_package_group_response = 
    sm_client.create_model_package_group(**model_package_group_input_dict)
print('ModelPackageGroup Arn :
    {}'.format(create_model_package_group_response['ModelPackageGroupArn']))
```

Create a Model Group (console)

To create a Model Group in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.
Studio

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. Choose Register, then choose Model group.
6. In the Register model group dialog box, enter the following information:
   - The name of the new Model Group in the Model group name field.
   - (Optional) A description for the Model Group in the Description field.
   - (Optional) Any key-value pairs you want to associate with the Model Group in the Tags field. For information about using tags, see Tagging AWS resources in the AWS General Reference.
7. Choose Register model group.
8. (Optional) In the Models page, choose the Registered models tab, then choose Model Groups. Confirm your newly-created Model Group appears in the list of Model Groups.

Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the Home icon ( ).
3. Choose Models, and then Model registry.
4. Choose Actions, then choose Create model group.
5. In the Create model group dialog box, enter the following information:
   - Enter the name of the new Model Group in the Model group name field.
   - (Optional) Enter a description for the Model Group in the Description field.
• (Optional) Enter any key-value pairs you want to associate with the Model Group in the **Tags** field. For information about using tags, see [Tagging AWS resources](https://docs.aws.amazon.com/general/latest/dg/aws-tagging-resource-elements.html) in the AWS General Reference.

• (Optional) Choose a project with which to associate the Model Group in the **Project** field. For information about projects, see [Automate MLOps with SageMaker Projects](https).

6. Choose **Create model group**.

---

### Delete a Model Group

This procedure demonstrates how to delete a Model Group in the Amazon SageMaker Studio console.

#### Delete a Model Group (console)

⚠️ **Important**

You can only delete an empty model group. Before you delete your model group, remove its model versions, if any.

To delete a Model Group in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](https).
2. In the left navigation pane, choose **Models**.
3. Choose the **Registered models** tab, if not selected already.
4. Immediately below the Registered models tab label, choose **Model Groups**, if not selected already.
5. From the model groups list, select the check box next to the name of the Model Group you want to delete.
6. Choose the vertical ellipsis above the top right corner of the model groups list, and choose **Delete**.
7. In the **Delete model group** dialog box, choose **Yes, delete the model group**.
8. Choose **Delete**.

9. Confirm that your deleted model groups no longer appear in your list of model groups.

### Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).

2. In the left navigation pane, choose the **Home** icon (↑).

3. Choose **Models**, and then **Model registry**. A list of your Model Groups appears.

4. From the model groups list, select the name of the Model Group you want to delete.

5. In the top right corner, choose **Remove**.

6. In the confirmation dialog box, enter **REMOVE**.

7. Choose **Remove**.

### Register a Model Version

You can register an Amazon SageMaker model by creating a model version that specifies the model group to which it belongs. A model version must include both the model artifacts (the trained weights of a model) and the inference code for the model.

An **inference pipeline** is a SageMaker model composed of a linear sequence of two to fifteen containers that process inference requests. You register an inference pipeline by specifying the containers and the associated environment variables. For more information on inference pipelines, see [Host models along with pre-processing logic as serial inference pipeline behind one endpoint](#).

You can register a model with an inference pipeline, by specifying the containers and the associated environment variables. To create a model version with an inference pipeline by using either the AWS SDK for Python (Boto3), the Amazon SageMaker Studio console, or by creating a step in a SageMaker model building pipeline, use the following steps.

### Topics

- [Register a Model Version (SageMaker Pipelines)](#)
- [Register a Model Version (Boto3)](#)
Register a Model Version (SageMaker Pipelines)

To register a model version by using a SageMaker model building pipeline, create a `RegisterModel` step in your pipeline. For information about creating a `RegisterModel` step as part of a pipeline, see [Step 8: Define a RegisterModel Step to Create a Model Package](#).

Register a Model Version (Boto3)

To register a model version by using Boto3, call the `create_model_package` API operation.

First, you set up the parameter dictionary to pass to the `create_model_package` API operation.

```python
# Specify the model source
model_url = "s3://your-bucket-name/model.tar.gz"

modelpackage_inference_specification = {
    "InferenceSpecification": {
        "Containers": [
            {
                "Image": image_uri,
                "ModelDataUrl": model_url
            }
        ],
        "SupportedContentTypes": [ "text/csv" ],
        "SupportedResponseMIMETypes": [ "text/csv" ],
    }
}

# Alternatively, you can specify the model source like this:
# modelpackage_inference_specification["InferenceSpecification"]["Containers"][0]["ModelDataUrl"] = model_url

create_model_package_input_dict = {
    "ModelPackageGroupName": model_package_group_name,
    "ModelPackageDescription": "Model to detect 3 different types of irises (Setosa, Versicolour, and Virginica)",
    "ModelApprovalStatus": "PendingManualApproval"
}

create_model_package_input_dict.update(modelpackage_inference_specification)
```
Then you call the `create_model_package` API operation, passing in the parameter dictionary that you just set up.

```python
create_model_package_response = sm_client.create_model_package(**create_model_package_input_dict)
model_package_arn = create_model_package_response['ModelPackageArn']
print('ModelPackage Version ARN : {}'.format(model_package_arn))
```

Register a Model Version (console)

To register a model version in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models from the menu.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. Choose Register, then choose Model version.
6. In the Register model version form, enter the following information:
   - In the Model group name dropdown, select the name of the model group to which your version belongs.
   - (Optional) Enter a description for your model version.
   - In the Model Approval Status dropdown, select the version approval status.
   - (Optional) In the Custom metadata field, choose + Add new and add custom tags as key-value pairs.
7. Choose Next.
8. In the Inference Specification form, enter the following information:
   - In Inference image location (ECR), enter your ECR inference image location.
   - In Model artifact location (S3), enter the Amazon S3 bucket location of your model data artifacts.
• To specify and input data configuration or environment variables, choose **Additional configuration** and enter this information.

• To add more containers, choose **+ Add container**.

• In **Realtime inference instance type**, enter the instance type to use for real-time inference.

• In **Transform inference instance type**, enter the instance type to use for batch transformations.

• In **Supported content types**, enter your input MIME types.

• In **Supported response content types**, enter your output MIME types.

9. Choose Next.

10. In the optional **Inference Recommendation** form, enter the following information:

   • For **Business problem**, choose the application the applies to your model.
   
   • For **Task**, choose the type of problem that applies to your model.
   
   • For **S3 bucket address**, enter the Amazon S3 bucket location of your sample payload.
   
   • For the first container, enter the following information:
      
     - For **Model name**, enter the model name as used in model zoos.
     
     - For **Framework**, choose a framework.
     
     - For **Framework version**, enter a framework version.
     
     • Repeat the previous step for all containers.

11. Choose Next.

12. Select the check box next to one or more of the displayed model metrics.

13. Choose Next.

14. Ensure the displayed settings are correct, and choose **Register model version**. If you subsequently see a modal window with an error message, choose View (next to the message) to view the source of the error.

15. Confirm your new model version appears in the parent model group page.

### Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
2. In the left navigation pane, choose the **Home** icon (  
   🏡).

3. Choose **Models**, and then **Model registry**.

4. Open the **Register Version** form. You can do this in one of two ways:
   - Choose **Actions**, and then choose **Create model version**.
   - Select the name of the model group for which you want to create a model version, then choose **Create model version**.

5. In the **Register model version** form, enter the following information:
   - In the **Model package group name** dropdown, select the model group name.
   - (Optional) Enter a description for your model version.
   - In the **Model Approval Status** dropdown, select the version approval status.
   - (Optional) In the **Custom metadata** field, add custom tags as key-value pairs.

6. Choose **Next**.

7. In the **Inference Specification** form, enter the following information:
   - Enter your inference image location.
   - Enter your model data artifacts location.
   - (Optional) Enter information about images to use for transform and real-time inference jobs, and supported input and output MIME types.

8. Choose **Next**.

9. (Optional) Provide details to aid endpoint recommendations.

10. Choose **Next**.

11. (Optional) Choose model metrics you want to include.

12. Choose **Next**.

13. Ensure the displayed settings are correct, and choose **Register model version**. If you subsequently see a modal window with an error message, choose **View** (next to the message) to view the source of the error.

14. Confirm your new model version appears in the parent model group page.
Register a Model Version from a Different Account

To register model versions with a Model Group created by a different AWS account, you must add a cross-account AWS Identity and Access Management resource policy to enable that account. For example, one AWS account in your organization is responsible for training models, and a different account is responsible for managing, deploying, and updating models. You create IAM resource policies and apply the policies to the specific account resource to which you want to grant access for this case. For more information about cross-account resource policies in AWS, see Cross-account policy evaluation logic in the AWS Identity and Access Management User Guide.

Note
You must also use a KMS key to encrypt the output data config action during training for cross-account model deployment.

To enable cross-account model registry in SageMaker, you have to provide a cross-account resource policy for the Model Group that contains the model versions. The following is an example that creates cross-account policies for the Model Group and applies these policies to that specific resource.

The following configuration must be set in the source account which registers models cross-account in a Model Group. In this example, the source account is the model training account which will train and then register the model cross-account into the Model Registry of the Model Registry account.

The example assumes that you previously defined the following variables:

- `model_package_group_name` – The Model Group to which you want to grant access.
- `model_package_group_arn` – The Model Group ARN to which you want to grant cross-account access.
- `bucket` – The Amazon S3 bucket where the model training artifacts are stored.

To be able to deploy a model created in a different account, the user must have a role that has access to SageMaker actions, such as a role with the AmazonSageMakerFullAccess managed policy. For information about SageMaker managed policies, see AWS Managed Policies for Amazon SageMaker.
Required IAM resource policies

The following diagram captures the policies required to allow cross-account model registration. As shown, these policies need to be active during model training to properly register the model into the Model Registry account.

Amazon ECR, Amazon S3, and AWS KMS policies are demonstrated in the following code samples.

Sample Amazon ECR policy

```json
{
```
Sample Amazon S3 policy

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "AddPerm",
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::\{model_registry_account\}:root"
            },
            "Action": [
                "s3:GetObject",
                "s3:GetBucketAcl",
                "s3:GetObjectAcl"
            ],
            "Resource": "arn:aws:s3:::bucket/*"
        }
    ]
}
```

Sample AWS KMS policy

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "AddPerm",
            "Effect": "Allow",
            "Principal": {
                "AWS": "arn:aws:iam::\{model_registry_account\}:root"
            },
            "Action": [
                "kms:Encrypt",
                "kms:Decrypt",
                "kms:ReEncrypt*,
                "kms:GenerateDataKey*",
                "kms:DescribeKey"
            ],
            "Resource": "arn:aws:kms::\{model_registry_account\}:key/\{kms_key_id\}"
        }
    ]
}
```
"Sid": "AddPerm",
"Effect": "Allow",
"Principal": {
    "AWS": "arn:aws:iam::\{model_registry_account\}:root"
},
"Action": [
    "kms:Decrypt",
    "kms:GenerateDataKey***"
],
"Resource": "***"
}
]
]

Apply resource policies to accounts

The following policy configuration applies the policies discussed in the previous section and must be put in the model training account.

```python
import json

# The Model Registry account id of the Model Group
model_registry_account = "111111111111"

# The model training account id where training happens
model_training_account = "222222222222"

# 1. Create a policy for access to the ECR repository
# in the model training account for the Model Registry account Model Group
ecr_repository_policy = {
    "Version": "2012-10-17",
    "Statement": [{
        "Sid": "AddPerm",
        "Effect": "Allow",
        "Principal": {
            "AWS": f"arn:aws:iam::\{model_registry_account\}:root"
        },
        "Action": [
            "ecr:BatchGetImage",
            "ecr:Describe***"
        ]
    }]
}

# Convert the ECR policy from JSON dict to string
```
ecr_repository_policy = json.dumps(ecr_repository_policy)

# Set the new ECR policy
ecr = boto3.client('ecr')
response = ecr.set_repository_policy(
    registryId = model_training_account,
    repositoryName = "decision-trees-sample",
    policyText = ecr_repository_policy
)

# 2. Create a policy in the model training account for access to the S3 bucket
# where the model is present in the Model Registry account Model Group
bucket_policy = {
    "Version": "2012-10-17",
    "Statement": [{
        "Sid": "AddPerm",
        "Effect": "Allow",
        "Principal": {
            "AWS": f"arn:aws:iam::{model_registry_account}:root"
        },
        "Action": [
            "s3:GetObject",
            "s3:GetBucketAcl",
            "s3:GetObjectAcl"
        ],
        "Resource": "arn:aws:s3:::{bucket}/*"
    }
]
}

# Convert the S3 policy from JSON dict to string
bucket_policy = json.dumps(bucket_policy)

# Set the new bucket policy
s3 = boto3.client("s3")
response = s3.put_bucket_policy(
    Bucket = bucket,
    Policy = bucket_policy
)

# 3. Create the KMS grant for the key used during training for encryption
# in the model training account to the Model Registry account Model Group
client = boto3.client("kms")

response = client.create_grant(
    GranteePrincipal=model_registry_account,
    KeyId=kms_key_id
    Operations=[
        "Decrypt",
    ]
)
The following configuration needs to be put in the Model Registry account where the Model Group exists.

```python
# The Model Registry account id of the Model Group
model_registry_account = "111111111111"

# 1. Create policy to allow the model training account to access the ModelPackageGroup
model_package_group_policy = {
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "AddPermModelPackageVersion",
            "Effect": "Allow",
            "Principal": {
                "AWS": f"arn:aws:iam::{model_training_account}:root"},
            "Action": ["sagemaker:CreateModelPackage"],
            "Resource": f"arn:aws:sagemaker:{region}:{model_registry_account}:model-package/" + model_package_group_name + "/*"
        }
    ]
}

# Convert the policy from JSON dict to string
model_package_group_policy = json.dumps(model_package_group_policy)

# Set the new policy
response = sm_client.put_model_package_group_policy(
    ModelPackageGroupName = model_package_group_name,
    ResourcePolicy = model_package_group_policy)
```

Finally, use the create_model_package action from the model training account to register the model package in the cross-account.

```python
# Specify the model source
model_url = "s3://{bucket}/model.tar.gz"

#Set up the parameter dictionary to pass to the create_model_package API operation
```
modelpackage_inference_specification = {
    "InferenceSpecification": {
        "Containers": [
            {
                "Image": f'{model_training_account}.dkr.ecr.us-east-2.amazonaws.com/decision-trees-sample:latest",
                "ModelDataUrl": model_url
            }
        ],
        "SupportedContentTypes": [ "text/csv" ],
        "SupportedResponseMIMETypes": [ "text/csv" ],
    }
}

# Alternatively, you can specify the model source like this:
# modelpackage_inference_specification["InferenceSpecification"]['Containers'][0]
# "ModelDataUrl"=model_url

create_model_package_input_dict = {
    "ModelPackageName" : model_package_group_arn,
    "ModelPackageDescription" : "Model to detect 3 different types of irises (Setosa, Versicolour, and Virginica)",
    "ModelApprovalStatus" : "PendingManualApproval"
}
create_model_package_input_dict.update(modelpackage_inference_specification)

# Create the model package in the Model Registry account
create_model_package_response = 
    sm_client.create_model_package(**create_model_package_input_dict)
model_package_arn = create_model_package_response['ModelPackageArn']
print('ModelPackage Version ARN : {}'.format(model_package_arn))

View Model Groups and Versions

Model Groups and versions help you organize your models. You can view a list of the model versions in a Model Group by using either the AWS SDK for Python (Boto3) (Boto3) or the Amazon SageMaker Studio console.

View a List of Model Versions in a Group

You can view all of the model versions that are associated with a Model Group. If a Model Group represents all models that you train to address a specific ML problem, you can view all of those related models.
View a List of Model Versions in a Group (Boto3)

To view model versions associated with a Model Group by using Boto3, call the list_model_packages API operation, and pass the name of the Model Group as the value of the ModelPackageGroupName parameter. The following code lists the model versions associated with the Model Group you created in Create a Model Group (Boto3).

```python
sm_client.list_model_packages(ModelPackageGroupName=model_package_group_name)
```

View a List of Model Versions in a Group (console)

To view a list of the model versions in a Model Group in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models from the menu.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the model groups list, choose the angle bracket to the left of the model group you want to view.
6. A list of the model versions in the model group appears.
7. (Optional) Choose View all, if shown, to view additional model versions.

**Studio Classic**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the Home icon (🏠).
3. Choose Models, and then Model registry.
4. From the model groups list, select the name of the Model Group you want to view.
5. A new tab appears with a list of the model versions in the Model Group.

View the Details of a Model Version

You can view details of a specific model version by using either the AWS SDK for Python (Boto3) or the Amazon SageMaker Studio console.

View the Details of a Model Version (Boto3)

To view the details of a model version by using Boto3, complete the following steps.

1. Call the `list_model_packages` API operation to view the model versions in a Model Group.

   ```python
   sm_client.list_model_packages(ModelPackageName="ModelGroup1")
   ```

   The response is a list of model package summaries. You can get the Amazon Resource Name (ARN) of the model versions from this list.

   ```json
   [{'ModelPackageName': 'AbaloneMPG-16039329888329896',
     'ModelVersion': 1,
     'ModelArn': 'arn:aws:sagemaker:us-east-2:123456789012:model-package/ModelGroup1/1',
     'ModelDescription': 'TestMe',
     'CreationTime': datetime.datetime(2020, 10, 29, 1, 27, 46, 46000, tzinfo=tzlocal()),
     'ModelStatus': 'Completed',
     'ModelApprovalStatus': 'Approved'}]
   ```

2. Call `describe_model_package` to see the details of the model version. You pass in the ARN of a model version that you got in the output of the call to `list_model_packages`.

   ```python
   sm_client.describe_model_package(ModelPackageName="arn:aws:sagemaker:us-east-2:123456789012:model-package/ModelGroup1/1")
   ```
The output of this call is a JSON object with the model version details.

```json
{"ModelPackageGroupName": 'ModelGroup1',
 'ModelPackageVersion': 1,
 ModelGroup/1',
 'ModelPackageDescription': 'Test Model',
 'CreationTime': datetime.datetime(2020, 10, 29, 1, 27, 46, 46000,
 tzinfo=tzlocal()),
 'InferenceSpecification': {'Containers': [{'Image': '257758044811.dkr.ecr.us-
 east-2.amazonaws.com/sagemaker-xgboost:1.0-1-cpu-py3',
 'ImageDigest': 'sha256:99fa602cff19aee33297a5926f8497ca7bcd2a391b7d600300204eef803bca66',
 'ModelDataUrl': 's3://sagemaker-us-east-2.amazonaws.com/sagemaker-xgboost/1.0-1-cpu-py3/model.tar.gz'}],
 'SupportedTransformInstanceTypes': ['ml.m5.xlarge'],
 'SupportedRealtimeInferenceInstanceTypes': ['ml.t2.medium', 'ml.m5.xlarge'],
 'SupportedContentTypes': ['text/csv'],
 'SupportedResponseMIMETypes': ['text/csv'],
 'ModelPackageStatus': 'Completed',
 'ModelPackageStatusDetails': {'ValidationStatuses': [],
 'ImageScanStatuses': []},
 'CertifyForMarketplace': False,
 'ModelApprovalStatus': 'PendingManualApproval',
 'LastModifiedTime': datetime.datetime(2020, 10, 29, 1, 28, 0, 438000,
 tzinfo=tzlocal()),
 'ResponseMetadata': {'RequestId': '12345678-abcd-1234-abcd-aabbccddeeff',
 'HTTPStatusCode': 200,
 'HTTPHeaders': {'x-amzn-requestid': '212345678-abcd-1234-abcd-aabbccddeeff',
 'content-type': 'application/x-amz-json-1.1',
 'content-length': '1038',
 'date': 'Mon, 23 Nov 2020 04:59:38 GMT'},
 'RetryAttempts': 0}}
```

**View the Details of a Model Version (console)**

To view the details of a model version in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.
1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.

2. In the left navigation pane, choose Models from the menu.

3. Choose the Registered models tab, if not selected already.

4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.

5. Select the name of the model group containing the model version to view.

6. In the list of model versions, select the model version to view.

7. To view details related to model training, choose the Training radio button. To view details related to inference, choose the Inference radio button.

The following tabs include the details related to model training:

- **Performance**: Statistical measurements to assess model performance, such as relative mean error (RME).
- **Evaluation**: Charts and metrics to describe bias and explainability.
- **Associations**: The resources that derived, are derived from, or are associated with the model version.
- **Activity**: The actions you performed with the model version, such as approval.
- **Tags**: The tags that belong to the model version.
- **Metadata**: The ARN information for model version and associated Identity and Access Management (IAM) roles.

The following tabs include details related to model inference:

- **Instances**: The instances on which your model is deployed.
- **Metadata**: The containers running inference with your model.

### Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the **Home** icon (  
\[home icon\]).

3. Choose **Models**, and then **Model registry**.

4. From the model groups list, select the name of the Model Group you want to view.

5. A new tab appears with a list of the model versions in the Model Group.

6. In the list of model versions, select the name of the model version for which you want to view details.

7. On the model version tab that opens, choose one of the following to see details about the model version:

   - **Activity**: Shows events for the model version, such as approval status updates.
   
   - **Model quality**: Reports metrics related to your Model Monitor model quality checks, which compare model predictions to Ground Truth. For more information about Model Monitor model quality checks, see [Monitor model quality](#).
   
   - **Explainability**: Reports metrics related to your Model Monitor feature attribution checks, which compare the relative rankings of your features in training data versus live data. For more information about Model Monitor explainability checks, see [Monitor Feature Attribution Drift for Models in Production](#).
   
   - **Bias**: Reports metrics related to your Model Monitor bias drift checks, which compare the distribution of live data to training data. For more information about Model Monitor bias drift checks, see [Monitor Bias Drift for Models in Production](#).
   
   - **Inference recommender**: Provides initial instance recommendations for optimal performance based on your model and sample payloads.
   
   - **Load test**: Runs load tests across your choice of instance types when you provide your specific production requirements, such as latency and throughput constraints.
   
   - **Inference specification**: Displays instance types for your real-time inference and transform jobs, and information about your Amazon ECR containers.
   
   - **Information**: Shows information such as the project with which the model version is associated, the pipeline that generated the model, the Model Group, and the model's location in Amazon S3.
Compare Model Versions

As you generate model versions, you might want to compare models versions by viewing relevant model quality metrics side-by-side. For example, you might want to track accuracy by comparing mean squared error (MSE) values, or you might decide to remove models that perform poorly on selected measures. The following procedure shows you how to set up model version comparison in Model Registry using the Amazon SageMaker Studio Classic console.

Compare Model Versions (Amazon SageMaker Studio Classic)

Note

You can only compare model versions the Amazon SageMaker Studio Classic console.

To compare model versions within a model group, complete the following steps:

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the left navigation pane, choose the Home icon (▲).
3. Choose Models, and then Model registry.
4. From the model groups list, select the name of the Model Group you want to view. A new tab opens with a list of the model versions in the Model Group.
5. In the list of model versions, check the boxes next to the model versions you want to compare.
6. Choose the Actions dropdown menu, then choose Compare. A listing of model quality metrics appears for your selected models.

View and Manage Model Group and Model Version Tags

Model Registry helps you view and manage tags related to your model groups. You can use tags to categorize model groups by purpose, owner, environment, or other criteria. The following instructions show you how to view, add, delete, and edit your tags in the Amazon SageMaker Studio console.
View and manage model group tags

Studio

To view a model group tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models to display a list of your model groups.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the Model Groups list, select the name of the Model Group you want to view.
6. In the model group page, choose the Tags tab. View the tags associated with your model group.

To add a model group tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models to display a list of your model groups.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the Model Groups list, select the name of the Model Group you want to edit.
6. In the model group page, choose the Tags tab.
7. Choose Add/Edit tags.
8. Above + Add new tag, enter your new key in the blank Key field.
9. (Optional) Enter your new value in the blank Value field.
10. Choose Confirm changes.
11. Confirm your new tag appears in the Tags section of the Information page.
To delete a model group tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models to display a list of your model groups.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the Model Groups list, select the name of the Model Group you want to edit.
6. In the model group page, choose the Tags tab.
7. Choose Add/Edit tags.
8. Choose the Trash icon next to the key-value pair you want to remove.
9. Choose Confirm changes.

To edit a model group tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models to display a list of your model groups.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the Model Groups list, select the name of the Model Group you want to edit.
6. In the model group page, choose the Tags tab.
7. Choose Add/Edit tags.
8. Enter a new value in the Value field of the key-pair you want to edit.
9. Choose Confirm changes.

Studio Classic

To view a model group tag, complete the following steps:

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the Home icon (🏠).
3. Choose Models, and then Model registry.
4. From the Model Groups list, select the name of the Model Group you want to edit.
5. Choose Information.
6. View your tags in the Tags section of the Information page.

To add a model group tag, complete the following steps:

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the left navigation pane, choose the Home icon (🏠).
3. Choose Models, and then Model registry.
4. From the Model Groups list, select the name of the Model Group you want to edit.
5. Choose Information.
6. If you don't have any tags, choose Add tags.
7. If you have pre-existing tags, choose Manage tags in the Tags section. A list of the model group's tags appears as key-value pairs.
8. Above Add new tag, enter your new key in the blank Key field.
9. (Optional) Enter your new value in the blank Value field.
10. Choose Confirm changes.
11. Confirm your new tag appears in the Tags section of the Information page.

To delete a model group tag, complete the following steps:

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the left navigation pane, choose the **Home** icon (  
   ![Home icon]  
).
3. Choose **Models**, and then **Model registry**.
4. From the Model Groups list, select the name of the Model Group you want to edit.
5. Choose **Information**.
6. In the **Tags** section, choose **Manage tags**. A list of the model group's tags appears as key-value pairs.
7. Choose the **Trash** icon to the right of the tag you want to remove.
8. Choose **Confirm changes**.
9. Confirm your removed tag does not appear in the **Tags** section of the **Information** page.

**To edit a model group tag, complete the following steps:**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Amazon SageMaker domain overview](#).
2. In the left navigation pane, choose the **Home** icon (  
   ![Home icon]  
).
3. Choose **Models**, and then **Model registry**.
4. From the Model Groups list, select the name of the Model Group you want to edit.
5. Choose **Information**.
6. In the **Tags** section, choose **Manage tags**. A list of the model group's tags appears as key-value pairs.
7. Edit any key or value.
8. Choose **Confirm changes**.
9. Confirm your tag contains your edits in the **Tags** section of the **Information** page.

**Manage model version tags**

To manage model version tags in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.
To add a model version tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models from the menu.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. Select the name of the model group containing the model version to update.
6. In the list of model versions, select the model version to update.
7. Under the Actions dropdown menu, choose Edit.
8. If you are adding your first tag, enter your new key-value pair in the blank Key and Value fields.
9. (Optional) To add another custom metadata key-value pair, choose Add New and enter your new key-value pair in the blank Name and Value fields.
10. Choose Save changes.

To delete a model version tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models from the menu.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. Select the name of the model group containing the model version to update.
6. In the list of model versions, select the model version to update.
7. Under the Actions dropdown menu, choose Edit.
8. Choose the Trash icon to the right of the tag that you want to remove.
9. Choose Confirm changes.
To edit a model version tag, complete the following steps:

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models from the menu.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. Select the name of the model group containing the model version to update.
6. In the list of model versions, select the model version to update.
7. Under the Actions dropdown menu, choose Edit.
8. Edit any key or value.
9. Choose Confirm changes.

Studio Classic

To add a model version tag, complete the following steps:

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the Home icon (🏠).
3. Choose Models, and then Model registry.
4. From the model groups list, select the name of the Model Group you want to view. A new tab opens with a list of the model versions in the Model Group.
5. In the list of model versions, select the name of the model version you want to update.
6. Under the Actions dropdown menu, choose Edit.
7. If you are adding your first tag, enter your new key-value pair in the blank Key and Value fields.
8. (Optional) To add another custom metadata key-value pair, choose Add New and enter your new key-value pair in the blank Name and Value fields.
9. Choose Save changes.
10. Confirm your new tag appears in the **Tags** section of the **Information** page.

**To delete a model version tag, complete the following steps:**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
2. In the left navigation pane, choose the **Home** icon (≡).
3. Choose **Models**, and then **Model registry**.
4. From the model groups list, select the name of the Model Group you want to view. A new tab opens with a list of the model versions in the Model Group.
5. In the list of model versions, select the name of the model version you want to update.
6. Under the **Actions** dropdown menu, choose **Edit**.
7. Choose the **Trash** icon to the right of the tag that you want to remove.
8. Choose **Confirm changes**.
9. Confirm your removed tag does not appear in the **Tags** section of the **Information** page.

**To edit a model version tag, complete the following steps:**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
2. In the left navigation pane, choose the **Home** icon (≡).
3. Choose **Models**, and then **Model registry**.
4. From the model groups list, select the name of the Model Group that you want to view. A new tab opens with a list of the model versions in the Model Group.
5. In the list of model versions, select the name of the model version you want to update.
6. Under the **Actions** dropdown menu, choose **Edit**.
7. Edit any key or value.
8. Choose **Confirm changes**.
9. Confirm your tag contains your edits in the **Tags** section of the **Information** page.
Share Models with SageMaker Canvas Users

**Note**
You can only share models with SageMaker Canvas in the Amazon SageMaker Studio Classic console.

You might have a model registered in your Model Registry that you want to share with a user in SageMaker Canvas. You can share a model that's been trained outside of SageMaker as long as it's registered in your Model Registry. With this functionality, SageMaker Canvas users can import models that you've trained and generate predictions with them. For more information about how to share a model with a SageMaker Canvas user, see [Bring your own model to SageMaker Canvas](#).

**Delete a Model Version**

This procedure demonstrates how to delete a model version in the Amazon SageMaker Studio console.

**Delete a Model Version (console)**

To delete a model version in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](#).
2. In the left navigation pane, choose **Models** to display a list of your model groups.
3. Choose the **Registered models** tab, if not selected already.
4. Immediately below the **Registered models** tab label, choose **Model Groups**, if not selected already.
5. From the model groups list, choose the angle bracket to the left of the model group that you want to view.
6. A list of the model versions in the model group appears. If you don't see the model version that you want to delete, choose **View all**.
7. Select the check boxes next to the model versions that you want to delete.
8. Choose the vertical ellipsis above the top right corner of the table, and choose **Delete** (or **Delete model version** if you are in the model group details page).

9. In the **Delete model version** dialog box, choose **Yes, delete the model version**.

10. Choose **Delete**.

11. Confirm that your deleted model versions no longer appear in the model group.

**Studio Classic**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).

2. In the left navigation pane, choose the **Home** icon ( ![Home](#) ).

3. Choose **Models**, and then **Model registry**. A list of your Model Groups appears.

4. From the model groups list, select the name of the Model Group of the model version that you want to delete.

5. From the list of model versions, select the name of the model version that you want to delete.

6. Choose the **Actions** dropdown menu, and choose **Remove**.

7. In the confirmation dialog box, enter **REMOVE**.

8. Choose **Remove**.

9. Confirm the model version you removed does not appear in the list of the model group's model versions.

**Update the Approval Status of a Model**

After you create a model version, you typically want to evaluate its performance before you deploy it to a production endpoint. If it performs to your requirements, you can update the approval status of the model version to **Approved**. Setting the status to **Approved** can initiate CI/CD deployment for the model. If the model version does not perform to your requirements, you can update the approval status to **Rejected**.
You can manually update the approval status of a model version after you register it, or you can create a condition step to evaluate the model when you create a SageMaker pipeline. For information about creating a condition step in a SageMaker pipeline, see Pipeline Steps.

When you use one of the SageMaker provided project templates and the approval status of a model version changes, the following action occurs. Only valid transitions are shown.

- PendingManualApproval to Approved – initiates CI/CD deployment for the approved model version
- PendingManualApproval to Rejected – No action
- Rejected to Approved – initiates CI/CD deployment for the approved model version
- Approved to Rejected – initiates CI/CD to deploy the latest model version with an Approved status

You can update the approval status of a model version by using the AWS SDK for Python (Boto3) or by using the Amazon SageMaker Studio console. You can also update the approval status of a model version as part of a condition step in a SageMaker pipeline. For information about using a model approval step in a SageMaker pipeline, see SageMaker Pipelines Overview.

**Update the Approval Status of a Model (Boto3)**

When you created the model version in Register a Model Version, you set the ModelApprovalStatus to PendingManualApproval. You update the approval status for the model by calling update_model_package. Note that you can automate this process by writing code that, for example, sets the approval status of a model depending on the result of an evaluation of some measure of the model's performance. You can also create a step in a pipeline that automatically deploys a new model version when it is approved. The following code snippet shows how to manually change the approval status to Approved.

```python
model_package_update_input_dict = {
    "ModelPackageArn" : model_package_arn,
    "ModelApprovalStatus" : "Approved"
}
model_package_update_response = 
    sm_client.update_model_package(**model_package_update_input_dict)
```
Update the Approval Status of a Model (console)

To manually change the approval status in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

Studio

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose the Models to display a list of your model groups.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the model groups list, choose the angle bracket to the left of the model group that you want to view.
6. A list of the model versions in the model group appears. If you don't see the model version that you want to delete, choose View all to display the complete list of model versions in the model group details page.
7. Select the name of the model version that you want to update.
8. Choose the vertical ellipsis at the top right, choose Update status, and then the final model status.
9. In the Update model status dialog box, insert an optional comment and choose Save and update.

Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the Home icon (🏠).
3. Choose Models, and then Model registry.
4. From the model groups list, select the name of the Model Group that you want to view. A new tab opens with a list of the model versions in the Model Group.
5. In the list of model versions, select the name of the model version that you want to update.

6. Under the **Actions** dropdown menu, you can choose one of two possible menu options to update the model version status.

- Using the **Update Status** option
  1. Under the **Actions** dropdown menu, choose the **Update Status** dropdown menu, and choose the new model version status.
  2. (Optional) In the **Comment** field, add additional details.
  3. Choose **Save and Update**.

- Using the **Edit** option
  1. Under the **Actions** dropdown menu, choose **Edit**.
  2. (Optional) In the **Comment** field, add additional details.
  3. Choose **Save changes**.

7. Confirm the model version status is updated to the correct value in the model version page.

### Deploy a Model from the Registry

After you register a model version and approve it for deployment, deploy it to a SageMaker endpoint for real-time inference. You can deploy your model by using the SageMaker SDK or the AWS SDK for Python (Boto3) (Boto3).

When you create a machine learning operations (MLOps) project and choose an MLOps project template that includes model deployment, approved model versions in the Model Registry are automatically deployed to production. For information about using SageMaker MLOps projects, see [Automate MLOps with SageMaker Projects](#).

You can also enable an AWS account to deploy model versions that were created in a different account by adding a cross-account resource policy. For example, one team in your organization might be responsible for training models, and a different team is responsible for deploying and updating models.

### Topics

- [Deploy a Model from the Registry (SageMaker SDK)](#)
- [Deploy a Model from the Registry (Boto3)](#)
- [Deploy a Model Version from a Different Account](#)
Deploy a Model from the Registry (SageMaker SDK)

To deploy a model version using the Amazon SageMaker Python SDK use the following code snippet:

```python
from sagemaker import ModelPackage
from time import gmtime, strftime

model_package_arn = 'arn:aws:sagemaker:us-east-2:12345678901:model-package/modeltest/1'
model = ModelPackage(role=role,
                     model_package_arn=model_package_arn,
                     sagemaker_session=sagemaker_session)
model.deploy(initial_instance_count=1, instance_type='ml.m5.xlarge')
```

Deploy a Model from the Registry (Boto3)

To deploy a model version using the AWS SDK for Python (Boto3), complete the following steps:

1. The following code snippet assumes you already created the SageMaker Boto3 client `sm_client` and a model version whose ARN is stored in the variable `model_version_arn`.

   Create a model object from the model version by calling the `create_model` API operation. Pass the Amazon Resource Name (ARN) of the model version as part of the `Containers` for the model object:

   ```python
   model_name = 'DEMO-modelregistry-model-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
   print("Model name : {}".format(model_name))
   container_list = [{'ModelPackageName': model_version_arn}]

   create_model_response = sm_client.create_model(
       ModelName = model_name,
       ExecutionRoleArn = role,
       Containers = container_list
   )
   print("Model arn : {}".format(create_model_response['ModelArn']))
   ```

2. Create an endpoint configuration by calling `create_endpoint_config`. The endpoint configuration specifies the number and type of Amazon EC2 instances to use for the endpoint.

   ```python
   endpoint_config_name = 'DEMO-modelregistry-EndpointConfig-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
   print(endpoint_config_name)
   ```
3. Create the endpoint by calling `create_endpoint`.

```python
# Create endpoint configuration
create_endpoint_config_response = sm_client.create_endpoint_config(
    EndpointConfigName = endpoint_config_name,
    ProductionVariants=[[
        'InstanceType':'ml.m4.xlarge',
        'InitialVariantWeight':1,
        'InitialInstanceCount':1,
        'ModelName':model_name,
        'VariantName':'AllTraffic']])

# Create endpoint
endpoint_name = 'DEMO-modelregistry-endpoint-' + strftime("%Y-%m-%d-%H-%M-%S",
    gmtime())
print("EndpointName={}".format(endpoint_name))
create_endpoint_response = sm_client.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name)
print(create_endpoint_response['EndpointArn'])
```

### Deploy a Model Version from a Different Account

You can permit an AWS account to deploy model versions that were created in a different account by adding a cross-account resource policy. For example, one team in your organization might be responsible for training models, and a different team is responsible for deploying and updating models. When you create these resource policies, you apply the policy to the specific resource to which you want to grant access. For more information about cross-account resource policies in AWS, see [Cross-account policy evaluation logic](https://docs.aws.amazon.com/IdentityAndAccessManagement/latest/UserGuide/cross-account-policy-eval-logic.html) in the *AWS Identity and Access Management User Guide*.

⚠️ **Note**

You must use a KMS key to encrypt the [output data config](https://docs.aws.amazon.com/AmazonSageMaker/latest/DG/how-to-deploy-endpoints-with-sagemaker-sdk.html) action during training for cross-account model deployment.

To enable cross-account model deployment in SageMaker, you have to provide a cross-account resource policy for the Model Group that contains the model versions you want to deploy, the Amazon ECR repository where the inference image for the Model Group resides, and the Amazon S3 bucket where the model versions are stored.
To be able to deploy a model that was created in a different account, you must have a role that has access to SageMaker actions, such as a role with the AmazonSageMakerFullAccess managed policy. For information about SageMaker managed policies, see AWS Managed Policies for Amazon SageMaker.

The following example creates cross-account policies for all three of these resources, and applies the policies to the resources. The example also assumes that you previously defined the following variables:

- **bucket** – The Amazon S3 bucket where the model versions are stored.
- **kms_key_id** – The KMS key used to encrypt the training output.
- **sm_client** – A SageMaker Boto3 client.
- **model_package_group_name** – The Model Group to which you want to grant cross-account access.
- **model_package_group_arn** – The Model Group ARN to which you want to grant cross-account access.

```python
import json

cross_account_id = "123456789012"

ecr_repository_policy = {
    'Version': '2012-10-17',
    'Statement': [
        {'Sid': 'AddPerm',
         'Effect': 'Allow',
         'Principal': {
             'AWS': f'arn:aws:iam::{cross_account_id}:root'
         },
         'Action': ['ecr:*']
    ]
}

ecr_repository_policy = json.dumps(ecr_repository_policy)

# Set the new ECR policy
ecr = boto3.client('ecr')
response = ecr.set_repository_policy(
    registryId = account,
    repositoryName = 'decision-trees-sample',
    policyText = ecr_repository_policy
)

# Create a policy for accessing the S3 bucket
bucket_policy = {
    'Version': '2012-10-17',
    'Statement': [
        {'Sid': 'AddPerm',
         'Effect': 'Allow',
         'Principal': {
             'AWS': f'arn:aws:iam::{cross_account_id}:root'
         },
         'Action': 's3:*',
         'Resource': f'arn:aws:s3:::{bucket}/*'
        }
    ]
}

# Convert the policy from JSON dict to string
bucket_policy = json.dumps(bucket_policy)

# Set the new policy
s3 = boto3.client('s3')
response = s3.put_bucket_policy(
    Bucket = bucket,
    Policy = bucket_policy)

# Create the KMS grant for encryption in the source account to the 
# Model Registry account Model Group
client = boto3.client('kms')

response = client.create_grant(
    GranteePrincipal=cross_account_id,
    KeyId=kms_key_id
    Operations=[
        'Decrypt',
        'GenerateDataKey',
    ],
)

# 3. Create a policy for access to the Model Group.
```python
model_package_group_policy = {
    'Version': '2012-10-17',
    'Statement': [
        {'Sid': 'AddPermModelPackageGroup',
         'Effect': 'Allow',
         'Principal': {
            'AWS': f'arn:aws:iam::{cross_account_id}:root'
         },
         'Action': ['sagemaker:DescribeModelPackageGroup'],
         'Resource': f'arn:aws:sagemaker:{region}:{account}:model-package-group/{model_package_group_name}'
        },
        {'Sid': 'AddPermModelPackageVersion',
         'Effect': 'Allow',
         'Principal': {
            'AWS': f'arn:aws:iam::{cross_account_id}:root'
         },
         'Action': ['sagemaker:DescribeModelPackage',
                    'sagemaker:ListModelPackages',
                    'sagemaker:UpdateModelPackage',
                    'sagemaker:CreateModel'],
         'Resource': f'arn:aws:sagemaker:{region}:{account}:model-package/{model_package_group_name}/*'
        }
    ]
}

# Convert the policy from JSON dict to string
model_package_group_policy = json.dumps(model_package_group_policy)

# Set the policy to the Model Group
response = sm_client.put_model_package_group_policy(
    ModelPackageName = model_package_group_name,
    ResourcePolicy = model_package_group_policy)

print('ModelPackageGroupArn : ' + create_model_package_group_response['ModelPackageGroupArn'])
print("First Versioned ModelPackageArn: " + model_package_arn)
print("Second Versioned ModelPackageArn: " + model_package_arn2)
print("Success! You are all set to proceed for cross-account deployment.")
```
View the Deployment History of a Model

To view the deployments for a model version in the Amazon SageMaker Studio console, complete the following steps based on whether you use Studio or Studio Classic.

Studio

View the deployment history for a model version

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models to display a list of your model groups.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Model Groups, if not selected already.
5. From the model groups list, choose the angle bracket to the left of the model group that you want to view.
6. A list of the model versions in the model group appears. If you don't see the model version that you want to delete, choose View all.
7. Select the name of the model version that you want to view.
8. Choose the Activity tab. Deployments for the model version appear as events in the activity list with an Event type of ModelDeployment.

Studio Classic

View the deployment history for a model version

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.
2. In the left navigation pane, choose the Home icon ( ).
3. Choose Models, and then Model registry.
4. From the model groups list, select the name of the Model Group that you want to view.
5. A new tab appears with a list of the model versions in the Model Group.
6. In the list of model versions, select the name of the model version for which you want to view details.

7. On the model version tab that opens, choose Activity. Deployments for the model version appear as events in the activity list with an Event type of ModelDeployment.

Model Registry Collections

You can use Collections to group registered models that are related to each other and organize them in hierarchies to improve model discoverability at scale. With Collections, you can organize registered models that are associated with one another. For example, you could categorize your models based on the domain of the problem they solve as Collections titled NLP-models, CV-models, or Speech-recognition-models. To organize your registered models in a tree structure, you can nest Collections within each other. Any operations you perform on a Collection, such as create, read, update, or delete, will not alter your registered models. You can use the Amazon SageMaker Studio UI or the Python SDK to manage your Collections.

The Collections tab in the Model Registry displays a list of all the Collections in your account. The following sections describe how you can use options in the Collections tab to do the following:

- Create Collections
- Add Model Groups to a Collection
- Move Model Groups between Collections
- Remove Model Groups or Collections from other Collections

Any operation you perform on your Collections does not affect the integrity of the individual Model Groups they contain—the underlying Model Group artifacts in Amazon S3 and Amazon ECR are not modified.

While Collections provide greater flexibility in organizing your models, the internal representation imposes some constraints on the size of your hierarchy. For a summary of these constraints, see Constraints.

The following topics show you how to create and work with Collections in the Model Registry.

**Topics**

- Prerequisites
- Create a Collection
• Add Model Groups to a Collection
• Remove Model Groups or Collections from a Collection
• Move a Model Group Between Collections
• View a Model Group's Parent Collection

Constraints

Prerequisites

Create a custom policy which includes the following required Resource Groups actions:

• resource-groups:CreateGroup
• resource-groups:DeleteGroup
• resource-groups:GetGroupQuery
• resource-groups:ListGroupResources
• resource-groups:Tag
• tag:GetResources

For instructions on how to add an inline policy, see Adding IAM identity permissions (console). When you choose the policy format, choose the JSON format and add the following policy:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "resource-groups:ListGroupResources"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "resource-groups:GetGroupQuery"
            ],
            "Resource": "arn:aws:resource-groups::*:*:group/*"
        }
    ]
}
```
Create a Collection

You can create a Collection in the SMMonarchlong; console. To create a Collection, complete the following steps based on whether you use Studio or Studio Classic.

Studio

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the **Registered models** tab label, choose **Collections**.

5. (Optional) To create a Collection inside another Collection, navigate to the hierarchy where you want to add your Collection. Otherwise, your Collection is created at the root level.

6. In the **Actions** dropdown menu in the top right, choose **Create new collection**.

7. Enter a name for your Collection in the **Name** field of the dialog box.

8. (Optional) To add Model Groups to your Collection, complete the following steps:
   
   a. Choose **Select model groups**.
   
   b. Select the Model Groups that you want to add. You can select up to 10.

9. Choose **Create**.

10. Check to make sure your Collection was created in the current hierarchy. If you do not immediately see your new Collection, choose **Refresh**.

**Studio Classic**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).

2. In the left navigation pane, choose the **Home** icon ( ).

3. Choose **Models**, and then **Model registry**.

4. Choose the **Collections** tab.

5. (Optional) To create a Collection inside another Collection, navigate to the hierarchy where you want to add your Collection. Otherwise, your Collection is created at the root level.

6. In the **Actions** dropdown menu in the top right, choose **Create new collection**.

7. Enter a name for your Collection in the **Name** field of the dialog box.
8. (Optional) To add Model Groups to your Collection, complete the following steps:
   a. Choose Select model groups.
   b. Select the Model Groups that you want to add. You can select up to 10.
9. Choose Create.
10. Check to make sure your Collection was created in the current hierarchy. If you do not immediately see your new Collection, choose Refresh.

Add Model Groups to a Collection

You can add model groups to a Collection in the Amazon SageMaker Studio console. To add Model Groups to a Collection, complete the following steps based on whether you use Studio or Studio Classic.

Studio

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.
2. In the left navigation pane, choose Models.
3. Choose the Registered models tab, if not selected already.
4. Immediately below the Registered models tab label, choose Models, if not selected already.
5. Select the check box next to the model groups that you want to add. You can select up to 10 Model Groups. If you select more than 10, the UI option to add your Model Groups to a Collection is inactive.
6. Choose the vertical ellipses next to Create, and choose Add to collection.
7. Select the radio button for the collection to which you want to add your selected Model Groups.
8. Choose **Add to collection**.
9. Check to make sure your Model Groups were added in to the collection. In the **Collections** column of the Model Groups you selected, you should see the name of collection to which you added the Model Groups.

**Studio Classic**

You can add Model Groups to a Collection from either the **Model Groups** or **Collections** tab.

To add one or more Model Groups to a Collection from the **Collections** tab, complete the following steps:

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
2. In the left navigation pane, choose the **Home** icon (🏠).
3. Choose **Models**, and then **Model registry**.
4. Choose the **Collections** tab.
5. Select the Collection to which you want to add Model Groups. If the desired Collection is not at root level, navigate to the hierarchy where you want to add your Model Groups.
6. In the **Actions** dropdown menu in the top right, choose **Add model groups**.
7. Select the Model Groups that you want to add. You can select up to 10 Model Groups. If you select more than 10, the UI option to add your Model Groups to a Collection is inactive.
8. Choose **Add to collection**.
9. Check to make sure your Model Groups were added in the current hierarchy. If you do not immediately see your new Model Groups, choose **Refresh**.

To add one or more Model Groups to a Collection from the **Model Groups** tab, complete the following steps:

1. Sign in to Studio Classic. For more information, see [Amazon SageMaker domain overview](#).
2. In the left navigation pane, choose the **Home** icon (🏠).
3. Choose **Models**, and then **Model registry**.

4. Choose the **Model Groups** tab.

5. Select the Model Groups that you want to add. You can select up to 10. If you select more than 10, the UI option to add your Model Groups to a Collection is inactive.

6. In the **Actions** dropdown menu in the top right, choose **Add to collection**.

7. In the pop-up dialog, choose the root path location **Collections**. This link to the root location appears above the table.

8. Navigate to the hierarchy which contains your destination Collection, or where you want to create a new Collection to which you add your models.

9. (Optional) To add your Model Groups to an existing Collection, complete the following steps:
   
   a. Select the destination Collection.
   
   b. Choose **Add to collection**.

10. (Optional) To add your Model Groups to a new Collection, complete the following steps:
   
   a. Choose **New collection**.
   
   b. Enter a name for your new Collection.
   
   c. Choose **Create**.

**Remove Model Groups or Collections from a Collection**

When you remove Model Groups or Collections from a Collection, you are removing them from a particular grouping and not from the Model Registry. You can remove Model Groups from a Collection in the Amazon SageMaker Studio console.

To remove one or more Model Groups or Collections from a Collection, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](#).

2. In the left navigation pane, choose **Models**.

3. Choose the **Registered models** tab, if not selected already.
4. Immediately below the **Registered models** tab label, choose **Collections**.

5. Navigate to the Collection which contains the Model Groups or Collections you want to remove.

6. Select the Model Groups or Collections that you want to remove. You can select up to 10. If you select more than 10 Model Groups or Collections, the UI option to remove them is inactive.

**Important**

You cannot simultaneously select Model Groups and Collections for removal. To remove both Model Groups and Collections, first remove Model Groups, and then remove Collections.

**Important**

You cannot remove non-empty Collections. To remove a non-empty Collection, first remove its contents.

7. In the **Actions** dropdown menu in the top right, choose **Remove X items from collection** (where X is the number of Model Groups that you selected).

8. Confirm that you want to remove the selected Model Groups.

**Studio Classic**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).

2. In the left navigation pane, choose the **Home** icon (  

3. Choose **Models**, and then **Model registry**.

4. Choose the **Collections** tab.

5. Navigate to the Collection which contains the Model Groups or Collections you want to remove.
6. Select the Model Groups or Collections that you want to remove. You can select up to 10. If you select more than 10 Model Groups or Collections, the UI option to remove them is inactive.

⚠️ Important
You cannot simultaneously select Model Groups and Collections for removal. To remove both Model Groups and Collections, first remove Model Groups, and then remove Collections.

⚠️ Important
You cannot remove non-empty Collections. To remove a non-empty Collection, first remove its contents.

7. In the Actions dropdown menu in the top right, choose Remove X items from collection (where X is the number of Model Groups you selected).

8. Confirm that you want to remove the selected Model Groups.

Move a Model Group Between Collections

You can move one or more Model Groups from one Collection to another in the Amazon SageMaker Studio console.

To move Model Groups, complete the following steps based on whether you use Studio or Studio Classic.

Studio

1. Open the SageMaker Studio console by following the instructions in Launch Amazon SageMaker Studio.

2. In the left navigation pane, choose Models.

3. Choose the Registered models tab, if not selected already.

4. Immediately below the Registered models tab label, choose Collections.

5. Navigate to the Collection which contains the Model Groups that you want to move.
6. Select the Model Groups that you want to move. You can select up to 10. If you select more than 10, the UI option to move your Model Groups is inactive.

7. In the Actions dropdown menu in the top right, choose Move to.

8. In the dialog box, choose the root path location Collections. This link to the root location appears above the table.

9. Navigate to the hierarchy which contains your destination Collection.

10. Select your destination Collection in the table.


Studio Classic

1. Sign in to Amazon SageMaker Studio Classic. For more information, see Launch Amazon SageMaker Studio Classic.

2. In the left navigation pane, choose the Home icon (🏠).

3. Choose Models, and then Model registry.

4. Choose the Collections tab.

5. Navigate to the Collection which contains the Model Groups that you want to move.

6. Select the Model Groups that you want to move. You can select up to 10. If you select more than 10, the UI option to move your Model Groups is inactive.

7. In the Actions dropdown menu in the top right, choose Move to.

8. In the dialog box, choose the root path location Collections. This link to the root location appears above the table.

9. Navigate to the hierarchy which contains your destination Collection.

10. Select your destination Collection in the table.


View a Model Group's Parent Collection

You can view the Collections which contain a particular Model Group in the Amazon SageMaker Studio console.
To view the Collections which contain a particular Model Group, complete the following steps based on whether you use Studio or Studio Classic.

**Studio**

1. Open the SageMaker Studio console by following the instructions in [Launch Amazon SageMaker Studio](#).
2. In the left navigation pane, choose **Models**.
3. Choose the **Registered models** tab, if not selected already.
4. Immediately below the **Registered models** tab label, choose **Model Groups**, if not selected already.
5. View the **Collection** column for your Model Group, which displays the name of the Collection which contains this Model Group. If multiple Collections contain this Model Group, choose the **Collection** column entry to display a pop-up listing the Collections which contain this Model Group.

**Studio Classic**

1. Sign in to Amazon SageMaker Studio Classic. For more information, see [Launch Amazon SageMaker Studio Classic](#).
2. In the left navigation pane, choose the **Home** icon (🏠).
3. Choose **Models**, and then **Model registry**.
4. Choose the **Model Groups** tab.
5. Find your Model Group in the table.
6. View the **Collection** column for your Model Group, which displays the name of the Collection which contains this Model Group. If multiple Collections contain this Model Group, choose the **Collection** column entry to display a pop-up listing the Collections which contain this Model Group.
Constraints

While using Collections, you may face issues related to tag length constraints or rate limits for Collection operations. Review the following list of caveats so you can avoid issues related to these limitations when you work with your Collections.

VPC constraints

- Collections are not supported in VPC mode.

Collection operation constraints

- You can add a maximum of 10 Model Groups to a Collection at a time.
- You can remove a maximum of 10 Model Groups from a Collection at a time.
- You can move a maximum of 10 Model Groups from one Collection to another at a time.
- You cannot delete a Collection unless it is empty.
- A Model Group can belong to multiple Collections, but a Collection can only belong to one Collection.

Tag-related constraints

- A Model Group can belong to a maximum of 48 Collections. For more details, see the following section Collection and Model Group tagging.
- A Collection’s absolute path can be a maximum of 256 characters long. Since Collection names are user-specified, you can control the path length. For more details, see the following section Collection and Model Group tagging.

Collection and Model Group tagging

The SageMaker Model Registry uses tag rules and tags to internally represent your Collection groupings and hierarchy. You can access these tag elements in the AWS Resource Access Manager, the SageMaker SDK, and the AWS CLI, but it is important that you do not alter or delete them.

⚠️ Important

Do not delete or alter any tag rules or tags which belong to your Collections or Model Groups. Doing so prevents you from performing Collection operations!
A tag rule is a key-value pair that SageMaker uses to identify a Collection's location in the hierarchy. In short, the key is the key of the parent Collection, and the value is the path of the Collection within the hierarchy. SageMaker allows tag values to be 256 characters or less, so if you have multiple nested hierarchies you are advised to keep your Collection names short.

⚠️ Important

Keep your Collection names short. The absolute path to any Collection must be 256 characters long or less.

Model Groups, on the other hand, do not have tag rules but use tags. A Model Group's tags include the tag rules for all the Collections which contain the Model Group. For example, if four Collections contain `model-group-1`, `model-group-1` has four tags. SageMaker allows a single AWS resource to have a maximum of 50 tags. Since two are pre-allocated for general purposes, a Model Group can have a maximum of 48 tags. In conclusion, a Model Group can belong to a maximum of 48 Collections.

Amazon SageMaker Model Registry FAQ

Use the following FAQ items to find answers to commonly asked questions about SageMaker Model Registry.

Q. How should I organize my models into Model Groups and model packages in the SageMaker Model Registry?

A model package is the actual model that is registered into the Model Registry as a versioned entity. Please note there are two ways you can use model packages in SageMaker. One is with SageMaker Marketplace — these model packages are not versioned. The other is with the SageMaker Model Registry, in which the model package must be versioned. The Model Registry receives every new model that you retrain, gives it a version, and assigns it to a Model Group inside the Model Registry. The following image shows an example of a Model Group with 25 consecutively-versioned models.
Q. How does the SageMaker Model Registry differ from Amazon Elastic Container Registry (Amazon ECR)?

The SageMaker Model Registry is a metadata store for your machine learning models. Amazon Elastic Container Registry is a repository that stores all of your containers. Within the Model Registry, models are versioned and registered as model packages within Model Groups. Each model package contains an Amazon S3 URI to the model files associated with the trained model and an Amazon ECR URI that points to the container used while serving the model.

Q. How do I tag model packages in the SageMaker Model Registry?

Model packages in the SageMaker Model Registry do not support tags—these are versioned model packages. Instead, you can add key value pairs using CustomerMetadataProperties. Model package groups in the model registry support tagging.

Q. How should I assign or tag model package groups in the SageMaker Model Registry to a project?

To assign or tag model groups to a project, complete the following steps:
1. Get tags with key `sagemaker:project-name` and `sagemaker:project-id` for the SageMaker project using the `ListTags` API.

2. To apply the tags to your model package group, choose one of the following methods:

   - If you create a new model package group and want to add tags, pass your tags from Step 1 to the `CreateModelPackageGroup` API.
   - If you want to add tags to an existing model package group, use the `AddTags` APIs.
   - If you create your model package group through SageMaker Pipelines, use the `pipeline.create()` or `pipeline.upsert()` methods, or pass your tags to the `RegisterModel` step.

### Model Deployment in SageMaker

Once you train and approve a model for production, use SageMaker to deploy your model to an endpoint for real-time inference. SageMaker provides multiple inference options so that you can pick the option that best suits your workload. You also configure your endpoint by choosing the instance type and number of instances you need for optimal performance. For details about model deployment, see [Deploy models for inference](#).

After you deploy your models to production, you might want to explore ways to further optimize model performance while maintaining availability of your current models. For example, you can set up a shadow test to try out a different model or model serving infrastructure before committing to the change. SageMaker deploys the new model, container, or instance in shadow mode and routes to it a copy of the inference requests in real time within the same endpoint. You can log the responses of the shadow variant for comparison. For details about shadow testing, see [Shadow tests](#). If you decide to go ahead and change your model, deployment guardrails help you control the switch from the current model to a new one. You can select such methods as blue/green or canary testing of the traffic shifting process to maintain granular control during the update. For information about deployment guardrails, see [Update models in production](#).

### SageMaker Model Monitor

Once a model is in production, you can monitor its performance in real time with Amazon SageMaker Model Monitor. Model Monitor helps you maintain model quality by detecting violations of user-defined thresholds for data quality, model quality, bias drift and feature...
attribution drift. In addition, you can configure alerts so you can troubleshoot violations as they arise and promptly initiate retraining. Model Monitor is integrated with SageMaker Clarify to improve visibility into potential bias.

To learn about SageMaker Model Monitor, see Monitor data and model quality.

Automate MLOps with SageMaker Projects

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the Studio Classic application. For information about using the updated Studio experience, see Amazon SageMaker Studio.

Create end-to-end ML solutions with CI/CD by using SageMaker projects.

Use SageMaker projects to create an MLOps solution to orchestrate and manage:

- Building custom images for processing, training, and inference
- Data preparation and feature engineering
- Training models
- Evaluating models
- Deploying models
- Monitoring and updating models

Topics

- What is a SageMaker Project?
- SageMaker Studio Permissions Required to Use Projects
- Create an MLOps Project using Amazon SageMaker Studio Classic
- MLOps Project Templates
- View Project Resources
- Update an MLOps Project in Amazon SageMaker Studio Classic
What is a SageMaker Project?

SageMaker Projects help organizations set up and standardize developer environments for data scientists and CI/CD systems for MLOps engineers. Projects also help organizations set up dependency management, code repository management, build reproducibility, and artifact sharing.

You can provision SageMaker Projects from the AWS Service Catalog using custom or SageMaker-provided templates. For information about the AWS Service Catalog, see What Is AWS Service Catalog. With SageMaker Projects, MLOps engineers and organization admins can define their own templates or use SageMaker-provided templates. The SageMaker-provided templates bootstrap the ML workflow with source version control, automated ML pipelines, and a set of code to quickly start iterating over ML use cases.

When Should You Use a SageMaker Project?

While notebooks are helpful for model building and experimentation, a team of data scientists and ML engineers sharing code needs a more scalable way to maintain code consistency and strict version control.

Every organization has its own set of standards and practices that provide security and governance for its AWS environment. SageMaker provides a set of first-party templates for organizations that want to quickly get started with ML workflows and CI/CD. The templates include projects that use AWS-native services for CI/CD, such as AWS CodeBuild, AWS CodePipeline, and AWS CodeCommit. The templates also offer the option to create projects that use third-party tools, such as Jenkins and GitHub. For a list of the project templates that SageMaker provides, see Use SageMaker-Provided Project Templates.

Organizations often need tight control over the MLOps resources that they provision and manage. Such responsibility assumes certain tasks, including configuring IAM roles and policies, enforcing resource tags, enforcing encryption, and decoupling resources across multiple accounts. SageMaker Projects can support all these tasks through custom template offerings where organizations use AWS CloudFormation templates to define the resources needed for an ML workflow. Data Scientists can choose a template to bootstrap and pre-configure their ML workflow. These custom templates are created as Service Catalog products and you can provision them in the Studio Classic UI under
**Organization Templates.** The Service Catalog is a service that helps organizations create and manage catalogs of products that are approved for use on AWS. For more information about creating custom templates, see [Build Custom SageMaker Project Templates – Best Practices](#).

SageMaker Projects can help you manage your Git repositories so that you can collaborate more efficiently across teams, ensure code consistency, and support CI/CD. SageMaker Projects can help you with the following tasks:

- Organize all entities of the ML lifecycle under one project.
- Establish a single-click approach to set up standard ML infrastructure for model training and deployment that incorporates best practices.
- Create and share templates for ML infrastructure to serve multiple use cases.
- Leverage SageMaker-provided pre-built templates to quickly start focusing on model building, or create custom templates with organization-specific resources and guidelines.
- Integrate with tools of your choice by extending the project templates. For an example, see [Create a SageMaker Project to integrate with GitLab and GitLab Pipelines](#).
- Organize all entities of the ML lifecycle under one project.

**What is in a SageMaker Project?**

Customers have the flexibility to set up their projects with the resources that best serve their use case. The example below showcases the MLOps setup for an ML workflow, including model training and deployment.
A typical project with a SageMaker-provided template might include the following:

- One or more repositories with sample code to build and deploy ML solutions. These are working examples that you can clone locally and modify for your needs. You own this code and can take advantage of the version-controlled repositories for your tasks.

- A SageMaker pipeline that defines steps for data preparation, training, model evaluation, and model deployment, as shown in the following diagram.

![SageMaker Pipelines Diagram](image)

- A CodePipeline or Jenkins pipeline that runs your SageMaker pipeline every time you check in a new version of the code. For information about CodePipeline, see [What is AWS CodePipeline](#). For information about Jenkins, see [Jenkins User Documentation](#).

- A model group that contains model versions. Every time you approve the resulting model version from a SageMaker pipeline run, you can deploy it to a SageMaker endpoint.

Each SageMaker project has a unique name and ID that are applied as tags to all of the SageMaker and AWS resources created in the project. With the name and ID, you can view all of the entities associated with your project. These include:

- Pipelines
- Registered models
- Deployed models (endpoints)
Do I Need to Create a Project to Use SageMaker Pipelines?

No. SageMaker pipelines are standalone entities just like training jobs, processing jobs, and other SageMaker jobs. You can create, update, and run pipelines directly within a notebook by using the SageMaker Python SDK without using a SageMaker project.

Projects provide an additional layer to help you organize your code and adopt operational best practices that you need for a production-quality system.

SageMaker Studio Permissions Required to Use Projects

Users can view SageMaker provided project templates and create projects with those templates when you grant Projects permissions for users. You can grant these permissions when you onboard or update Amazon SageMaker Studio Classic. There are two permissions to grant.

1. Grant Projects permissions for the Studio Classic administrator to permit the Studio Classic administrator to view the SageMaker-provided templates in the Service Catalog console. The administrator can see what other Studio Classic users create if you grant them permission to use SageMaker projects. The administrator can also view the AWS CloudFormation template that the SageMaker-provided project templates define in the Service Catalog console. For information about using the Service Catalog console, see What Is Service Catalog in the Service Catalog User Guide.

2. Allow Studio Classic users who are configured to use the same execution role as the domain to create projects. This grants Studio Classic users permission to use the SageMaker-provided project templates to create a project from within Studio Classic.

⚠️ Important

Do not manually create your roles. Always create roles through Studio Classic Settings using the steps described in the following procedure.
For users who use any role other than the domain's execution role to view and use SageMaker-provided project templates, you need to grant Projects permissions to the individual user profiles.

The following procedures show how to grant Projects permissions after you onboard to Studio Classic. For more information about onboarding to Studio Classic, see Amazon SageMaker domain overview.

To grant Projects permissions for the administrator and domain execution role users

1. Open the SageMaker console.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Choose Create domain.
5. If you choose Quick setup to set up your SageMaker Domain, you have permissions to use project templates by default.
6. If you choose Standard setup to set up your SageMaker Domain, make sure you turn on the following options when you configure Studio Classic settings:
   - Enable Amazon SageMaker project templates and Amazon SageMaker JumpStart for this account
   - Enable Amazon SageMaker project templates and Amazon SageMaker JumpStart for Studio Classic users
7. To confirm that your SageMaker Domain has active project template permissions:
   a. Open the SageMaker console.
   b. On the left navigation pane, choose Admin configurations.
   c. Under Admin configurations, choose domains.
   d. Select your domain.
   e. Choose the Domain Settings tab.
   f. Under SageMaker Projects and JumpStart, make sure the following options are turned on:
      - Enable Amazon SageMaker project templates and Amazon SageMaker JumpStart for this account
      - Enable Amazon SageMaker project templates and Amazon SageMaker JumpStart for Studio Classic users
8. To view a list of your roles:
   a. Open the SageMaker console.
   b. On the left navigation pane, choose Admin configurations.
   c. Under Admin configurations, choose domains.
   d. Select your domain.
   e. Choose the Domain Settings tab.
   f. A list of your roles appears in the Apps card under the Studio tab.

⚠️ Important
As of July 25, we require additional roles to use project templates. Here is the complete list of roles you should see under Projects:

- AmazonSageMakerServiceCatalogProductsLaunchRole
- AmazonSageMakerServiceCatalogProductsUseRole
- AmazonSageMakerServiceCatalogProductsApiGatewayRole
- AmazonSageMakerServiceCatalogProductsCloudformationRole
- AmazonSageMakerServiceCatalogProductsCodeBuildRole
- AmazonSageMakerServiceCatalogProductsCodePipelineRole
- AmazonSageMakerServiceCatalogProductsEventsRole
- AmazonSageMakerServiceCatalogProductsFirehoseRole
- AmazonSageMakerServiceCatalogProductsGlueRole
- AmazonSageMakerServiceCatalogProductsLambdaRole
- AmazonSageMakerServiceCatalogProductsExecutionRole

For descriptions of these roles, see AWS Managed Policies for SageMaker projects and JumpStart.

Create an MLOps Project using Amazon SageMaker Studio Classic

This procedure demonstrates how to create an MLOps project using Amazon SageMaker Studio Classic.

Prerequisites

- An IAM account or IAM Identity Center to sign in to Studio Classic. For information, see Amazon SageMaker domain overview.
• Permission to use SageMaker-provided project templates. For information, see SageMaker Studio Permissions Required to Use Projects.

• Basic familiarity with the Studio Classic user interface. For information, see Amazon SageMaker Studio Classic UI Overview.

To create a project in Studio Classic

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.

2. In the Studio Classic sidebar, choose the Home icon ( ).

3. Select Deployments from the menu, and then select Projects.

4. Choose Create project.

   The Create project tab opens displaying a list of available templates.

5. If not selected already, choose SageMaker templates. For more information about project templates, see MLOps Project Templates.

6. Choose the template Model building, training, and deployment.

7. Choose Select project template.

   The Create project tab changes to display Project details.

8. Enter the following information:

   • For Project details, enter a name and description for your project.
   • Optionally, add tags, which are key-value pairs that you can use to track your projects.

9. Choose Create project and wait for the project to appear in the Projects list.

MLOps Project Templates

An Amazon SageMaker project template automates the setup and implementation of MLOps for your projects. A SageMaker project template is an Service Catalog product that SageMaker makes available to Amazon SageMaker Studio Classic users. These Service Catalog products are visible in your Service Catalog console after you enable permissions when you onboard or update Amazon SageMaker Studio Classic. For information about enabling permissions to use SageMaker project templates, see SageMaker Studio Permissions Required to Use Projects. Use SageMaker project templates to create a project that is an end-to-end MLOps solution.
If you are an administrator, you can create custom project templates from scratch or modify one of the project templates provided by SageMaker. Studio Classic users in your organization can use these custom project templates to create their projects.

Topics
- Use SageMaker-Provided Project Templates
- Create Custom Project Templates

Use SageMaker-Provided Project Templates

Amazon SageMaker provides project templates that create the infrastructure you need to create an MLOps solution for continuous integration and continuous deployment (CI/CD) of ML models. Use these templates to process data, extract features, train and test models, register the models in the SageMaker model registry, and deploy the models for inference. You can customize the seed code and the configuration files to suit your requirements.

⚠️ Important
As of July 25, 2022, we require additional roles to use project templates. For a complete list of required roles and instructions on how to create them, see SageMaker Studio Permissions Required to Use Projects. If you do not have the new roles, you will get the error message CodePipeline is not authorized to perform AssumeRole on role arn:aws:iam::xxx:role/service-role/AmazonSageMakerServiceCatalogProductsCodePipelineRole when you try to create a new project and cannot proceed.

SageMaker project templates offer you the following choice of code repositories, workflow automation tools, and pipeline stages:

- **Code repository**: AWS CodeCommit or third-party Git repositories such as GitHub and Bitbucket
- **CI/CD workflow automation**: AWS CodePipeline or Jenkins
- **Pipeline stages**: Model building and training, model deployment, or both

The following discussion provides an overview of each template you can choose when you create your SageMaker project. You can also view the available templates in Studio Classic by following Step 1: Create the Project of the Project walkthrough.
For step-by-step instructions on how to create a real project, you can follow one of the project walkthroughs:

- If you want to use the template **MLOps template for model building, training, and deployment**, see [SageMaker MLOps Project Walkthrough](#).
- If you want to use the template **MLOps template for model building, training, and deployment with third-party Git repositories using CodePipeline**, see [SageMaker MLOps Project Walkthrough Using Third-party Git Repos](#).
- If you want to use the template **MLOps template for model building, training, and deployment with third-party Git repositories using Jenkins**, see [Create Amazon SageMaker projects using third-party source control and Jenkins](#).

**Topics**

- **MLOps template for model building, training, and deployment**
- **MLOps template for model building, training, deployment, and Amazon SageMaker Model Monitor**
- **MLOps template for image building, model building, and model deployment**
- **MLOps template for model building, training, and deployment with third-party Git repositories using CodePipeline**
- **MLOps template for model building, training, and deployment with third-party Git repositories using Jenkins**
- **Model deployment for Salesforce**
- **Update SageMaker Projects to Use Third-Party Git Repositories**

**MLOps template for model building, training, and deployment**

This template is a combination of the following two templates, each of which can be used independently, and contains all of the resources provided in those templates.

- **Code repository**: AWS CodeCommit
- **CI/CD workflow automation**: AWS CodePipeline
MLOps template for model building and training

Use this template when you want an MLOps solution to process data, extract features, train and test models, and register the models in the SageMaker model registry.

This template provides the following resources:

- An AWS CodeCommit repository that contains sample code that creates an Amazon SageMaker pipeline in Python code and shows how to create and update the SageMaker pipeline. This repository also has a sample Python notebook that you can open and run in Studio Classic.

- An AWS CodePipeline pipeline that has source and build steps. The source step points to the CodeCommit repository. The build step gets the code from that repository, creates and updates the SageMaker pipeline, starts a pipeline execution, and waits for the pipeline execution to complete.

- An Amazon S3 bucket to store artifacts, including CodePipeline and CodeBuild artifacts, and any artifacts generated from the SageMaker pipeline runs.

The following diagram illustrates the workflow and AWS resources used by this template to help you build and train your models.
MLOps template for model deployment

Use this template to automate the deployment of models in the SageMaker model registry to SageMaker endpoints for real-time inference. This template recognizes changes in the model registry. When a new model version is registered and approved, it automatically initiates a deployment.

The template provisions a CodeCommit repository with configuration files to specify the model deployment steps, AWS CloudFormation templates to define endpoints as infrastructure, and seed code for testing the endpoint.

This template provides the following resources:

- An AWS CodeCommit repository that contains sample code that deploys models to endpoints in staging and production environments.
- An AWS CodePipeline pipeline that has source, build, deploy-to-staging, and deploy-to-production steps. The source step points to the CodeCommit repository, and the build step gets the code from that repository and generates CloudFormation stacks to deploy. The deploy-to-staging and deploy-to-production steps deploy the CloudFormation stacks to their respective environments. There is a manual approval step between the staging and production build steps, so that a MLOps engineer must approve the model before it is deployed to production.

  There is also a programmatic approval step with placeholder tests in the example code in the CodeCommit repository. You can add additional tests to replace the placeholders tests.

- An Amazon S3 bucket to store artifacts, including CodePipeline and CodeBuild artifacts, and any artifacts generated from the SageMaker pipeline runs.
- A CloudWatch event to initiate the pipeline when a model package version is approved or rejected.

The following diagram illustrates the workflow and AWS resources used by this template to help you deploy your models.
As previously mentioned, see [Project Walkthrough](#) for a demonstration that uses this template to create a real project.

**MLOps template for model building, training, deployment, and Amazon SageMaker Model Monitor**

This template is an extension of the MLOps template for model building, training, and deployment. It includes both the model building, training, and deployment components of the template, and an additional Amazon SageMaker Model Monitor template that provides the following types of monitoring:

- **Data Quality** – Monitor drift in data quality.
- **Model Quality** – Monitor drift in model quality metrics, such as accuracy.
- **Bias Drift for Models in Production** – Monitor bias in a model's predictions.

- **Code repository**: AWS CodeCommit
- **CI/CD workflow automation**: AWS CodePipeline
MLOps template for Amazon SageMaker Model Monitor

You can use this template for an MLOps solution to deploy one or more of the Amazon SageMaker data quality, model quality, model bias, and model explainability monitors to monitor a deployed model on a SageMaker inference endpoint.

This template provides the following resources:

- An AWS CodeCommit repository that contains sample Python code that gets the baselines used by the monitors from the SageMaker Model Registry, and updates the template's parameters for the staging and production environments. It also contains a AWS CloudFormation template to create the Amazon SageMaker Model Monitors.

- An AWS CodePipeline pipeline that has source, build, and deploy steps. The source step points to the CodePipeline repository. The build step gets the code from that repository, gets the baseline from the Model Registry, and updates template parameters for the staging and production environments. The deploy steps deploy the configured monitors into the staging and production environments. The manual approval step, within the DeployStaging stage, requires you to verify that the production SageMaker endpoint is InService before approving and moving to the DeployProd stage.

- The template uses the same S3 bucket created by the MLOps template for model building, training, and deployment to store the monitors' outputs.

- Two Amazon EventBridge events rules initiate the Amazon SageMaker Model Monitor AWS CodePipeline every time the staging SageMaker endpoint is updated, or a code change is committed to the CodePipeline repository.

MLOps template for image building, model building, and model deployment

This template is an extension of the MLOps template for model building, training, and deployment. It includes both the model building, training, and deployment components of that template and the following options:

- Include processing image–building pipeline
- Include training image–building pipeline
- Include inference image–building pipeline

For each of the components selected during project creation, the following are created by using the template:
• An Amazon ECR repository

• **A SageMaker Image**

• A CodeCommit repository containing a Dockerfile that you can customize

• A CodePipeline that is initiated by changes to the CodePipeline repository

• A CodeBuild project that builds a Docker image and registers it in the Amazon ECR repository

• An EventBridge rule that initiates the CodePipeline on a schedule

When the CodePipeline is initiated, it builds a new Docker container and registers it with an Amazon ECR repository. When a new container is registered with the Amazon ECR repository, a new ImageVersion is added to the SageMaker image. This initiates the model building pipeline, which in turn initiates the deployment pipeline.

The newly created image is used in the model building, training, and deployment portions of the workflow where applicable.

**MLOps template for model building, training, and deployment with third-party Git repositories using CodePipeline**

• **Code repository**: Third-party Git. Establish the AWS CodeStar connection from your AWS account to your GitHub user or organization. Add a tag with the key `sagemaker` and value `true` to this AWS CodeStar connection.

• **CI/CD workflow automation**: AWS CodePipeline

This template provides the following resources:

• Associations with one or more customer-specified Git repositories.

• An AWS CodePipeline pipeline that has source, build, deploy-to-staging, and deploy-to-production steps. The source step points to the third-party Git repository and the build step gets the code from that repository and generates CloudFormation stacks to deploy. The deploy-to-staging and deploy-to-production steps deploy the CloudFormation stacks to their respective environments. There is a manual approval step between the staging and production build steps, so that a MLOps engineer must approve the model before it is deployed to production.

• An AWS CodeBuild project to populate the Git repositories with the seed code information. This requires an AWS CodeStar connection from your AWS account to your account on the Git repository host.
• An Amazon S3 bucket to store artifacts, including CodePipeline and CodeBuild artifacts, and any artifacts generated from the SageMaker pipeline runs.

As previously mentioned, see Project Walkthrough Using Third-party Git Repos for a demonstration that uses this template to create a real project.

MLOps template for model building, training, and deployment with third-party Git repositories using Jenkins

• Code repository: Third-party Git. Establish the AWS CodeStar connection from your AWS account to your GitHub user or organization. Add a tag with the key sagemaker and value true to this AWS CodeStar connection.

• CI/CD workflow automation: Jenkins

This template provides the following resources:

• Associations with one or more customer-specified Git repositories.

• Seed code to generate Jenkins pipelines that have source, build, deploy-to-staging, and deploy-to-production steps. The source step points to the customer-specified Git repository. The build step gets the code from that repository and generates two CloudFormation stacks. The deploy steps deploy the CloudFormation stacks to their respective environments. There is an approval step between the staging step and the production step.

• An AWS CodeBuild project to populate the Git repositories with the seed code information. This requires an AWS CodeStar connection from your AWS account to your account on the Git repository host.

• An Amazon S3 bucket to store artifacts of the SageMaker project and SageMaker pipeline.

The template creates the association between your project and the source control repositories, but you need to perform additional manual steps to establish communication between your AWS account and Jenkins. For the detailed steps, see Create Amazon SageMaker projects using third-party source control and Jenkins.

The instructions help you build the architecture shown in the following diagram, with GitHub as the source control repository in this example. As shown, you are attaching your Git repository to the project to check in and manage code versions. Jenkins initiates the model build pipeline when it detects changes to the model build code in the Git repository. You are also connecting the
project to Jenkins to orchestrate your model deployment steps, which start when you approve the model registered in the model registry, or when Jenkins detects changes to the model deployment code.

In summary, the steps guide you through the following tasks:

1. Establish the connection between your AWS and GitHub accounts.
2. Create the Jenkins account and import needed plugins.
3. Create the Jenkins IAM user and permissions policy.
4. Set the AWS credentials for the Jenkins IAM user on your Jenkins server.
5. Create an API token for communication with your Jenkins server.
6. Use a CloudFormation template to set up an EventBridge rule to monitor the model registry for newly-approved models.
7. Create the SageMaker project, which seeds your GitHub repositories with model build and deploy code.
8. Create your Jenkins model build pipeline with the model build seed code.
9. Create your Jenkins model deploy pipeline with the model deploy seed code.
Model deployment for Salesforce

- **Code repository**: AWS CodeCommit
- **CI/CD workflow automation**: AWS CodePipeline

This template provides the following resources:

- An AWS CodeCommit repository that contains sample code that creates an Amazon SageMaker pipeline in Python code and shows how to create and update the pipeline. This repository also has a Python Jupyter Notebook that you can open and run in Studio Classic.
- An AWS CodePipeline pipeline that has source and build steps. The source step points to the CodeCommit repository. The build step gets the code from the repository, creates and updates the SageMaker pipeline, starts a pipeline run, and waits for the pipeline run to complete.
- An Amazon S3 bucket to store artifacts, including CodePipeline and CodeBuild artifacts, and any artifacts generated from the SageMaker pipeline runs.

Your admin may need to perform additional setup to enable data access from Salesforce Data Cloud to SageMaker Studio to build AI/ML models. See the solution overview in the blog post [Use the Amazon SageMaker and Salesforce Data Cloud integration to power your Salesforce apps with AI/ML](https://example.com) for detailed information and instructions.

The following diagram illustrates the high-level workflow used by this template to help you build and train your models. After you set up a connection between the Salesforce Data Cloud to Data Wrangler and preprocess your data, use the Model deployment for Salesforce project template to automate model training and deployment. The template provides customizable model deploy code and a sample notebook from AWS CodePipeline to train your model and register it into the SageMaker model registry. Once you approve the model, the endpoint is exposed to Salesforce as an API through API Gateway, and customers can start making predictions with the deployed model from within Salesforce.

**Note**

This template permits Transport Layer Security (TLS) policy versions 1.0 and 1.1 for API Gateway setup. You can make this configuration more secure with custom domain names. For details, see [Setting up custom domain names for REST APIs](https://example.com).
The blog post [Use the Amazon SageMaker and Salesforce Data Cloud integration to power your Salesforce apps with AI/ML](#) provides detailed instructions to guide you through the following steps:

1. Select the project template **Model deployment for Salesforce**, and provide the secret manager name.
2. Clone the repository to use the customizable SageMaker-provided sample notebook and model deploy code.
3. Preprocess your data with Data Wrangler.
   a. Create a connection to the Salesforce Data Cloud and import data into Data Wrangler.
   b. Use Data Wrangler to prepare the data with some example transformations.
   c. Initiate a processing job to process the data using your Data Wrangler configuration.
4. Train the model.
5. Register your model in the model registry.
6. Approve your model in model registry.
7. View your endpoint in the SageMaker console.
8. Invoke the API URL from Salesforce Einstein Studio to register and use the model inferences in Einstein Studio.

The following diagram shows in greater detail the workflow and AWS resources used by the SageMaker project template with Salesforce Data Cloud Integration.

Update SageMaker Projects to Use Third-Party Git Repositories

The managed policy attached to the AmazonSageMakerServiceCatalogProductsUseRole role was updated on July 27, 2021 for use with the third-party Git templates. Users who onboard to Amazon SageMaker Studio Classic after this date and enable project templates use the new policy. Users who onboarded prior to this date must update the policy to use these templates. Use one of the following options to update the policy:

- Delete role and toggle Studio Classic settings
1. In the IAM console, delete AmazonSageMakerServiceCatalogProductsUseRole.
2. In the Studio Classic control panel, choose Edit Settings.
3. Toggle both settings and then choose Submit.

- In the IAM console, add the following permissions to AmazonSageMakerServiceCatalogProductsUseRole:

```json
{
  "Effect": "Allow",
  "Action": [
    "codestar-connections:UseConnection"
  ],
  "Resource": "arn:aws:codestar-connections:*:*:connection/*",
  "Condition": {
    "StringEqualsIgnoreCase": {
      "aws:ResourceTag/sagemaker": "true"
    }
  }
},
{
  "Effect": "Allow",
  "Action": [
    "s3:PutObjectAcl"
  ],
  "Resource": [
    "arn:aws:s3:::sagemaker-*"
  ]
}
```

**Create Custom Project Templates**

If the SageMaker-provided templates do not meet your needs (for example, you want to have more complex orchestration in the CodePipeline with multiple stages or custom approval steps), create your own templates.

We recommend starting by using SageMaker-provided templates to understand how to organize your code and resources and build on top of it. To do this, after you enable administrator access to the SageMaker templates, log in to the [https://console.aws.amazon.com/servicecatalog/](https://console.aws.amazon.com/servicecatalog/), choose Portfolios, then choose Imported. For information about Service Catalog, see [Overview of Service Catalog](https://docs.aws.amazon.com/servicecatalog/latest/adminguide/servicecatalog-overview.html) in the [Service Catalog User Guide](https://docs.aws.amazon.com/servicecatalog/latest/adminguide/servicecatalog-overview.html).
Create your own project templates to customize your MLOps project. SageMaker project templates are Service Catalog–provisioned products to provision the resources for your MLOps project.

To create a custom project template, complete the following steps.

1. Create a portfolio. For information, see Step 3: Create an Service Catalog Portfolio.
2. Create a product. A product is a CloudFormation template. You can create multiple versions of the product. For information, see Step 4: Create an Service Catalog Product.

For the product to work with SageMaker projects, add the following parameters to your product template.

```yaml
SageMakerProjectName:
  Type: String
  Description: Name of the project

SageMakerProjectId:
  Type: String
  Description: Service generated Id of the project.
```

⚠️ Important

We recommend that you wrap the CodeCommit repository into the SageMaker code repository for the project's repositories to be visible in VPC mode. The sample template and required addition are shown in the following code samples.

Original (sample) template:

```yaml
ModelBuildCodeCommitRepository:
  Type: AWS::CodeCommit::Repository
  Properties:
    # Max allowed length: 100 chars
    RepositoryName: !Sub sagemaker-${SageMakerProjectName}-${SageMakerProjectId}-modelbuild # max: 10+33+15+10=68
    RepositoryDescription: !Sub SageMaker Model building workflow infrastructure as code for the Project ${SageMakerProjectName}
    Code:
      S3:
        Bucket: SEEDCODE_BUCKETNAME
        Key: toolchain/model-building-workflow-v1.0.zip
        BranchName: main
```
Additional content to add in VPC mode:

```yaml
SageMakerRepository:
  Type: AWS::SageMaker::CodeRepository
  Properties:
    GitConfig:
      RepositoryUrl: !GetAtt ModelBuildCodeCommitRepository.CloneUrlHttp
      Branch: main
```

3. Add a launch constraint. A launch constraint designates an IAM role that Service Catalog assumes when a user launches a product. For information, see Step 6: Add a Launch Constraint to Assign an IAM Role.

4. Provision the product on https://console.aws.amazon.com/servicecatalog/ to test the template. If you are satisfied with your template, continue to the next step to make the template available in Studio Classic.

5. Grant access to the Service Catalog portfolio that you created in step 1 to your Studio Classic execution role. Use either the Studio Classic domain execution role or a user role that has Studio Classic access. For information about adding a role to the portfolio, see Step 7: Grant End Users Access to the Portfolio.

6. To make your project template available in your Organization templates list in Studio Classic, create a tag with the following key and value to the Service Catalog product you created in step 2.
   - **key**: sagemaker:studio-visibility
   - **value**: true

After you complete these steps, Studio Classic users in your organization can create a project with the template you created by following the steps in Create an MLOps Project using Amazon SageMaker Studio Classic and choosing Organization templates when you choose a template.

**View Project Resources**

After you create a project, view the resources associated with the project in Amazon SageMaker Studio Classic.
To create a project in Studio Classic

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the Studio Classic sidebar, choose the Home icon (🏠).
3. Select Deployments from the menu, and then select Projects.
4. Select the name of the project for which you want to view details.

A tab with the project details appears.

On the project details tab, you can view the following entities associated with the project.

- Repositories: Code repositories (repos) associated with this project. If you use a SageMaker-provided template when you create your project, it creates a AWS CodeCommit repo or a third-party Git repo. For more information about CodeCommit, see What is AWS CodeCommit.
- Pipelines: SageMaker ML pipelines that define steps to prepare data, train, and deploy models. For information about SageMaker ML pipelines, see Create and Manage SageMaker Pipelines.
- Experiments: One or more Amazon SageMaker Autopilot experiments associated with the project. For information about Autopilot, see SageMaker Autopilot.
- Model groups: Groups of model versions that were created by pipeline executions in the project. For information about model groups, see Create a Model Group.
- Endpoints: SageMaker endpoints that host deployed models for real-time inference. When a model version is approved, it is deployed to an endpoint.
- Settings: Settings for the project. This includes the name and description of the project, information about the project template and SourceModelPackageGroupName, and metadata about the project.

Update an MLOps Project in Amazon SageMaker Studio Classic

This procedure demonstrates how to update an MLOps project in Amazon SageMaker Studio Classic. You can update the Description, template version, and template parameters.

Prerequisites

- An IAM account or IAM Identity Center to sign in to Studio Classic. For information, see Amazon SageMaker domain overview.
• Basic familiarity with the Studio Classic user interface. For information, see Amazon SageMaker Studio Classic UI Overview.

• Add the following custom inline policies to the specified roles:

User-created role having AmazonSageMakerFullAccess

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": ["servicecatalog:CreateProvisionedProductPlan",
                  "servicecatalog:DescribeProvisionedProductPlan",
                  "servicecatalog:DeleteProvisionedProductPlan"
                  ],
      "Resource": "*"
    }
  ]
}
```

AmazonSageMakerServiceCatalogProductsLaunchRole

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": ["cloudformation:CreateChangeSet",
                  "cloudformation:DeleteChangeSet",
                  "cloudformation:DescribeChangeSet"
                  ],
      "Resource": "arn:aws:cloudformation::*::*:stack/SC-*"
    },
    {
      "Effect": "Allow",
      "Action": ["codecommit:PutRepositoryTriggers"
                  ],
      "Resource": "arn:aws:codecommit::*::*:sagemaker-*"
    }
  ]
}
```
To update a project in Studio Classic

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the Studio Classic sidebar, choose the Home icon (  ).
3. Select Deployments from the menu, and then select Projects. A list of your projects appears.
4. Select the name of the project you want to update in the projects list.
5. Choose Update from the Actions menu in the upper-right corner of the project tab.
6. In the Update project dialog box, you can edit the Description and listed template parameters.
7. Choose View difference.

A dialog box displays your original and updated project settings. Any change in your project settings can modify or delete resources in the current project. The dialog box displays these changes as well.
8. You may need to wait a few minutes for the Update button to become active. Choose Update.
9. The project update may take a few minutes to complete. Select Settings in the project tab and ensure the parameters have been updated correctly.

Delete an MLOps Project using Amazon SageMaker Studio Classic

This procedure demonstrates how to delete an MLOps project using Amazon SageMaker Studio Classic.

Prerequisites

⚠️ Note

You can only delete projects in Studio Classic that you have created. This condition is part of the service catalog permission servicecatalog:TerminateProvisionedProduct in the AmazonSageMakerFullAccess policy. If needed, you can update this policy to remove this condition.
An IAM account or IAM Identity Center to sign in to Studio Classic. For information, see Amazon SageMaker domain overview.

• Basic familiarity with the Studio Classic user interface. For information, see Amazon SageMaker Studio Classic UI Overview.

To delete a project in Amazon SageMaker Studio Classic

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the Studio Classic sidebar, choose the Home icon ( ).
3. Select Deployments from the menu, and then select Projects.
4. Select the target project from the dropdown list. If you don't see your project, type the project name and apply the filter to find your project.
5. Once you've found your project, select the project name to view details.
6. Choose Delete from the Actions menu.
7. Confirm your choice by choosing Delete from the Delete Project window.

SageMaker MLOps Project Walkthrough

This walkthrough uses the template MLOps template for model building, training, and deployment to demonstrate using MLOps projects to create a CI/CD system to build, train, and deploy models.

Prerequisites

To complete this walkthrough, you need:

• An IAM account or IAM Identity Center to sign in to Studio Classic. For information, see Amazon SageMaker domain overview.

• Permission to use SageMaker-provided project templates. For information, see SageMaker Studio Permissions Required to Use Projects.

• Basic familiarity with the Studio Classic user interface. For information, see Amazon SageMaker Studio Classic UI Overview.

Topics
• Step 1: Create the Project
• Step 2: Clone the Code Repository
• Step 3: Make a Change in the Code
• Step 4: Approve the Model
• (Optional) Step 5: Deploy the Model Version to Production
• Step 6: Clean Up Resources

Step 1: Create the Project

In this step, you create a SageMaker MLOps project by using a SageMaker-provided project template to build, train, and deploy models.

To create the SageMaker MLOps project

1. Sign in to Studio Classic. For more information, see Amazon SageMaker domain overview.
2. In the Studio Classic sidebar, choose the Home icon ( ).
3. Select Deployments from the menu, and then select Projects.
4. Choose Create project.
   The Create project tab appears.
5. If not selected already, choose SageMaker templates, then choose MLOps template for model building, training, and deployment.
6. For Project details, enter a name and description for your project.

When the project appears in the Projects list with a Status of Create completed, move on to the next step.

⚠️ Important

As of July 25, 2022, we require additional roles to use project templates. If you see the error message CodePipeline is not authorized to perform AssumeRole on role arn:aws:iam::xxx:role/service-role/AmazonSageMakerServiceCatalogProductsCodePipelineRole, see Steps 5-6 of SageMaker Project walkthrough.
Step 2: Clone the Code Repository

After you create the project, two CodeCommit repositories are created in the project. One of the repositories contains code to build and train a model, and one contains code to deploy the model. In this step, you clone the repository to your local SageMaker project that contains the code to build and train the model to the local Studio Classic environment so that you can work with the code.

To clone the code repository

1. In the Studio Classic sidebar, choose the Home icon (``).
2. Select Deployments from the menu, and then select Projects.
3. Select the project you created in the previous step to open the project tab for your project.
4. In the project tab, choose Repositories, and in the Local path column for the repository that ends with modelbuild, choose clone repo....
5. In the dialog box that appears, accept the defaults and choose Clone repository.

![Clone Repository](image-url)
When clone of the repository is complete, the local path appears in the **Local path** column. Choose the path to open the local folder that contains the repository code in Studio Classic.

**Step 3: Make a Change in the Code**

Now make a change to the pipeline code that builds the model and check in the change to initiate a new pipeline run. The pipeline run registers a new model version.

**To make a code change**

1. In Studio Classic, choose the file browser icon (￼), and navigate to the pipelines/abalone folder. Double-click pipeline.py to open the code file.

2. In the pipeline.py file, find the line that sets the training instance type.

   ```python
   training_instance_type = ParameterString(
       name="TrainingInstanceType", default_value="ml.m5.xlarge"
   )
   ```

   Change `ml.m5.xlarge` to `ml.m5.large`, then type Ctrl+S to save the change.

3. Choose the **Git** icon (￼). Stage, commit, and push the change in pipeline.py. Also, enter a summary in the **Summary** field and an optional description in the **Description** field. For information about using Git in Studio Classic, see [Clone a Git Repository in SageMaker Studio Classic](#).
After pushing your code change, the MLOps system initiates a run of the pipeline that creates a new model version. In the next step, you approve the new model version to deploy it to production.

**Step 4: Approve the Model**

Now you approve the new model version that was created in the previous step to initiate a deployment of the model version to a SageMaker endpoint.

To approve the model version

1. In the Studio Classic sidebar, choose the **Home** icon ( ).
2. Select **Deployments** from the menu, and then select **Projects**.
3. Select the name of the project you created in the first step to open the project tab for your project.
4. In the project tab, choose **Model groups**, then double-click the name of the model group that appears.

   The model group tab appears.
5. In the model group tab, double-click **Version 1**. The **Version 1** tab opens. Choose **Update status**.
6. In the model **Update model version status** dialog box, in the **Status** dropdown list, select **Approve**, then choose **Update status**.

   Approving the model version causes the MLOps system to deploy the model to staging. To view the endpoint, choose the **Endpoints** tab on the project tab.

**(Optional) Step 5: Deploy the Model Version to Production**

Now you can deploy the model version to the production environment.

**Note**

To complete this step, you need to be an administrator in your Studio Classic domain. If you are not an administrator, skip this step.
To deploy the model version to the production environment

1. Log in to the CodePipeline console at https://console.aws.amazon.com/codepipeline/
2. Choose Pipelines, then choose the pipeline with the name sagemaker-projectname-projectid-modeldeploy, where projectname is the name of your project, and projectid is the ID of your project.
3. In the DeployStaging stage, choose Review.
4. In the Review dialog box, choose Approve.

Approving the DeployStaging stage causes the MLOps system to deploy the model to production. To view the endpoint, choose the Endpoints tab on the project tab in Studio Classic.

Step 6: Clean Up Resources

To stop incurring charges, clean up the resources that were created in this walkthrough. To do this, complete the following steps.

Note
To delete the AWS CloudFormation stack and the Amazon S3 bucket, you need to be an administrator in Studio Classic. If you are not an administrator, ask your administrator to complete those steps.

1. In the Studio Classic sidebar, choose the Home icon ().
2. Select Deployments from the menu, and then select Projects.
3. Select the target project from the dropdown list. If you don’t see your project, type the project name and apply the filter to find your project.
4. You can delete a Studio Classic project in one of the following ways:
   a. You can delete the project from the projects list.
      
      Right-click the target project and choose Delete from the dropdown list.
b. **You can delete a project from the Project details section.**
   
i. When you've found your project, double-click it to view its details in the main panel.
   
ii. Choose **Delete** from the **Actions** menu.

5. Confirm your choice by choosing **Delete** from the **Delete Project** window.

   This deletes the Service Catalog provisioned product that the project created. This includes the CodeCommit, CodePipeline, and CodeBuild resources created for the project.

6. Delete the AWS CloudFormation stacks that the project created. There are two stacks, one for staging and one for production. The names of the stacks are `sagemaker-projectname-project-id-deploy-staging` and `sagemaker-projectname-project-id-deploy-prod`, where `projectname` is the name of your project, and `project-id` is the ID of your project.

   For information about how to delete a AWS CloudFormation stack, see [Deleting a stack on the AWS CloudFormation console](https://aws.amazon.com/documentation/cloudformation/reference/glossary.html) in the **AWS CloudFormation User Guide**.

7. Delete the Amazon S3 bucket that the project created. The name of the bucket is `sagemaker-project-project-id`, where `project-id` is the ID of your project.

---

### SageMaker MLOps Project Walkthrough Using Third-party Git Repos

This walkthrough uses the template [MLOps template for model building, training, and deployment with third-party Git repositories using CodePipeline](https://aws.amazon.com/sagemaker/) to demonstrate how to use MLOps projects to create a CI/CD system to build, train, and deploy models.

#### Prerequisites

To complete this walkthrough, you need:

- An IAM or IAM Identity Center account to sign in to Studio Classic. For information, see [Amazon SageMaker domain overview](https://aws.amazon.com/sagemaker/).
• Permission to use SageMaker-provided project templates. For information, see SageMaker Studio Permissions Required to Use Projects.

• Basic familiarity with the Studio Classic user interface. For information, see Amazon SageMaker Studio Classic UI Overview.

• Two GitHub repositories initialized with a README. You input these repositories into the project template, which will seed these repos with model build and deploy code.

Topics

• **Step 1: Set up the GitHub connection**
• **Step 2: Create the Project**
• **Step 3: Make a Change in the Code**
• **Step 4: Approve the Model**
• **(Optional) Step 5: Deploy the Model Version to Production**
• **Step 6: Clean Up Resources**

**Step 1: Set up the GitHub connection**

In this step, you connect to your GitHub repositories using an AWS CodeStar connection. The SageMaker project uses this connection to access your source code repositories.

**To set up the GitHub connection:**

1. Log in to the CodePipeline console at https://console.aws.amazon.com/codepipeline/
2. Under **Settings** in the navigation pane, choose **Connections**.
3. Choose **Create connection**.
4. For **Select a provider**, select GitHub.
5. For **Connection name**, enter a name.
6. Choose **Connect to GitHub**.
7. If the AWS Connector GitHub app isn't previously installed, choose **Install new app**.
   
   This displays a list of all the GitHub personal accounts and organizations to which you have access.
8. Choose the account where you want to establish connectivity for use with SageMaker projects and GitHub repositories.
9. Choose **Configure**.

10. You can optionally select your specific repositories or choose **All repositories**.

11. Choose **Save**. When the app is installed, you’re redirected to the **Connect to GitHub** page and the installation ID is automatically populated.

12. Choose **Connect**.

13. Add a tag with the key `sagemaker` and value `true` to this AWS CodeStar connection.

14. Copy the connection ARN to save for later. You use the ARN as a parameter in the project creation step.

**Step 2: Create the Project**

In this step, you create a SageMaker MLOps project by using a SageMaker-provided project template to build, train, and deploy models.

**To create the SageMaker MLOps project**

1. Sign in to Studio Classic. For more information, see [Amazon SageMaker domain overview](#).

2. In the Studio Classic sidebar, choose the **Home** icon (  

3. Select **Deployments** from the menu, and then select **Projects**.

4. Choose **Create project**.

   The **Create project** tab appears.

5. For **SageMaker project templates**, choose **MLOps template for model building, training, and deployment with third-party Git repositories**.

6. Choose **Select project template**.

7. Under **ModelBuild CodeRepository Info**, provide the following parameters:

   - For **URL**, enter the URL of your Git repository for the model build code in `https://git-url.git` format.
   - For **Branch**, enter the branch to use from your Git repository for pipeline activities.
   - For **Full Repository Name**, enter the Git repository name in the format of `username/repository name` or `organization/repository name`.
   - For **Codestar Connection ARN**, enter the ARN of the AWS CodeStar connection you created in Step 1.
The **Sample Code** toggle switch lets you choose whether to populate the repository with model build seed code. We can leave it on for this demo.

8. Under **ModelDeploy CodeRepository Info**, provide the following parameters:

- For **URL**, enter the URL of your Git repository for the model deploy code in https://git-url.git format.
- For **Branch**, enter the branch to use from your Git repository for pipeline activities.
- For **Full Repository Name**, enter the Git repository name in the format of `username/repository name` or `organization/repository name`.
- For **Codestar Connection ARN**, enter the ARN of the AWS CodeStar connection you created in Step 1.
- The **Sample Code** toggle switch lets you choose whether to populate the repository with model deployment seed code. We can leave it on for this demo.

9. Choose **Create Project**.

The project appears in the **Projects** list with a **Status** of **Created**.

**Step 3: Make a Change in the Code**

Now make a change to the pipeline code that builds the model and commit the change to initiate a new pipeline run. The pipeline run registers a new model version.

**To make a code change**

1. In your model build GitHub repo, navigate to the `pipelines/abalone` folder. Double-click `pipeline.py` to open the code file.
2. In the `pipeline.py` file, find the line that sets the training instance type.

   ```python
   training_instance_type = ParameterString(
       name="TrainingInstanceType", default_value="ml.m5.xlarge"
   )
   ```

   Open the file for editing, change `ml.m5.xlarge` to `ml.m5.large`, then commit.

After you commit your code change, the MLOps system initiates a run of the pipeline that creates a new model version. In the next step, you approve the new model version to deploy it to production.
Step 4: Approve the Model

Now you approve the new model version that was created in the previous step to initiate a deployment of the model version to a SageMaker endpoint.

To approve the model version

1. In the Studio Classic sidebar, choose the Home icon (🏠).
2. Select Deployments from the menu, and then select Projects.
3. Find the name of the project you created in the first step and double-click on it to open the project tab for your project.
4. In the project tab, choose Model groups, then double-click the name of the model group that appears.

   The model group tab appears.
5. In the model group tab, double-click Version 1. The Version 1 tab opens. Choose Update status.
6. In the model Update model version status dialog box, in the Status dropdown list, select Approve and then choose Update status.

   Approving the model version causes the MLOps system to deploy the model to staging. To view the endpoint, choose the Endpoints tab on the project tab.

(Optional) Step 5: Deploy the Model Version to Production

Now you can deploy the model version to the production environment.

⚠️ Note

To complete this step, you need to be an administrator in your Studio Classic domain. If you are not an administrator, skip this step.

To deploy the model version to the production environment

1. Log in to the CodePipeline console at https://console.aws.amazon.com/codepipeline/
2. Choose **Pipelines**, then choose the pipeline with the name  
   `sagemaker-projectname-projectid-modeldeploy`, where `projectname` is the name of  
   your project, and `projectid` is the ID of your project.

3. In the **DeployStaging** stage, choose **Review**.

4. In the **Review** dialog box, choose **Approve**.

   Approving the **DeployStaging** stage causes the MLOps system to deploy the model to  
   production. To view the endpoint, choose the **Endpoints** tab on the project tab in Studio  
   Classic.

### Step 6: Clean Up Resources

To stop incurring charges, clean up the resources that were created in this walkthrough.

**Note**

To delete the AWS CloudFormation stack and the Amazon S3 bucket, you need to be an  
administrator in Studio Classic. If you are not an administrator, ask your administrator to  
complete those steps.

1. In the Studio Classic sidebar, choose the **Home** icon (  
   ![Home icon](image)
   ).

2. Select **Deployments** from the menu, and then select **Projects**.

3. Select the target project from the dropdown list. If you don’t see your project, type the project  
   name and apply the filter to find your project.

4. Select your project to view its details in the main panel.

5. Choose **Delete** from the **Actions** menu.

6. Confirm your choice by choosing **Delete** from the **Delete Project** window.

   This deletes the Service Catalog provisioned product that the project created. This includes the  
   CodeCommit, CodePipeline, and CodeBuild resources created for the project.

7. Delete the AWS CloudFormation stacks that the project created. There  
   are two stacks, one for staging and one for production. The names of the  
   stacks are `sagemaker-projectname-project-id-deploy-staging` and  
   `sagemaker-projectname-project-id-deploy-production`.
sagemaker-projectname-project-id-deploy-prod, where projectname is the name of your project, and project-id is the ID of your project.

For information about how to delete a AWS CloudFormation stack, see Deleting a stack on the AWS CloudFormation console in the AWS CloudFormation User Guide.

8. Delete the Amazon S3 bucket that the project created. The name of the bucket is sagemaker-project-project-id, where project-id is the ID of your project.

Amazon SageMaker MLOps FAQ

Use the following FAQ items to find answers to commonly asked questions about MLOps in SageMaker.

**Q. Do I need to use the SageMaker Python SDK to create a SageMaker pipeline?**

No, the SageMaker Python SDK is not required to create a SageMaker pipeline. You can also use boto3 or AWS CloudFormation. Creating a pipeline requires a pipeline definition, which is a JSON object that defines each step of the pipeline. The SageMaker SDK offers a simple way to construct the pipeline definition, which you can use with any of the APIs previously mentioned to create the pipeline itself. Without using the SDK, users have to write the raw JSON definition to create the pipeline without any of the error checks provided by the SageMaker Python SDK. To see the schema for the pipeline JSON definition, see SageMaker Pipeline Definition JSON Schema. The following code sample shows an example of a SageMaker pipeline definition JSON object:

```json
{
   'Version': '2020-12-01',
   'Metadata': {},
   'Parameters': [{
      'Name': 'ProcessingInstanceType',
      'Type': 'String',
      'DefaultValue': 'ml.m5.xlarge'},
   [{
      'Name': 'ProcessingInstanceCount', 'Type': 'Integer', 'DefaultValue': 1},
   [{
      'Name': 'TrainingInstanceType',
      'Type': 'String',
      'DefaultValue': 'ml.m5.xlarge'},
   [{
      'Name': 'ModelApprovalStatus',
      'Type': 'String',
      'DefaultValue': 'PendingManualApproval'},
   [{
      'Name': 'ProcessedData',
       'Type': 'String',
      'DefaultValue': 'S3_URL'},
```
Q. Why do I see a repack step in my SageMaker pipeline?

Model repacking happens when the pipeline needs to include a custom script in the compressed model file (model.tar.gz) to be uploaded to Amazon S3 and used to deploy a model to a SageMaker endpoint. When SageMaker pipeline trains a model and registers it to the model registry, it introduces a repack step if the trained model output from the training job needs to include a custom inference script. The repack step uncompresses the model, adds a new script, and recompresses the model. Running the pipeline adds the repack step as a training job.

Q. Can I use SageMaker Experiments with SageMaker Pipelines?

Yes. SageMaker Pipelines is natively integrated with SageMaker Experiments. You can use PipelineExperimentConfig when creating a pipeline and set your own SageMaker Experiment name. Each run of the pipeline creates a trial, and each step in the pipeline corresponds to a TrialComponent within the trial. If no trial name is specified in the experiment config, the pipeline execution ID is used as the trial name.
Q. SageMaker Project templates have a model deploy repository that uses CloudFormation (CFN) to create an endpoint. Are there ways to deploy the model without using CloudFormation?

You can customize the deploy repository in the project template to deploy the model from the model registry any way you like. The template uses CloudFormation to create a real-time endpoint, as an example. You can update the deployment to use the SageMaker SDK, boto3, or any other API that can create endpoints instead of CFN. If you need to update the CodeBuild steps as part of the deployment pipeline, you can create a custom template.

Q. How do we pass the model file Amazon S3 URL from the train step to the model register step in a SageMaker pipeline at run time?

You can reference the model location as a property of the training step, as shown in the end-to-end example CustomerChurn pipeline in Github.

Q. If I am extending a prebuilt container to train an estimator or for a ProcessingStep on SageMaker Pipelines, is it necessary to copy the script to the container in the Dockerfile?

No, you can either copy the script to the container or pass it via the entry_point argument (of your estimator entity) or code argument (of your processor entity), as demonstrated in the following code sample.

```python
tf
step_process = ProcessingStep(
    name="PreprocessAbaloneData",
    processor=sklearn_processor,
    inputs = [
        ProcessingInput(
            name = "preprocess_abalone_data",
            s3_data = "s3://bucket/model_data/abalone.csv",
            s3_input_mode = "stream"))
)
```
Q. What’s the recommended way to manage dependencies for different SageMaker Pipelines steps?

You can use a SageMaker Projects template to implement image-building CI/CD. With this template, you can automate the CI/CD of images that are built and pushed to Amazon ECR. Changes in the container files in your project’s source control repositories initiate the ML pipeline and deploy the latest version for your container. For more information, see the blog [Create Amazon SageMaker projects with image building CI/CD pipelines](#).
Q. How do I provide SageMaker Project access to specific user profiles in Amazon SageMaker Studio Classic?

Since SageMaker Projects is backed by Service Catalog, you must add each role that requires access to SageMaker Projects to the Amazon SageMaker Solutions and ML Ops products Portfolio in the service catalog. You can do this on the Groups, roles, and users tab, as shown in the following image. If each user profile in Studio Classic has a different role, you should add each of those roles to the service catalog. You can also do this while creating a user profile in Studio Classic.

Q. Where do I see the properties associated with each SageMaker pipeline step so that I can use them in subsequent steps?

Each step in the pipeline uses the underlying SageMaker APIs for the corresponding jobs. For example, TrainingStep invokes the CreateTrainingJob API and the step properties correspond to the response from DescribeTrainingJob. The response output can be found in the API reference link for DescribeTrainingJob. You can follow the same procedure to get the properties for TransformStep, ProcessingStep, TuningStep, and CreateModelStep. For more information about pipeline steps, see Pipeline Steps.
Q. In SageMaker Pipelines, can I specify a unique output path for a pipeline step so that its output data will not be overridden by future runs?

Yes, you can use `ExecutionVariables` and the `Join` function to specify your output location. `ExecutionVariables` is resolved at runtime. For instance, `ExecutionVariables.PIPELINE_EXECUTION_ID` is resolved to the ID of the current execution, which can be used as a unique identifier across different runs.

```python
from sagemaker.workflow.execution_variables import ExecutionVariables

processor_run_args = sklearn_processor.run(
    outputs=[
        ProcessingOutput(
            output_name="train",
            source="/opt/ml/processing/train",
            destination=Join(on="/",
                              values=[
                                      "s3:/",
                                      default_bucket,
                                      base_job_prefix,
                                      ExecutionVariables.PIPELINE_EXECUTION_ID,
                                      "PreprocessData"],
                              ),
                      ),
        ProcessingOutput(
            output_name="validation",
            source="/opt/ml/processing/validation",
            destination=Join(on="/",
                              values=[
                                      "s3:/",
                                      default_bucket,
                                      base_job_prefix,
                                      ExecutionVariables.PIPELINE_EXECUTION_ID,
                                      "PreprocessData"],
                              ),
                      ),
        ProcessingOutput(
            output_name="test",
            source="/opt/ml/processing/test",
        ),
    ]
)
```
Q. What’s the best way to reproduce my model in SageMaker?

SageMaker’s Lineage Tracking service works in the backend to track all the metadata associated with your model training and deployment workflows. This includes your training jobs, datasets used, pipelines, endpoints, and the actual models. You can query the lineage service at any point to find the exact artifacts used to train a model. Using those artifacts, you can recreate the same ML workflow to reproduce the model as long as you have access to the exact dataset that was used. A trial component tracks the training job. This trial component has all the parameters used as part of the training job. If you don’t need to rerun the entire workflow, you can reproduce the training job to derive the same model.

Q. If I try to delete a SageMaker project created from a SageMaker template and receive an error due to non-empty Amazon S3 buckets or Amazon ECR repositories, how can I delete the project?

If you try to delete your SageMaker project and get one of the following error messages:

- The bucket you tried to delete is not empty
- The repository with name 'repository-name' in registry

MLOps FAQ
then you have non-empty Amazon S3 buckets or Amazon ECR repositories which you need to manually delete before you delete the SageMaker project. AWS CloudFormation does not automatically delete non-empty Amazon S3 buckets or Amazon ECR repositories for you.
Monitor data and model quality

Amazon SageMaker Model Monitor monitors the quality of Amazon SageMaker machine learning models in production. You can set up continuous monitoring with a real-time endpoint (or a batch transform job that runs regularly), or on-schedule monitoring for asynchronous batch transform jobs. With Model Monitor, you can set alerts that notify you when there are deviations in the model quality. Early and proactive detection of these deviations enables you to take corrective actions, such as retraining models, auditing upstream systems, or fixing quality issues without having to monitor models manually or build additional tooling. You can use Model Monitor prebuilt monitoring capabilities that do not require coding. You also have the flexibility to monitor models by coding to provide custom analysis.

Model Monitor provides the following types of monitoring:

- **Monitor data quality** - Monitor drift in data quality.
- **Monitor model quality** - Monitor drift in model quality metrics, such as accuracy.
- **Monitor Bias Drift for Models in Production** - Monitor bias in your model's predictions.
- **Monitor Feature Attribution Drift for Models in Production** - Monitor drift in feature attribution.

Topics

- Monitoring a Model in Production
- How Model Monitor Works
- Capture data
- Monitor data quality
- Monitor model quality
- Monitor Bias Drift for Models in Production
- Monitor Feature Attribution Drift for Models in Production
- Schedule monitoring jobs
- Amazon SageMaker Model Monitor prebuilt container
- Interpret results
- Visualize results for real-time endpoints in Amazon SageMaker Studio
- Advanced topics
- Model Monitor FAQs
Monitoring a Model in Production

After you deploy a model into your production environment, use Amazon SageMaker model monitor to continuously monitor the quality of your machine learning models in real time. Amazon SageMaker model monitor enables you to set up an automated alert triggering system when there are deviations in the model quality, such as data drift and anomalies. Amazon CloudWatch Logs collects log files of monitoring the model status and notifies when the quality of your model hits certain thresholds that you preset. CloudWatch stores the log files to an Amazon S3 bucket you specify. Early and pro-active detection of model deviations through AWS model monitor products enables you to take prompt actions to maintain and improve the quality of your deployed model.

For more information about SageMaker model monitoring products, see Monitor data and model quality.

To start your machine learning journey with SageMaker, sign up for an AWS account at Set Up SageMaker.

How Model Monitor Works

Amazon SageMaker Model Monitor automatically monitors machine learning (ML) models in production and notifies you when quality issues arise. Model Monitor uses rules to detect drift in your models and alerts you when it happens. The following figure shows how this process works in the case that your model is deployed to a real-time endpoint.
You can also use Model Monitor to monitor a batch transform job instead of a real-time endpoint. In this case, instead of receiving requests to an endpoint and tracking the predictions, Model Monitor will monitor inference inputs and outputs. The following figure diagrams the process of monitoring a batch transform job.
To enable model monitoring, you take the following steps, which follow the path of the data through the various data collection, monitoring, and analysis processes:

- For a real-time endpoint, enable the endpoint to capture data from incoming requests to a trained ML model and the resulting model predictions.
- For a batch transform job, enable data capture of the batch transform inputs and outputs.
- Create a baseline from the dataset that was used to train the model. The baseline computes metrics and suggests constraints for the metrics. Real-time or batch predictions from your model are compared to the constraints, and are reported as violations if they are outside the constrained values.
- Create a monitoring schedule specifying what data to collect, how often to collect it, how to analyze it, and which reports to produce.
- Inspect the reports, which compare the latest data with the baseline, and watch for any violations reported and for metrics and notifications from Amazon CloudWatch.
Notes

- Model Monitor computes model metrics and statistics on tabular data only. For example, an image classification model that takes images as input and outputs a label based on that image can still be monitored. Model Monitor would be able to calculate metrics and statistics for the output, not the input.

- Model Monitor currently supports only endpoints that host a single model and does not support monitoring multi-model endpoints. For information on using multi-model endpoints, see Host multiple models in one container behind one endpoint.

- Model Monitor supports monitoring inference pipelines, but capturing and analyzing data is done for the entire pipeline, not for individual containers in the pipeline.

- To prevent impact to inference requests, Data Capture stops capturing requests at high levels of disk usage. It is recommended you keep your disk utilization below 75% in order to ensure data capture continues capturing requests.

- If you launch SageMaker Studio in a custom Amazon VPC, you need to create VPC endpoints to enable Model Monitor to communicate with Amazon S3 and CloudWatch. For information about VPC endpoints, see VPC endpoints in the Amazon Virtual Private Cloud User Guide. For information about launching SageMaker Studio in a custom VPC, see Connect SageMaker Studio Notebooks in a VPC to External Resources.

Model Monitor Sample Notebooks

For a sample notebook that takes you through the full end-to-end workflow using Model Monitor with your real-time endpoint, see Introduction to Amazon SageMaker Model Monitor.

For a sample notebook that visualizes the statistics.json file for a selected execution in a monitoring schedule, see the Model Monitor Visualization.

For instructions that show you how to create and access Jupyter notebook instances that you can use to run the example in SageMaker, see Amazon SageMaker Notebook Instances. After you have created a notebook instance and opened it, choose the SageMaker Examples tab to see a list of all the SageMaker samples. To open a notebook, choose the notebook's Use tab and choose Create copy.
Capture data

To log the inputs to your endpoint and the inference outputs from your deployed model to Amazon S3, you can enable a feature called Data Capture. Data Capture is commonly used to record information that can be used for training, debugging, and monitoring. Amazon SageMaker Model Monitor automatically parses this captured data and compares metrics from this data with a baseline that you create for the model. For more information about Model Monitor see Monitor data and model quality.

You can implement Data Capture for both real-time and batch model-monitor modes using the AWS SDK for Python (Boto) or the SageMaker Python SDK. For a real-time endpoint, you will specify your Data Capture configuration when you create your endpoint. Due to the persistent nature of your real-time endpoint, you can configure additional options to turn data capturing on or off at certain times, or change the sampling frequency. You can also choose to encrypt your inference data.

For a batch transform job, you can enable Data Capture if you want to run on-schedule model monitoring or continuous model-monitoring for regular, periodic batch transform jobs. You will specify your Data Capture configuration when you create your batch transform job. Within this configuration, you have the option to turn on encryption or generate the inference ID with your output, which helps you match your captured data to Ground Truth data.

Capture data from real-time endpoint

⚠️ Note

To prevent impact to inference requests, Data Capture stops capturing requests at high levels of disk usage. It is recommended you keep your disk utilization below 75% in order to ensure data capture continues capturing requests.

To capture data for your real-time endpoint, you must deploy a model using SageMaker hosting services. This requires that you create a SageMaker model, define an endpoint configuration, and create an HTTPS endpoint.

The steps required to turn on data capture are similar whether you use the AWS SDK for Python (Boto) or the SageMaker Python SDK. If you use the AWS SDK, define the DataCaptureConfig dictionary, along with required fields, within the CreateEndpointConfig method to turn on data capture. If you use the SageMaker Python SDK, import the DataCaptureConfig Class and initialize...
an instance from this class. Then, pass this object to the DataCaptureConfig parameter in the sagemaker.model.Model.deploy() method.

To use the proceeding code snippets, replace the *italicized placeholder text* in the example code with your own information.

**How to enable data capture**

Specify a data capture configuration. You can capture the request payload, the response payload, or both with this configuration. The proceeding code snippet demonstrates how to enable data capture using the AWS SDK for Python (Boto) and the SageMaker Python SDK.

### Note

You do not need to use Model Monitor to capture request or response payloads.

AWS SDK for Python (Boto)

Configure the data you want to capture with the `DataCaptureConfig` dictionary when you create an endpoint using the `CreateEndpointConfig` method. Set `EnableCapture` to the boolean value `True`. In addition, provide the following mandatory parameters:

- **EndpointConfigName**: the name of your endpoint configuration. You will use this name when you make a `CreateEndpoint` request.
- **ProductionVariants**: a list of models you want to host at this endpoint. Define a dictionary data type for each model.
- **DataCaptureConfig**: dictionary data type where you specify an integer value that corresponds to the initial percentage of data to sample (`InitialSamplingPercentage`), the Amazon S3 URI where you want captured data to be stored, and a capture options (`CaptureOptions`) list. Specify either `Input` or `Output` for `CaptureMode` within the `CaptureOptions` list.

You can optionally specify how SageMaker should encode captured data by passing key-value pair arguments to the `CaptureContentTypeHeader` dictionary.

```python
# Create an endpoint config name.
endpoint_config_name = '<endpoint-config-name>'
```
# The name of the production variant.
variant_name = '<name-of-production-variant>'

# The name of the model that you want to host.
# This is the name that you specified when creating the model.
model_name = '<The_name_of_your_model>'

instance_type = '<instance-type>'
#instance_type='ml.m5.xlarge' # Example

# Number of instances to launch initially.
initial_instance_count = <integer>

# Sampling percentage. Choose an integer value between 0 and 100
initial_sampling_percentage = <integer>

# The S3 URI containing the captured data
s3_capture_upload_path = 's3://<bucket-name>/<data_capture_s3_key>'

# Specify either Input, Output, or both
capture_modes = [ "Input", "Output" ]
#capture_mode = [ "Input"] # Example - If you want to capture input only

capture_data_from_real-time_endpoint
For more information about other endpoint configuration options, see the CreateEndpointConfig API in the Amazon SageMaker Service API Reference Guide.

SageMaker Python SDK

Import the DataCaptureConfig Class from sagemaker.model_monitor module. Enable data capture by setting EnableCapture to the boolean value True.

Optionally provide arguments for the following parameters:

- **SamplingPercentage**: an integer value that corresponds to percentage of data to sample. If you do not provide a sampling percentage, SageMaker will sample a default of 20 (20%) of your data.
- **DestinationS3Uri**: the Amazon S3 URI SageMaker will use to store captured data. If you do not provide one, SageMaker will store captured data in "s3://<default-session-bucket>/ model-monitor/data-capture".

```python
from sagemaker.model_monitor import DataCaptureConfig

# Set to True to enable data capture
enable_capture = True

# Optional - Sampling percentage. Choose an integer value between 0 and 100
sampling_percentage = <int>
# sampling_percentage = 30 # Example 30%

# Optional - The S3 URI of stored captured-data location
s3_capture_upload_path = 's3://<bucket-name>/<data_capture_s3_key>'

# Specify either Input, Output or both.
capture_modes = ['REQUEST','RESPONSE'] # In this example, we specify both
# capture_mode = ['REQUEST'] # Example - If you want to only capture input.

data_capture_config = DataCaptureConfig(
)```
Deploy your model

Deploy your model and create an HTTPS endpoint with DataCapture enabled.

AWS SDK for Python (Boto3)

Provide the endpoint configuration to SageMaker. The service launches the ML compute instances and deploys the model or models as specified in the configuration.

Once you have your model and endpoint configuration, use the `CreateEndpoint` API to create your endpoint. The endpoint name must be unique within an AWS Region in your AWS account.

The following creates an endpoint using the endpoint configuration specified in the request. Amazon SageMaker uses the endpoint to provision resources and deploy models.

```python
# The name of the endpoint. The name must be unique within an AWS Region in your AWS account.
endpoint_name = '<endpoint-name>'

# The name of the endpoint configuration associated with this endpoint.
endpoint_config_name='<endpoint-config-name>'

create_endpoint_response = sagemaker_client.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name)
```

For more information, see the `CreateEndpoint` API.

SageMaker Python SDK

Define a name for your endpoint. This step is optional. If you do not provide one, SageMaker will create a unique name for you:

```python
from datetime import datetime
```

Capture data from real-time endpoint
endpoint_name = f"DEMO-{datetime.utcnow():%Y-%m-%d-%H%M}"  
print("EndpointName =", endpoint_name)

Deploy your model to a real-time, HTTPS endpoint with the Model object's built-in deploy() method. Provide the name of the Amazon EC2 instance type to deploy this model to in the instance_type field along with the initial number of instances to run the endpoint on for the initial_instance_count field:

initial_instance_count=<integer>  
# initial_instance_count=1 # Example

instance_type='<instance-type>'  
# instance_type='ml.m4.xlarge' # Example

# Uncomment if you did not define this variable in the previous step  
data_capture_config = <name-of-data-capture-configuration>

model.deploy(  
    initial_instance_count=initial_instance_count,  
    instance_type=instance_type,  
    endpoint_name=endpoint_name,  
    data_capture_config=data_capture_config  
)

**View Captured Data**

Create a predictor object from the SageMaker Python SDK **Predictor** Class. You will use the object returned by the Predictor Class to invoke your endpoint in a future step. Provide the name of your endpoint (defined earlier as endpoint_name), along with serializer and deserializer objects for the serializer and deserializer, respectively. For information about serializer types, see the **Serializers** Class in the SageMaker Python SDK.

```python
from sagemaker.predictor import Predictor
from sagemaker.serializers import <Serializer>  
from sagemaker.deserializers import <Deserializer>

predictor = Predictor(endpoint_name=endpoint_name,  
        serializer = <Serializer_Class>,  
        deserializer = <Deserializer_Class>)
```
In the proceeding code example scenario we invoke the endpoint with sample validation data that we have stored locally in a CSV file named validation_with_predictions. Our sample validation set contains labels for each input.

The first few lines of the with statement first opens the validation set CSV file, then splits each row within the file by the comma character ",", and then stores the two returned objects into a label and input_cols variables. For each row, the input (input_cols) is passed to the predictor variable's (predictor) objects built-in method Predictor.predict().

Suppose the model returns a probability. Probabilities range between integer values of 0 and 1.0. If the probability returned by the model is greater than 80% (0.8) we assign the prediction an integer value label of 1. Otherwise, we assign the prediction an integer value label of 0.
Amazon SageMaker

Developer Guide

print(".", end="", flush=True)
sleep(0.5)
print()
print("Done!")

Because you enabled the data capture in the previous steps, the request and response payload,
along with some additional meta data, is saved in the Amazon S3 location that you speciﬁed in
DataCaptureConfig. The delivery of capture data to Amazon S3 can require a couple of minutes.
View captured data by listing the data capture ﬁles stored in Amazon S3. The format of the
Amazon S3 path is: s3:///{endpoint-name}/{variant-name}/yyyy/mm/dd/hh/
filename.jsonl.
Expect to see diﬀerent ﬁles from diﬀerent time periods, organized based on the hour when the
invocation occurred. Run the following to print out the contents of a single capture ﬁle:
print("\n".join(capture_file[-3:-1]))

This will return a SageMaker speciﬁc JSON-line formatted ﬁle. The following is a response sample
taken from a real-time endpoint that we invoked using csv/text data:

{"captureData":{"endpointInput":{"observedContentType":"text/csv","mode":"INPUT",
"data":"69,0,153.7,109,194.0,105,256.1,114,14.1,6,1,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0
"encoding":"CSV"},"endpointOutput":{"observedContentType":"text/csv;
charset=utf-8","mode":"OUTPUT","data":"0.0254181120544672","encoding":"CSV"}},
"eventMetadata":{"eventId":"aaaaaaaa-bbbb-cccc-ddddeeeeeeeeeeee","inferenceTime":"2022-02-14T17:25:49Z"},"eventVersion":"0"}
{"captureData":{"endpointInput":{"observedContentType":"text/csv","mode":"INPUT",
"data":"94,23,197.1,125,214.5,136,282.2,103,9.5,5,4,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0
"encoding":"CSV"},"endpointOutput":{"observedContentType":"text/csv;
charset=utf-8","mode":"OUTPUT","data":"0.07675473392009735","encoding":"CSV"}},
"eventMetadata":{"eventId":"aaaaaaaa-bbbb-cccc-ddddeeeeeeeeeeee","inferenceTime":"2022-02-14T17:25:49Z"},"eventVersion":"0"}

In the proceeding example, the capture_file object is a list type. Index the ﬁrst element of the
list to view a single inference request.
# The capture_file object is a list. Index the first element to view a single inference
request
print(json.dumps(json.loads(capture_file[0]), indent=2))

Capture data from real-time endpoint

5160


Capture data from batch transform job

The steps required to turn on data capture for your batch transform job are similar whether you use the AWS SDK for Python (Boto) or the SageMaker Python SDK. If you use the AWS SDK, define the `DataCaptureConfig` dictionary, along with required fields, within the `CreateTransformJob` method to turn on data capture. If you use the SageMaker Python SDK, import the `BatchDataCaptureConfig` class and initialize an instance from this class. Then, pass this object to the `batch_data_capture_config` parameter of your transform job instance.

To use the following code snippets, replace the italicized placeholder text in the example code with your own information.

How to enable data capture

Specify a data capture configuration when you launch a transform job. Whether you use the AWS SDK for Python (Boto3) or the SageMaker Python SDK, you must provide the `DestinationS3Uri`
argument, which is the directory where you want the transform job to log the captured data. Optionally, you can also set the following parameters:

- **KmsKeyId**: The AWS KMS key used to encrypt the captured data.
- **GenerateInferenceId**: A Boolean flag that, when capturing the data, indicates if you want the transform job to append the inference ID and time to your output. This is useful for model quality monitoring, where you need to ingest the Ground Truth data. The inference ID and time help to match the captured data with your Ground Truth data.

**AWS SDK for Python (Boto3)**

Configure the data you want to capture with the `DataCaptureConfig` dictionary when you create a transform job using the `CreateTransformJob` method.

```python
input_data_s3_uri = "s3://input_S3_uri"
output_data_s3_uri = "s3://output_S3_uri"
data_capture_destination = "s3://captured_data_S3_uri"

model_name = "model_name"

sm_client.create_transform_job(
    TransformJobName="transform_job_name",
    MaxConcurrentTransforms=2,
    ModelName=model_name,
    TransformInput={
        "DataSource": {
            "S3DataSource": {
                "S3DataType": "S3Prefix",
                "S3Uri": input_data_s3_uri,
            }
        },
        "ContentType": "text/csv",
        "CompressionType": "None",
        "SplitType": "Line",
    },
    TransformOutput={
        "S3OutputPath": output_data_s3_uri,
        "Accept": "text/csv",
        "AssembleWith": "Line",
    },
    TransformResources={
```
"InstanceType": "ml.m4.xlarge",
"InstanceCount": 1,
},
DataCaptureConfig={
  "DestinationS3Uri": data_capture_destination,
  "KmsKeyId": "kms_key",
  "GenerateInferenceId": True,
}
)}

SageMaker Python SDK

Import the BatchDataCaptureConfig class from the sagemaker.model_monitor.

```python
from sagemaker.transformer import Transformer
from sagemaker.inputs import BatchDataCaptureConfig

# Optional - The S3 URI of where to store captured data in S3
data_capture_destination = "s3://captured_data_S3_uri"

model_name = "model_name"

transformer = Transformer(model_name=model_name, ...)
transform_arg = transformer.transform(
    batch_data_capture_config=BatchDataCaptureConfig(
        destination_s3_uri=data_capture_destination,
        kms_key_id="kms_key",
        generate_inference_id=True,
    ),
    ...
)
```

How to view data captured

Once the transform job completes, the captured data gets logged under the DestinationS3Uri you provided with the data capture configuration. There are two subdirectories under DestinationS3Uri, /input and /output. If DestinationS3Uri is s3://my-data-capture, then the transform job creates the following directories:

- s3://my-data-capture/input: The captured input data for the transform job.
- s3://my-data-capture/output: The captured output data for the transform job.
To avoid data duplication, the captured data under the preceding two directories are manifests. Each manifest is a JSONL file that contains the Amazon S3 locations of the source objects. A manifest file may look like the following example:

```json
# under "/input" directory
[
   {"prefix":"s3://input_S3_uri/"},
   "dummy_0.csv",
   "dummy_1.csv",
   "dummy_2.csv",
   ...
]

# under "/output" directory
[
   {"prefix":"s3://output_S3_uri/"},
   "dummy_0.csv.out",
   "dummy_1.csv.out",
   "dummy_2.csv.out",
   ...
]
```

The transform job organizes and labels these manifests with a `yyyy/mm/dd/hh` S3 prefix to indicate when they were captured. This helps the model monitor determine the appropriate portion of data to analyze. For example, if you start your transform job at 2022-8-26 13PM UTC, then the captured data is labeled with a `2022/08/26/13/` prefix string.

**InferenceId Generation**

When you configure `DataCaptureConfig` for a transform job, you can turn on the Boolean flag `GenerateInferenceId`. This is particularly useful when you need to run model quality and model bias monitoring jobs, for which you need user-ingested Ground Truth data. Model monitor relies on an inference ID to match the captured data and the Ground Truth data. For addition details about Ground Truth ingestion, see [Ingest Ground Truth Labels and Merge Them With Predictions](#). When `GenerateInferenceId` is on, the transform output appends an inference ID (a random UUID) as well as the transform job start time in UTC for each record. You need these two values to run model quality and model bias monitoring. When you construct the Ground Truth data, you need to provide the same inference ID to match the output data. Currently, this feature supports transform outputs in CSV, JSON, and JSONL formats.

If your transform output is in CSV format, the output file looks like the following example:
Amazon SageMaker

Developer Guide

0, 1f1d57b1-2e6f-488c-8c30-db4e6d757861,2022-08-30T00:49:15Z
1, 22445434-0c67-45e9-bb4d-bd1bf26561e6,2022-08-30T00:49:15Z
...

The last two columns are inference ID and the transform job start time. Do not modify them. The
remaining columns are your transform job outputs.
If your transform output is in JSON or JSONL format, the output ﬁle looks like the following
example:
{"output": 0, "SageMakerInferenceId": "1f1d57b1-2e6f-488c-8c30-db4e6d757861",
"SageMakerInferenceTime": "2022-08-30T00:49:15Z"}
{"output": 1, "SageMakerInferenceId": "22445434-0c67-45e9-bb4d-bd1bf26561e6",
"SageMakerInferenceTime": "2022-08-30T00:49:15Z"}
...

There are two appended ﬁelds that are reserved, SageMakerInferenceId and
SageMakerInferenceTime. Do not modify these ﬁelds if you need to run model quality or model
bias monitoring — you need them for merge jobs.

Monitor data quality
Data quality monitoring automatically monitors machine learning (ML) models in production and
notiﬁes you when data quality issues arise. ML models in production have to make predictions on
real-life data that is not carefully curated like most training datasets. If the statistical nature of the
data that your model receives while in production drifts away from the nature of the baseline data
it was trained on, the model begins to lose accuracy in its predictions. Amazon SageMaker Model
Monitor uses rules to detect data drift and alerts you when it happens. To monitor data quality,
follow these steps:
• Enable data capture. This captures inference input and output from a real-time inference
endpoint or batch transform job and stores the data in Amazon S3. For more information, see
Capture data.
• Create a baseline. In this step, you run a baseline job that analyzes an input dataset that you
provide. The baseline computes baseline schema constraints and statistics for each feature using
Deequ, an open source library built on Apache Spark, which is used to measure data quality in
large datasets. For more information, see Create a Baseline.
Monitor data quality

5165


• Define and schedule data quality monitoring jobs. For specific information and code samples of data quality monitoring jobs, see Schedule data quality monitoring jobs. For general information about monitoring jobs, see Schedule monitoring jobs.

• Optionally use preprocessing and postprocessing scripts to transform the data coming out of your data quality analysis. For more information, see Preprocessing and Postprocessing.

• View data quality metrics. For more information, see Schema for Statistics (statistics.json file).

• Integrate data quality monitoring with Amazon CloudWatch. For more information, see CloudWatch Metrics.

• Interpret the results of a monitoring job. For more information, see Interpret results.

• Use SageMaker Studio to enable data quality monitoring and visualize results if you are using a real-time endpoint. For more information, see Visualize results for real-time endpoints in Amazon SageMaker Studio.

Note
Model Monitor computes model metrics and statistics on tabular data only. For example, an image classification model that takes images as input and outputs a label based on that image can still be monitored. Model Monitor would be able to calculate metrics and statistics for the output, not the input.

Topics
• Create a Baseline
• Schedule data quality monitoring jobs
• Schema for Statistics (statistics.json file)
• CloudWatch Metrics
• Schema for Violations (constraint_violations.json file)

Create a Baseline

The baseline calculations of statistics and constraints are needed as a standard against which data drift and other data quality issues can be detected. Model Monitor provides a built-in container that provides the ability to suggest the constraints automatically for CSV and flat JSON input. This sagemaker-model-monitor-analyzer container also provides you with a range of model monitoring
capabilities, including constraint validation against a baseline, and emitting Amazon CloudWatch metrics. This container is based on Spark version 3.3.0 and is built with Deequ version 2.0.2. All column names in your baseline dataset must be compliant with Spark. For column names, use only lowercase characters, and _ as the only special character.

The training dataset that you used to train the model is usually a good baseline dataset. The training dataset data schema and the inference dataset schema should exactly match (the number and order of the features). Note that the prediction/output columns are assumed to be the first columns in the training dataset. From the training dataset, you can ask SageMaker to suggest a set of baseline constraints and generate descriptive statistics to explore the data. For this example, upload the training dataset that was used to train the pretrained model included in this example. If you already stored the training dataset in Amazon S3, you can point to it directly.

**To Create a baseline from a training dataset**

When you have your training data ready and stored in Amazon S3, start a baseline processing job with DefaultModelMonitor.suggest_baseline(..) using the Amazon SageMaker Python SDK. This uses an Amazon SageMaker Model Monitor prebuilt container that generates baseline statistics and suggests baseline constraints for the dataset and writes them to the output_s3_uri location that you specify.

```python
from sagemaker.model_monitor import DefaultModelMonitor
from sagemaker.model_monitor.dataset_format import DatasetFormat

my_default_monitor = DefaultModelMonitor(
    role=role,
    instance_count=1,
    instance_type='ml.m5.xlarge',
    volume_size_in_gb=20,
    max_runtime_in_seconds=3600,
)

my_default_monitor.suggest_baseline(
    baseline_dataset=baseline_data_uri+'/training-dataset-with-header.csv',
    dataset_format=DatasetFormat.csv(header=True),
    output_s3_uri=baseline_results_uri,
    wait=True
)
```
Note

If you provide the feature/column names in the training dataset as the first row and set the `header=True` option as shown in the previous code sample, SageMaker uses the feature name in the constraints and statistics file.

The baseline statistics for the dataset are contained in the `statistics.json` file and the suggested baseline constraints are contained in the `constraints.json` file in the location you specify with `output_s3_uri`.

Output Files for Tabular Dataset Statistics and Constraints

<table>
<thead>
<tr>
<th>File Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>statistics.json</code></td>
<td>This file is expected to have columnar statistics for each feature in the dataset that is analyzed. For more information about the schema for this file, see [Schema for Statistics](statistics.json file).</td>
</tr>
<tr>
<td><code>constraints.json</code></td>
<td>This file is expected to have the constraints on the features observed. For more information about the schema for this file, see [Schema for Constraints](constraints.json file).</td>
</tr>
</tbody>
</table>

The Amazon SageMaker Python SDK provides convenience functions described to generate the baseline statistics and constraints. But if you want to call processing job directly for this purpose instead, you need to set the `Environment` map as shown in the following example:

```json
"Environment": {
    "dataset_format": "{"csv": { "header": true}}",
    "dataset_source": "/opt/ml/processing/sm_input",
    "output_path": "/opt/ml/processing/sm_output",
    "publish_cloudwatch_metrics": "Disabled",
}
```
Schedule data quality monitoring jobs

After you create your baseline, you can call the `create_monitoring_schedule()` method of your `DefaultModelMonitor` class instance to schedule an hourly data quality monitor. The following sections show you how to create a data quality monitor for a model deployed to a real-time endpoint as well as for a batch transform job.

⚠️ **Important**

You can specify either a batch transform input or an endpoint input, but not both, when you create your monitoring schedule.

Data quality monitoring for models deployed to real-time endpoints

To schedule a data quality monitor for a real-time endpoint, pass your `EndpointInput` instance to the `endpoint_input` argument of your `DefaultModelMonitor` instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

data_quality_model_monitor = DefaultModelMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

schedule = data_quality_model_monitor.create_monitoring_schedule(
    monitor_schedule_name=schedule_name,
    post_analytics_processor_script=s3_code_postprocessor_uri,
    output_s3_uri=s3_report_path,
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    statistics=data_quality_model_monitor.baseline_statistics(),
    constraints=data_quality_model_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
    endpoint_input=EndpointInput(
        endpoint_name=endpoint_name,
        destination="/opt/ml/processing/input/endpoint",
    ),
)
```
Data quality monitoring for batch transform jobs

To schedule a data quality monitor for a batch transform job, pass your BatchTransformInput instance to the batch_transform_input argument of your DefaultModelMonitor instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

data_quality_model_monitor = DefaultModelMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

schedule = data_quality_model_monitor.create_monitoring_schedule(
    monitor_schedule_name=mon_schedule_name,
    batch_transform_input=BatchTransformInput(
        data_captured_destination_s3_uri=s3_capture_upload_path,
        destination="/opt/ml/processing/input",
        dataset_format=MonitoringDatasetFormat.csv(header=False),
    ),
    output_s3_uri=s3_report_path,
    statistics= statistics_path,
    constraints = constraints_path,
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
)

Schema for Statistics (statistics.json file)

Amazon SageMaker Model Monitor prebuilt container computes per column/feature statistics. The statistics are calculated for the baseline dataset and also for the current dataset that is being analyzed.

```json
{
    "version": 0,
    # dataset level stats
    "dataset": {
        "item_count": number
    },
    # feature level stats
    "features": [
        
```
"name": "feature-name",
"inferred_type": "Fractional" | "Integral",
"numerical_statistics": {
  "common": {
    "num_present": number,
    "num_missing": number
  },
  "mean": number,
  "sum": number,
  "std_dev": number,
  "min": number,
  "max": number,
  "distribution": {
    "kll": {
      "buckets": [
        {
          "lower_bound": number,
          "upper_bound": number,
          "count": number
        }
      ],
      "sketch": {
        "parameters": {
          "c": number,
          "k": number
        },
        "data": [
          [num, num, num, num],
          [num, num]
        ]
      }
    }
  }
}
Note the following:

- The prebuilt containers compute KLL sketch, which is a compact quantiles sketch.
- By default, we materialize the distribution in 10 buckets. This is not currently configurable.

**CloudWatch Metrics**

You can use the built-in Amazon SageMaker Model Monitor container for CloudWatch metrics. When the emit_metrics option is Enabled in the baseline constraints file, SageMaker emits these metrics for each feature/column observed in the dataset in the following namespace:

- For real-time endpoints: /aws/sagemaker/Endpoints/data-metric namespace with EndpointName and ScheduleName dimensions.
For batch transform jobs: `/aws/sagemaker/ModelMonitoring/data-metric` namespace with `MonitoringSchedule` dimension.

For numerical fields, the built-in container emits the following CloudWatch metrics:

- Metric: Max → query for MetricName: `feature_data_{feature_name}`, Stat: Max
- Metric: Min → query for MetricName: `feature_data_{feature_name}`, Stat: Min
- Metric: Sum → query for MetricName: `feature_data_{feature_name}`, Stat: Sum
- Metric: SampleCount → query for MetricName: `feature_data_{feature_name}`, Stat: SampleCount
- Metric: Average → query for MetricName: `feature_data_{feature_name}`, Stat: Average

For both numerical and string fields, the built-in container emits the following CloudWatch metrics:

- Metric: Completeness → query for MetricName: `feature_non_null_{feature_name}`, Stat: Sum
- Metric: Baseline Drift → query for MetricName: `feature_baseline_drift_{feature_name}`, Stat: Sum

**Schema for Violations (constraint_violations.json file)**

The violations file is generated as the output of a `MonitoringExecution`, which lists the results of evaluating the constraints (specified in the `constraints.json` file) against the current dataset that was analyzed. The Amazon SageMaker Model Monitor prebuilt container provides the following violation checks.

```json
{
    "violations": [
        {
            "feature_name" : "string",
            "constraint_check_type" : 
                "data_type_check",
            "completeness_check",
            "baseline_drift_check",
            "missing_column_check",
            "extra_column_check",
            "categorical_values_check"
        }
    ]
}
```
### Types of Violations Monitored

<table>
<thead>
<tr>
<th>Violation Check Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data_type_check</td>
<td>If the data types in the current execution are not the same as in the baseline dataset, this violation is flagged. During the baseline step, the generated constraints suggest the inferred data type for each column. The <code>monitoring_config.datatype_check_threshold</code> parameter can be tuned to adjust the threshold on when it is flagged as a violation.</td>
</tr>
<tr>
<td>completeness_check</td>
<td>If the completeness (% of non-null items) observed in the current execution exceeds the threshold specified in <code>completeness_threshold</code> specified per feature, this violation is flagged. During the baseline step, the generated constraints suggest a completeness value.</td>
</tr>
<tr>
<td>baseline_drift_check</td>
<td>If the calculated distribution distance between the current and the baseline datasets is more than the threshold specified in <code>monitoring_config.comparison_threshold</code>, this violation is flagged.</td>
</tr>
<tr>
<td>missing_column_check</td>
<td>If the number of columns in the current dataset is less than the number in the baseline dataset, this violation is flagged.</td>
</tr>
</tbody>
</table>
## Monitor model quality

Model quality monitoring jobs monitor the performance of a model by comparing the predictions that the model makes with the actual Ground Truth labels that the model attempts to predict. To do this, model quality monitoring merges data that is captured from real-time or batch inference with actual labels that you store in an Amazon S3 bucket, and then compares the predictions with the actual labels.

To measure model quality, model monitor uses metrics that depend on the ML problem type. For example, if your model is for a regression problem, one of the metrics evaluated is mean square error (mse). For information about all of the metrics used for the different ML problem types, see [Model Quality Metrics](#).

Model quality monitoring follows the same steps as data quality monitoring, but adds the additional step of merging the actual labels from Amazon S3 with the predictions captured from the real-time inference endpoint or batch transform job. To monitor model quality, follow these steps:

- Enable data capture. This captures inference input and output from a real-time inference endpoint or batch transform job and stores the data in Amazon S3. For more information, see [Capture data](#).
- Create a baseline. In this step, you run a baseline job that compares predictions from the model with Ground Truth labels in a baseline dataset. The baseline job automatically creates baseline statistical rules and constraints that define thresholds against which the model performance is evaluated. For more information, see [Create a Model Quality Baseline](#).
• Define and schedule model quality monitoring jobs. For specific information and code samples of model quality monitoring jobs, see [Schedule Model Quality Monitoring Jobs](#). For general information about monitoring jobs, see [Schedule monitoring jobs](#).

• Ingest Ground Truth labels that model monitor merges with captured prediction data from a real-time inference endpoint or batch transform job. For more information, see [Ingest Ground Truth Labels and Merge Them With Predictions](#).

• Integrate model quality monitoring with Amazon CloudWatch. For more information, see [Model Quality CloudWatch Metrics](#).

• Interpret the results of a monitoring job. For more information, see [Interpret results](#).

• Use SageMaker Studio to enable model quality monitoring and visualize results. For more information, see [Visualize results for real-time endpoints in Amazon SageMaker Studio](#).

Topics

• [Create a Model Quality Baseline](#)

• [Schedule Model Quality Monitoring Jobs](#)

• [Ingest Ground Truth Labels and Merge Them With Predictions](#)

• [Model Quality Metrics](#)

• [Model Quality CloudWatch Metrics](#)

Create a Model Quality Baseline

Create a baseline job that compares your model predictions with ground truth labels in a baseline dataset that you have stored in Amazon S3. Typically, you use a training dataset as the baseline dataset. The baseline job calculates metrics for the model and suggests constraints to use to monitor model quality drift.

To create a baseline job, you need to have a dataset that contains predictions from your model along with labels that represent the Ground Truth for your data.

To create a baseline job use the `ModelQualityMonitor` class provided by the SageMaker Python SDK, and complete the following steps.

**To create a model quality baseline job**

1. First, create an instance of the `ModelQualityMonitor` class. The following code snippet shows how to do this.
from sagemaker import get_execution_role, session, Session
from sagemaker.model_monitor import ModelQualityMonitor

role = get_execution_role()
session = Session()

model_quality_monitor = ModelQualityMonitor(
    role=role,
    instance_count=1,
    instance_type='ml.m5.xlarge',
    volume_size_in_gb=20,
    max_runtime_in_seconds=1800,
    sagemaker_session=session
)

2. Now call the `suggest_baseline` method of the `ModelQualityMonitor` object to run a baseline job. The following code snippet assumes that you have a baseline dataset that contains both predictions and labels stored in Amazon S3.

```python
baseline_job_name = "MyBaselineJob"
job = model_quality_monitor.suggest_baseline(
    job_name=baseline_job_name,
    baseline_dataset=baseline_dataset_uri, # The S3 location of the validation dataset.
    dataset_format=DatasetFormat.csv(header=True),
    output_s3_uri = baseline_results_uri, # The S3 location to store the results.
    problem_type='BinaryClassification',
    inference_attribute= "prediction", # The column in the dataset that contains predictions.
    probability_attribute= "probability", # The column in the dataset that contains probabilities.
    ground_truth_attribute= "label" # The column in the dataset that contains ground truth labels.
)
job.wait(logs=False)
```

3. After the baseline job finishes, you can see the constraints that the job generated. First, get the results of the baseline job by calling the `latest_baselining_job` method of the `ModelQualityMonitor` object.

```python
baseline_job = model_quality_monitor.latest_baselining_job
```
4. The baseline job suggests constraints, which are thresholds for metrics that model monitor measures. If a metric goes beyond the suggested threshold, Model Monitor reports a violation. To view the constraints that the baseline job generated, call the suggested_constraints method of the baseline job. The following code snippet loads the constraints for a binary classification model into a Pandas dataframe.

```python
import pandas as pd
pd.DataFrame(baseline_job.suggested_constraints().body_dict["binary_classification_constraints"]).T
```

We recommend that you view the generated constraints and modify them as necessary before using them for monitoring. For example, if a constraint is too aggressive, you might get more alerts for violations than you want.

If your constraint contains numbers expressed in scientific notation, you will need to convert them to float. The following python preprocessing script example shows how to convert numbers in scientific notation to float.

```python
import csv

def fix_scientific_notation(col):
    try:
        return format(float(col), "f")
    except:
        return col

def preprocess_handler(csv_line):
    reader = csv.reader([csv_line])
    csv_record = next(reader)
    #skip baseline header, change HEADER_NAME to the first column's name
    if csv_record[0] == "HEADER_NAME":
        return []
    return { str(i).zfill(20) : fix_scientific_notation(d) for i, d in enumerate(csv_record)}
```

You can add your pre-processing script to a baseline or monitoring schedule as a record_preprocessor_script, as defined in the Model Monitor documentation.

5. When you are satisfied with the constraints, pass them as the constraints parameter when you create a monitoring schedule. For more information, see Schedule Model Quality Monitoring Jobs.
The suggested baseline constraints are contained in the constraints.json file in the location you specify with `output_s3_uri`. For information about the schema for this file in the `Schema for Constraints (constraints.json file)`.

**Schedule Model Quality Monitoring Jobs**

After you create your baseline, you can call the `create_monitoring_schedule()` method of your `ModelQualityMonitor` class instance to schedule an hourly model quality monitor. The following sections show you how to create a model quality monitor for a model deployed to a real-time endpoint as well as for a batch transform job.

⚠️ **Important**

You can specify either a batch transform input or an endpoint input, but not both, when you create your monitoring schedule.

Unlike data quality monitoring, you need to supply Ground Truth labels if you want to monitor model quality. However, Ground Truth labels could be delayed. To address this, specify offsets when you create your monitoring schedule.

**Model monitor offsets**

Model quality jobs include `StartTimeOffset` and `EndTimeOffset`, which are fields of the `ModelQualityJobInput` parameter of the `create_model_quality_job_definition` method that work as follows:

- **StartTimeOffset** - If specified, jobs subtract this time from the start time.
- **EndTimeOffset** - If specified, jobs subtract this time from the end time.

The format of the offsets are, for example, `-PT7H`, where 7H is 7 hours. You can use `-PT#H` or `-P#D`, where H=hours, D=days, and M=minutes, and # is the number. In addition, the offset should be in ISO 8601 duration format.

For example, if your Ground Truth starts coming in after 1 day, but is not complete for a week, set `StartTimeOffset` to `-P8D` and `EndTimeOffset` to `-P1D`. Then, if you schedule a job to run at 2020-01-09T13:00, it analyzes data from between 2020-01-01T13:00 and 2020-01-08T13:00.
Important

The schedule cadence should be such that one execution finishes before the next execution starts, which allows the Ground Truth merge job and monitoring job from the execution to complete. The maximum runtime of an execution is divided between the two jobs, so for an hourly model quality monitoring job, the value of `MaxRuntimeInSeconds` specified as part of `StoppingCondition` should be no more than 1800.

Model quality monitoring for models deployed to real-time endpoints

To schedule a model quality monitor for a real-time endpoint, pass your `EndpointInput` instance to the `endpoint_input` argument of your `ModelQualityMonitor` instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

model_quality_model_monitor = ModelQualityMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

schedule = model_quality_model_monitor.create_monitoring_schedule(
    monitor_schedule_name=schedule_name,
    post_analytics_processor_script=s3_code_postprocessor_uri,
    output_s3_uri=s3_report_path,
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    statistics=model_quality_model_monitor.baseline_statistics(),
    constraints=model_quality_model_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
    endpoint_input=EndpointInput(
        endpoint_name=endpoint_name,
        destination="/opt/ml/processing/input/endpoint",
        start_time_offset="-PT2D",
        end_time_offset="-PT1D",
    )
)
```
Model quality monitoring for batch transform jobs

To schedule a model quality monitor for a batch transform job, pass your BatchTransformInput instance to the batch_transform_input argument of your ModelQualityMonitor instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

model_quality_model_monitor = ModelQualityMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

schedule = model_quality_model_monitor.create_monitoring_schedule(
    monitor_schedule_name=mon_schedule_name,
    batch_transform_input=BatchTransformInput(
        data_captured_destination_s3_uri=s3_capture_upload_path,
        destination="/opt/ml/processing/input",
        dataset_format=MonitoringDatasetFormat.csv(header=False),
        # the column index of the output representing the inference probablity
        probability_attribute="0",
        # the threshold to classify the inference probablity to class 0 or 1 in
        # binary classification problem
        probability_threshold_attribute=0.5,
        # look back 6 hour for transform job outputs.
        start_time_offset="-PT6H",
        end_time_offset="-PT0H"
    ),
    ground_truth_input=gt_s3_uri,
    output_s3_uri=s3_report_path,
    problem_type="BinaryClassification",
    constraints = constraints_path,
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
)
```

Ingest Ground Truth Labels and Merge Them With Predictions

Model quality monitoring compares the predictions your model makes with ground truth labels to measure the quality of the model. For this to work, you periodically label data captured by your endpoint or batch transform job and upload it to Amazon S3.
To match Ground Truth labels with captured prediction data, there must be a unique identifier for each record in the dataset. The structure of each record for ground truth data is as follows:

```json
{
    "groundTruthData": {
        "data": "1",
        "encoding": "CSV" # only CSV supported at launch, we assume "data" only consists of label
    },
    "eventMetadata": {
        "eventId": "aaaa-bbbb-cccc"
    },
    "eventVersion": "0"
}
```

In the `groundTruthData` structure, `eventId` can be one of the following:

- `eventId` – This ID is automatically generated when a user invokes the endpoint.
- `inferenceId` – The caller supplies this ID when they invoke the endpoint.

If `inferenceId` is present in captured data records, Model Monitor uses it to merge captured data with Ground Truth records. You are responsible for making sure that the `inferenceId` in the Ground Truth records match the `inferenceId` in the captured records. If `inferenceId` is not present in captured data, model monitor uses `eventId` from the captured data records to match them with a Ground Truth record.

You must upload Ground Truth data to an Amazon S3 bucket that has the same path format as captured data, which is of the following form:

```
s3://bucket/prefix/yyyy/mm/dd/hh
```

The date in this path is the date when the Ground Truth label is collected, and does not have to match the date when the inference was generated.

After you create and upload the Ground Truth labels, include the location of the labels as a parameter when you create the monitoring job. If you are using AWS SDK for Python (Boto3), do this by specifying the location of Ground Truth labels as the `S3Uri` field of the `GroundTruthS3Input` parameter in a call to the `create_model_quality_job_definition` method. If you are using the SageMaker Python SDK, specify the location of the
Ground Truth labels as the `ground_truth_input` parameter in the call to the `create_monitoring_schedule` of the `ModelQualityMonitor` object.

**Model Quality Metrics**

Model quality monitoring jobs compute different metrics depending on the ML problem type. The following sections list the metrics analyzed for each ML problem type.

**Note**

Standard deviation for metrics are provided only when at least 200 samples are available. Model Monitor computes standard deviation by randomly sampling 80% of the data 5 times, computing the metric, and taking the standard deviation for those results.

**Regression Metrics**

The following shows an example of the metrics that model quality monitor computes for a regression problem.

```json
"regression_metrics" : {
   "mae" : {
      "value" : 0.3711832061068702,
      "standard_deviation" : 0.0037566388129940394
   },
   "mse" : {
      "value" : 0.3711832061068702,
      "standard_deviation" : 0.0037566388129940524
   },
   "rmse" : {
      "value" : 0.609248066149471,
      "standard_deviation" : 0.003079253267651125
   },
   "r2" : {
      "value" : -1.3766111872212665,
      "standard_deviation" : 0.022653980022771227
   }
}
```
Binary Classification Metrics

The following shows an example of the metrics that model quality monitor computes for a binary classification problem.

```
"binary_classification_metrics" : {
    "confusion_matrix" : {
        "0" : {
            "0" : 1,
            "1" : 2
        },
        "1" : {
            "0" : 0,
            "1" : 1
        }
    },
    "recall" : {
        "value" : 1.0,
        "standard_deviation" : "NaN"
    },
    "precision" : {
        "value" : 0.3333333333333333,
        "standard_deviation" : "NaN"
    },
    "accuracy" : {
        "value" : 0.5,
        "standard_deviation" : "NaN"
    },
    "recall_best_constant_classifier" : {
        "value" : 1.0,
        "standard_deviation" : "NaN"
    },
    "precision_best_constant_classifier" : {
        "value" : 0.25,
        "standard_deviation" : "NaN"
    },
    "accuracy_best_constant_classifier" : {
        "value" : 0.25,
        "standard_deviation" : "NaN"
    },
    "true_positive_rate" : {
        "value" : 1.0,
        "standard_deviation" : "NaN"
    }
}
```
"true_negative_rate": {
  "value": 0.33333333333333337,
  "standard_deviation": "NaN"
},
"false_positive_rate": {
  "value": 0.6666666666666666,
  "standard_deviation": "NaN"
},
"false_negative_rate": {
  "value": 0.0,
  "standard_deviation": "NaN"
},
"receiver_operating_characteristic_curve": {
  "false_positive_rates": [0.0, 0.0, 0.0, 0.0, 0.0, 1.0],
  "true_positive_rates": [0.0, 0.25, 0.5, 0.75, 1.0, 1.0]
},
"precision_recall_curve": {
  "precisions": [1.0, 1.0, 1.0, 1.0, 1.0]
},
"auc": {
  "value": 1.0,
  "standard_deviation": "NaN"
},
"f0_5": {
  "value": 0.3846153846153846,
  "standard_deviation": "NaN"
},
"f1": {
  "value": 0.5,
  "standard_deviation": "NaN"
},
"f2": {
  "value": 0.7142857142857143,
  "standard_deviation": "NaN"
},
"f0_5_best_constant_classifier": {
  "value": 0.29411764705882354,
  "standard_deviation": "NaN"
},
"f1_best_constant_classifier": {
  "value": 0.4,
  "standard_deviation": "NaN"}
Multiclass Metrics

The following shows an example of the metrics that model quality monitor computes for a multiclass classification problem.

```
"multiclass_classification_metrics" : {
    "confusion_matrix" : {
        "0" : {
            "0" : 1180,
            "1" : 510
        },
        "1" : {
            "0" : 268,
            "1" : 138
        }
    },
    "accuracy" : {
        "value" : 0.6288167938931297,
        "standard_deviation" : 0.00375663881299405
    },
    "weighted_recall" : {
        "value" : 0.6288167938931297,
        "standard_deviation" : 0.003756638812994008
    },
    "weighted_precision" : {
        "value" : 0.6983172269629505,
        "standard_deviation" : 0.006195912915307507
    },
    "weighted_f0_5" : {
        "value" : 0.6803947317178771,
        "standard_deviation" : 0.005328406973561699
    },
    "weighted_f1" : {
        "value" : 0.6571162346664904,
        "standard_deviation" : 0.004385008075019733
    }
}
```
"weighted_f2" : {
    "value" : 0.6384024354394601,
    "standard_deviation" : 0.003867109755267757
},
"accuracy_best_constant_classifier" : {
    "value" : 0.19370229007633588,
    "standard_deviation" : 0.0032049848450732355
},
"weighted_recall_best_constant_classifier" : {
    "value" : 0.19370229007633588,
    "standard_deviation" : 0.0032049848450732355
},
"weighted_precision_best_constant_classifier" : {
    "value" : 0.03752057718081697,
    "standard_deviation" : 0.001241536088657851
},
"weighted_f0_5_best_constant_classifier" : {
    "value" : 0.04473443104152011,
    "standard_deviation" : 0.0014460485504284792
},
"weighted_f1_best_constant_classifier" : {
    "value" : 0.06286421244683643,
    "standard_deviation" : 0.0019113576884608862
},
"weighted_f2_best_constant_classifier" : {
    "value" : 0.10570313141262414,
    "standard_deviation" : 0.002734216826748117
}

**Model Quality CloudWatch Metrics**

If you set the value of the `enable_cloudwatch_metrics` to True when you create the monitoring schedule, model quality monitoring jobs send all metrics to Amazon CloudWatch.

Model quality metrics appear in the following namespace:

- For real-time endpoints: `aws/sagemaker/Endpoints/model-metrics`
- For batch transform jobs: `aws/sagemaker/ModelMonitoring/model-metrics`

For a list of the metrics that are emitted, see [Model Quality Metrics](#).
You can use CloudWatch metrics to create an alarm when a specific metric doesn't meet the threshold you specify. For instructions about how to create CloudWatch alarms, see Create a CloudWatch Alarm Based on a Static Threshold in the Amazon CloudWatch User Guide.

Monitor Bias Drift for Models in Production

Amazon SageMaker Clarify bias monitoring helps data scientists and ML engineers monitor predictions for bias on a regular basis. As the model is monitored, customers can view exportable reports and graphs detailing bias in SageMaker Studio and configure alerts in Amazon CloudWatch to receive notifications if bias beyond a certain threshold is detected. Bias can be introduced or exacerbated in deployed ML models when the training data differs from the data that the model sees during deployment (that is, the live data). These kinds of changes in the live data distribution might be temporary (for example, due to some short-lived, real-world events) or permanent. In either case, it might be important to detect these changes. For example, the outputs of a model for predicting home prices can become biased if the mortgage rates used to train the model differ from current, real-world mortgage rates. With bias detection capabilities in Model Monitor, when SageMaker detects bias beyond a certain threshold, it automatically generates metrics that you can view in SageMaker Studio and through Amazon CloudWatch alerts.

In general, measuring bias only during the train-and-deploy phase might not be sufficient. It is possible that after the model has been deployed, the distribution of the data that the deployed model sees (that is, the live data) is different from data distribution in the training dataset. This change might introduce bias in a model over time. The change in the live data distribution might be temporary (for example, due to some short-lived behavior like the holiday season) or permanent. In either case, it might be important to detect these changes and take steps to reduce the bias when appropriate.

To detect these changes, SageMaker Clarify provides functionality to monitor the bias metrics of a deployed model continuously and raise automated alerts if the metrics exceed a threshold. For example, consider the DPPL bias metric. Specify an allowed range of values $A=(a_{\min}, a_{\max})$, for instance an interval of $(-0.1, 0.1)$, that DPPL should belong to during deployment. Any deviation from this range should raise a bias detected alert. With SageMaker Clarify, you can perform these checks at regular intervals.

For example, you can set the frequency of the checks to 2 days. This means that SageMaker Clarify computes the DPPL metric on data collected during a 2-day window. In this example, $D_{\text{win}}$ is the data that the model processed during last 2-day window. An alert is issued if the DPPL value $b_{\text{win}}$ computed on $D_{\text{win}}$ falls outside of an allowed range $A$. This approach to checking if
$b_{\text{win}}$ is outside of $A$ can be somewhat noisy. $D_{\text{win}}$ might consist of very few samples and might not be representative of the live data distribution. The small sample size means that the value of bias $b_{\text{win}}$ computed over $D_{\text{win}}$ might not be a very robust estimate. In fact, very high (or low) values of $b_{\text{win}}$ may be observed purely due to chance. To ensure that the conclusions drawn from the observed data $D_{\text{win}}$ are statistically significant, SageMaker Clarify makes use of confidence intervals. Specifically, it uses the Normal Bootstrap Interval method to construct an interval $C=(c_{\text{min}},c_{\text{max}})$ such that SageMaker Clarify is confident that the true bias value computed over the full live data is contained in $C$ with high probability. Now, if the confidence interval $C$ overlaps with the allowed range $A$, SageMaker Clarify interprets it as “it is likely that the bias metric value of the live data distribution falls within the allowed range”. If $C$ and $A$ are disjoint, SageMaker Clarify is confident that the bias metric does not lie in $A$ and raises an alert.

**Model Monitor Sample Notebook**

Amazon SageMaker Clarify provides the following sample notebook that shows how to capture inference data for a real-time endpoint, create a baseline to monitor evolving bias against, and inspect the results:

- [Monitoring bias drift and feature attribution drift Amazon SageMaker Clarify](#) – Use Amazon SageMaker Model Monitor to monitor bias drift and feature attribution drift over time.

This notebook has been verified to run in Amazon SageMaker Studio only. If you need instructions on how to open a notebook in Amazon SageMaker Studio, see [Create or Open an Amazon SageMaker Studio Classic Notebook](#). If you’re prompted to choose a kernel, choose [Python 3 (Data Science)](#). The following topics contain the highlights from the last two steps, and they contain code examples from the example notebook.

**Topics**

- [Create a Bias Drift Baseline](#)
- [Bias Drift Violations](#)
- [Configure Parameters to Monitor Bias Drift](#)
- [Schedule Bias Drift Monitoring Jobs](#)
- [Inspect Reports for Data Bias Drift](#)
- [CloudWatch Metrics for Bias Drift Analysis](#)
Create a Bias Drift Baseline

After you have configured your application to capture real-time or batch transform inference data, the first task to monitor for bias drift is to create a baseline. This involves configuring the data inputs, which groups are sensitive, how the predictions are captured, and the model and its post-training bias metrics. Then you need to start the baselining job.

Model bias monitor can detect bias drift of ML models on a regular basis. Similar to the other monitoring types, the standard procedure of creating a model bias monitor is first baselining and then establishing a monitoring schedule.

```python
model_bias_monitor = ModelBiasMonitor(
    role=role,
    sagemaker_session=sagemaker_session,
    max_runtime_in_seconds=1800,
)
```

DataConfig stores information about the dataset to be analyzed (for example, the dataset file), its format (that is, CSV or JSON Lines), headers (if any) and label.

```python
model_bias_baselining_job_result_uri = f"{baseline_results_uri}/model_bias"
model_bias_data_config = DataConfig(
    s3_data_input_path=validation_dataset,
    s3_output_path=model_bias_baselining_job_result_uri,
    label=label_header,
    headers=all_headers,
    dataset_type=dataset_type,
)
```

BiasConfig is the configuration of the sensitive groups in the dataset. Typically, bias is measured by computing a metric and comparing it across groups. The group of interest is called the facet. For post-training bias, you should also take the positive label into account.

```python
model_bias_config = BiasConfig(
    label_values_or_threshold=[1],
    facet_name="Account Length",
    facet_values_or_threshold=[100],
)
```
ModelPredictedLabelConfig specifies how to extract a predicted label from the model output. In this example, the 0.8 cutoff has been chosen in anticipation that customers will turn over frequently. For more complicated outputs, there are a few more options, like "label" is the index, name, or JMESPath to locate predicted label in endpoint response payload.

```python
model_predicted_label_config = ModelPredictedLabelConfig(
    probability_threshold=0.8,
)
```

ModelConfig is the configuration related to the model to be used for inferencing. In order to compute post-training bias metrics, the computation needs to get inferences for the model name provided. To accomplish this, the processing job uses the model to create an ephemeral endpoint (also known as *shadow endpoint*). The processing job deletes the shadow endpoint after the computations are completed. This configuration is also used by the explainability monitor.

```python
model_config = ModelConfig(
    model_name=model_name,
    instance_count=endpoint_instance_count,
    instance_type=endpoint_instance_type,
    content_type=dataset_type,
    accept_type=dataset_type,
)
```

Now you can start the baselining job.

```python
model_bias_monitor.suggest_baseline(
    model_config=model_config,
    data_config=model_bias_data_config,
    bias_config=model_bias_config,
    model_predicted_label_config=model_predicted_label_config,
)
print(f"ModelBiasMonitor baselining job: {model_bias_monitor.latest_baselining_job_name}")
```

The scheduled monitor automatically picks up baselining job name and waits for it before monitoring begins.
Bias Drift Violations

Bias drift jobs evaluate the baseline constraints provided by the baseline configuration against the analysis results of current MonitoringExecution. If violations are detected, the job lists them to the constraint_violations.json file in the execution output location, and marks the execution status as Interpret results.

Here is the schema of the bias drift violations file.

- facet – The name of the facet, provided by the monitoring job analysis configuration facet name_or_index.
- facet_value – The value of the facet, provided by the monitoring job analysis configuration facet value_or_threshold.
- metric_name – The short name of the bias metric. For example, "CI" for class imbalance. See Measure Pre-training Bias for the short names of each of the pre-training bias metrics and Measure Post-training Data and Model Bias for the short names of each of the post-training bias metrics.
- constraint_check_type – The type of violation monitored. Currently only bias_drift_check is supported.
- description – A descriptive message to explain the violation.

```json
{
  "version": "1.0",
  "violations": [
    {
      "facet": "string",
      "facet_value": "string",
      "metric_name": "string",
      "constraint_check_type": "string",
      "description": "string"
    }
  ]
}
```

A bias metric is used to measure the level of equality in a distribution. A value close to zero indicates that the distribution is more balanced. If the value of a bias metric in the job analysis results file (analysis.json) is worse than its corresponding value in the baseline constraints file, a violation is logged. As an example, if the baseline constraint for the DPPL bias metric is 0.2, and the analysis result is 0.1, no violation is logged because 0.1 is closer to 0 than 0.2. However,
if the analysis result is -0.3, a violation is logged because it is farther from 0 than the baseline constraint of 0.2.

```
{
    "version": "1.0",
    "violations": [ [
        "facet": "Age",
        "facet_value": "40",
        "metric_name": "CI",
        "constraint_check_type": "bias_drift_check",
        "description": "Value 0.0751544567666083 does not meet the constraint requirement"
    ], [
        "facet": "Age",
        "facet_value": "40",
        "metric_name": "DPPL",
        "constraint_check_type": "bias_drift_check",
        "description": "Value -0.0791244970125596 does not meet the constraint requirement"
    ]
}
```

**Configure Parameters to Monitor Bias Drift**

Amazon SageMaker Clarify bias monitoring reuses a subset of the parameters used in the analysis configuration of Configure the Analysis. After describing the configuration parameters, this topic provides examples of JSON files. These files are used to configure CSV and JSON Lines datasets to monitor them for bias drift when machine learning models are in production.

The following parameters must be provided in a JSON file. The path to this JSON file must be provided in the ConfigUri parameter of the ModelBiasAppSpecification API.

- "**version**" – (Optional) Schema version of the configuration file. If not provided, the latest supported version is used.
- "**headers**" – (Optional) A list of column names in the dataset. If the dataset_type is "application/jsonlines" and "label" is specified, then the last header becomes the header of the label column.
- "**label**" – (Optional) Target attribute for the model to be used for bias metrics. Specified either as a column name, or an index (if dataset format is CSV), or as a JMESPath (if dataset format is JSON Lines).
• "label_values_or_threshold" – (Optional) List of label values or threshold. Indicates positive outcome used for bias metrics.

• "facet" – (Optional) A list of features that are sensitive attributes, referred to as facets. Facets are used for bias metrics in the form of pairs, and include the following:
  • "name_or_index" – Facet column name or index.
  • "value_or_threshold" – (Optional) List of values or threshold that the facet column can take. Indicates the sensitive group, such as the group that is used to measure bias against. If not provided, bias metrics are computed as one group for every unique value (rather than all values). If the facet column is numeric, this threshold value is applied as the lower bound to select the sensitive group.
  • "group_variable" – (Optional) A column name or index to indicate the group variable to be used for the bias metric Conditional Demographic Disparity.

The other parameters should be provided in EndpointInput (for real-time endpoints) or BatchTransformInput (for batch transform jobs) of the ModelBiasJobInput API.

• FeaturesAttribute – This parameter is required if endpoint input data format is "application/jsonlines". It is the JMESPath used to locate the feature columns if the dataset format is JSON Lines.

• InferenceAttribute – Index or JMESPath location in the model output for the target attribute to be used for monitored for bias using bias metrics. If it is not provided in the CSV accept_type case, then it is assumed that the model output is a single numeric value corresponding to a score or probability.

• ProbabilityAttribute – Index or JMESPath location in the model output for probabilities. If the model output is JSON Lines with a list of labels and probabilities, for example, then the label that corresponds to the maximum probability is selected for bias computations.

• ProbabilityThresholdAttribute – (Optional) A float value to indicate the threshold to select the binary label, in the case of binary classification. The default value is 0.5.

Example JSON Configuration Files for CSV and JSON Lines Datasets

Here are examples of the JSON files used to configure CSV and JSON Lines datasets to monitor them for bias drift.

Topics
CSV Datasets

Consider a dataset that has four feature columns and one label column, where the first feature and the label are binary, as in the following example.

```
0, 0.5814568701544718, 0.6651538910132964, 0.3138080342665499, 0
1, 0.6711642728531724, 0.7466687034026017, 0.1215477472819713, 1
0, 0.0453256543003371, 0.6377430803264152, 0.3558625219713576, 1
1, 0.4785191813363956, 0.0265841045263860, 0.0376935084990697, 1
```

Assume that the model output has two columns, where the first one is the predicted label and the second one is the probability, as in the following example.

```
1, 0.5385257417814224
```

Then the following JSON configuration file shows an example of how this CSV dataset can be configured.

```
{
   "headers": [
      "feature_0",
      "feature_1",
      "feature_2",
      "feature_3",
      "target"
   ],
   "label": "target",
   "label_values_or_threshold": [1],
   "facet": [{
      "name_or_index": "feature_1",
      "value_or_threshold": [1]
   }]
}
```

The predicted label is selected by the "InferenceAttribute" parameter. Zero-based numbering is used, so 0 indicates the first column of the model output,
Alternatively, you can use different parameters to convert probability values to binary predicted labels. Zero-based numbering is used: 1 indicates the second column; the `ProbabilityThresholdAttribute` value of 0.6 indicates that a probability greater than 0.6 predicts the binary label as 1.

```
"EndpointInput": {
  ...
  "InferenceAttribute": 0
  ...
}
```

### JSON Lines Datasets

Consider a dataset that has four feature columns and one label column, where the first feature and the label are binary, as in the following example.

```
{"features":[0, 0.5814568701544718, 0.6651538910132964, 0.3138080342665499], "label":0}
{"features":[1, 0.6711642728531724, 0.7466687034026017, 0.1215477472819713], "label":1}
{"features":[0, 0.0453256543003371, 0.6377430803264152, 0.3558625219713576], "label":1}
{"features":[1, 0.4785191813363956, 0.0265841045263860, 0.0376935084990697], "label":1}
```

Assume that the model output has two columns, where the first is a predicted label and the second is a probability.

```
{"predicted_label":1, "probability":0.5385257417814224}
```

The following JSON configuration file shows an example of how this JSON Lines dataset can be configured.

```
{
  "headers": [
    "feature_0",
    "feature_1",
    ...
  ]
}
Then, the "features" parameter value in EndpointInput (for real-time endpoints) or BatchTransformInput (for batch transform jobs) is used to locate the features in the dataset, and the "predicted_label" parameter value selects the predicted label from the model output.

```json
"EndpointInput": {
  ...
  "FeaturesAttribute": "features",
  "InferenceAttribute": "predicted_label"
  ...
}
```

Alternatively, you can convert probability values to predicted binary labels using the ProbabilityThresholdAttribute parameter value. A value of 0.6, for example, indicates that a probability greater than 0.6 predicts the binary label as 1.

```json
"EndpointInput": {
  ...
  "FeaturesAttribute": "features",
  "ProbabilityAttribute": "probability",
  "ProbabilityThresholdAttribute": 0.6
  ...
}
```

## Schedule Bias Drift Monitoring Jobs

After you create your baseline, you can call the `create_monitoring_schedule()` method of your `ModelBiasModelMonitor` class instance to schedule an hourly bias drift monitor. The following sections show you how to create bias drift monitor for a model deployed to a real-time endpoint as well as for a batch transform job.
Important

You can specify either a batch transform input or an endpoint input, but not both, when you create your monitoring schedule.

Unlike data quality monitoring, you need to supply Ground Truth labels if you want to monitor model quality. However, Ground Truth labels could be delayed. To address this, specify offsets when you create your monitoring schedule. For details about how to create time offsets, see Model monitor offsets.

If you have submitted a baselining job, the monitor automatically picks up analysis configuration from the baselining job. If you skip the baselining step or the capture dataset has a different nature from the training dataset, you must provide the analysis configuration.

Bias drift monitoring for models deployed to real-time endpoint

To schedule a bias drift monitor for a real-time endpoint, pass your EndpointInput instance to the endpoint_input argument of your ModelBiasModelMonitor instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

model_bias_monitor = ModelBiasModelMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

model_bias_analysis_config = None
if not model_bias_monitor.latest_baselining_job:
    model_bias_analysis_config = BiasAnalysisConfig(
        model_bias_config,
        headers=all_headers,
        label=label_header,
    )

model_bias_monitor.create_monitoring_schedule(
    monitor_schedule_name=schedule_name,
    post_analytics_processor_script=s3_code_postprocessor_uri,
    output_s3_uri=s3_report_path,
    statistics=model_bias_monitor.baseline_statistics(),
)```
Bias drift monitoring for batch transform jobs

To schedule a bias drift monitor for a batch transform job, pass your `BatchTransformInput` instance to the `batch_transform_input` argument of your `ModelBiasModelMonitor` instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

model_bias_monitor = ModelBiasModelMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

model_bias_analysis_config = None
if not model_bias_monitor.latest_baselining_job:
    model_bias_analysis_config = BiasAnalysisConfig(
        model_bias_config,
        headers=all_headers,
        label=label_header,
    )

schedule = model_bias_monitor.create_monitoring_schedule(
    monitor_schedule_name=schedule_name,
    post_analytics_processor_script=s3_code_postprocessor_uri,
    output_s3_uri=s3_report_path,
    statistics=model_bias_monitor.baseline_statistics(),
    constraints=model_bias_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
    analysis_config=model_bias_analysis_config,
)"
batch_transform_input=BatchTransformInput(
    destination="opt/ml/processing/input",
    data_captured_destination_s3_uri=s3_capture_path,
    start_time_offset="-PT1H",
    end_time_offset="-PT0H",
    probability_threshold_attribute=0.8
),
)

**Inspect Reports for Data Bias Drift**

If you are not able to inspect the results of the monitoring in the generated reports in SageMaker Studio, you can print them out as follows:

```
schedule_desc = model_bias_monitor.describe_schedule()
execution_summary = schedule_desc.get("LastMonitoringExecutionSummary")
if execution_summary and execution_summary["MonitoringExecutionStatus"] in
    ["Completed", "CompletedWithViolations"]:
    last_model_bias_monitor_execution = model_bias_monitor.list_executions()[-1]
    last_model_bias_monitor_execution_report_uri =
    last_model_bias_monitor_execution.output.destination
    print(f'Report URI: {last_model_bias_monitor_execution_report_uri}')
    last_model_bias_monitor_execution_report_files =
    sorted(S3Downloader.list(last_model_bias_monitor_execution_report_uri))
    print("Found Report Files:")
    print("\n ".join(last_model_bias_monitor_execution_report_files))
else:
    last_model_bias_monitor_execution = None
    print("====STOP==== 
 No completed executions to inspect further. Please wait till
 an execution completes or investigate previously reported failures.")
```

If there are violations compared to the baseline, they are listed here:

```
if last_model_bias_monitor_execution:
    model_bias_violations = last_model_bias_monitor_execution.constraint_violations()
    if model_bias_violations:
        print(model_bias_violations.body_dict)
```

If your model is deployed to a real-time endpoint, you can see visualizations in SageMaker Studio of the analysis results and CloudWatch metrics by choosing the **Endpoints** tab, and then double-clicking the endpoint.
CloudWatch Metrics for Bias Drift Analysis

This guide shows CloudWatch metrics and their properties that you can use for bias drift analysis in SageMaker Clarify. Bias drift monitoring jobs compute both pre-training bias metrics and post-training bias metrics, and publish them to the following CloudWatch namespace:

- For real-time endpoints: aws/sagemaker/Endpoints/bias-metrics
- For batch transform jobs: aws/sagemaker/ModelMonitoring/bias-metrics

The CloudWatch metric name appends the metric's short name to bias_metric.

For example, bias_metric_CI is the bias metric for class imbalance (CI).

Note

+-infinity is published as the floating point number +/- 2.348543e108, and errors including null values are not published.

Each metric has the following properties:

- Endpoint: The name of the monitored endpoint, if applicable.
- MonitoringSchedule: The name of the schedule for the monitoring job.
- BiasStage: The name of the stage of the bias drift monitoring job. Choose either Pre-training or Post-Training.
- Label: The name of the target feature, provided by the monitoring job analysis configuration label.
- LabelValue: The value of the target feature, provided by the monitoring job analysis configuration label_values_or_threshold.
- Facet: The name of the facet, provided by the monitoring job analysis configuration facet name_of_index.
- FacetValue: The value of the facet, provided by the monitoring job analysis configuration facet nvalue_or_threshold.

To stop the monitoring jobs from publishing metrics, set publish_cloudwatch_metrics to Disabled in the Environment map of model bias job definition.
Monitor Feature Attribution Drift for Models in Production

A drift in the distribution of live data for models in production can result in a corresponding drift in the feature attribution values, just as it could cause a drift in bias when monitoring bias metrics. Amazon SageMaker Clarify feature attribution monitoring helps data scientists and ML engineers monitor predictions for feature attribution drift on a regular basis. As the model is monitored, customers can view exportable reports and graphs detailing feature attributions in SageMaker Studio and configure alerts in Amazon CloudWatch to receive notifications if it is detected that the attribution values drift beyond a certain threshold.

To illustrate this with a specific situation, consider a hypothetical scenario for college admissions. Assume that we observe the following (aggregated) feature attribution values in the training data and in the live data:

**College Admission Hypothetical Scenario**

<table>
<thead>
<tr>
<th>Feature</th>
<th>Attribution in training data</th>
<th>Attribution in live data</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAT score</td>
<td>0.70</td>
<td>0.10</td>
</tr>
<tr>
<td>GPA</td>
<td>0.50</td>
<td>0.20</td>
</tr>
<tr>
<td>Class rank</td>
<td>0.05</td>
<td>0.70</td>
</tr>
</tbody>
</table>

The change from training data to live data appears significant. The feature ranking has completely reversed. Similar to the bias drift, the feature attribution drifts might be caused by a change in the live data distribution and warrant a closer look into the model behavior on the live data. Again, the first step in these scenarios is to raise an alarm that a drift has happened.

We can detect the drift by comparing how the ranking of the individual features changed from training data to live data. In addition to being sensitive to changes in ranking order, we also want to be sensitive to the raw attribution score of the features. For instance, given two features that fall in the ranking by the same number of positions going from training to live data, we want to be more sensitive to the feature that had a higher attribution score in the training data. With these properties in mind, we use the Normalized Discounted Cumulative Gain (NDCG) score for comparing the feature attributions rankings of training and live data.

Specifically, assume we have the following:
• $F=[f_1, \ldots, f_m]$ is the list of features sorted with respect to their attribution scores in the training data where $m$ is the total number of features. For instance, in our case, $F=[\text{SAT Score}, \text{GPA}, \text{Class Rank}]$.

• $a(f)$ is a function that returns the feature attribution score on the training data given a feature $f$. For example, $a(\text{SAT Score}) = 0.70$.

• $F'=[f'_1, \ldots, f'_m]$ is the list of features sorted with respect to their attribution scores in the live data. For example, $F'=[\text{Class Rank}, \text{GPA}, \text{SAT Score}]$.

Then, we can compute the NDCG as:

$$\text{NDCG} = \frac{\text{DCG}}{\text{iDCG}}$$

with

• $\text{DCG} = \sum_{1}^{m} a(f'_i)/\log_2(i+1)$

• $\text{iDCG} = \sum_{1}^{m} a(f_i)/\log_2(i+1)$

The quantity DCG measures whether features with high attribution in the training data are also ranked higher in the feature attribution computed on the live data. The quantity iDCG measures the ideal score and it's just a normalizing factor to ensure that the final quantity resides in the range $[0, 1]$, with 1 being the best possible value. A NDCG value of 1 means that the feature attribution ranking in the live data is the same as the one in the training data. In this particular example, because the ranking changed by quite a bit, the NDCG value is 0.69.

In SageMaker Clarify, if the NDCG value is below 0.90, we automatically raise an alert.

**Model Monitor Example Notebook**

SageMaker Clarify provides the following example notebook that shows how to capture inference data for a real-time endpoint, create a baseline to monitor evolving bias against, and inspect the results:

• **Monitoring bias drift and feature attribution drift Amazon SageMaker Clarify** – Use Amazon SageMaker Model Monitor to monitor bias drift and feature attribution drift over time.

This notebook has been verified to run in SageMaker Studio only. If you need instructions on how to open a notebook in SageMaker Studio, see [Create or Open an Amazon SageMaker Studio Classic](#).
Notebook. If you're prompted to choose a kernel, choose **Python 3 (Data Science)**. The following topics contain the highlights from the last two steps, and they contain code examples from the example notebook.

**Topics**
- **Create a SHAP Baseline for Models in Production**
- **Model Feature Attribution Drift Violations**
- **Configure Parameters to Monitor Attribution Drift**
- **Schedule Feature Attribute Drift Monitoring Jobs**
- **Inspect Reports for Feature Attribute Drift in Production Models**
- **CloudWatch Metrics for Feature Drift Analysis**

**Create a SHAP Baseline for Models in Production**

Explanations are typically contrastive, that is, they account for deviations from a baseline. For information on explainability baselines, see [SHAP Baselines for Explainability](#).

In addition to providing explanations for per-instance inferences, SageMaker Clarify also supports global explanation for ML models that helps you understand the behavior of a model as a whole in terms of its features. SageMaker Clarify generates a global explanation of an ML model by aggregating the Shapley values over multiple instances. SageMaker Clarify supports the following different ways of aggregation, which you can use to define baselines:

- **mean_abs** – Mean of absolute SHAP values for all instances.
- **median** – Median of SHAP values for all instances.
- **mean_sq** – Mean of squared SHAP values for all instances.

After you have configured your application to capture real-time or batch transform inference data, the first task to monitor for drift in feature attribution is to create a baseline to compare against. This involves configuring the data inputs, which groups are sensitive, how the predictions are captured, and the model and its posttraining bias metrics. Then you need to start the baselining job. Model explainability monitor can explain the predictions of a deployed model that's producing inferences and detect feature attribution drift on a regular basis.

```python
model_explainability_monitor = ModelExplainabilityMonitor(
```
In this example, the explainability baselining job shares the test dataset with the bias baselining job, so it uses the same DataConfig, and the only difference is the job output URI.

```python
model_explainability_baselining_job_result_uri = f"{baseline_results_uri}/model_explainability"
model_explainability_data_config = DataConfig(
    s3_data_input_path=validation_dataset,
    s3_output_path=model_explainability_baselining_job_result_uri,
    label=label_header,
    headers=all_headers,
    dataset_type=dataset_type,
)
```

Currently the SageMaker Clarify explainer offers a scalable and efficient implementation of SHAP, so the explainability config is SHAPConfig, including the following:

- **baseline** – A list of rows (at least one) or S3 object URI to be used as the baseline dataset in the Kernel SHAP algorithm. The format should be the same as the dataset format. Each row should contain only the feature columns/values and omit the label column/values.

- **num_samples** – Number of samples to be used in the Kernel SHAP algorithm. This number determines the size of the generated synthetic dataset to compute the SHAP values.

- **agg_method** – Aggregation method for global SHAP values. Following are valid values:
  - **mean_abs** – Mean of absolute SHAP values for all instances.
  - **median** – Median of SHAP values for all instances.
  - **mean_sq** – Mean of squared SHAP values for all instances.
  - **use_logit** – Indicator of whether the logit function is to be applied to the model predictions. Default is False. If use_logit is True, the SHAP values will have log-odds units.
  - **save_local_shap_values** (bool) – Indicator of whether to save the local SHAP values in the output location. Default is False.

```python
# Here use the mean value of test dataset as SHAP baseline
test_dataframe = pd.read_csv(test_dataset, header=None)
```
shap_baseline = [list(test_dataframe.mean())]

shap_config = SHAPConfig(
    baseline=shap_baseline,
    num_samples=100,
    agg_method="mean_abs",
    save_local_shap_values=False,
)

Start a baselining job. The same model_config is required because the explainability baselining job needs to create a shadow endpoint to get predictions for the generated synthetic dataset.

model_explainability_monitor.suggest_baseline(
    data_config=model_explainability_data_config,
    model_config=model_config,
    explainability_config=shap_config,
)

print(f"ModelExplainabilityMonitor baselining job:
    {model_explainability_monitor.latest_baselining_job_name}"
)

**Model Feature Attribution Drift Violations**

Feature attribution drift jobs evaluate the baseline constraints provided by the baseline configuration against the analysis results of current MonitoringExecution. If violations are detected, the job lists them to the constraint_violations.json file in the execution output location, and marks the execution status as Interpret results.

Here is the schema of the feature attribution drift violations file.

- **label** – The name of the label, job analysis configuration label_headers or a placeholder such as "label0".
- **metric_name** – The name of the explainability analysis method. Currently only shap is supported.
- **constraint_check_type** – The type of violation monitored. Currently only feature_attribution_drift_check is supported.
- **description** – A descriptive message to explain the violation.

```json
{
    "version": "1.0",
    "label": "label0",
    "metric_name": "shap",
    "constraint_check_type": "feature_attribution_drift_check",
    "description": "Violation description goes here.
```
For each label in the explanations section, the monitoring jobs calculate the nDCG score of its global SHAP values in the baseline constraints file and in the job analysis results file (analysis.json). If the score is less than 0.9, then a violation is logged. The combined global SHAP value is evaluated, so there are no “feature” fields in the violation entry. The following output provides an example of several logged violations.

```json
{
  "version": "1.0",
  "violations": [[
    "label": "label0",
    "metric_name": "shap",
    "constraint_check_type": "feature_attribution_drift_check",
    "description": "Feature attribution drift 0.7639720923277322 exceeds threshold 0.9"
  ], [
    "label": "label1",
    "metric_name": "shap",
    "constraint_check_type": "feature_attribution_drift_check",
    "description": "Feature attribution drift 0.7323763972092327 exceeds threshold 0.9"
  ]]
}
```

**Configure Parameters to Monitor Attribution Drift**

Amazon SageMaker Clarify explainability monitor reuses a subset of the parameters used in the analysis configuration of Configure the Analysis. The following parameters must be provided in a JSON file and the path must be provided in the ConfigUri parameter of ModelExplainabilityAppSpecification.

- "version" – (Optional) Schema version of the configuration file. If not provided, the latest supported version is used.
• "headers" – (Optional) A list of feature names in the dataset. Explainability analysis does not require labels.

• "methods" – A list of methods and their parameters for the analyses and reports. If any section is omitted, then it is not computed.

• "shap" – (Optional) Section on SHAP value computation.
  • "baseline" – (Optional) A list of rows (at least one), or an Amazon Simple Storage Service Amazon S3 object URI. To be used as the baseline dataset (also known as a background dataset) in the Kernel SHAP algorithm. The format should be the same as the dataset format. Each row should contain only the feature columns (or values). Before you send each row to the model, omit any column that must be excluded.
  • "num_samples" – Number of samples to be used in the Kernel SHAP algorithm. This number determines the size of the generated synthetic dataset to compute the SHAP values. If not provided, then a SageMaker Clarify job chooses the value based on a count of features.
  • "agg_method" – Aggregation method for global SHAP values. Valid values are as follows:
    • "mean_abs" – Mean of absolute SHAP values for all instances.
    • "median" – Median of SHAP values for all instances.
    • "mean_sq" – Mean of squared SHAP values for all instances.
  • "use_logit" – (Optional) Boolean value to indicate if the logit function is to be applied to the model predictions. If "use_logit" is true, then the SHAP values have log-odds units. The default value is false.
  • "save_local_shap_values" – (Optional) Boolean value to indicate if local SHAP values are to be saved in the output location. Use true to save them. Use false to not save them. The default is false.

• "predictor" – (Optional for real-time endpoint, required for batch transform) Section on model parameters, required if "shap" and "post_training_bias" sections are present.
  • "model_name" – Model name created by CreateModel API, with container mode as SingleModel.
  • "instance_type" – Instance type for the shadow endpoint.
  • "initial_instance_count" – Instance count for the shadow endpoint.
  • "content_type" – (Optional) The model input format to be used for getting inferences with the shadow endpoint. Valid values are "text/csv" for CSV, "application/jsonlines" for JSON Lines, application/x-parquet for Apache Parquet, and application/x-image to
enable Computer Vision explainability. The default value is the same as the dataset_type format.

- "accept_type" – (Optional) The model output format to be used for getting inferences with the shadow endpoint. Valid values are "text/csv" for CSV, "application/jsonlines" for JSON Lines. If omitted, SageMaker Clarify uses the response data type of the captured data.

- "content_template" – (Optional) A template string used to construct the model input from dataset instances. It is only used when "content_type" is "application/jsonlines". The template should have only one placeholder, $features, which is replaced by the features list at runtime. For example, given "content_template": "{\"myfeatures\": $features}"; if an instance (no label) is 1, 2, 3, then model input becomes JSON Lines '{"myfeatures": [1, 2, 3]}'.

- "label_headers" – (Optional) A list of values that the "label" takes in the dataset. Associates the scores returned by the model endpoint or batch transform job with their corresponding label values. If it is provided, then the analysis report uses the headers instead of placeholders like "label0".

The other parameters should be provided in EndpointInput (for real-time endpoints) or BatchTransformInput (for batch transform jobs) of the ModelExplainabilityJobInput API.

- FeaturesAttribute – This parameter is required if endpoint or batch job input data format is "application/jsonlines". It is the JMESPath used to locate the feature columns if the dataset format is JSON Lines.

- ProbabilityAttribute – Index or JMESPath location in the model output for probabilities. If the model output is JSON Lines with a list of labels and probabilities, for example, then the label that corresponds to the maximum probability is selected for bias computations.

### Example JSON Configuration Files for CSV and JSON Lines Datasets

Here are examples of the JSON files used to configure CSV and JSON Lines datasets to monitor them for feature attribution drift.

#### Topics

- **CSV Datasets**
- **JSON Lines Datasets**
CSV Datasets

Consider a dataset that has three numerical feature columns, as in the following example.

- 0.5814568701544718, 0.6651538910132964, 0.3138080342665499
- 0.6711642728531724, 0.7466687034026017, 0.1215477472819713
- 0.0453256543003371, 0.6377430803264152, 0.3558625219713576
- 0.4785191813363956, 0.0265841045263860, 0.0376935084990697

Assume that the model output has two columns, where the first one is the predicted label and the second one is the probability, as in the following example.

- 1, 0.5385257417814224

The following example JSON configuration file shows how this CSV dataset can be configured.

```json
{
    "headers": [
        "feature_1",
        "feature_2",
        "feature_3"
    ],
    "methods": {
        "shap": {
            "baseline": [
                [0.4441164946610942, 0.5190374448171748, 0.20722795300473712]
            ],
            "num_samples": 100,
            "agg_method": "mean_abs"
        }
    },
    "predictor": {
        "model_name": "my_model",
        "instance_type": "ml.m5.xlarge",
        "initial_instance_count": 1
    }
}
```

The predicted label is selected by the "ProbabilityAttribute" parameter. Zero-based numbering is used, so 1 indicates the second column of the model output.
"EndpointInput": {
    ...
    "ProbabilityAttribute": 1
    ...
}

JSON Lines Datasets

Consider a dataset that has four feature columns and one label column, where the first feature and
the label are binary, as in the following example.

{"features":[0, 0.5814568701544718, 0.6651538910132964, 0.3138080342665499], "label":0}
{"features":[1, 0.6711642728531724, 0.7466687034026017, 0.1215477472819713], "label":1}
{"features":[0, 0.0453256543003371, 0.6377430803264152, 0.3558625219713576], "label":1}
{"features":[1, 0.4785191813363956, 0.0265841045263860, 0.0376935084990697], "label":1}

The model input is the same as the dataset format, and the model output are JSON Lines, as in the
following example.

{"predicted_label":1, "probability":0.5385257417814224}

In the following example, the JSON configuration file shows how this JSON Lines dataset can be
configured.

{
    "headers": [
        "feature_1",
        "feature_2",
        "feature_3"
    ],
    "methods": {
        "shap": {
            "baseline": [
                "features":[0.4441164946610942, 0.5190374448171748, 0.2072279530473712]
            ],
            "num_samples": 100,
            "agg_method": "mean_abs"
        }
    }
}
Then the "features" parameter value in EndpointInput (for real-time endpoints) or BatchTransformInput (for batch transform jobs) is used to locate the features in the dataset, and the "probability" parameter value selects the probability value from model output.

Schedule Feature Attribute Drift Monitoring Jobs

After you create your SHAP baseline, you can call the create_monitoring_schedule() method of your ModelExplainabilityMonitor class instance to schedule an hourly model explainability monitor. The following sections show you how to create a model explainability monitor for a model deployed to a real-time endpoint as well as for a batch transform job.

⚠️ Important

You can specify either a batch transform input or an endpoint input, but not both, when you create your monitoring schedule.

If a baselining job has been submitted, the monitor automatically picks up analysis configuration from the baselining job. However, if you skip the baselining step or the capture dataset has a different nature from the training dataset, you have to provide the analysis configuration. ModelConfig is required by ExplainabilityAnalysisConfig for the same reason that it's required for the baselining job. Note that only features are required for computing feature attribution, so you should exclude Ground Truth labeling.
Feature attribution drift monitoring for models deployed to real-time endpoint

To schedule a model explainability monitor for a real-time endpoint, pass your EndpointInput instance to the endpoint_input argument of your ModelExplainabilityMonitor instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

model_exp_model_monitor = ModelExplainabilityMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

schedule = model_exp_model_monitor.create_monitoring_schedule(
    monitor_schedule_name=schedule_name,
    post_analytics_processor_script=s3_code_postprocessor_uri,
    output_s3_uri=s3_report_path,
    statistics=model_exp_model_monitor.baseline_statistics(),
    constraints=model_exp_model_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
    endpoint_input=EndpointInput(
        endpoint_name=endpoint_name,
        destination="/opt/ml/processing/input/endpoint",
    ),
)
```

Feature attribution drift monitoring for batch transform jobs

To schedule a model explainability monitor for a batch transform job, pass your BatchTransformInput instance to the batch_transform_input argument of your ModelExplainabilityMonitor instance, as shown in the following code sample:

```python
from sagemaker.model_monitor import CronExpressionGenerator

model_exp_model_monitor = ModelExplainabilityMonitor(
    role=sagemaker.get_execution_role(),
    ...
)

schedule = model_exp_model_monitor.create_monitoring_schedule(
    monitor_schedule_name=schedule_name,
)
```
post_analytics_processor_script=s3_code_postprocessor_uri,
output_s3_uri=s3_report_path,
statistics=model_exp_model_monitor.baseline_statistics(),
constraints=model_exp_model_monitor.suggested_constraints(),
schedule_cron_expression=CronExpressionGenerator.hourly(),
enable_cloudwatch_metrics=True,
batch_transform_input=BatchTransformInput(
    destination="opt/ml/processing/data",
    model_name="batch-fraud-detection-model",
    input_manifests_s3_uri="s3://my-bucket/batch-fraud-detection/on-schedule-
    monitoring/in/",
    excludeFeatures="0",
)
)

Inspect Reports for Feature Attribute Drift in Production Models

After the schedule that you set up is started by default, you need to wait for the its first execution to start, and then stop the schedule to avoid incurring charges.

To inspect the reports, use the following code:

```
schedule_desc = model_explainability_monitor.describe_schedule()
execution_summary = schedule_desc.get("LastMonitoringExecutionSummary")
if execution_summary and execution_summary["MonitoringExecutionStatus"] in
["Completed", "CompletedWithViolations"]:
    last_model_explainability_monitor_execution =
    model_explainability_monitor.list_executions()[-1]
    last_model_explainability_monitor_execution_report_uri =
    last_model_explainability_monitor_execution.output.destination
    print(f'Report URI: {last_model_explainability_monitor_execution_report_uri}')
    last_model_explainability_monitor_execution_report_files =
    sorted(S3Downloader.list(last_model_explainability_monitor_execution_report_uri))
    print("Found Report Files:")
    print("\n ".join(last_model_explainability_monitor_execution_report_files))
else:
    last_model_explainability_monitor_execution = None
    print("====STOP==== 
 No completed executions to inspect further. Please wait till an execution completes or investigate previously reported failures.")
```

If there are any violations compared to the baseline, they are listed here:

```
if last_model_explainability_monitor_execution:
```

Inspect Reports for Feature Attribute Drift

5214
model_explainabilityViolations =
last_model_explainability_monitor_execution.constraint_violations()
if model_explainabilityViolations:
    print(model_explainabilityViolations.body_dict)

If your model is deployed to a real-time endpoint, you can see visualizations in SageMaker Studio of the analysis results and CloudWatch metrics by choosing the **Endpoints** tab, and then double-clicking the endpoint.

### CloudWatch Metrics for Feature Drift Analysis

This guide shows CloudWatch metrics and their properties that you can use for feature attribute drift analysis in SageMaker Clarify. Feature attribute drift monitoring jobs compute and publish two types of metrics:

- The global SHAP value of each feature.

  **Note**
  
  The name of this metric appends the feature name provided by the job analysis configuration to `feature_`. For example, `feature_X` is the global SHAP value for feature X.

- The `ExpectedValue` of the metric.

These metrics are published to the following CloudWatch namespace:

- For real-time endpoints: `aws/sagemaker/Endpoints/explainability-metrics`
- For batch transform jobs: `aws/sagemaker/ModelMonitoring/explainability-metrics`

Each metric has the following properties:

- **Endpoint**: The name of the monitored endpoint, if applicable.
- **MonitoringSchedule**: The name of the schedule for the monitoring job.
- **ExplainabilityMethod**: The method used to compute Shapley values. Choose `KernelShap`.
- **Label**: The name provided by job analysis configuration `label_headers`, or a placeholder like `label0`. 
• **ValueType**: The type of the value returned by the metric. Choose either `GlobalShapValues` or `ExpectedValue`.

To stop the monitoring jobs from publishing metrics, set `publish_cloudwatch_metrics` to `Disabled` in the `Environment` map of [model explainability job](#).

## Schedule monitoring jobs

Amazon SageMaker Model Monitor provides you the ability to monitor the data collected from your real-time endpoints. You can monitor your data on a recurring schedule, or you can monitor it one time, immediately. You can create a monitoring schedule with the `CreateMonitoringSchedule` API.

With a monitoring schedule, SageMaker can start processing jobs to analyze the data collected during a given period. In the processing job, SageMaker compares the dataset for the current analysis with the baseline statistics and constraints that you provide. Then, SageMaker generate a violations report. In addition, CloudWatch metrics are emitted for each feature under analysis.

SageMaker provides a prebuilt container for performing analysis on tabular datasets. Alternatively, you could choose to bring your own container as outlined in the [Bring Your Own Containers](#) topic.

You can create a model monitoring schedule for your real-time endpoint or batch transform job. Use the baseline resources (constraints and statistics) to compare against the real-time traffic or batch job inputs.

### Example baseline assignments

In the following example, the training dataset used to train the model was uploaded to Amazon S3. If you already have it in Amazon S3, you can point to it directly.

```python
# copy over the training dataset to Amazon S3 (if you already have it in Amazon S3, you could reuse it)
baseline_prefix = prefix + '/baselining'
baseline_data_prefix = baseline_prefix + '/data'
baseline_results_prefix = baseline_prefix + '/results'

baseline_data_uri = 's3://{}/{}'.format(bucket, baseline_data_prefix)
baseline_results_uri = 's3://{}/{}'.format(bucket, baseline_results_prefix)
print('Baseline data uri: {}'.format(baseline_data_uri))
```

---

*Schedule monitoring jobs*
print('Baseline results uri: {}'.format(baseline_results_uri))

training_data_file = open("test_data/training-dataset-with-header.csv", 'rb')
s3_key = os.path.join(baseline_prefix, 'data', 'training-dataset-with-header.csv')
boto3.Session().resource('s3').Bucket(bucket).Object(s3_key).upload_fileobj(training_data_file)

Example schedule for recurring analysis

If you are scheduling a model monitor for a real-time endpoint, use the baseline constraints and statistics to compare against real-time traffic. The following code snippet shows the general format you use to schedule a model monitor for a real-time endpoint. This example schedules the model monitor to run hourly.

```python
from sagemaker.model_monitor import CronExpressionGenerator
from time import gmtime, strftime

mon_schedule_name = 'my-model-monitor-schedule-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
my_default_monitor.create_monitoring_schedule(
    monitor_schedule_name=mon_schedule_name,
    endpoint_input=EndpointInput(
        endpoint_name=endpoint_name,
        destination="/opt/ml/processing/input/endpoint"
    ),
    post_analytics_processor_script=s3_code_postprocessor_uri,
    output_s3_uri=s3_report_path,
    statistics=my_default_monitor.baseline_statistics(),
    constraints=my_default_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
)
```

Example schedule for one-time analysis

You can also schedule the analysis to run once without recurring by passing arguments like the following to the create_monitoring_schedule method:

```python
schedule_cron_expression=CronExpressionGenerator.now(),
data_analysis_start_time="-PT1H",
data_analysis_end_time="-PT0H",
```
In these arguments, the schedule_cron_expression parameter schedules the analysis to run once, immediately, with the value CronExpressionGenerator.now(). For any schedule with this setting, the data_analysis_start_time and data_analysis_end_time parameters are required. These parameters set the start time and end time of an analysis window. Define these times as offsets that are relative to the current time, and use ISO 8601 duration format. In this example, the times -PT1H and -PT0H define a window between one hour in the past and the current time. With this schedule, the analysis evaluates only the data that was collected during the specified window.

**Example schedule for a batch transform job**

The following code snippet shows the general format you use to schedule a model monitor for a batch transform job.

```python
from sagemaker.model_monitor import (  
    CronExpressionGenerator,  
    BatchTransformInput,  
    MonitoringDatasetFormat,  
)
from time import gmtime, strftime
mon_schedule_name = 'my-model-monitor-schedule-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
my_default_monitor.create_monitoring_schedule(  
    monitor_schedule_name=mon_schedule_name,  
    batch_transform_input=BatchTransformInput(  
        destination="opt/ml/processing/input",  
        data_captured_destination_s3_uri=s3_capture_upload_path,  
        dataset_format=MonitoringDatasetFormat.csv(header=False),  
    ),  
    post_analytics_processor_script=s3_code_postprocessor_uri,  
    output_s3_uri=s3_report_path,  
    statistics=my_default_monitor.baseline_statistics(),  
    constraints=my_default_monitor.suggested_constraints(),  
    schedule_cron_expression=CronExpressionGenerator.hourly(),  
    enable_cloudwatch_metrics=True,
)

desc_schedule_result = my_default_monitor.describe_schedule()
print('Schedule status: {}'.format(desc_schedule_result['MonitoringScheduleStatus']))
```
The cron expression for monitoring schedule

To provide details for the monitoring schedule, use **ScheduleConfig**, which is a **cron** expression that describes details about the monitoring schedule.

Amazon SageMaker Model Monitor supports the following **cron** expressions:

- To set the job to start every hour, use the following:
  ```yaml
  Hourly: cron(0 * ? * *)
  ```

- To run the job daily, use the following:
  ```yaml
  cron(0 [00-23] ? * * *)
  ```

- The run the job one time, immediately, use the following keyword:
  ```yaml
  NOW
  ```

For example, the following are valid **cron** expressions:

- Daily at 12 PM UTC: `cron(0 12 ? * *)`
- Daily at 12 AM UTC: `cron(0 0 ? * *)`

To support running every 6, 12 hours, Model Monitor supports the following expression:

```yaml
cron(0 [00-23]/[01-24] ? * *)
```

For example, the following are valid **cron** expressions:

- Every 12 hours, starting at 5 PM UTC: `cron(0 17/12 ? * *)`
- Every two hours, starting at 12 AM UTC: `cron(0 0/2 ? * *)`

---

**Notes**

- Although the **cron** expression is set to start at 5 PM UTC, note that there could be a delay of 0-20 minutes from the actual requested time to run the execution.
- If you want to run on a daily schedule, don't provide this parameter. SageMaker picks a time to run every day.
Currently, SageMaker only supports hourly integer rates between 1 hour and 24 hours.

### Configuring service control policies for monitoring schedules

You have to specify the parameters of a monitoring job when you create or update a schedule for it with the [CreateMonitoringSchedule](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateMonitoringSchedule.html) API or the [UpdateMonitoringSchedule](https://docs.aws.amazon.com/sagemaker/latest/dg/API_UpdateMonitoringSchedule.html) API, respectively. Depending on your use case, you can do this in one of the following ways:

- You can specify the `MonitoringJobDefinition` field of `MonitoringScheduleConfig`, when you invoke `CreateMonitoringSchedule` or `UpdateMonitoringSchedule`. You can use this only to create or update a schedule for a data quality monitoring job.

- You can specify the name of a monitoring job definition, that you have already created, for the `MonitoringJobDefinitionName` field of `MonitoringScheduleConfig`, when you invoke `CreateMonitoringSchedule` or `UpdateMonitoringSchedule`. You can use this for any job definition that you create with one of the following APIs:
  - [CreateDataQualityJobDefinition](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateDataQualityJobDefinition.html)
  - [CreateModelQualityJobDefinition](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateModelQualityJobDefinition.html)
  - [CreateModelBiasJobDefinition](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateModelBiasJobDefinition.html)
  - [CreateModelExplainabilityJobDefinition](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateModelExplainabilityJobDefinition.html)

If you want to use the SageMaker Python SDK to create or update schedules, then you have to use this process.

The aforementioned processes are mutually exclusive, that is, you can either specify the `MonitoringJobDefinition` field or the `MonitoringJobDefinitionName` field when creating or updating monitoring schedules.

When you create a monitoring job definition, or specify one in the `MonitoringJobDefinition` field, you can set security parameters, such as `NetworkConfig` and `VolumeKmsKeyId`. As an administrator, you might want that these parameters are always set to certain values, so that the monitoring jobs always run in a secure environment. To ensure this, set up appropriate [Service control policies](https://docs.aws.amazon.com/solutions/latest/dg/amazon-solutions-service-control-policies.html) (SCPs). SCPs are a type of organization policy that you can use to manage permissions in your organization.

The following example shows a SCP that you can use to ensure that infrastructure parameters are properly set when creating or updating schedules for monitoring jobs.
Configuring SCPs for monitoring schedules

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Deny",
            "Action": [
                "sagemaker:CreateDataQualityJobDefinition",
                "sagemaker:CreateModelBiasJobDefinition",
                "sagemaker:CreateModelExplainabilityJobDefinition",
                "sagemaker:CreateModelQualityJobDefinition"
            ],
            "Resource": "arn:*:sagemaker:*:*:*",
            "Condition": {
                "Null": {
                    "sagemaker:VolumeKmsKey": "true",
                    "sagemaker:VpcSubnets": "true",
                    "sagemaker:VpcSecurityGroupIds": "true"
                }
            }
        },
        {
            "Effect": "Deny",
            "Action": [
                "sagemaker:CreateDataQualityJobDefinition",
                "sagemaker:CreateModelBiasJobDefinition",
                "sagemaker:CreateModelExplainabilityJobDefinition",
                "sagemaker:CreateModelQualityJobDefinition"
            ],
            "Resource": "arn:*:sagemaker:*:*:*",
            "Condition": {
                "Bool": {
                    "sagemaker:InterContainerTrafficEncryption": "false"
                }
            }
        },
        {
            "Effect": "Deny",
            "Action": [
                "sagemaker:CreateMonitoringSchedule",
                "sagemaker:UpdateMonitoringSchedule"
            ],
            "Resource": "arn:*:sagemaker:*:*:monitoring-schedule/*",
            "Condition": {
                "Null": {
                    "sagemaker:VolumeKmsKey": "true",
                    "sagemaker:VpcSubnets": "true",
                    "sagemaker:VpcSecurityGroupIds": "true"
                }
            }
        }
    ]
}
```
The first two rules in the example, ensure that the security parameters are always set for monitoring job definitions. The final rule requires that anyone, in your organization, creating or updating a schedule, have to always specify the `MonitoringJobDefinitionName` field. This ensures that no one in your organization, can set insecure values for the security parameters by specifying the `MonitoringJobDefinition` field, when creating or updating schedules.

**Amazon SageMaker Model Monitor prebuilt container**

SageMaker provides a built-in image called `sagemaker-model-monitor-analyzer` that provides you with a range of model monitoring capabilities, including constraint suggestion, statistics generation, constraint validation against a baseline, and emitting Amazon CloudWatch metrics. This image is based on Spark version 3.3.0 and is built with [Deequ](https://deeplearning.ai/deequ) version 2.0.2.

ℹ️ **Note**

You can not pull the built-in `sagemaker-model-monitor-analyzer` image directly. You can use the `sagemaker-model-monitor-analyzer` image when you submit a baseline processing or monitoring job using one of the AWS SDKs.

Use the SageMaker Python SDK (see `image_uris.retrieve` in the [SageMaker Python SDK reference guide](https://boto3.amazonaws.com/v1/documentation/api/latest/reference/services/sagemaker.html#SageMaker.Client.image_uris.retrieve)) to generate the ECR image URI for you, or specify the ECR image URI directly. The prebuilt image for SageMaker Model Monitor can be accessed as follows:

```
<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/sagemaker-model-monitor-analyzer
```

For example: `159807026194.dkr.ecr.us-west-2.amazonaws.com/sagemaker-model-monitor-analyzer`
If you are in an AWS region in China, the prebuilt images for SageMaker Model Monitor can be accessed as follows:

\(<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com.cn/sagemaker-model-monitor-analyzer\)

For account IDs and AWS Region names, see Docker Registry Paths and Example Code.

To write your own analysis container, see the container contract described in Customize monitoring.

**Interpret results**

After you run a baseline processing job and obtained statistics and constraint for your dataset, you can execute monitoring jobs that calculate statistics and list any violations encountered relative to the baseline constraints. Amazon CloudWatch metrics are also reported in your account by default. For information on viewing the results of monitoring in Amazon SageMaker Studio, see Visualize results for real-time endpoints in Amazon SageMaker Studio.

**List Executions**

The schedule starts monitoring jobs at the specified intervals. The following code lists the latest five executions. If you are running this code after creating the hourly schedule, the executions might be empty, and you might have to wait until you cross the hour boundary (in UTC) to see the executions start. The following code includes the logic for waiting.

```python
mon_executions = my_default_monitor.list_executions()
print("We created a hourly schedule above and it will kick off executions ON the hour (plus 0 - 20 min buffer.\nWe will have to wait till we hit the hour...")

while len(mon_executions) == 0:
    print("Waiting for the 1st execution to happen...")
    time.sleep(60)
    mon_executions = my_default_monitor.list_executions()
```

**Inspect a Specific Execution**

In the previous step, you picked up the latest completed or failed scheduled execution. You can explore what went right or wrong. The terminal states are:
- **Completed** – The monitoring execution completed and no issues were found in the violations report.
- **CompletedWithViolations** – The execution completed, but constraint violations were detected.
- **Failed** – The monitoring execution failed, possibly due to client error (for example, a role issues) or infrastructure issues. To identify the cause, see the `FailureReason` and `ExitMessage`.

```python
latest_execution = mon_executions[-1]  # latest execution's index is -1, previous is -2 and so on..
time.sleep(60)
latest_execution.wait(logs=False)

print("Latest execution status: {}".format(latest Execution.describe() ['ProcessingJobStatus']))
print("Latest execution result: {}".format(latest Execution.describe() ['ExitMessage']))

latest_job = latest_execution.describe()
if (latest_job['ProcessingJobStatus'] != 'Completed'):
    print("====STOP==== \n No completed executions to inspect further. Please wait till an execution completes or investigate previously reported failures."")

report_uri=latest_execution.output.destination
print('Report Uri: {}'.format(report_uri))
```

### List Generated Reports

Use the following code to list the generated reports.

```python
from urllib.parse import urlparse
s3uri = urlparse(report_uri)
report_bucket = s3uri.netloc
report_key = s3uri.path.lstrip('/')
print('Report bucket: {}'.format(report_bucket))
print('Report key: {}'.format(report_key))

s3_client = boto3.Session().client('s3')
result = s3_client.list_objects(Bucket=report_bucket, Prefix=report_key)
report_files = [report_file.get('Key') for report_file in result.get('Contents')]
print("Found Report Files:")
```
print("\n \n\nViolations Report

If there are violations compared to the baseline, they are generated in the violations report. Use the following code to list the violations.

```python
violations = my_default_monitor.latest_monitoring_constraint_violations()
pd.set_option('display.max_colwidth', -1)
constraints_df = pd.io.json.json_normalize(violations.body_dict['violations'])
constraints_df.head(10)
```

This applies only to datasets that contain tabular data. The following schema files specify the statistics calculated and the violations monitored for.

**Output Files for Tabular Datasets**

<table>
<thead>
<tr>
<th>File Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>statistics.json</td>
<td>Contains columnar statistics for each feature in the dataset that is analyzed. See the schema of this file in the next topic.</td>
</tr>
<tr>
<td>constraint_violations.json</td>
<td>Contains a list of violations found in this current set of data as compared to the baseline statistics and constraints file specified in the baseline_constraints and baseline_statistics paths.</td>
</tr>
</tbody>
</table>

The Amazon SageMaker Model Monitor prebuilt container saves a set of Amazon CloudWatch metrics for each feature by default.
The container code can emit CloudWatch metrics in this location: `/opt/ml/output/metrics/cloudwatch`.

**Visualize results for real-time endpoints in Amazon SageMaker Studio**

If you are monitoring a real-time endpoint, you can also visualize the results in Amazon SageMaker Studio. You can view the details of any monitoring job run, and you can create charts that show the baseline and captured values for any metric that the monitoring job calculates.

**To view the detailed results of a monitoring job**

1. Sign in to Studio. For more information, see [Amazon SageMaker domain overview](#).
2. In the left navigation pane, choose the *Components and registries* icon (🔍).
3. Choose **Endpoints** in the drop-down menu.

![Endpoints in Amazon SageMaker Studio](image)

4. On the endpoint tab, choose the monitoring type for which you want to see job details.
5. Choose the name of the monitoring job run for which you want to view details from the list of monitoring jobs.

6. The **MONITORING JOB DETAILS** tab opens with a detailed report of the monitoring job.
You can create a chart that displays the baseline and captured metrics for a time period.

**To create a chart in SageMaker Studio to visualize monitoring results**

1. Sign in to Studio. For more information, see Amazon SageMaker domain overview.
2. In the left navigation pane, choose the **Components and registries** icon (🔗).
3. Choose **Endpoints** in the drop-down menu.

4. On the **Endpoint** tab, choose the monitoring type you want to create a chart for. This example shows a chart for the **Model quality** monitoring type.

5. Choose **Add chart**.
6. On the **CHART PROPERTIES** tab, choose the time period, statistic, and metric that you want to chart. This example shows a chart for a **Timeline** of 1 week, the **Average Statistic** of, and the **F1 Metric**.
7. The chart that shows the baseline and current metric statistic you chose in the previous step shows up in the Endpoint tab.
Advanced topics

The following sections contain more advanced tasks that explain how to customize monitoring using preprocessing and postprocessing scripts, how to build your own container, and how to use AWS CloudFormation to create a monitoring schedule.

Topics

- Customize monitoring
- Create a Monitoring Schedule for a Real-time Endpoint with an AWS CloudFormation Custom Resource

Customize monitoring

In addition to using the built-in monitoring mechanisms, you can create your own custom monitoring schedules and procedures using preprocessing and postprocessing scripts or by using or building your own container.

Topics

- Preprocessing and Postprocessing
- Bring Your Own Containers

Preprocessing and Postprocessing

You can use custom preprocessing and postprocessing Python scripts to transform the input to your model monitor or extend the code after a successful monitoring run. Upload these scripts to Amazon S3 and reference them when creating your model monitor.

The following example shows how you can customize monitoring schedules with preprocessing and postprocessing scripts. Replace user placeholder text with your own information.

```python
import boto3, os
from sagemaker import get_execution_role, Session
from sagemaker.model_monitor import CronExpressionGenerator, DefaultModelMonitor

# Upload pre and postprocessor scripts
session = Session()
bucket = boto3.Session().resource("s3").Bucket(session.default_bucket())
```
prefix = "demo-sagemaker-model-monitor"
pre_processor_script = bucket.Object(os.path.join(prefix, "preprocessor.py")).upload_file("preprocessor.py")
post_processor_script = bucket.Object(os.path.join(prefix, "postprocessor.py")).upload_file("postprocessor.py")

# Get execution role
role = get_execution_role() # can be an empty string

# Instance type
instance_type = "instance-type"
# instance_type = "ml.m5.xlarge" # Example

# Create a monitoring schedule with pre and postprocessing
my_default_monitor = DefaultModelMonitor(
    role=role,
    instance_count=1,
    instance_type=instance_type,
    volume_size_in_gb=20,
    max_runtime_in_seconds=3600,
)

s3_report_path = "s3://{}/{}".format(bucket, "reports")
monitor_schedule_name = "monitor-schedule-name"
endpoint_name = "endpoint-name"

my_default_monitor.create_monitoring_schedule(
    post_analytics_processor_script=post_processor_script,
    record_preprocessor_script=pre_processor_script,
    monitor_schedule_name=monitor_schedule_name,
    # use endpoint_input for real-time endpoint
    endpoint_input=endpoint_name,
    # or use batch_transform_input for batch transform jobs
    # batch_transform_input=batch_transform_name,
    output_s3_uri=s3_report_path,
    statistics=my_default_monitor.baseline_statistics(),
    constraints=my_default_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
)

Topics

- Preprocessing Script
Preprocessing Script

Use preprocessing scripts when you need to transform the inputs to your model monitor.

For example, suppose the output of your model is an array \([1.0, 2.1]\). The Amazon SageMaker Model Monitor container only works with tabular or flattened JSON structures, like \{"prediction0": 1.0, "prediction1": 2.1\}. You could use a preprocessing script like the following to transform the array into the correct JSON structure.

```python
def preprocess_handler(inference_record):
    input_data = inference_record.endpoint_input.data
    output_data = inference_record.endpoint_output.data.rstrip("
")
data = output_data + "," + input_data
return { str(i).zfill(20) : d for i, d in enumerate(data.split(",")) }
```

In another example, suppose your model has optional features and you use \(-1\) to denote that the optional feature has a missing value. If you have a data quality monitor, you may want to remove the \(-1\) from the input value array so that it isn't included in the monitor's metric calculations. You could use a script like the following to remove those values.

```python
def preprocess_handler(inference_record):
    input_data = inference_record.endpoint_input.data
    return {i : None if x == -1 else x for i, x in enumerate(input_data.split(","))}
```

Your preprocessing script receives an `inference_record` as its only input. The following code snippet shows an example of an `inference_record`.

```json
{
    "captureData": {
        "endpointInput": {
            "observedContentType": "text/csv",
            "mode": "INPUT",
            "data": "132,25,113.2,96,269.9,107,,0,0,0,0,0,0,1,0,1,0,0,1",
            "encoding": "CSV"
        }
    }
}
```
The following code snippet shows the full class structure for an `inference_record`.

```python
KEY_EVENT_METADATA = "eventMetadata"
KEY_EVENT_METADATA_EVENT_ID = "eventId"
KEY_EVENT_METADATA_EVENT_TIME = "inferenceTime"
KEY_EVENT_METADATA_CUSTOM_ATTR = "customAttributes"

KEY_EVENTDATA_ENCODING = "encoding"
KEY_EVENTDATA_DATA = "data"

KEY_GROUND_TRUTH_DATA = "groundTruthData"

KEY_EVENTDATA = "captureData"
KEY_EVENTDATA_ENDPOINT_INPUT = "endpointInput"
KEY_EVENTDATA_ENDPOINT_OUTPUT = "endpointOutput"

KEY_EVENTDATA_BATCH_OUTPUT = "batchTransformOutput"
KEY_EVENTDATA_OBSERVED_CONTENT_TYPE = "observedContentType"
KEY_EVENTDATA_MODE = "mode"

KEY_EVENT_VERSION = "eventVersion"

class EventConfig:
    def __init__(self, endpoint, variant, start_time, end_time):
        self.endpoint = endpoint
        self.variant = variant
        self.start_time = start_time
        self.end_time = end_time
```
class EventMetadata:
    def __init__(self, event_metadata_dict):
        self.event_id = event_metadata_dict.get(KEY_EVENT_METADATA_EVENT_ID, None)
        self.event_time = event_metadata_dict.get(KEY_EVENT_METADATA_EVENT_TIME, None)
        self.custom_attribute = event_metadata_dict.get(KEY_EVENT_METADATA_CUSTOM_ATTR, None)

class EventData:
    def __init__(self, data_dict):
        self.encoding = data_dict.get(KEY_EVENTDATA_ENCODING, None)
        self.data = data_dict.get(KEY_EVENTDATA_DATA, None)
        self.observedContentType = data_dict.get(KEY_EVENTDATA_OBSERVED_CONTENT_TYPE, None)
        self.mode = data_dict.get(KEY_EVENTDATA_MODE, None)

def as_dict(self):
    ret = {
        KEY_EVENTDATA_ENCODING: self.encoding,
        KEY_EVENTDATA_DATA: self.data,
        KEY_EVENTDATA_OBSERVED_CONTENT_TYPE: self.observedContentType,
    }
    return ret

class CapturedData:
    def __init__(self, event_dict):
        self.event_metadata = None
        self.endpoint_input = None
        self.endpoint_output = None
        self.batch_transform_output = None
        self.ground_truth = None
        self.event_version = None
        self.event_dict = event_dict
        self._event_dict_postprocessed = False

        if KEY_EVENT_METADATA in event_dict:
            self.event_metadata = EventMetadata(event_dict[KEY_EVENT_METADATA])
        if KEY_EVENTDATA in event_dict:
            if KEY_EVENTDATA_ENDPOINT_INPUT in event_dict[KEY_EVENTDATA]:
                self.endpoint_input = EventData(event_dict[KEY_EVENTDATA][KEY_EVENTDATA_ENDPOINT_INPUT])
if KEY_EVENTDATA_ENDPOINT_OUTPUT in event_dict[KEY_EVENTDATA]:
    self.endpoint_output = EventData(event_dict[KEY_EVENTDATA][KEY_EVENTDATA_ENDPOINT_OUTPUT])

if KEY_EVENTDATA_BATCH_OUTPUT in event_dict[KEY_EVENTDATA]:
    self.batch_transform_output = EventData(event_dict[KEY_EVENTDATA][KEY_EVENTDATA_BATCH_OUTPUT])

if KEY_GROUND_TRUTH_DATA in event_dict:
    self.ground_truth = EventData(event_dict[KEY_GROUND_TRUTH_DATA])

if KEY_EVENT_VERSION in event_dict:
    self.event_version = event_dict[KEY_EVENT_VERSION]

def as_dict(self):
    if self._event_dict_postprocessed is True:
        return self.event_dict

    if KEY_EVENTDATA in self.event_dict:
        if KEY_EVENTDATA_ENDPOINT_INPUT in self.event_dict[KEY_EVENTDATA]:
            self.event_dict[KEY_EVENTDATA][KEY_EVENTDATA_ENDPOINT_INPUT] =
            self.endpoint_input.as_dict()

        if KEY_EVENTDATA_ENDPOINT_OUTPUT in self.event_dict[KEY_EVENTDATA]:
            self.event_dict[KEY_EVENTDATA][KEY_EVENTDATA_ENDPOINT_OUTPUT] =
            self.endpoint_output.as_dict()

        if KEY_EVENTDATA_BATCH_OUTPUT in self.event_dict[KEY_EVENTDATA]:
            self.event_dict[KEY_EVENTDATA][KEY_EVENTDATA_BATCH_OUTPUT] =
            self.batch_transform_output.as_dict()

    self._event_dict_postprocessed = True
    return self.event_dict

def __str__(self):
    return str(self.as_dict())

Custom Sampling

You can also apply a custom sampling strategy in your preprocessing script. To do this, configure Model Monitor's first-party, pre-built container to ignore a percentage of the records according to your specified sampling rate. In the following example, the handler samples 10 percent of the records by returning the record in 10 percent of handler calls and an empty list otherwise.

import random
def preprocess_handler(inference_record):
    # we set up a sampling rate of 0.1
    if random.random() > 0.1:
        # return an empty list
        return []
    input_data = inference_record.endpoint_input.data
    return {i : None if x == -1 else x for i, x in enumerate(input_data.split(','))}

Custom logging for preprocessing script

If your preprocessing script returns an error, check the exception messages logged to CloudWatch to debug. You can access the logger on CloudWatch through the preprocess_handler interface. You can log any information you need from your script to CloudWatch. This can be useful when debug your preprocessing script. The following example shows how you can use the preprocess_handler interface to log to CloudWatch

def preprocess_handler(inference_record, logger):
    logger.info(f'I\'m a processing record: {inference_record}\n')
    logger.debug(f'I\'m debugging a processing record: {inference_record}\n')
    logger.warning(f'I\'m processing record with missing value: {inference_record}\n')
    logger.error(f'I\'m a processing record with bad value: {inference_record}\n')
    return inference_record

Postprocessing Script

Use a postprocessing script when you want to extend the code following a successful monitoring run.

def postprocess_handler():
    print("Hello from post-proc script!")

Bring Your Own Containers

Amazon SageMaker Model Monitor provides a prebuilt container with ability to analyze the data captured from endpoints or batch transform jobs for tabular datasets. If you would like to bring your own container, Model Monitor provides extension points which you can leverage.

Under the hood, when you create a MonitoringSchedule, Model Monitor ultimately kicks off processing jobs. Hence the container needs to be aware of the processing job contract documented
in the Build Your Own Processing Container (Advanced Scenario) topic. Note that Model Monitor kicks off the processing job on your behalf per the schedule. While invoking, Model Monitor sets up additional environment variables for you so that your container has enough context to process the data for that particular execution of the scheduled monitoring. For additional information on container inputs, see the Container Contract Inputs.

In the container, using the above environment variables/context, you can now analyze the dataset for the current period in your custom code. After this analysis is complete, you can chose to emit your reports to be uploaded to an S3 bucket. The reports that the prebuilt container generates are documented in Container Contract Outputs. If you would like the visualization of the reports to work in SageMaker Studio, you should follow the same format. You can also choose to emit completely custom reports.

You also emit CloudWatch metrics from the container by following the instructions in CloudWatch Metrics for Bring Your Own Containers.

Topics
• Container Contract Inputs
• Container Contract Outputs
• CloudWatch Metrics for Bring Your Own Containers

Container Contract Inputs

The Amazon SageMaker Model Monitor platform invokes your container code according to a specified schedule. If you choose to write your own container code, the following environment variables are available. In this context, you can analyze the current dataset or evaluate the constraints if you choose and emit metrics, if applicable.

The available environment variables are the same for real-time endpoints and batch transform jobs, except for the dataset_format variable. If you are using a real-time endpoint, the dataset_format variable supports the following options:

```json
{"sagemakerCaptureJson": {"captureIndexNames": ["endpointInput","endpointOutput"]}}
```

If you are using a batch transform job, the dataset_format supports the following options:

```json
{"csv": {"header": ["true","false"]}}
```
The following code sample shows the complete set of environment variables available for your container code (and uses the dataset_format format for a real-time endpoint).

"Environment": {
  "dataset_format": "{\"sagemakerCaptureJson\": {\"captureIndexNames\": [\"endpointInput\", \"endpointOutput\"]}}",
  "dataset_source": "/opt/ml/processing/endpointdata",
  "end_time": "2019-12-01T16: 20: 00Z",
  "output_path": "/opt/ml/processing/resultdata",
  "publish_cloudwatch_metrics": "Disabled",
  "sagemaker_endpoint_name": "endpoint-name",
  "sagemaker_monitoring_schedule_name": "schedule-name",
  "start_time": "2019-12-01T15: 20: 00Z"
}

### Parameters

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dataset_format</td>
<td>For a job started from a MonitoringSchedule backed by an Endpoint, this is sageMakerCaptureJson with the capture indices endpointInput, or endpointOutput, or both. For a batch transform job, this specifies the data format, whether CSV, JSON, or Parquet.</td>
</tr>
<tr>
<td>dataset_source</td>
<td>If you are using a real-time endpoint, the local path in which the data corresponding to the monitoring period, as specified by start_time and end_time, are available. At this path, the data is available in /{endpoint-name}/{variant-name}/yyyy/mm/dd/hh.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>output_path</td>
<td>We sometimes download more than what is specified by the start and end times. It is up to the container code to parse the data as required.</td>
</tr>
<tr>
<td>publish_cloudwatch_metrics</td>
<td>For a job launched by CreateMonitoringSchedule, this parameter is set to Enabled. The container can choose to write the Amazon CloudWatch output file at [filepath].</td>
</tr>
<tr>
<td>sagemaker_endpoint_name</td>
<td>If you are using a real-time endpoint, the name of the Endpoint that this scheduled job was launched for.</td>
</tr>
<tr>
<td>sagemaker_monitoring_schedule_name</td>
<td>The name of the MonitoringSchedule that launched this job.</td>
</tr>
<tr>
<td><em>sagemaker_endpoint_datacapture_prefix</em></td>
<td>If you are using a real-time endpoint, the prefix specified in the DataCaptureConfig parameter of the Endpoint. The container can use this if it needs to directly access more data than already downloaded by SageMaker at the dataset_source path.</td>
</tr>
<tr>
<td>Parameter Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>start_time, end_time</td>
<td>The time window for this analysis run. For example, for a job scheduled to run at 05:00 UTC and a job that runs on 20/02/2020, start_time: is 2020-02-19T06:00:00Z and end_time: is 2020-02-20T05:00:00Z</td>
</tr>
<tr>
<td>baseline_constraints:</td>
<td>The local path of the baseline constraint file specified in BaselineConfig.ConstraintResource.S3Uri. This is available only if this parameter was specified in the CreateMonitoringSchedule request.</td>
</tr>
<tr>
<td>baseline_statistics</td>
<td>The local path to the baseline statistics file specified in BaselineConfig.StatisticsResource.S3Uri. This is available only if this parameter was specified in the CreateMonitoringSchedule request.</td>
</tr>
</tbody>
</table>

**Container Contract Outputs**

The container can analyze the data available in the *dataset_source* path and write reports to the path in *output_path*. The container code can write any reports that suit your needs.

If you use the following structure and contract, certain output files are treated specially by SageMaker in the visualization and API. This applies only to tabular datasets.

**Output Files for Tabular Datasets**

<table>
<thead>
<tr>
<th>File Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>statistics.json</td>
<td>This file is expected to have columnar statistics for each feature in the dataset that is analyzed. The schema for this file is available in the next section.</td>
</tr>
<tr>
<td>File Name</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>constraints.json</td>
<td>This file is expected to have the constraints on the features observed. The schema for this file is available in the next section.</td>
</tr>
<tr>
<td>constraints_violations.json</td>
<td>This file is expected to have the list of violations found in this current set of data as compared to the baseline statistics and constraints file specified in the baseline_constraints and baseline_statistics path.</td>
</tr>
</tbody>
</table>

In addition, if the publish_cloudwatch_metrics value is "Enabled" container code can emit Amazon CloudWatch metrics in this location: /opt/ml/output/metrics/cloudwatch. The schema for these files is described in the following sections.

Topics

- [Schema for Statistics (statistics.json file)](#)
- [Schema for Constraints (constraints.json file)](#)

Schema for Statistics (statistics.json file)

The schema defined in the statistics.json file specifies the statistical parameters to be calculated for the baseline and data that is captured. It also configures the bucket to be used by KLL, a very compact quantiles sketch with lazy compaction scheme.

```json
{
    "version": 0,
    # dataset level stats
    "dataset": {
        "item_count": number
    },
    # feature level stats
    "features": [
        {
            "name": "feature-name",
            "inferred_type": "Fractional" | "Integral",
```
"numerical_statistics": {
    "common": {
        "num_present": number,
        "num_missing": number
    },
    "mean": number,
    "sum": number,
    "std_dev": number,
    "min": number,
    "max": number,
    "distribution": {
        "kll": {
            "buckets": [
                {
                    "lower_bound": number,
                    "upper_bound": number,
                    "count": number
                }
            ],
            "sketch": {
                "parameters": {
                    "c": number,
                    "k": number
                },
                "data": [
                    [
                        num, num, num, num
                    ],
                    [
                        num, num
                    ],
                    [num, num]
                ]
            }
        }
    }
},
"num_stats"}
Notes

- The specified metrics are recognized by SageMaker in later visualization changes. The container can emit more metrics if required.
- KLL sketch is the recognized sketch. Custom containers can write their own representation, but it won't be recognized by SageMaker in visualizations.
- By default, the distribution is materialized in 10 buckets. You can't change this.

Schema for Constraints (constraints.json file)

A constraints.json file is used to express the constraints that a dataset must satisfy. Amazon SageMaker Model Monitor containers can use the constraints.json file to evaluate datasets against. Prebuilt containers provide the ability to generate the constraints.json file automatically for a baseline dataset. If you bring your own container, you can provide it with similar abilities or you can...
create the constraints.json file in some other way. Here is the schema for the constraint file that the prebuilt container uses. Bring your own containers can adopt the same format or enhance it as required.

```json
{
    "version": 0,
    "features": [
        {
            "name": "string",
            "inferred_type": "Integral" | "Fractional" | "String" | "Unknown",
            "completeness": number,
            "num_constraints": {
                "is_non_negative": boolean
            },
            "string_constraints": {
                "domains": [
                    "list of",
                    "observed values",
                    "for small cardinality"
                ]
            },
            "monitoringConfigOverrides": {}
        }
    ],
    "monitoring_config": {
        "evaluate_constraints": "Enabled",
        "emit_metrics": "Enabled",
        "datatype_check_threshold": 0.1,
        "domain_content_threshold": 0.1,
        "distribution_constraints": {
            "perform_comparison": "Enabled",
            "comparison_threshold": 0.1,
            "comparison_method": "Simple" || "Robust",
            "categorical_comparison_threshold": 0.1,
            "categorical_drift_method": "LInfinity" || "ChiSquared"
        }
    }
}
```
The `monitoring_config` object contains options for monitoring job for the feature. The following table describes each option.

### Monitoring Constraints

<table>
<thead>
<tr>
<th>Constraint</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>evaluate_constraints</code></td>
<td>When Enabled, evaluates whether the current dataset being analyzed satisfies the constraints specified in the <code>constraints.json</code> file taken as a baseline. Valid values: Enabled or Disabled. Default: Enabled.</td>
</tr>
<tr>
<td><code>emit_metrics</code></td>
<td>When Enabled, emits CloudWatch metrics for the data contained in the file. Valid values: Enabled or Disabled. Default: Enabled.</td>
</tr>
<tr>
<td><code>datatype_check_threshold</code></td>
<td>If the threshold is above the value of the specified <code>datatype_check_threshold</code>, this causes a failure that is treated as a violation in the violation report. If the data types in the current execution are not the same as in the baseline dataset, this threshold is used to evaluate if it needs to be flagged as a violation. During the baseline step, the generated constraints suggest the inferred data type for each column. The <code>datatype_check_threshold</code> parameter can be tuned to adjust...</td>
</tr>
<tr>
<td>Constraint</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>the threshold on when it is flagged as a violation.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default: 0.1</td>
</tr>
<tr>
<td>domain_content_threshold</td>
<td>If there are more unknown values for a String field in the current dataset than in the baseline dataset, this threshold can be used to dictate if it needs to be flagged as a violation.</td>
</tr>
<tr>
<td></td>
<td>Valid values: float</td>
</tr>
<tr>
<td></td>
<td>Default: 0.1</td>
</tr>
<tr>
<td>distribution_constraints</td>
<td>perform_comparison</td>
</tr>
<tr>
<td></td>
<td>When Enabled, this flag instructs the code to perform a distribution comparison between the baseline distribution and the distribution observed for the current dataset.</td>
</tr>
<tr>
<td></td>
<td>Valid values: Enabled or Disabled</td>
</tr>
<tr>
<td></td>
<td>Default: Enabled</td>
</tr>
<tr>
<td>Constraint</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>comparison_threshold</td>
<td>If the threshold is above the value set for the <code>comparison_threshold</code>, this causes a failure that is treated as a violation in the violation report. The distance is calculated by getting the maximum absolute difference between the cumulative distribution functions of two distributions. Valid values: float. Default: 0.1</td>
</tr>
<tr>
<td>comparison_method</td>
<td>Whether to calculate <code>linf_simple</code> or <code>linf_robust</code>. The <code>linf_simple</code> is based on the maximum absolute difference between the cumulative distribution functions of two distributions. Calculating <code>linf_robust</code> is based on <code>linf_simple</code>, but is used when there are not enough samples. The <code>linf_robust</code> formula is based on the Two-sample Kolmogorov–Smirnov test. Valid values: <code>linf_simple</code> or <code>linf_robust</code></td>
</tr>
<tr>
<td>Constraint</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>categorical_comparison_threshold</td>
<td>Optional. Sets a threshold for categorical features. If the value in the dataset exceeds the threshold that you set, a violation is recorded in the violation report. Valid values: float Default: The value assigned to the comparison_threshold parameter</td>
</tr>
<tr>
<td>categorical_drift_method</td>
<td>Optional. For categorical features, specifies the computation method used to detect distribution drift. If you don't set this parameter, the K-S (LInfinity) test is used. Valid Values: LInfinity or ChiSquared Default: LInfinity</td>
</tr>
</tbody>
</table>

**CloudWatch Metrics for Bring Your Own Containers**

If the publish_cloudwatch_metrics value is Enabled in the Environment map in the `/opt/ml/processing/processingjobconfig.json` file, the container code emits Amazon CloudWatch metrics in this location: `/opt/ml/output/metrics/cloudwatch`

The schema for this file is closely based on the CloudWatch PutMetrics API. The namespace is not specified here. It defaults to the following:

- For real-time endpoints: `/aws/sagemaker/Endpoint/data-metrics`
- For batch transform jobs: `/aws/sagemaker/ModelMonitoring/data-metrics`

However, you can specify dimensions. We recommend you add the following dimensions at minimum:
• Endpoint and MonitoringSchedule for real-time endpoints
• MonitoringSchedule for batch transform jobs

The following JSON snippets show how to set your dimensions.

For a real-time endpoint, see the following JSON snippet which includes the Endpoint and MonitoringSchedule dimensions:

```json
{
  "MetricName": "", # Required
  "Timestamp": "2019-11-26T03:00:00Z", # Required
  "Dimensions" : ["Endpoint":"endpoint_0",
  "MonitoringSchedule":"schedule_0"],
  "Value": Float,
  # Either the Value or the StatisticValues field can be populated and not both.
  "StatisticValues": {
    "SampleCount": Float,
    "Sum": Float,
    "Minimum": Float,
    "Maximum": Float
  },
  "Unit": "Count", # Optional
}
```

For a batch transform job, see the following JSON snippet which includes the MonitoringSchedule dimension:

```json
{
  "MetricName": "", # Required
  "Timestamp": "2019-11-26T03:00:00Z", # Required
  "Dimensions" : ["MonitoringSchedule":"schedule_0"],
  "Value": Float,
  # Either the Value or the StatisticValues field can be populated and not both.
  "StatisticValues": {
    "SampleCount": Float,
    "Sum": Float,
    "Minimum": Float,
    "Maximum": Float
  },
  "Unit": "Count", # Optional
}
```
Create a Monitoring Schedule for a Real-time Endpoint with an AWS CloudFormation Custom Resource

If you are using a real-time endpoint, you can use an AWS CloudFormation custom resource to create a monitoring schedule. The custom resource is in Python. To deploy it, see [Python Lambda deployment](#).

**Custom Resource**

Start by adding a custom resource to your AWS CloudFormation template. This points to an AWS Lambda function that you create in the next step.

This resource enables you to customize the parameters for the monitoring schedule. You can add or remove more parameters by modifying the AWS CloudFormation resource and the Lambda function in the following example resource.

```json
{
   "AWSTemplateFormatVersion": "2010-09-09",
   "Resources": {
      "MonitoringSchedule": {
         "Type": "Custom::MonitoringSchedule",
         "Version": "1.0",
         "Properties": {
            "ScheduleName": "YourScheduleName",
            "EndpointName": "YourEndpointName",
            "BaselineConstraintsUri": "s3://your-baseline-constraints/constraints.json",
            "BaselineStatisticsUri": "s3://your-baseline-stats/statistics.json",
            "PostAnalyticsProcessorSourceUri": "s3://your-post-processor/postprocessor.py",
            "RecordPreprocessorSourceUri": "s3://your-preprocessor/preprocessor.py",
            "InputLocalPath": "/opt/ml/processing/endpointdata",
            "OutputLocalPath": "/opt/ml/processing/localpath",
            "OutputS3URI": "s3://your-output-uri",
            "ImageURI": "11111111111.dkr.ecr.us-west-2.amazonaws.com/your-image",
            "ScheduleExpression": "cron(0 * ? * * *)",
            "PassRoleArn": "arn:aws:iam::111111111111:role/AmazonSageMaker-ExecutionRole"
         }
      }
   }
}
```
Lambda Custom Resource Code

This AWS CloudFormation custom resource uses the Custom Resource Helper AWS library, which you can install with pip using `pip install crhelper`.

This Lambda function is invoked by AWS CloudFormation during the creation and deletion of the stack. This Lambda function is responsible for creating and deleting the monitoring schedule and using the parameters defined in the custom resource described in the preceding section.

```python
import boto3
import botocore
import logging
from crhelper import CfnResource
from botocore.exceptions import ClientError

logger = logging.getLogger(__name__)
sm = boto3.client('sagemaker')

# cfnhelper makes it easier to implement a CloudFormation custom resource
helper = CfnResource()

# CFN Handlers

def handler(event, context):
    helper(event, context)

@helper.create
def create_handler(event, context):
    """
    Called when CloudFormation custom resource sends the create event
    """
    create_monitoring_schedule(event)

@helper.delete
def delete_handler(event, context):
```

Called when CloudFormation custom resource sends the delete event

```python
schedule_name = get_schedule_name(event)
delete_monitoring_schedule(schedule_name)
```

@helper.poll_create
def poll_create(event, context):
    
    Return true if the resource has been created and false otherwise so CloudFormation polls again.

    ```python
    schedule_name = get_schedule_name(event)
    logger.info('Polling for creation of schedule: %s', schedule_name)
    return is_schedule_ready(schedule_name)
    ```

@helper.update
def noop():
    
    Not currently implemented but crhelper will throw an error if it isn't added

    ```python
    pass
    ```

# Helper Functions
def get_schedule_name(event):
    return event['ResourceProperties']['ScheduleName']

def create_monitoring_schedule(event):
    schedule_name = get_schedule_name(event)
    monitoring_schedule_config = create_monitoring_schedule_config(event)

    logger.info('Creating monitoring schedule with name: %s', schedule_name)
    ```python
    sm.create_monitoring_schedule(
        MonitoringScheduleName=schedule_name,
        MonitoringScheduleConfig=monitoring_schedule_config)
    ```

def is_schedule_ready(schedule_name):
    is_ready = False

    schedule = sm.describe_monitoring_schedule(MonitoringScheduleName=schedule_name)
    status = schedule['MonitoringScheduleStatus']
if status == 'Scheduled':
    logger.info('Monitoring schedule (%s) is ready', schedule_name)
    is_ready = True
elif status == 'Pending':
    logger.info('Monitoring schedule (%s) still creating, waiting and polling again...', schedule_name)
else:
    raise Exception('Monitoring schedule ({}) has unexpected status:
{}' .format(schedule_name, status))

return is_ready

def create_monitoring_schedule_config(event):
    props = event['ResourceProperties']

    return {
        "ScheduleConfig": {
            "ScheduleExpression": props["ScheduleExpression"],
        },
        "MonitoringJobDefinition": {
            "BaselineConfig": {
                "ConstraintsResource": {
                    "S3Uri": props['BaselineConstraintsUri'],
                },
                "StatisticsResource": {
                    "S3Uri": props['BaselineStatisticsUri'],
                }
            },
            "MonitoringInputs": [
                {
                    "EndpointInput": {
                        "EndpointName": props['EndpointName'],
                        "LocalPath": props['InputLocalPath'],
                    }
                }
            ],
            "MonitoringOutputConfig": {
                "MonitoringOutputs": [
                    {
                        "S3Output": {
                            "S3Uri": props['OutputS3URI'],
                            "LocalPath": props['OutputLocalPath'],
                        }
                    }
                ]
            }
        }
    }
def delete_monitoring_schedule(schedule_name):
    logger.info('Deleting schedule: %s', schedule_name)
    try:
        sm.delete_monitoring_schedule(MonitoringScheduleName=schedule_name)
    except ClientError as e:
        if e.response['Error'] ['Code'] == 'ResourceNotFound':
            logger.info('Resource not found, nothing to delete')
        else:
            logger.error('Unexpected error while trying to delete monitoring schedule')
            raise e

### Model Monitor FAQs

Refer to the following FAQs for more information about Amazon SageMaker Model Monitor.

**Q: How do Model Monitor and SageMaker Clarify help customers monitor model behavior?**
Amazon SageMaker Model Monitor automates model monitoring alleviating the need to monitor the models manually or building any additional tooling. In order to automate the process, Model Monitor provides you with the ability to create a set of baseline statistics and constraints using the data with which your model was trained, then set up a schedule to monitor the predictions made on your endpoint. Model Monitor uses rules to detect drift in your models and alerts you when it happens. The following steps describe what happens when you enable model monitoring:

- **Enable model monitoring**: For a real-time endpoint, you have to enable the endpoint to capture data from incoming requests to a deployed ML model and the resulting model predictions. For a batch transform job, enable data capture of the batch transform inputs and outputs.

- **Baseline processing job**: You then create a baseline from the dataset that was used to train the model. The baseline computes metrics and suggests constraints for the metrics. For example, the recall score for the model shouldn't regress and drop below 0.571, or the precision score shouldn't fall below 1.0. Real-time or batch predictions from your model are compared to the constraints and are reported as violations if they are outside the constrained values.

- **Monitoring job**: Then, you create a monitoring schedule specifying what data to collect, how often to collect it, how to analyze it, and which reports to produce.

- **Merge job**: This is only applicable if you are leveraging Amazon SageMaker Ground Truth. Model Monitor compares the predictions your model makes with Ground Truth labels to measure the quality of the model. For this to work, you periodically label data captured by your endpoint or batch transform job and upload it to Amazon S3.

After you create and upload the Ground Truth labels, include the location of the labels as a parameter when you create the monitoring job.

When you use Model Monitor to monitor a batch transform job instead of a real-time endpoint, instead of receiving requests to an endpoint and tracking the predictions, Model Monitor monitors inference inputs and outputs. In a Model Monitor schedule, the customer provides the count and
type of instances that are to be used in the processing job. These resources remain reserved until the schedule is deleted irrespective of the status of current execution.

**Q: What is Data Capture, why is it required, and how can I enable it?**

In order to log the inputs to the model endpoint and the inference outputs from the deployed model to Amazon S3, you can enable a feature called Data Capture. For more details about how to enable it for a real-time endpoint and batch transform job, see Capture data from real-time endpoint and Capture data from batch transform job.

**Q: Does enabling Data Capture impact the performance of a real-time endpoint?**

Data Capture happens asynchronously without impacting production traffic. After you have enabled the data capture, then the request and response payload, along with some additional metadata, is saved in the Amazon S3 location that you specified in the DataCaptureConfig. Note that there can be a delay in the propagation of the captured data to Amazon S3.

You can also view the captured data by listing the data capture files stored in Amazon S3. The format of the Amazon S3 path is: s3://{endpoint-name}/{variant-name}/yyyy/mm/dd/hh/filename.jsonl. Amazon S3 Data Capture should be in the same region as the Model Monitor schedule. You should also ensure that the column names for the baseline dataset only have lowercase letters and an underscore (_) as the only separator.

**Q: Why is Ground Truth needed for model monitoring?**

Ground Truth labels are required by the following features of Model Monitor:

- **Model quality monitoring** compares the predictions your model makes with Ground Truth labels to measure the quality of the model.
- **Model bias monitoring** monitors predictions for bias. One way bias can be introduced in deployed ML models is when the data used in training differs from the data used to generate predictions. This is especially pronounced if the data used for training changes over time (such as fluctuating mortgage rates), and the model prediction is not as accurate unless the model is retrained with updated data. For example, a model for predicting home prices can be biased if the mortgage rates used to train the model differ from the most current real-world mortgage rate.

**Q: For customers leveraging Ground Truth for labeling, what are the steps I can take to monitor the quality of the model?**
Model quality monitoring compares the predictions your model makes with Ground Truth labels to measure the quality of the model. For this to work, you periodically label data captured by your endpoint or batch transform job and upload it to Amazon S3. Besides captures, model bias monitoring execution also requires Ground Truth data. In real use cases, Ground Truth data should be regularly collected and uploaded to the designated Amazon S3 location. To match Ground Truth labels with captured prediction data, there must be a unique identifier for each record in the dataset. For the structure of each record for Ground Truth data, see Ingest Ground Truth Labels and Merge Them With Predictions.

The following code example can be used for generating artificial Ground Truth data for a tabular dataset.

```python
import random

def ground_truth_with_id(inference_id):
    random.seed(inference_id)  # to get consistent results
    rand = random.random()

    # format required by the merge container
    return {
        "groundTruthData": {
            "data": "1" if rand < 0.7 else "0",  # randomly generate positive labels
        },
        "encoding": "CSV",
        "eventId": str(inference_id),
        "eventVersion": "$0$",
    }

def upload_ground_truth(upload_time):
    records = [ground_truth_with_id(i) for i in range(test_dataset_size)]
    fake_records = [json.dumps(r) for r in records]
    data_to_upload = "\n".join(fake_records)
    target_s3_uri = f"{ground_truth_upload_path}/{upload_time:%Y/%m/%d/%H%M%S}.jsonl"
    print(f"Uploading {len(fake_records)} records to", target_s3_uri)
    S3Uploader.upload_string_as_file_body(data_to_upload, target_s3_uri)

    # Generate data for the last hour
    upload_ground_truth(datetime.utcnow() - timedelta(hours=1))
    # Generate data once a hour
    def generate_fake_ground_truth(terminate_event):
```
upload_ground_truth(datetime.utcnow())
for _ in range(0, 60):
    time.sleep(60)
    if terminate_event.is_set():
        break

ground_truth_thread = WorkerThread(do_run=generate_fake_ground_truth)
ground_truth_thread.start()

The following code example shows how to generate artificial traffic to send to the model endpoint. Notice the inferenceId attribute used above to invoke. If this is present, it is used to join with Ground Truth data (otherwise, the eventId is used).

```python
import threading

class WorkerThread(threading.Thread):
    def __init__(self, do_run, *args, **kwargs):
        super(WorkerThread, self).__init__(*args, **kwargs)
        self.__do_run = do_run
        self.__terminate_event = threading.Event()

    def terminate(self):
        self.__terminate_event.set()

    def run(self):
        while not self.__terminate_event.is_set():
            self.__do_run(self.__terminate_event)

def invoke_endpoint(terminate_event):
    with open(test_dataset, "r") as f:
        i = 0
        for row in f:
            payload = row.rstrip("\n")
            response = sagemaker_runtime_client.invoke_endpoint(
                EndpointName=endpoint_name,
                ContentType="text/csv",
                Body=payload,
                InferenceId=str(i),  # unique ID per row
            )
            i += 1
            response["Body"].read()
            time.sleep(1)
            if terminate_event.is_set():
```
# Keep invoking the endpoint with test data
invoke_endpoint_thread = WorkerThread(do_run=invoke_endpoint)
invoke_endpoint_thread.start()

You must upload Ground Truth data to an Amazon S3 bucket that has the same path format as captured data, which is in the following format: s3://<bucket>/<prefix>/yyyy/mm/dd/hh

**Note**

The date in this path is the date when the Ground Truth label is collected. It doesn't have to match the date when the inference was generated.

Q: How can customers customize monitoring schedules?

In addition to using the built-in monitoring mechanisms, you can create your own custom monitoring schedules and procedures using pre-processing and post-processing scripts, or by using or building your own container. It's important to note that pre-processing and post-processing scripts only work with data and model quality jobs.

Amazon SageMaker provides the capability for you to monitor and evaluate the data observed by the model endpoints. For this, you have to create a baseline with which you compare the real-time traffic. After a baseline is ready, set up a schedule to continuously evaluate and compare against the baseline. While creating a schedule, you can provide the pre-processing and post-processing script.

The following example shows how you can customize monitoring schedules with pre-processing and post-processing scripts.

```python
import boto3, os
from sagemaker import get_execution_role, Session
from sagemaker.model_monitor import CronExpressionGenerator, DefaultModelMonitor

# Upload pre and postprocessor scripts
session = Session()
bucket = boto3.Session().resource("s3").Bucket(session.default_bucket())
prefix = "demo-sagemaker-model-monitor"
pre_processor_script = bucket.Object(os.path.join(prefix, "preprocessor.py")).upload_file("preprocessor.py")
```
post_processor_script = bucket.Object(os.path.join(prefix, "postprocessor.py")).upload_file("postprocessor.py")
# Get execution role
role = get_execution_role() # can be an empty string
# Instance type
instance_type = "instance-type"
# instance_type = "ml.m5.xlarge" # Example
# Create a monitoring schedule with pre and post-processing
my_default_monitor = DefaultModelMonitor(
    role=role,
    instance_count=1,
    instance_type=instance_type,
    volume_size_in_gb=20,
    max_runtime_in_seconds=3600,
)

s3_report_path = "s3://{}/{}".format(bucket, "reports")
monitor_schedule_name = "monitor-schedule-name"
endpoint_name = "endpoint-name"
my_default_monitor.create_monitoring_schedule(
    post_analytics_processor_script=post_processor_script,
    record_preprocessor_script=pre_processor_script,
    monitor_schedule_name=monitor_schedule_name,
    # use endpoint_input for real-time endpoint
    endpoint_input=endpoint_name,
    # or use batch_transform_input for batch transform jobs
    output_s3_uri=s3_report_path,
    statistics=my_default_monitor.baseline_statistics(),
    constraints=my_default_monitor.suggested_constraints(),
    schedule_cron_expression=CronExpressionGenerator.hourly(),
    enable_cloudwatch_metrics=True,
)

Q: What are some of the scenarios or use cases where I can leverage a pre-processing script?

You can use pre-processing scripts when you need to transform the inputs to your model monitor. Consider the following example scenarios:

1. Pre-processing script for data transformation.

Suppose the output of your model is an array: [1.0, 2.1]. The Model Monitor container only works with tabular or flattened JSON structures, such as{"prediction0": 1.0,
“prediction1” : 2.1}. You could use a pre-processing script like the following example to transform the array into the correct JSON structure.

```python
def preprocess_handler(inference_record):
    input_data = inference_record.endpoint_input.data
    output_data = inference_record.endpoint_output.data.rstrip("\n")
    data = output_data + "," + input_data
    return { str(i).zfill(20) : d for i, d in enumerate(data.split(","))) }
```

2. Exclude certain records from Model Monitor's metric calculations.

Suppose your model has optional features and you use -1 to denote that the optional feature has a missing value. If you have a data quality monitor, you may want to remove the -1 from the input value array so that it isn't included in the monitor's metric calculations. You could use a script like the following to remove those values.

```python
def preprocess_handler(inference_record):
    input_data = inference_record.endpoint_input.data
    return {i : None if x == -1 else x for i, x in enumerate(input_data.split(","))) }
```

3. Apply a custom sampling strategy.

You can also apply a custom sampling strategy in your pre-processing script. To do this, configure Model Monitor's first-party, pre-built container to ignore a percentage of the records according to your specified sampling rate. In the following example, the handler samples 10% of the records by returning the record in 10% of handler calls and an empty list otherwise.

```python
import random
def preprocess_handler(inference_record):
    # we set up a sampling rate of 0.1
    if random.random() > 0.1:
        # return an empty list
        return []
    input_data = inference_record.endpoint_input.data
    return {i : None if x == -1 else x for i, x in enumerate(input_data.split(","))) }
```

4. Use custom logging.
You can log any information you need from your script to Amazon CloudWatch. This can be useful when debugging your pre-processing script in case of an error. The following example shows how you can use the `preprocess_handler` interface to log to CloudWatch.

```python
def preprocess_handler(inference_record, logger):
    logger.info(f"I'm a processing record: {inference_record}")
    logger.debug(f"I'm debugging a processing record: {inference_record}")
    logger.warning(f"I'm processing record with missing value: {inference_record}")
    logger.error(f"I'm a processing record with bad value: {inference_record}")
    return inference_record
```

**Note**

When the pre-processing script is run on batch transform data, the input type is not always the `CapturedData` object. For CSV data, the type is a string. For JSON data, the type is a Python dictionary.

**Q: When can I leverage a post-processing script?**

You can leverage a post-processing script as an extension following a successful monitoring run. The following is a simple example, but you can perform or call any business function that you need to perform after a successful monitoring run.

```python
def postprocess_handler():
    print("Hello from the post-processing script!")
```

**Q: When should I consider bringing my own container for model monitoring?**

SageMaker provides a pre-built container for analyzing data captured from endpoints or batch transform jobs for tabular datasets. However, there are scenarios where you might want to create your own container. Consider the following scenarios:

- You have regulatory and compliance requirements to only use the containers that are created and maintained internally in your organization.
- If you want to include a few third-party libraries, you can place a `requirements.txt` file in a local directory and reference it using the `source_dir` parameter in the SageMaker.
estimator, which enables library installation at run-time. However, if you have lots of libraries or dependencies that increase the installation time while running the training job, you might want to leverage BYOC.

- Your environment forces no internet connectivity (or Silo), which prevents package download.
- You want to monitor data that's in data formats other than tabular, such as NLP or CV use cases.
- When you require additional monitoring metrics than the ones supported by Model Monitor.

Q: I have NLP and CV models. How do I monitor them for data drift?

Amazon SageMaker's prebuilt container supports tabular datasets. If you want to monitor NLP and CV models, you can bring your own container by leveraging the extension points provided by Model Monitor. For more details about the requirements, see Bring your own containers. Some of the following are more examples:

- For a detailed explanation of how to use Model Monitor for a computer vision use case, see Detecting and Analyzing incorrect predictions.
- For a scenario where Model Monitor can be leveraged for a NLP use case, see Detect NLP data drift using custom Amazon SageMaker Model Monitor.

Q: I want to delete the model endpoint for which Model Monitor was enabled, but I’m unable to do so since the monitoring schedule is still active. What should I do?

If you want to delete an inference endpoint hosted in SageMaker which has Model Monitor enabled, first you must delete the model monitoring schedule (with the DeleteMonitoringSchedule CLI or API). Then, delete the endpoint.

Q: Does SageMaker Model Monitor calculate metrics and statistics for input?

Model Monitor calculates metrics and statistics for output, not input.

Q: Does SageMaker Model Monitor support multi-model endpoints?

No, Model Monitor only supports endpoints that host a single model and doesn't support monitoring multi-model endpoints.

Q: Does SageMaker Model Monitor provide monitoring data about individual containers in an inference pipeline?
Model Monitor supports monitoring inference pipelines, but capturing and analyzing data is done for the entire pipeline, not for individual containers in the pipeline.

**Q: What can I do to prevent impact to inference requests when data capture is set up?**

To prevent impact to inference requests, Data Capture stops capturing requests at high levels of disk usage. It is recommended you keep your disk utilization below 75% in order to ensure data capture continues capturing requests.

**Q: Can Amazon S3 Data Capture be in a different AWS region than the region in which the monitoring schedule was set up?**

No, Amazon S3 Data Capture must be in the same region as the monitoring schedule.

**Q: What is a baseline, and how do I create one? Can I create a custom baseline?**

A baseline is used as a reference to compare real-time or batch predictions from the model. It computes statistics and metrics along with constraints on them. During monitoring, all of these are used in conjunction to identify violations.

To use the default solution of Amazon SageMaker Model Monitor, you can leverage the Amazon SageMaker Python SDK. Specifically, use the `suggest_baseline` method of the `ModelMonitor` or the `ModelQualityMonitor` class to trigger a processing job that computes the metrics and constraints for the baseline.

The result of a baseline job are two files: `statistics.json` and `constraints.json`. [Schema for statistics](#) and [schema for constraints](#) contain the schema of the respective files. You can review the generated constraints and modify them before using them for monitoring. Based on your understanding of the domain and business problem, you can make a constraint more aggressive, or relax it to control the number and nature of the violations.

**Q: What are the guidelines to create a baseline dataset?**

The primary requirement for any kind of monitoring is to have a baseline dataset that is used to compute metrics and constraints. Typically, this is the training dataset used by the model, but in some cases you might choose to use some other reference dataset.

The column names of the baseline dataset should be compatible with Spark. In order to maintain the maximum compatibility between Spark, CSV, JSON and parquet it is advisable to use only lowercase letters, and only use _ as the separator. Special characters including “” can cause issues.
Q: What are the StartTimeOffset and EndTimeOffset parameters, and when are they used?

When Amazon SageMaker Ground Truth is required for monitoring jobs like model quality, you need to ensure that a monitoring job only uses data for which Ground Truth is available. The start_time_offset and end_time_offset parameters of EndpointInput can be used to select the data that the monitoring job uses. The monitoring job uses the data in the time window that is defined by start_time_offset and end_time_offset. These parameters need to be specified in the ISO 8601 duration format. The following are some examples:

- If your Ground Truth results arrive 3 days after the predictions have been made, set start_time_offset="-P3D" and end_time_offset="-P1D", which is 3 days and 1 day respectively.
- If the Ground Truth results arrive 6 hours after the predictions and you have an hourly schedule, set start_time_offset="-PT6H" and end_time_offset="-PT1H", which is 6 hours and 1 hour.

Q: Can I run ‘on demand’ monitoring jobs?

Yes, you can run ‘on demand’ monitoring jobs by running a SageMaker Processing job. For Batch Transform, SageMaker Pipelines has a MonitorBatchTransformStep that you can use to create a SageMaker pipeline that runs monitoring jobs on demand. The SageMaker examples repository has code samples for running data quality and model quality monitoring jobs on demand.

Q: How do I set up Model Monitor?

You can set up Model Monitor in the following ways:

- **Amazon SageMaker Python SDK** – There is a Model Monitor module which contains classes and functions that assist in suggesting baselines, creating monitoring schedules, and more. See the Amazon SageMaker Model Monitor notebook examples for detailed notebooks that leverage the SageMaker Python SDK for setting up Model Monitor.
- **SageMaker Pipelines** – SageMaker Pipelines are integrated with Model Monitor through the QualityCheck Step and ClarifyCheckStep APIs. You can create a SageMaker pipeline that contains these steps and that can be used to run monitoring jobs on demand whenever the pipeline is executed.
- **Amazon SageMaker Studio Classic** – You can create a data or model quality monitoring schedule along with model bias and explainability schedules directly from the UI by selecting an
endpoint from the list of deployed model endpoints. Schedules for other types of monitoring can be created by selecting the relevant tab in the UI.

- **SageMaker Model Dashboard** – You can enable monitoring on endpoints by selecting a model that has been deployed to an endpoint. In the following screenshot of the SageMaker console, a model named group1 has been selected from the **Models** section of the **Model dashboard**. On this page, you can create a monitoring schedule, and you can edit, activate or deactivate existing monitoring schedules and alerts. For a step by step guide on how to view alerts and model monitor schedules, see [View Model Monitor schedules and alerts](#).

![](image)

**Q: How does Model Monitor Integrate with SageMaker Model Dashboard**

[SageMaker Model Dashboard](#) gives you unified monitoring across all your models by providing automated alerts about deviations from expected behavior and troubleshooting to inspect models and analyze factors impacting model performance over time.
Evaluate, explain, and detect bias in models

Amazon SageMaker offers features to improve your machine learning (ML) models by detecting potential bias and helping explain the predictions that models make. It helps you identify various types of bias in pre-training data and in post-training that can emerge during model training or when the model is in production. You can also evaluate a language model for model quality and responsibility metrics using foundation model evaluations.

The following topics give information about how to evaluate, explain, and detect bias with Amazon SageMaker.

Topics

- Use SageMaker Clarify to evaluate large language models
- Use SageMaker Clarify to explain and detect bias
- Use SageMaker Clarify explainability with SageMaker Autopilot

Use SageMaker Clarify to evaluate large language models

⚠️ Important

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see Migrating from Amazon SageMaker Studio Classic. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

SageMaker Clarify offers Foundation Model Evaluations (FMEval) as a single place to evaluate and compare model quality and responsibility metrics for any large language model (LLM). Use FMEval to evaluate pre-trained and fine-tuned language models, and text classifiers. A foundation model serves as a starting point, from which you can develop downstream natural language processing (NLP) applications.
You can create a model evaluation job using pre-trained, text-based foundation models available in SageMaker JumpStart or use SageMaker JumpStart model that you have previously deployed to an endpoint.

**What are foundation model evaluations?**

FMEval can help you quantify model risks, such as inaccurate, toxic, or biased content. Evaluating your LLM helps you comply with international guidelines around responsible generative AI, such as the [ISO 42001](https://www.iso.org) AI Management System Standard and the NIST AI Risk Management Framework.

Model evaluation jobs can evaluate LLMs that generate responses for the following tasks:

- **Open-ended generation** – The production of natural human responses to text that does not have a pre-defined structure.
- **Text summarization** – The generation of a concise and condensed summary while retaining the meaning and key information that's contained in larger text.
- **Question answering** – The generation of a relevant and accurate response to a prompt.
- **Classification** – Assigning a category, such as a label or score to text, based on its content.
- **Custom** – Allows you to define custom evaluation dimensions for your model evaluation job.

Each of these tasks have specific metrics associated with them that you can use to evaluate your model.

**Automatic model evaluation jobs**

Automatic model evaluation jobs use metrics based on benchmarks to measure toxic, harmful, or otherwise poor responses to your customers. Model responses are scored using either built-in datasets specific to the task or you can specify your own custom prompt dataset. To create an automatic model evaluation job, you can use Amazon SageMaker Studio or the `fmeval` library in your own code that you can customize for your use case.

To help ensure that the SageMaker JumpStart model you select performs well against all prompts SageMaker Clarify automatically formats your input prompts to be in a format that works best for the **Evaluation dimensions** you select. To see how SageMaker Clarify will modify the prompts, choose **prompt template** under an **Evaluation dimensions** you have added to the model evaluation job.
The toggle allows you to turn off or to turn on the automatic prompt templating support that SageMaker Clarify provides. Turning off the automatic prompt templating allows you to specify your own custom prompts that will be applied to all prompts in your dataset for that evaluation dimension. The following is an example using `meta-textgenerationneuron-llama-2-7b-f`.

**Example Example of an updated prompt template using Prompt template**

Imagine a simple scenario where you have a simple dataset made up of only two prompts, and you want to evaluate them using `meta-textgenerationneuron-llama-2-7b-f`.

```json
{
  "model_input": "Is himalaya the highest mountain in the world?",
  "target_output": "False, Mt. Everest is the highest mountain in the world",
  "category": "Geography"
},
{
  "model_input": "Is Olympia the capital of Washington?",
  "target_output": "True",
  "category": "Capitals"
}
```

Because your prompts are question and answer pairs, you choose the **Question Answering (Q&A)** task type.

By choosing **Prompt template** in Studio you can see how SageMaker Clarify will format your prompts to match the requirements of the `meta-textgenerationneuron-llama-2-7b-f` SageMaker JumpStart model.

```inst
[INST]<<<SYS>>Respond to the following question. Valid answers are "True" or "False".<<<SYS>>Is himalaya the highest mountain in the world?[/INST]
```

For this model SageMaker Clarify will supplement your prompts to contain the correct prompt format by adding the [INST] and <<<SYS>>tags. It will also augment your initial request by adding Respond to the following question. Valid answers are "True" or "False". to help the model respond better.

The SageMaker Clarify provided text might not be well suited for your use case. To turn off the default prompt templates, slide the **Dataset default prompt templates** toggle to **Off**.

You can edit the prompt template to be aligned with your use case.
Respond to the following question with a short response. Is himalaya the highest mountain in the world?

Now all built-in or custom prompt datasets under the specified Evaluation dimension will use the prompt template you specified.

To better understand your results, use the generated report. The report includes visualizations and examples.

To use a model not publicly available in SageMaker JumpStart, you must use the fmeval library to run the automatic model evaluation job. For a list of SageMaker JumpStart models, see Explore the latest foundation models.

Model evaluation jobs that use humans workers

You can also employ human workers to manually evaluate your model responses for more subjective dimensions, such as helpfulness or style. To create a model evaluation job that uses human workers, you must use Studio.

In a model evaluation job that uses human workers you can compare the responses for up to two SageMaker JumpStart models. Optionally, you can also specify responses from models outside of AWS. All model evaluation jobs that use human workers require that you create a custom prompt dataset, and store it in Amazon S3.

In Studio, you can define the criteria that your human workforce uses to evaluate responses from models. You can also document evaluation instructions using a template available in Studio. Furthermore, you can create a work team in Studio. The work team are people who you want to participate in your model evaluation job.

Get started with model evaluations

Important

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update
A large language model (LLM) is a machine learning model that can analyze and generate natural language text. If you want to evaluate an LLM, SageMaker provides the following three options that you can choose:

- Set up manual evaluations for a human workforce using Studio.
- Evaluate your model with an algorithm using Studio.
- Automatically evaluate your model with a customized work flow using the fmeval library.

You can either use an algorithm to automatically evaluate your foundation model or ask a human work team to evaluate the models' responses.

Human work teams can evaluate and compare up to two models concurrently using metrics that indicate preference for one response over another. The work flow, metrics, and instructions for a human evaluation can be tailored to fit a particular use case. Humans can also provide a more refined evaluation than an algorithmic evaluation.

You can also use an algorithm to evaluate your LLM using benchmarks to rapidly score your model responses in Studio. Studio provides a guided work flow to evaluate responses from a SageMaker JumpStart model using pre-defined metrics. These metrics are specific to generative AI tasks. This guided flow uses built-in or custom datasets to evaluate your LLM.

Alternatively, you can use the fmeval library to create a more customized work flow using automatic evaluations than what is available in Studio. Using Python code and the fmeval library, you can evaluate any text-based LLM, including models that were created outside of SageMaker JumpStart.

The following topics provide an overview of foundation model evaluations, a summary of the automatic and human Foundation Model Evaluation (FMEval) work flows, how to run them, and how to view an analysis report of your results. The automatic evaluation topic shows how to configure and run both a starting and customized evaluation.

Topics

- Foundation model evaluation overview
In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see Migrating from Amazon SageMaker Studio Classic. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

The following sections provide an overview of how to use automatic and human-based model evaluation jobs.

Model evaluation tasks

In a model evaluation job, an evaluation task is a task you want the model to perform based on information found in the prompts.

You can choose one task type per model evaluation job. Use the following sections to learn more about each task type. Each section also includes a list of available built-in datasets and their corresponding metrics that can be used only in automatic model evaluation jobs.

Open-ended generation

Open-ended text generation is a foundation model task that generates natural language responses to prompts that don't have a pre-defined structure, such as general-purpose queries to a chatbot. For open-ended text generation, Foundation Model Evaluations (FMEval) can evaluate your model along the following dimensions.

- **Factual knowledge** – Evaluates how well your model encodes factual knowledge. FMEval can measure your model against your own custom dataset or use a built-in dataset based on the TREX open source dataset.
• **Semantic robustness** – Evaluates how much your model output changes as the result of small, semantic-preserving changes in the input. FMEval measures how your model output changes as a result of keyboard typos, random changes to uppercase, and random additions or deletions of white spaces.

• **Prompt stereotyping** – Measures the probability of your model encoding biases in its response. These biases include those for race, gender, sexual orientation, religion, age, nationality, disability, physical appearance, and socioeconomic status. FMEval can measure your model responses against your own custom dataset or use a built-in dataset based on the CrowS-Pairs open source challenge dataset.

• **Toxicity** – Evaluates text using toxicity detection models. FMEval checks your model for sexual references, rude, unreasonable, hateful or aggressive comments, profanity, insults, flirtations, attacks on identities, and threats. FMEval can measure your model against your own custom dataset or use built-in datasets based on the RealToxicityPrompts, RealToxicityPromptsChallenging, and BOLD datasets.

RealToxicityPromptsChallenging is a subset of RealToxicityPrompts that is used to test the limits of a large language model (LLM). It also identifies areas where LLMs are vulnerable to generating toxic text.

You can evaluate your model with the following toxicity detectors:

• **UnitaryAI Detoxify-unbiased** – A multi-label text classifier trained on Toxic Comment Classification Challenge and Jigsaw Unintended Bias in Toxicity Classification. The model provides 7 scores for the following classes: toxicity, severe toxicity, obscenity, threat, insult, sexual explicit and identity attack.

• **Toxigen-roberta** – A binary RoBERTa-based text classifier fine-tuned on the ToxiGen dataset. The ToxiGen dataset contains sentences with subtle and implicit toxicity pertaining to minority groups.

**Text summarization**

Text summarization is used for tasks, such as creating summaries of news, legal documents, academic papers, content previews, and content curation. The following can influence the quality of responses: ambiguity, coherence, bias, fluency of the text used to train the foundation model, and information loss, accuracy, relevance, or context mismatch. FMEval can evaluate your model against your own custom dataset or use built-in datasets based on the Government Report.
Dataset, and Gigaword datasets. For text summarization, FMEval can evaluate your model for the following:

- **Accuracy** – A numerical score indicating the similarity of the summarization to a reference summary that is accepted as a gold standard. A high numerical score indicates that the summary is of high quality. A low numerical score indicates a poor summary. The following metrics are used to evaluate the accuracy of a summarization:
  - **ROUGE-N** – Computes N-gram overlaps between the reference and model summary.
  - **Meteor** – Computes the word overlap between the reference and model summary while also accounting for rephrasing.
  - **BERTScore** – Computes and compares sentence embeddings for the summarization and reference. FMEval uses the roberta-large-mnli or microsoft/deberta-xlarge-mnli models to compute the embeddings.

- **Toxicity** – Scores for generated summaries that are calculated using a toxicity detector model. For additional information, see the Toxicity section in the previous for Open-ended generation task for details.

- **Semantic robustness** – A measure of how much the quality of your model's text summary changes as the result of small, semantic-preserving changes in the input. Examples of these changes include keyboard typos, the inaccurate conversion of numbers to words, random changes to uppercase, and random additions or deletions of white spaces. Semantic robustness uses the absolute difference in accuracy between a text summary that is unperturbed and one that is perturbed. The accuracy algorithm uses the ROUGE-N, Meteor, and BERTScore metrics, as detailed previously in this section.

### Question answering

Question answering is used for tasks such as generating automatic help-desk responses, information retrieval, and e-learning. FMEval can evaluate your model against your own custom dataset or use built-in datasets based on the BoolQ, TriviaQA, and Natural Questions datasets. For question answering, FMEval can evaluate your model for the following:

- **Accuracy** – An average score comparing the generated response to the question answer pairs given in the references. The score is averaged from the following methods:
  - **Exact match** – A binary score of 1 is assigned to an exact match, and 0 otherwise.
  - **Quasi-exact match** – A binary score of 1 is assigned to a match with grammatical articles (such as the, a, and) after punctuation has been removed.
• **F1 over words** – The F1 score, or harmonic mean of precision and recall between the normalized response and reference. The F1 score is equal to twice precision multiplied by recall divided by the sum of precision (P) and recall (R), or \( F1 = \frac{2*P*R}{P + R} \).

In the previous calculation, precision is defined as the number of true positives (TP) divided by the sum of true positives and false positives (FP), or \( P = \frac{TP}{TP+FP} \).

Recall is defined as the number of true positives divided by the sum of true positives and false negatives (FN), or \( R = \frac{TP}{TP+FN} \).

A higher F1 over words score indicates higher quality responses.

• **Semantic robustness** – A measure of how much the quality of your model's text summary changes as the result of small, semantic-preserving changes in the input. Examples of these changes include keyboard typos, the inaccurate conversion of numbers to words, random changes to uppercase, and random additions or deletions of white spaces. Semantic robustness uses the absolute difference in accuracy between a text summary that is unperturbed and one that is perturbed. Accuracy is measured using exact-match, quasi-exact match and F1 over words, as described previously.

• **Toxicity** – Scores evaluate generated answers using a toxicity detector model. For additional information, see the **Toxicity** section in the previous for **Open-ended generation** task for details.

**Classification**

Classification is used to categorize text into pre-defined categories. Applications that use text classification include content recommendation, spam detection, language identification and trend analysis on social media. Imbalanced, ambiguous, noisy data, bias in labeling are some issues that can cause errors in classification. FMEval evaluates your model against a built-in dataset based on the **Women's ECommerce Clothing Reviews** dataset, and/or against your own prompt datasets for the following.

• **Accuracy** – A score that compares the predicted class to its label. Accuracy is measured using the following metrics:

  • **Classification accuracy** – A binary score of 1 if the predicted label equals the true label, and 0 otherwise.

  • **Precision** – The ratio of true positives to all positives, calculated over the entire dataset. Precision is an appropriate measure when reducing false positives is important.
The score for each data point can be aggregated using the following values for the `multiclass_average_strategy` parameter:

- **micro** (default) – The sum of true positive values across all classes divided by the sum of all positives across all classes. This aggregation type gives a measure of the overall positive predictive accuracy of your model. It does this by using equal weights for each response without regard to its class. For example, this aggregation can assess your model's ability to predict a correct diagnosis for each patient.

- **macro** – The sum of precision values calculated for each class divided by the number of classes. This aggregation type gives a measure of the overall positive predictive accuracy of your model, which gives equal weight to each class. For example, this aggregation can assess your model's ability to predict a correct disease using data from a large number of patients.

- **samples** (multi-class classification only) – The ratio of the sum of true positives for all samples to the sum of true positives and false positives for all samples. For multi-class classification, a sample consists of a set of predicted responses for each class. This aggregation type gives a granular measure of each sample's precision for multi-class problems. For example, because aggregating by samples treats each sample equally, this aggregation can assess your model's ability to predict a correct diagnosis for a patient with a rare disease, and highlighting false positives.

- **weighted** – The weight for one class multiplied by the precision for the same class, summed over all classes. This aggregation type provides a measure of overall precision while accommodating varying importance values among classes. For example, this aggregation can assess your model's ability to predict a correct diagnosis for a patient and give a higher weight to diseases that are life-threatening.

- **binary** – The precision calculated for the class that is specified by the value `pos_label`. This aggregation type ignores the unspecified class, and gives the overall predictive accuracy for a single class. For example, this aggregation can assess your model's ability to detect a rare disease changes over time.

- **none** – The precision calculated for each class. Class-specific precision can help you focus on the positive predictive value for each class. For example, this aggregation can help you decide which medical conditions your model is able to predict well, and which conditions may warrant additional focus.

- **Recall** – the ratio of true positives to the sum of true positives and false negatives, calculated over the entire dataset. Recall is an appropriate measure when reducing false negatives is important. The scores for each data point can be aggregated using the following values for the `multiclass_average_strategy` parameter.
- **micro** (default) – The sum of the true positives divided by the sum of true positives and false negatives for all classes. This aggregation type gives a measure of the overall predictive accuracy of your model, while considering all classes equally. For example, this aggregation can assess your model's ability to correctly classify patients with any disease including rare diseases, because it gives equal weight to all classes.

- **macro** – The sum of recall values calculated for each class divided by the number of classes. This aggregation type gives a measure of the predictive accuracy of your model for each class, with equal weight to each class. For example, this aggregation can assess your model's ability to predict all diseases, regardless of the prevalence or rarity of each condition.

- **samples** (multi-class classification only) – The ratio of the sum of true positives over all samples to the sum of true positives and false negatives for all samples. For multi-class classification, a sample consists of a set of predicted responses for each class. This aggregation type gives a granular measure of each sample's recall for multi-class problems. For example, because aggregating by samples treats each sample equally, this aggregation can assess your model's ability to predict a correct diagnosis for a patient with a rare disease while also minimizing false negatives.

- **weighted** – The weight for one class multiplied by the recall for the same class, summed over all classes. This aggregation type provides a measure of overall recall while accommodating varying importances among classes. For example, this aggregation can assess your model's ability to predict a correct diagnosis for a patient and give a higher weight to diseases that are life-threatening.

- **binary** – The recall calculated for the class that is specified by the value `pos_label`. This aggregation type ignores the unspecified class, and gives overall predictive accuracy for a single class. For example, this aggregation can assess your model's ability to screen a population for a specific highly contagious life-threatening disease.

- **none** – The recall calculated for each class. Class-specific recall can help you address class imbalances in your data when the penalty for error varies significantly between classes. For example, this aggregation can assess how well your model can identify all patients that may have a specific disease.

- **Balanced classification accuracy (BCA)** – The sum of recall and the true negative rate divided by 2 for binary classification. The true negative rate is the number of true negatives divided by the sum of true negatives and false positives. For multi-class classification, BCA is calculated as the sum of recall values for each class divided by the number of classes. BCA can help when the penalty for predicting both false positives and false negatives is high. For example, BCA
can assess how well your model can predict a number of highly contagious lethal diseases with intrusive treatments.

- **Semantic robustness** – Evaluates how much your model output changes as the result of small, semantic-preserving changes in the input. FMEval measures your model output as a result of keyboard typos, random changes to uppercase, and random additions or deletions of white spaces. Semantic robustness scores the absolute difference in accuracy between a text summary that is unperturbed and one that is perturbed.

**Types of foundation model evaluations**

The following sections provide details about both human and algorithmic types of evaluations for your foundation model.

**Human evaluations**

To evaluate your model by a human, you must define the metrics and associated metric types. If you want to evaluate more than one model, you can use a comparative or individual rating mechanism. If you want to evaluate one model, you must use an individual rating mechanism. The following rating mechanisms can be applied to any text-related task:

- (Comparative) **Likert scale - comparison** – A human evaluator will indicate their preference between two responses on a 5-point Likert scale according to your instructions. In the final report, the results will be shown as a histogram of ratings by preference strength over your whole dataset. Define the important points of the 5-point scale in your instructions so that your evaluators know how to rate the responses according to your expectations.

- (Comparative) **Choice buttons** – Allows a human evaluator to indicate one preferred response over another response using radio buttons, according to your instructions. The results in the final report will be shown as a percentage of responses that workers preferred for each model. Explain your evaluation method clearly in the instructions.

- (Comparative) **Ordinal rank** – Allows a human evaluator to rank their preferred responses to a prompt in order, starting at 1, and according to your instructions. In the final report, the results display as a histogram of the rankings from the evaluators over the whole dataset. Make sure that you define what a rank of 1 means in your instructions.

- (Individual) **Thumbs up/down** – Allows a human evaluator to rate each response from a model as acceptable or unacceptable according to your instructions. In the final report, the results show a percentage of the total number of ratings by evaluators that received a thumbs up rating for each model. You can use this rating method to evaluate one or more models. If you use this in
an evaluation that contains two models, the UI presents your work team with a thumbs up or down option for each model response. The final report will show the aggregated results for each model individually. Define what is an acceptable response in your instructions to your work team.

- **(Individual) Likert scale - individual** – Allows a human evaluator to indicate how strongly they approve of the model response, based on your instructions, on a 5-point Likert scale. In the final report, the results display a histogram of the 5-point ratings from the evaluators over your whole dataset. You can use this rating method for an evaluation containing one or more models. If you select this rating method in an evaluation that contains more than one model, a 5-point Likert scale is presented to your work team for each model response. The final report will show the aggregated results for each model individually. Define the important points on the 5-point scale in your instructions so that your evaluators know how to rate the responses according to your expectations.

**Automatic evaluations**

Automatic evaluations can leverage built-in datasets and algorithms, or you can bring your own dataset of prompts that are specific to your use case. The built-in datasets vary for each task and are listed in the following sections. For a summary of tasks and their associated metrics and datasets, see the table in the following **Foundation model summary evaluation** section.

**Foundation model evaluation summary**

The following table summarizes all of the evaluation tasks, metrics, and built-in datasets for both human and automatic evaluations.

<table>
<thead>
<tr>
<th>Task</th>
<th>Human evaluations</th>
<th>Human metrics</th>
<th>Automatic evaluations</th>
<th>Automatic metrics</th>
<th>Automatic built-in datasets</th>
</tr>
</thead>
<tbody>
<tr>
<td>Open-ended generation</td>
<td>Fluency, Coherence, Toxicity, Accuracy, Consistency, Relevance, User-defined</td>
<td>Preference rate, Preference strength, Preference rank, Approval rate,</td>
<td>Factual knowledge</td>
<td></td>
<td>TREX</td>
</tr>
<tr>
<td>Task</td>
<td>Human evaluations</td>
<td>Human metrics</td>
<td>Automatic evaluations</td>
<td>Automatic metrics</td>
<td>Automatic built-in datasets</td>
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<td>Approval strength</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Semantic robustness</td>
<td></td>
<td></td>
<td></td>
<td>TREX</td>
<td></td>
</tr>
<tr>
<td>BOLD</td>
<td></td>
<td></td>
<td></td>
<td>WikiText</td>
<td></td>
</tr>
<tr>
<td>Prompt stereotyping</td>
<td></td>
<td></td>
<td></td>
<td>CrowS-Pairs</td>
<td></td>
</tr>
<tr>
<td>Toxicity</td>
<td></td>
<td></td>
<td></td>
<td>RealToxicityPrompts</td>
<td></td>
</tr>
<tr>
<td>BOLD</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Text summarization</td>
<td>Accuracy</td>
<td>ROUGE-N</td>
<td>Government Report Dataset</td>
<td></td>
<td>BERTScore Gigaword</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Government Report Dataset</td>
<td></td>
<td>Gigaword</td>
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### Updating inference parameters to change your model's output

Inference parameters are a way to influence a model's output without having to retrain or fine-tune a model.

In automatic model evaluation job, you can change the model's Temperature, Top P, and the Max new tokens.

**Temperature**

Changes the amount of randomness in the model's responses. Lower the default temperature to decrease the amount of randomness, and increase to have more.

**Top P**

During inference, the model is generating text and choosing from a list of words to place the next word. Updating Top P changes the number of words in that list based on a percentage. Decreasing Top P results in more deterministic samples, while a higher value will allow for more variability and creativity in the generated text.
Max new tokens

Changes the length of response the model can provide.

You can update the inference parameters in Studio after adding the model to your model evaluation job.

Use a human evaluation

Important

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see Migrating from Amazon SageMaker Studio Classic. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

To create a model evaluation job that uses human workers you must set up your environment to have the correct permissions. Then, you can use the model evaluation job wizard in Studio to select the models you want to use, and then define the parameters and the workforce you want to use in the model evaluation job.

When the job is complete you can, view a report to understand how your workforce evaluated the models you selected. The results are also saved in Amazon S3 as a jsonlines output file.

In model evaluation job that uses human workers, you have the ability to bring inference data from models hosted outside of SageMaker and models hosted outside of AWS. To learn more, see Using your own inference data for model evaluation jobs that use human workers.

Set up your environment

Prerequisites

To run a model evaluation in the Amazon SageMaker Studio UI, your AWS Identity and Access Management (IAM) role and any input datasets must have the correct permissions. If you do not have a SageMaker Domain or IAM role, follow the steps from the top of Onboard from the console through all of Step 1. General Settings.
Setting up your permissions

The following section shows you how to create an Amazon S3 bucket and how to specify the correct Cross-origin resource sharing (CORS) permissions.

To create an Amazon S3 bucket and specify the CORS permissions

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the navigation pane, enter S3 into the search bar at the top of the page.
3. Choose S3 under Services.
4. Choose Buckets from the navigation pane.
5. In the General purpose buckets section, under Name, choose the name of the S3 bucket that you want to use to store your model input and output in the console. If you do not have an S3 bucket, do the following.
   1. Select Create bucket to open a new Create bucket page.
   2. In the General configuration section, under AWS Region, select the AWS region where your foundation model is located.
   3. Name your S3 bucket in the input box under Bucket name.
   4. Accept all of the default choices.
   5. Select Create bucket.
   6. In the General purpose buckets section, under Name, select the name of the S3 bucket that you created.
6. Choose the Permissions tab.
7. Scroll to the Cross-origin resource sharing (CORS) section at the bottom of the window. Choose Edit.
8. The following is the minimum required CORS policy that you must add to your Amazon S3 bucket. Copy and paste the following into the input box.

```json
[
  {
    "AllowedHeaders": ["*"],
    "AllowedMethods": [
      "GET",
      "HEAD",
      "PUT"
    ],
    "ExposeHeaders": []
  }
]
```
To add permissions to your IAM policy

You may want to consider the level of permissions to attach to your IAM role.

- You can create a custom IAM policy that allows the minimum required permissions tailored to this service.
- You can attach the existing AmazonSageMakerFullAccess and AmazonS3FullAccess policies to your existing IAM role, which is more permissive. For more information about the AmazonSageMakerFullAccess policy, see AmazonSageMakerFullAccess.

If you wish to attach the existing policies to your IAM role, you may skip the instructions set here and continue following the instructions under To add permissions to your IAM role.

The following instructions creates a custom IAM policy that is tailored to this service with minimum permissions.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the search bar at the top of the page, enter IAM.
3. Under Services, select Identity and Access Management (IAM).
4. Choose Policies from the navigation pane.
5. Choose Create policy. When the Policy editor opens, choose JSON.
6. Ensure that the following permissions appear in the Policy editor. You can also copy and paste the following into the Policy editor.

```json
{
    "Version": "2012-10-17",
    "AllowedOrigins": [
        "*
    ],
    "ExposeHeaders": [
        "Access-Control-Allow-Origin"
    ],
    "MaxAgeSeconds": 3000
}
```

9. Choose Save changes.
"Statement":
  [{
    "Effect": "Allow",
    "Action": ["
      "s3:GetObject",
      "s3:PutObject",
      "s3:ListBucket"
    ],
    "Resource": [
      "arn:aws:s3:::{input_bucket}/*",
      "arn:aws:s3:::{input_bucket}",
      "arn:aws:s3:::{output_bucket}/*",
      "arn:aws:s3:::{output_bucket}",
      "arn:aws:s3:::jumpstart-cache-prod-{region}/*",
      "arn:aws:s3:::jumpstart-cache-prod-{region}"
    ]
  },
  {
    "Effect": "Allow",
    "Action": [
      "sagemaker:CreateEndpoint",
      "sagemaker:DeleteEndpoint",
      "sagemaker:CreateEndpointConfig",
      "sagemaker:DeleteEndpointConfig"
    ],
    "Resource": [
      "arn:aws:sagemaker:{region}:{account-id}:endpoint/sm-margaret-*",
      "arn:aws:sagemaker:{region}:{account-id}:endpoint-config/sm-
margaret-**"
    ],
    "Condition": {
      "ForAnyValue:StringEquals": {
        "aws:TagKeys": "sagemaker-sdk:jumpstart-model-id"
      }
    }
  },
  {
    "Effect": "Allow",
    "Action": [
      "sagemaker:DescribeProcessingJob",
      "sagemaker:DescribeEndpoint",
      "sagemaker:InvokeEndpoint"
    ],
    "Resource": "*"
  }]
```
{
  "Effect": "Allow",
  "Action": [
    "sagemaker:DescribeInferenceComponent",
    "sagemaker:AddTags",
    "sagemaker:CreateModel",
    "sagemaker:DeleteModel"
  ],
  "Resource": "arn:aws:sagemaker:{region}:{account-id}:model/*",
  "Condition": {
    "ForAnyValue:StringEquals": {
      "aws:TagKeys": "sagemaker-sdk:jumpstart-model-id"
    }
  }
},
{
  "Effect": "Allow",
  "Action": [
    "sagemaker:DescribeFlowDefinition",
    "sagemaker:StartHumanLoop",
    "sagemaker:DescribeHumanLoop"
  ],
  "Resource": "*"
},
{
  "Effect": "Allow",
  "Action": [
    "logs:CreateLogStream",
    "logs:PutLogEvents",
    "logs:CreateLogGroup",
    "logs:DescribeLogStreams"
  ],
  "Resource": "arn:aws:logs:{region}:{account-id}:log-group:/aws/sagemaker/ProcessingJobs:*"
},
{
  "Effect": "Allow",
  "Action": [
    "cloudwatch:PutMetricData"
  ],
  "Resource": "*"
}
```
7. Choose Next.

8. Enter a policy name in the Policy details section, under Policy name. You can also enter an optional description. You will search for this policy name when you assign it to a role.

9. Choose Create policy.
To add permissions to your IAM role

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the search bar at the top of the page, enter IAM.
3. Under Services, select Identity and Access Management (IAM).
4. Choose Roles in the navigation pane.
5. If you are creating a new role:
   a. Choose Create role.
   b. On the Select trusted entity step, under Trusted entity type choose Custom trust policy.
   c. In the Custom trust policy editor, next to Add principal choose Add.
   d. On the Add principal pop-up box, under Principal type select AWS services from the dropdown list of options.
   e. Under ARN replace {ServiceName} with sagemaker.
   f. Choose Add principal.
   g. Choose Next.
   h. (Optional) Under Permissions policies select the policies you would like to add to your role.
   i. (Optional) Under Set permissions boundary - optional choose your permission boundary setting.
   j. Choose Next.
   k. On the Name, review, and create step, under Role details fill in your Role name and Description.
   l. (Optional) Under Add tags - optional, you can add tags by choosing Add new tag and enter a Key and Value - optional pair.
   m. Review your settings.
   n. Choose Create role.
6. If you are adding the policy to an existing role:
   a. Select the name of the role under Role name. The main window changes to show information about your role.
   b. In the Permissions policies section, choose the down arrow next to Add permissions.
   c. From the options that appear, choose Attach policies.
d. From the list of policies that appear, search for and select the policy that you created under **To add permissions to your IAM policy** and select the check box next to your policy's name. If you did not create a custom IAM policy, search for and select the check boxes next to the AWS provided **AmazonSageMakerFullAccess** and **AmazonS3FullAccess** policies. You may want to consider the level of permissions to attach to your IAM role. The instructions for the custom IAM policy is less permissive, while the latter is more permissive. For more information about the AmazonSageMakerFullAccess policy, see [AmazonSageMakerFullAccess](https://console.aws.amazon.com/sagemaker/).

e. Choose **Add permissions**. A banner at the top of the page should state **Policy was successfully attached to role**. when completed.

### To add trust policy to your IAM role

The following trust policy makes it so administrators can allow SageMaker to assume the role. You need to add the policy to your IAM role. Use the following steps to do so.

2. In the search bar at the top of the page, enter **IAM**.
3. Under **Services**, select **Identity and Access Management (IAM)**.
4. Choose **Roles** in the navigation pane.
5. Select the name of the role under **Role name**. The main window changes to show information about your role.
6. Choose the **Trust relationship** tab.
7. Choose **Edit trust policy**.
8. Ensure that the following policy appears under **Edit trust policy**. You can also copy and paste the following into the editor.

```json
{
   "Version": "2012-10-17",
   "Statement": [
   {
      "Sid": "",
      "Effect": "Allow",
      "Principal": {
         "Service": [
            "sagemaker.amazonaws.com"
         ]
   }
   ]
}
```
9. Choose **Update policy**. A banner at the top of the page should state **Trust policy updated**. when completed.

### Creating a model evaluation job that uses human workers

You can create a human evaluation job using a text-based model that is available in SageMaker JumpStart or you can use a SageMaker JumpStart model that you've previously deployed to an endpoint.

**To launch SageMaker JumpStart**

2. In the search bar at the top of the page, enter **SageMaker**.
3. Under **Services**, select **Amazon SageMaker**.
4. Choose **Studio** from the navigation pane.
5. Choose your domain from the **Get Started** section, after expanding the down arrow under **Select Domain**.
6. Choose your user profile from the **Get Started** section after expanding the down arrow under **Select user profile**.
7. Choose **Open Studio** to open the landing page for Studio.
8. Choose **Jobs** from the navigation pane.

**To set up an evaluation job**

1. On the Model evaluation home page, choose **Evaluate a model**
2. Specify job details.
   
   a. Enter the **Evaluation name** of your model evaluation. This name helps you identify your model evaluation job after it is submitted.
   b. Enter a **Description** to add more context to the name.
   c. Choose **Next**.
3. Set up evaluation

a. Under **Choose an evaluation type**, select the radio button next to **Human**.

b. Under **Choose the model(s) you want to evaluate**, choose **Add model to evaluation**. You can evaluate up to two models for each evaluation.

1. To use a pre-trained SageMaker JumpStart model, choose **Pre-trained SageMaker JumpStart foundation model**. If you want to use a SageMaker JumpStart model that you have previously deployed to an endpoint, choose **Endpoints with JumpStart foundation models**.

2. If the model requires a legal agreement, select the check box to confirm that you agree.

3. If you want to add another model, repeat the previous step.

c. To change how the model behave during inference choose, **Set parameters**.

Set parameters contains a list of inference parameters that affect the degree of randomness in your model's output, the length of your model's output, and what words the model will choose next.

d. Next, select an **Task type**. You can select any of the following:

   - **Text Summarization**
   - **Question Answering (Q&A)**
   - **Text classification**
   - **Open-ended Generation**
   - **Custom**

e. In the **Evaluation metrics** section, choose an **Evaluation dimension** and enter additional context about the dimension in the text box under **Description**. You can choose from the following dimensions:

   - **Fluency** – Measures the linguistic quality of a generated text.
   - **Coherence** – Measures the organization and structure of a generated text.
   - **Toxicity** – Measures the harmfulness of a generated text.
   - **Accuracy** – Indicates the accuracy of a generated text.
   - A custom evaluation dimension that you can define the name and description of for your work team.

To add a custom evaluation dimension, do the following:
• Choose **Add an evaluation dimension**.

• In the text box containing **Provide evaluation dimension**, input the name of your custom dimension.

• In the text box containing **Provide description for this evaluation dimension**, input a description so that your work team understands how to evaluate your custom dimension.

Under each of these metrics are reporting metrics that you can choose from the **Choose a metric type** down arrow. If you have two models to evaluate, you can choose either comparative or individual reporting metrics. If you have one model to evaluate, you can choose only individual reporting metrics. You can choose the following reporting metrics types for each of the above metrics.

• (Comparative) **Likert scale - comparison** – A human evaluator will indicate their preference between two responses on a 5-point Likert scale according to your instructions. The results in the final report will be shown as a histogram of preference strength ratings from the evaluators over your whole dataset. Define the important points of the 5-point scale in your instructions so that your evaluators know how to rate the responses according to your expectations.

• (Comparative) **Choice buttons** – Allows a human evaluator to indicate their one preferred response over another response. Evaluators indicate their preference between two responses according to your instructions using radio buttons. The results in the final report will be shown as a percentage of responses that workers preferred for each model. Explain your evaluation method clearly in your instructions.

• (Comparative) **Ordinal Rank** – Allows a human evaluator to rank their preferred responses to a prompt in order, starting at 1, according to your instructions. The results in the final report will be shown as a histogram of the rankings from the evaluators over the whole dataset. Define the what a rank of 1 means in your instructions.

• (Individual) **Thumbs up/down** – Allows a human evaluator to rate each response from a model as acceptable or unacceptable according to your instructions. The results in the final report will be shown as a percentage of the total number of ratings by evaluators that received a thumbs up rating for each model. You may use this rating method for an evaluation one or more models. If you use this in an evaluation that contains two models, a thumbs up or down will be presented to your work team for each model response and the final report will show the aggregated results for each
model individually. Define what is acceptable as a thumbs up or thumbs down rating in your instructions.

- **(Individual) Likert scale - individual** – Allows a human evaluator to indicate how strongly they approve of the model response based on your instructions on a 5-point Likert scale. The results in the final report will be shown as a histogram of the 5-point ratings from the evaluators over your whole dataset. You may use this scale for an evaluation containing one or more models. If you select this rating method in an evaluation that contains more than one model, a 5-point Likert scale will be presented to your work team for each model response and the final report will show the aggregated results for each model individually. Define the important points on the 5-point scale in your instructions so that your evaluators know how to rate the responses according to your expectations.

f. **Choose a Prompt dataset.** This dataset is required and will be used by your human work team to evaluate responses from your model. Provide the S3 URI to an Amazon S3 bucket that contains your prompt dataset in the text box under **S3 URI for your input dataset file.** Your dataset must be in jsonlines format and contain the following keys to identify which parts of your dataset the UI will use to evaluate your model:

- **prompt** – The request that you want your model to generate a response to.
- **(Optional) category** – The category labels for your prompt. The category key is used to categorize your prompts so you can filter your evaluation results later by category for a deeper understanding of the evaluation results. It does not participate in the evaluation itself, and workers do not see it on the evaluation UI.
- **(Optional) referenceResponse** – The reference answer for your human evaluators. The reference answer is not rated by your workers, but can be used to understand what responses are acceptable or unacceptable, based on your instructions.
- **(Optional) responses** – Used to specify inferences from a model outside of SageMaker or outside of AWS.

This object **requires** two additional key value pairs "modelIdentifier" which is a string that identifies the model, and "text" which is the model's inference.

If you specify a "responses" key in any input of the custom prompt dataset it must be specified in all inputs.
• The following JSON code example shows the accepted key-value pairs in a custom prompt dataset. The **Bring your own inference** check box must be checked if responses key is provided.

```json
{
    "prompt": {
        "text": "String",
    },
    "category": "String", // optional
    "referenceResponse": {
        "text": "String" // optional
    },
    "responses": { // optional
        // All inference must come from a single model
        [{
            "modelIdentifier": "String" // optional - A name you want to use to ID the model
            "text": "String" // optional - The inference reference answer
        }]
    }
}
```

**g.** Input an S3 bucket location where you want to save the output evaluation results in the text box under **Choose an S3 location to save your evaluation results.** The output file written to this S3 location will be in JSON format, ending in the extension, `.json`.

**h.**

> **Note**

If you want to include bring your own inference data in the model evaluation job, you can only use a single model.

(Optional) Choose the check box under **Bring your own inference** to indicate that your prompt dataset contains the `responses` key. If you specify the `responses` key as part of any prompts it must be present in all of them.

**i.** Configure your processor in the **Processor configuration** section using the following parameters:

- **Use Instance count** to specify the number of compute instances to use to run your model. If you use more than 1 instance, your model will run in parallel instances.
• **Use Instance type** to choose the kind of compute instance you want to use to run your model. AWS has general compute instances and instances that are optimized for computing and memory. For more information about instance types, see [Available Studio Classic Instance Types](#).

• If you want SageMaker to use your own AWS Key Management Service (AWS KMS) encryption key instead of the default AWS managed service key, toggle to select **On** under **Volume KMS key**, and input the AWS KMS key. SageMaker will use your AWS KMS key to encrypt data on the storage volume. For more information about keys, see [AWS Key Management Service](#).

• If you want SageMaker to use your own AWS Key Management Service (AWS KMS) encryption key instead of the default AWS managed service key, toggle to select **On** under **Output KMS key** and input the AWS KMS key. SageMaker will use your AWS KMS key to encrypt the processing job output.

• Use an IAM role to specify the access and permissions for the default processor. Input the IAM role that you set up in the section **Set up your IAM role** in this **Run a human evaluation** section.

  j. After you specify your model and criteria, select **Next**.

Your work team consists of the people that are evaluating your model. After your work team is created, it persists indefinitely and you cannot change its attributes. The following shows how to get started with your work team.

**Set up your work team**

1. Choose an existing team or **Create a new team** in the **Select team** input text box.

2. Specify a name of your organization in **Organization name**. This field only appears when you create the first work team in the account.

3. Specify a **contact email**. Your workers will use this email to communicate with you about the evaluation task that you will provide to them. This field only appears when you create the first work team in the account.

4. Specify a **Team name**. You cannot change this name later.

5. Specify a list of **Email addresses** for each of your human workers that will evaluate your large language model (LLM). When you specify the email addresses for your team, they are notified of a new job only when they are newly added to a work team. If you use the same team for a subsequent job, you must notify them manually.
6. Then, specify the **Number of workers per prompt**

**Provide instructions for your work team**

1. Provide detailed instructions to your human workforce so that they can evaluate your model to your metrics and standards. A template in the main window shows sample instructions that you can provide. For more information about how to give instructions, see [Creating good worker instructions](#).

2. To minimize bias in your human evaluation, select the check box next to **Randomize response positions**.

3. Select **Next**.

You can review the summary of the selections that you have made for your human job. If you must change your job, choose **Previous** to go back to an earlier selection.

**Submit your evaluation job request and view job progress**

1. To submit your evaluation job request, choose **Create resource**.

2. To see the status of all of your jobs, choose **Jobs** in the navigation pane. Then, choose **Model evaluation**. The evaluation status displays as **Completed**, **Failed**, or **In progress**.

The following also displays:

- Sample notebooks to run a model evaluation in SageMaker and Amazon Bedrock.
- Links to additional information including documentation, videos, news, and blogs about the model evaluation process.
- The URL to your **Private worker portal** is also available.

3. Select your model evaluation under **Name** to view a summary of your evaluation.

   - The summary gives information about the status of the job, what kind of evaluation task you ran on which model, and when it ran. Following the summary, the human evaluation scores are sorted and summarized by metric.

**View the report card of your model evaluation job that uses human workers**

1. To see the report for your jobs, choose **Jobs** in the navigation pane.
2. Then, choose **Model evaluation**. One the **Model evaluations** home page, use the table to find your model evaluation job. Once the job status has changed to **Completed** you can view your report card.

3. Choose the name of the model evaluation job to its report card.

**View your human analysis results Amazon S3**

The output from your human model evaluation is stored in the S3 location that you specified to save the output evaluation results while creating the model evaluation job.

You can view a model evaluation's job while it is still in progress. The output data appears in the S3 bucket when the evaluation job is complete. It may take a few minutes for output data to appear.

The contents of your output json file depends on your specific job description, metrics and model. The following output sample was generated for a single prompt in inputRecord, where a human rated the model response (modelResponses) as unacceptable ("result":false):

```json
{
   "output": [{
       "humanAnswers": [{
           "acceptanceTime": "2023-11-09T19:17:43.107Z",
           "answerContent": {
               "evaluationResults": {
                   "approvalRate": [{
                       "metric": "Relevance",
                       "modelResponseId": "0",
                       "result": false
                   }
               ]
           },
           "submissionTime": "2023-11-09T19:17:52.101Z",
           "timeSpentInSeconds": 8.994,
           "workerId": "444455556666",
           "workerMetadata": {
               "identityData": {
                   "identityProviderType": "Cognito",
                   "sub": "12345678-1234-1234-1234-1234"
               }
           }
       },
   },
   "submissionTime": "2023-11-09T19:17:52.101Z",
   "timeSpentInSeconds": 8.994,
   "workerId": "444455556666",
   "workerMetadata": {
       "identityData": {
           "identityProviderType": "Cognito",
           "sub": "12345678-1234-1234-1234-1234"
       }
   }
}
```
"humanLoopName": "1234567890abcdefghijklmnopqrstuvwxyz",
"inputRecord": {
  "prompt": "What does vitamin C serum do for skin?",
  "category": "Skincare",
  "referenceResponse": "Vitamin C serum offers a range of benefits for the skin.
Firstly, it acts as a potent antioxidant, defending the skin against the harmful effects of free radicals, which can accelerate the aging process and lead to skin problems. Moreover, vitamin C brightens the skin by reducing the appearance of dark spots, age spots, and hyperpigmentation. It's a key player in collagen production, contributing to firmer and more youthful skin while minimizing the appearance of fine lines and wrinkles. Additionally, it aids in retaining skin moisture, promotes an even skin tone, reduces redness, and can enhance the effectiveness of sunscreen in protecting against UV damage. Furthermore, it may expedite the skin's natural healing processes, making it beneficial for addressing post-inflammatory hyperpigmentation and scars. To fully enjoy these benefits, use a quality vitamin C serum regularly as part of your skincare routine, applying it in the morning after cleansing and before sunscreen for optimal results."
},
"modelResponses": [
  {
    "text": "Vitamin C serums are widely known for their"
  }
],
"flowDefinitionArn": "arn:aws:sagemaker:us-west-2:111122223333:flow-definition/flow-definition-name"
... 
],

The previous output sample uses the following parameters:
- **flowDefinitionArn** – The Amazon Resource Number (ARN) of the human review work flow (flow definition) that's used to create the human loop.
- **humanAnswers** – A list of json objects that contain worker responses in `answerContent`.
  
  ```json
  metric:"Relevance" – A value that is based on the **Metric type** that you selected when the model evaluation job was created.
  ```
- **humanLoopName** – The name of the human loop.
- **inputRecord** – A json object that contains an entry prompt from the input dataset.
- **modelResponses** – The individual responses from the models.
- **input** – The input content sent to SageMaker in the request to StartHumanLoop.

When your analysis is complete, you can see how your model performed against the dataset that you provided using the following steps:

1. From the Studio navigation pane, select **Jobs**, and then select **Model Evaluation**.
2. In the **Model Evaluations** page, successfully submitted jobs appear in a list. The list includes job name, status, model name, evaluation type, and the date it was created.
3. If your model evaluation completed successfully, you can click on the job name to see a summary of the evaluation results.
4. To view your human analysis report, select the name of the job that you want to examine.

An analysis report section that identifies your job by the name, status, model name, evaluation type, and the date it was created appears at the top of the main window after you select the name of a job that you want to examine.

The next analysis report section contains the type of task that your model performed, and the evaluation results for that task.

The following analysis report section contains the evaluation results for each model that you evaluated.

The next analysis report section shows the evaluation job configuration. It includes the resources that are used, the model that is evaluated, the location of the evaluation results, and your evaluation dimensions.

The last section contains a copy of the instructions that you provided to your workforce.
Using your own inference data for model evaluation jobs that use human workers

When you create a model evaluation job that uses human workers you have the option to bring your own inference data, and have your human workers compare that inference data to data produced by one other SageMaker JumpStart model or to SageMaker JumpStart model that you have previously deployed to an endpoint.

This topic describes the format required for the inference data, and a simplified procedure for how to add that data to your model evaluation job.

Choose a Prompt dataset. This dataset is required and will be used by your human work team to evaluate responses from your model. Provide the S3 URI to an Amazon S3 bucket that contains your prompt dataset in the text box under Choose an S3 location to save your evaluation results. Your dataset must be in .jsonl format. Each record must be a valid JSON object, and contain the following required keys:

- prompt – A JSON object that contains the text to be passed into the model.
- (Optional) category – The category labels for your prompt. The category key is used to categorize your prompts so you can filter your evaluation results later by category for a deeper understanding of the evaluation results. It does not participate in the evaluation itself, and workers do not see it on the evaluation UI.
- (Optional) referenceResponse – a JSON object that contains the reference answer for your human evaluators. The reference answer is not rated by your workers, but can be used to understand what responses are acceptable or unacceptable, based on your instructions.
- (Optional) responses – Used to specify inferences from a model outside of SageMaker or outside of AWS.

This object requires to additional key value pairs "modelIdentifier which is a string that identified the model, and "text" which is the model's inference.

If you specify a "responses" key in any input of the of custom prompt dataset it must specified in all inputs.

The following json code example shows the accepted key-value pairs in a custom prompt dataset.

```json
{
    "prompt": {
        "text": String,
    }
}
To get started launch Studio, and under choose Model evaluation under Jobs in the primary navigation.

To add your own inference data to a human model evaluation job.

1. **In Step 1: Specify job details** add the name of your model evaluation job, and an optional description.

2. **In Step 2: Set up evaluation** choose Human.

3. Next, under Choose the model(s) you want to evaluate you can choose the model that you want to use. You can use either a SageMaker JumpStart model that has already deployed or you can choose a Pre-trained Jumpstart foundation model.

4. Then, choose a Task type.

5. Next, you can add Evaluation metrics.

6. Next, under Prompt dataset choose the check box under Bring your own inference to indicate that your prompts have response keys in it.

7. Then continue setting up your model evaluation job.

To learn more about how the responses from your model evaluation job that uses human workers are saved, see [View your human analysis results Amazon S3](#).
Use an automatic evaluation

⚠️ Important

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see [Migrating from Amazon SageMaker Studio Classic](#). For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).

You can create an automatic model evaluation in Studio or by using the `fmeval` library inside your own code. Studio uses a wizard to create the model evaluation job. The `fmeval` library provides tools to customize your workflow further. The following sections show you how to use both types of automatic evaluations.

Both types of automatic model evaluation jobs support the use of publicly available SageMaker JumpStart models, and SageMaker JumpStart models that you previously deployed to an endpoint. If you use a SageMaker JumpStart that has *not* been previously deployed, SageMaker will handle creating the necessary resource, and shutting them down once the model evaluation job has finished.

To use text based LLMs from other AWS service or a model hosted outside of AWS, you must use the `fmeval` library.

Creating an automatic model evaluation job in Studio

The wizard available in Studio guides you through choosing a model to evaluate, selecting a task type, choosing metrics and datasets, and configuring any required resources. The following topics show you how to format an optional custom input dataset, set up your environment, and create the model evaluation job in Studio.

Format your input dataset

If you use a built-in dataset to evaluate your model in Studio, the dataset is formatted correctly. To use your own custom prompt dataset, it must be a `jsonlines` file, where each line is a valid JSON object. Each JSON object *must* contain a single prompt.
To help ensure that the SageMaker JumpStart model you select performs well, SageMaker Clarify automatically formats all prompt datasets to be in format that work best for the **Model Evaluation dimensions** you select. For built-in prompt datasets, SageMaker Clarify will also augment your prompt with additional instructional text. To see how SageMaker Clarify will modify the prompts, choose [prompt template](#) under an **Evaluation dimensions** you have added to the model evaluation job. To see an example of how you can modify a prompt template, see [Prompt template example](#).

The toggle allows you to turn off or to turn on the automatic prompt templating support that SageMaker Clarify provides for built-in datasets. Turning off the automatic prompt templating allows, you to specify your own custom prompt templates that will be applied to all prompts in your dataset.

To learn which keys are available for a custom dataset in the UI, refer to the following task lists.

- **model_input** – Required to indicate the input for the following tasks.
  - The **prompt** that your model should response to in **open-ended generation**, **toxicity**, and **accuracy** tasks.
  - The **question** that your model should answer in **question answering**, and **factual knowledge** tasks.
  - The **text** that your model should summarize in **text summarization** tasks.
  - The **text** that your model should classify in **classification** tasks.
  - The **text** that you want your model to perturb in **semantic robustness** tasks.
- **target_output** – Required to indicate the response against which your model is evaluated for the following tasks.
  - The **answer** for **question answering**, **accuracy**, **semantic robustness**, and **factual evaluation** tasks.
  - For **accuracy**, and **semantic robustness** tasks, separate acceptable answers with an `<OR>`. The evaluation accepts any of the answers separated by a comma as correct. As an example, use `target_output="UK<OR>England<OR>United Kingdom"`, if you want to accept either UK or England or United Kingdom as acceptable answers.
- **category** – Generates evaluation scores reported for each category.
- **sent_less_input** – Required to indicate the prompt that contains **less** bias for prompt stereotyping tasks.
• **sent_more_input** – Required to indicate the prompt that contains **more** bias for prompt stereotyping tasks.

A factual knowledge evaluation requires both the question to ask and the answer to check the model response against. Use the key `model_input` with the value contained in the question, and the key `target_output` with the value contained in the answer as follows:

```json
{"model_input": "Bobigny is the capital of", "target_output": "Seine-Saint-Denis", "category": "Capitals"}
```

The previous example is a single valid JSON object that makes up one record in a jsonlines input file. Each JSON object is sent to your model as a request. To make multiple requests, include multiple lines. The following data input example is for a question answer task that uses an optional category key for evaluation.

```json
{"target_output": "Cantal", "category": "Capitals", "model_input": "Aurillac is the capital of"}
{"target_output": "Bamiyan Province", "category": "Capitals", "model_input": "Bamiyan city is the capital of"}
{"target_output": "Abkhazia", "category": "Capitals", "model_input": "Sokhumi is the capital of"}
```

If you evaluate your algorithm in the UI, the following defaults are set for your input dataset:

- The number of records that the evaluation uses is fixed. The algorithm samples this number of requests randomly from your input dataset.
  - **To change this number:** Use the `fmeval` library as described in **Customize your work flow using the fmeval library**, and set the parameter `num_records` to your desired number of samples, or `-1` to specify the entire dataset. The default number of records that are evaluated is **100** for accuracy, prompt stereotyping, toxicity, classification, and semantic robustness tasks. The default number of records for a factual knowledge task is **300**.

- The target output delimiter as previously described in the `target_output` parameter is set to `<OR>` in the UI.
  - **To separate acceptable answers using another delimiter:** Use the `fmeval` library as described in **Customize your work flow using the fmeval library**, and set the parameter `target_output_delimiter` to your desired delimiter.
• You must use a text-based SageMaker JumpStart language model that is available for model evaluation. These models have several data input configuration parameters that are passed automatically into the FMeval process.

• **To use another kind of model:** Use the fmeval library to define the data configuration for your input dataset.

**Set up your environment**

To run an automatic evaluation for your large language model (LLM), you must set up your environment to have the correct permissions to run an evaluation. Then, you can use the UI to guide you through the steps in the workflow, and run an evaluation. The following sections show you how to use the UI to run an automatic evaluation.

**Prerequisites**

• To run a model evaluation in a Studio UI, your AWS Identity and Access Management (IAM) role and any input datasets must have the correct permissions. If you do not have a SageMaker Domain or IAM role, follow the steps from the top of [Onboard from the console](#) through all of **Step 1. General Settings**.

**To set permissions for your S3 bucket**

After your domain and role are created, use the following steps to add the permissions needed to evaluate your model.

2. In the navigation pane, enter **S3** into the search bar at the top of the page.
3. Choose **S3** under **Services**.
4. Choose **Buckets** from the navigation pane.
5. In the **General purpose buckets** section, under **Name**, choose the name of the Amazon S3 bucket that you want to use to store your custom prompt dataset, and where you want the results of your model evaluation job saved. Your Amazon S3 bucket must be in the same AWS Region as your Studio instance. If you don't have an Amazon S3 bucket, do the following.

   1. Select **Create bucket** to open a new **Create bucket** page.
   2. In the **General configuration** section, under **AWS Region**, select the AWS region where your foundation model is located.
3. Name your S3 bucket in the input box under **Bucket name**.
4. Accept all of the default choices.
5. Select **Create bucket**.
6. In the **General purpose buckets** section, under **Name**, select the name of the S3 bucket that you created.
6. Choose the **Permissions** tab.
7. Scroll to the **Cross-origin resource sharing (CORS)** section at the bottom of the window. Choose **Edit**.
8. To add the CORS permissions to your bucket copy the following code into the input box.

```json
[
  {
    "AllowedHeaders": [
      "*
    ],
    "AllowedMethods": [
      "GET",
      "PUT",
      "POST",
      "DELETE"
    ],
    "AllowedOrigins": [
      "*
    ],
    "ExposeHeaders": [
      "Access-Control-Allow-Origin"
    ]
  }
]
```
9. Choose **Save changes**.

**To add permissions to your IAM policy**

1. In the search bar at the top of the page, enter **IAM**.
2. Under **Services**, select **Identity and Access Management (IAM)**.
3. Choose **Policies** from the navigation pane.
4. Choose **Create policy**. When the **Policy editor** opens, choose **JSON**.
5. Choose **Next**.

6. Ensure that the following permissions appear in the **Policy editor**. You can also copy and paste the following into the **Policy editor**.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "cloudwatch:PutMetricData",
            "logs:CreateLogStream",
            "logs:PutLogEvents",
            "logs:CreateLogGroup",
            "logs:DescribeLogStreams",
            "s3:GetObject",
            "s3:PutObject",
            "s3:ListBucket",
            "ecr:GetAuthorizationToken",
            "ecr:BatchCheckLayerAvailability",
            "ecr:GetDownloadUrlForLayer",
            "ecr:BatchGetImage"
         ],
         "Resource": "*"
      },
      {
         "Effect": "Allow",
         "Action": [
            "sagemaker:Search",
            "sagemaker:CreateProcessingJob",
            "sagemaker:DescribeProcessingJob"
         ],
         "Resource": "*"
      }
   ]
}
```

7. Choose **Next**.

8. Enter a policy name in the **Policy details** section, under **Policy name**. You can also enter an optional description. You will search for this policy name when you assign it to a role.

9. Choose **Create policy**.
To add permissions to your IAM role

1. Choose **Roles** in the navigation pane. Input the name of the role that you want to use.
2. Select the name of the role under **Role name**. The main window changes to show information about your role.
3. In the **Permissions** policies section, choose the down arrow next to **Add permissions**.
4. From the options that appear, choose **Attach policies**.
5. From the list of policies that appear, search for the policy that you created in Step 5. Select the check box next to your policy's name.
6. Choose the down arrow next to **Actions**.
7. From the options that appear, select **Attach**.
8. Search for the name of the role that you created. Select the check box next to the name.
9. Choose **Add permissions**. A banner at the top of the page should state **Policy was successfully attached to role**.

Create an automatic model evaluation job in Studio

When you create an automatic model evaluation job, you can choose from available text-based SageMaker JumpStart models or you can use a text based SageMaker JumpStart model that you've previously deployed to an endpoint.

To create a automatic model evaluation job use the following procedure.

To launch an automatic model evaluation job in Studio.

2. In the search bar at the top of the page, enter **SageMaker**.
3. Under **Services**, select **Amazon SageMaker**.
4. Choose **Studio** from the navigation pane.
5. Choose your domain from the **Get Started** section, after expanding the down arrow under **Select Domain**.
6. Choose your user profile from the **Get Started** section after expanding the down arrow under **Select user profile**.
7. Choose Open Studio to open the landing page for Studio.

8. Choose Jobs from the primary navigation pane.

9. Then, choose Model evaluation.

To set up an evaluation job

1. Next, choose Evaluate a model.

2. In Step 1: Specify job details do the following:
   a. Enter the Name of your model evaluation. This name helps you identify your model evaluation job after it is submitted.
   b. Enter a Description to add more context to the name.
   c. Choose Next.

3. In Step 2: Set up evaluation do the following:
   b. Then, choose Add model to evaluation
   c. In the Add model modal you can choose to use either a Pre-trained Jumpstart foundation model or SageMaker endpoint. If you've already deployed SageMaker JumpStart model choose SageMaker endpoint otherwise choose Pre-trained Jumpstart foundation model.
   d. Then, choose Save.
   e. (Optional) After adding your model choose Prompt template to see the expected input format for prompts based on the model you selected.
   f. Then, under Task type choose a task type.

For more information about tasks types and the associated evaluation dimensions, see the Automatic evaluation in Foundation model evaluation overview.

4. In the Evaluation metrics section, choose an Evaluation dimension. The text box under Description contains additional context about the dimension.

   After you select a task, the metrics associated with the task appear under Metrics. In this section, do the following.
   h. Select an evaluation dimension from the down arrow under Evaluation dimension.
Choose an evaluation dataset. You can choose to use your own dataset or use a built-in dataset. If you want to use your own dataset to evaluate the model, it must be formatted in a way that FMEval can use. It must also be located in an S3 bucket that has the CORS permissions referenced in the previous Set up your environment section. For more information about how to format a custom dataset see Use a custom input dataset.

Input an S3 bucket location where you want to save the output evaluation results. This file is in jsonlines (.jsonl) format.

Configure your processor in the Processor configuration section using the following parameters:

- Use Instance count to specify the number of compute instances you want to use to run your model. If you use more than 1 instance, your model is run in parallel instances.
- Use Instance type to choose the kind of compute instance you want to use to run your model. For more information about instance types, see Available Studio Classic Instance Types.
- Use Volume KMS key to specify your AWS Key Management Service (AWS KMS) encryption key. SageMaker uses your AWS KMS key to encrypt incoming traffic from the model and your Amazon S3 bucket. For more information about keys, see AWS Key Management Service.
- Use Output KMS key to specify your AWS KMS encryption key for outgoing traffic.
- Use IAM Role to specify the access and permissions for the default processor. Enter the IAM role that you set up in Set up your environment.

After you specify your model and criteria, choose Next. The main window skips to Step 5 Review and Save.

Review and run your evaluation job

1. Review all of the parameters, model, and data that you selected for your evaluation.
2. Choose Create resource to run your evaluation.
3. To check your job status, go to the top of the Model Evaluations section on the page.

View analysis results from your automatic evaluation

This section lists three outputs for an automatic evaluation that is run in the UI.
1. The output.json file contains aggregate scores for your dataset. An example json output follows.

```json
{
    "evaluations": [
    {
        "evaluation_name": "factual_knowledge",
        "dataset_name": "trex",
        "prompt_template": "<s>[INST] <<SYS>>Answer the question at the end in as few words as possible. Do not repeat the question. Do not answer in complete sentences.</SYS> Question: $feature [/INST]",
        "dataset_scores": [
            {
                "name": "factual_knowledge",
                "value": 0.2966666666666667
            }
        ],
        "category_scores": [
            {
                "name": "Author",
                "scores": [
                    {
                        "name": "factual_knowledge",
                        "value": 0.4117647058823529
                    }
                ]
            },
            ...
            {
                "name": "Capitals",
                "scores": [
                    {
                        "name": "factual_knowledge",
                        "value": 0.2857142857142857
                    }
                ]
            }
        ]
    }
}
```
In the previous output example, the model scored on average of 0.2966666666666667. Average scores for each category are listed following the aggregate score.

2. One `evaluation_name_dataset_name`.jsonl file containing instance-wise results for each jsonlines request. If you had 300 requests in your jsonlines input data, this jsonlines output file contains 300 responses. The output file contains the request made to your model followed by the score for that evaluation. An example instance-wide output follows.

3. An Evaluation Report that contains the results of your foundation model evaluation. The content of the evaluation report depends on the kind of task you used to evaluate your model. Each report contains the following sections:
   a. The overall scores for each successful evaluation under the evaluation task. As an example of one evaluation with one dataset, if you evaluated your model for a classification task for Accuracy and Semantic Robustness, then a table summarizing the evaluation results for Accuracy and Accuracy Semantic Robustness appears at the top of your report. Other evaluations with other datasets may be structured differently.
   b. The configuration for your evaluation job including the model name, type, which evaluation methods were used and what datasets your model was evaluated against.
   c. A Detailed Evaluation Results section that summarizes the evaluation algorithm, provides information about and links to any built-in datasets, how scores are calculated, and tables showing some sample data with their associated scores.
   d. A Failed Evaluations section that contains a list of evaluations that did not complete. If no evaluations failed, this section of the report is omitted.

Use the fmeval library to run an automatic evaluation

Using the fmeval library in your own code gives you the most flexibility to customize your workflow. You can use the fmeval library to evaluate any LLM, and also to have more flexibility with your custom input datasets. The following steps show you how to set up your environment and how to run both a starting and a customized workflow using the fmeval library.

Get started using the fmeval library

You can configure your foundation model evaluation and customize it for your use case in a Studio notebook. Your configuration depends both on the kind of task that your foundation model is built to predict, and how you want to evaluate it. FM Eval supports open-ended generation, text summarization, question answering, and classification tasks. The steps in this section show you
how to set up a starting work flow. This starting work flow includes setting up your environment and running an evaluation algorithm using either a SageMaker JumpStart or an Amazon Bedrock foundation model with built-in datasets. If you must use a custom input dataset and work flow for a more specific use case, see Customize your workflow using the fmeval library.

Set up your environment

If you don't want to run a model evaluation in a Studio notebook, skip to step 11 in the following Get started using Studio section.

Prerequisites

- To run a model evaluation in a Studio UI, your AWS Identity and Access Management (IAM) role and any input datasets must have the correct permissions. If you do not have a SageMaker Domain or IAM role, follow the steps from the top of Onboard from the console through all of Step 1. General Settings.

To set permissions for your Amazon S3 bucket

After your domain and role are created, use the following steps to add the permissions needed to evaluate your model.

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/
2. In the navigation pane, enter S3 into the search bar at the top of the page.
3. Choose S3 under Services.
4. Choose Buckets from the navigation pane.
5. In the General purpose buckets section, under Name, choose the name of the S3 bucket that you want to use to store your model input and output in the console. If you do not have an S3 bucket, do the following:
   1. Select Create bucket to open a new Create bucket page.
   2. In the General configuration section, under AWS Region, select the AWS region where your foundation model is located.
   3. Name your S3 bucket in the input box under Bucket name.
   4. Accept all of the default choices.
   5. Select Create bucket.
6. In the **General purpose buckets** section, under **Name**, select the name of the S3 bucket that you created.

6. Choose the **Permissions** tab.

7. Scroll to the **Cross-origin resource sharing (CORS)** section at the bottom of the window. Choose **Edit**.

8. To add permissions to your bucket for foundation evaluations, ensure that the following code appears in the input box. You can also copy and paste the following into the input box.

```json
[
  {
    "AllowedHeaders": [
      "*"
    ],
    "AllowedMethods": [
      "GET",
      "PUT",
      "POST",
      "DELETE"
    ],
    "AllowedOrigins": [
      "*"
    ],
    "ExposeHeaders": [
      "Access-Control-Allow-Origin"
    ]
  }
]
```

9. Choose **Save changes**.

**To add permissions to your IAM policy**

1. In the search bar at the top of the page, enter **IAM**.

2. Under **Services**, select **Identity and Access Management (IAM)**.

3. Choose **Policies** from the navigation pane.

4. Input [AmazonSageMakerFullAccess](#) into the search bar. Select the radio button next to the policy that appears. The **Actions** button can now be selected.

5. Choose the down arrow next to **Actions**. Two options appear.
6. Choose **Attach**.

7. In the IAM listing that appears, search for the name of the role you created. Select the check box next to the name.

8. Choose **Attach policy**.

**Get started using Studio**

1. In the search bar at the top of the page, enter **SageMaker**.
2. Under **Services**, select **Amazon SageMaker**.
3. Choose **Studio** from the navigation pane.
4. Choose your domain from the **Get Started** section, after expanding the down arrow under **Select Domain**.
5. Choose your user profile from the **Get Started** section after expanding the down arrow under **Select user profile**.
6. Choose **Open Studio** to open the landing page for Studio.
7. Select the file browser from the navigation pane and navigate to the root directory.
8. Select **Create notebook**.
9. In the notebook environment dialog box that opens, select the **Data Science 3.0** image.
10. Choose **Select**.
11. Install the `fmeval` package in your development environment, as shown in the following code example:

   ```
   !pip install fmeval
   ```

   **Note**

   Install the `fmeval` library into an environment that uses Python 3.10. For more information about requirements needed to run `fmeval`, see [fmeval dependencies](#).

**Configure ModelRunner**

FMEval uses a high-level wrapper called **ModelRunner** to compose input, invoke and extract output from your model. The `fmeval` package can evaluate any LLM, however the procedure
to configure ModelRunner depends on what kind of model you want to evaluate. This section explains how to configure ModelRunner for a SageMaker JumpStart or Amazon Bedrock model. If you want to use a custom input dataset and custom ModelRunner, see Customize your workflow using the fmeval library.

**Use a SageMaker JumpStart model**

To use ModelRunner to evaluate a SageMaker JumpStart model, create or provide an endpoint, define the model and the built-in dataset, configure, and test ModelRunner.

**Define a SageMaker JumpStart model and configure a ModelRunner**

1. Provide an endpoint by doing either of the following:
   - Specify the **EndpointName** to an existing SageMaker JumpStart endpoint, the model_id, and model_version.
   - Specify the model_id and model_version for your model, and create a SageMaker JumpStart endpoint.

The following code example shows how create an endpoint for a Llama 2 foundation model that's available through SageMaker JumpStart.

```python
import sagemaker
from sagemaker.jumpstart.model import JumpStartModel

# JumpStart model and version
model_id, model_version = "meta-textgeneration-llama-2-7b-f", "*"

my_model = JumpStartModel(model_id=model_id)
predictor = my_model.deploy()
endpoint_name = predictor.endpoint_name

# Accept the EULA, and test the endpoint to make sure it can predict.
predictor.predict({"inputs": [[{"role":"user", "content": "Hello how are you?"}]},
custom_attributes='accept_eula=true')
```

The previous code example refers to EULA, which stands for end-use-license-agreement (EULA). The EULA can be found in the model card description of the model that you are using. To use some SageMaker JumpStart models, you must specify accept_eula=true, as shown
in the previous call to predict. For more information about EULA, see the Licenses and model sources section in Model sources and license agreements.

You can find a list of available SageMaker JumpStart models at Built-in Algorithms with pre-trained Model Table.

2. Configure ModelRunner by using the JumpStartModelRunner, as shown in the following configuration example:

```python
from fmeval.model_runners.sm_jumpstart_model_runner import JumpStartModelRunner

js_model_runner = JumpStartModelRunner(
    endpoint_name=endpoint_name,
    model_id=model_id,
    model_version=model_version
)
```

In the previous configuration example, use the same values for endpoint_name, model_id, and model_version that you used to create the endpoint.

3. Test your ModelRunner. Send a sample request to your model as shown in the following code example:

```python
js_model_runner.predict("What is the capital of London")
```

Use an Amazon Bedrock model

To evaluate an Amazon Bedrock model, you must define the model and built-in dataset, and configure ModelRunner.

Define an Amazon Bedrock model and configure a ModelRunner

1. To define and print model details, use the following code example for a Titan model that is available through Amazon Bedrock:

```python
import boto3
import json
bedrock = boto3.client(service_name='bedrock')
bedrock_runtime = boto3.client(service_name='bedrock-runtime')

model_id = "amazon.titan-tg1-large"
```
In the previous code example, the `accept` parameter specifies the format of the data that you want to use to evaluate your LLM. The `contentType` specifies the format of the input data in the request. Only MIME_TYPE_JSON is supported for `accept` and `contentType` for Amazon Bedrock models. For more information about these parameters, see `InvokeModelWithResponseStream`.

2. To configure `ModelRunner`, use the `BedrockModelRunner`, as shown in the following configuration example:

```python
from fmeval.model_runners.bedrock_model_runner import BedrockModelRunner

bedrock_model_runner = BedrockModelRunner(
    model_id=model_id,
    output='results[0].outputText',
    content_template='{"inputText": $prompt, "textGenerationConfig": {
        "maxTokenCount": 4096, "stopSequences": [], "temperature": 1.0, "topP": 1.0}}',
)
```

Parametrize the `ModelRunner` configuration as follows.

- Use the same values for `model_id` that you used to deploy the model.
- Use `output` to specify the format of the generated JSON response. As an example, if your LLM provided the response `"results": "this is the output"`, then `output='results[0].outputText'` returns `this is the output`.
- Use `content_template` to specify how your LLM interacts with requests. The following configuration template is detailed solely to explain the previous configuration example, and it's not required.
  - In the previous configuration example, the variable `inputText` specifies the prompt, which captures the request made by the user.
  - The variable `textGenerationConfig` specifies how the LLM generates responses as follows:
• The parameter `maxTokenCount` is used to limit the length of the response by limiting the number of tokens returned by the LLM.

• The parameter `stopSequences` is used to specify a list of character sequences that tell your LLM to stop generating a response. The model output is stopped the first time any of the listed strings are encountered in the output. As an example, you can use a carriage return sequence to limit the model response to a single line.

• The parameter `topP` controls the randomness by limiting the set of tokens to consider when generating the next token. This parameter accepts values between 0.0 and 1.0. Higher values of topP allow for a set containing a broader vocabulary and lower values restrict the set of tokens to more probable words.

• The parameter `temperature` controls the randomness of the generated text, and accepts positive values. Higher values of `temperature` instruct the model to generate more random and diverse responses. Lower values generate responses that are more predictable. Typical ranges for `temperature` lie between 0.2 and 2.0.

For more information about parameters for a specific Amazon Bedrock foundation model, see [Inference parameters for foundation models](#).

The format of the `content_template` parameter depends on the inputs and parameters supported by your LLM. For example, Anthropic’s Claude 2 model can support the following `content_template`:

```json
"content_template": "{\"prompt\": $prompt, \"max_tokens_to_sample\": 500}"
```

As another example, the Falcon 7b model can support the following `content_template`.

```json
"content_template": "{\"inputs\": $prompt, \"parameters\":{\"max_new_tokens\": \" 10, \"top_p\": 0.9, \"temperature\": 0.8}}"
```

Lastly, test your ModelRunner. Send a sample request to your model as shown in the following code example:

```python
bedrock_model_runner.predict("What is the capital of London?")
```
Evaluate your model

After you configure your data and ModelRunner, you can run an evaluation algorithm on the responses generated by your LLM. To see a list of all of the available evaluation algorithms, run the following code:

```python
from fmeval.eval_algo_mapping import EVAL_ALGORITHMS
print(EVAL_ALGORITHMS.keys())
```

Each algorithm has both an evaluate and an evaluate_sample method. The evaluate method computes a score for the entire dataset. The evaluate_sample method evaluates the score for a single instance.

The evaluate_sample method returns EvalScore objects. EvalScore objects contain aggregated scores of how well your model performed during evaluation. The evaluate_sample method has the following optional parameters:

- `model_output` – The model response for a single request.
- `model_input` – A prompt containing the request to your model.
- `target_output` – The expected response from the prompt contained in `model_input`.

The following code example shows how to use the evaluate_sample:

```python
#Evaluate your custom sample
model_output = model_runner.predict("London is the capital of?")[0]
eval_algo.evaluate_sample(target_output="UK<OR>England<OR>United Kingdom",
model_output=model_output)
```

The evaluate method has the following optional parameters:

- `model` – An instance of ModelRunner using the model that you want to evaluate.
- `dataset_config` – The dataset configuration. If `dataset_config` is not provided, the model is evaluated using all of the built-in datasets that are configured for this task.
- `prompt_template` – A template used to generate prompts. If `prompt_template` is not provided, your model is evaluated using a default prompt template.
- `save` – If set to True, record-wise prompt responses and scores are saved to the file EvalAlgorithmInterface.EVAL_RESULTS_PATH. Defaults to False.
- `num_records` – The number of records that are sampled randomly from the input dataset for evaluation. Defaults to 300.

The evaluate algorithm returns a list of `EvalOutput` objects that can include the following:

- `eval_name` – The name of the evaluation algorithm.

- `dataset_name` – The name of dataset used by the evaluation algorithm.

- `prompt_template` – A template used to compose prompts that is consumed if the parameter `model_output` is not provided in the dataset. For more information, see `prompt_template` in the Configure a SageMaker JumpStart ModelRunner section.

- `dataset_scores` – An aggregated score computed across the whole dataset.

- `category_scores` – A list of `CategoryScore` objects that contain the scores for each category in the dataset.

- `output_path` – The local path to the evaluation output. This output contains prompt-responses with record-wise evaluation scores.

- `error` – A string error message for a failed evaluation job.

The following dimensions are available for model evaluation:

- Accuracy
- Factual knowledge
- Prompt stereotyping
- Semantic robustness
- Toxicity

**Accuracy**

You can run an accuracy algorithm for a question answering, text summarization, or classification task. The algorithms are different for each task in order to accommodate the different data input types and problems as follows:

- For question answering tasks, run the QAAccuracy algorithm with a QAAccuracyConfig file.
• For text summarization tasks, run the `SummarizationAccuracy` algorithm with a `SummarizationAccuracyConfig`.

• For classification tasks, run the `ClassificationAccuracy` algorithm with a `ClassificationAccuracyConfig`.

The `QAAccuracy` algorithm returns a list of `EvalOutput` objects that contains one accuracy score for each sample. To run the question answer accuracy algorithm, instantiate a `QAAccuracyConfig` and pass in either `<OR>` or `None` as the `target_output_delimiter`. The question answer accuracy algorithm compares the response that your model generates with a known response. If you pass in `<OR>` as the target delimiter, then the algorithm scores the response as correct if it generates any of the content separated by `<OR>` in the answer. If you pass `None` or an empty string as the `target_output_delimiter`, the code throws an error.

Call the `evaluate` method and pass in your desired parameters as shown in the following code example:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.qa_accuracy import QAAccuracy, QAAccuracyConfig

eval_algo = QAAccuracy(QAAccuracyConfig(target_output_delimiter="<OR>"))
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config,
                                 prompt_template="$feature", save=True)
```

The `SummarizationAccuracy` algorithm returns a list of `EvalOutput` objects that contain scores for **ROUGE-N**, **Meteor**, and **BERTScore**. For more information about these scores, see the Text summarization section in [Foundation model evaluation overview](#). To run the text summarization accuracy algorithm, instantiate a `SummarizationAccuracyConfig` and pass in the following:

• Specify the type of **ROUGE** metric you want to use in your evaluation to `rouge_type`. You can choose `rouge1`, `rouge2`, or `rougeL`. These metrics compare generated summaries to reference summaries. ROUGE-1 compares the generated summaries and reference summaries using overlapping unigrams (sequences of one item such as “the”, “is”). ROUGE-2 compares the generated and reference summaries using bigrams (groups of two sequences such as “the large”, “is home”). ROUGE-L compares the longest matching sequence of words. For more information about ROUGE, see [ROUGE: A Package for Automatic Evaluation of Summaries](#).
• Set `use_stemmer_for_rouge` to True or False. A stemmer removes affixes from words before comparing them. For example, a stemmer removes the affixes from “swimming” and “swam” so that they are both “swim” after stemming.

• Set `model_type_for_bertscore` to the model that you want to use to calculate a BERTScore. You can choose `ROBERTA_MODEL` or the more advanced `MICROSOFT_DEBERTA_MODEL`.

Lastly, call the evaluate method and pass in your desired parameters as shown in the following code example:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.summarization_accuracy import SummarizationAccuracy, SummarizationAccuracyConfig

eval_algo = SummarizationAccuracy(SummarizationAccuracyConfig(rouge_type="rouge1",model_type_for_bertscore="MICROSOFT_DEBERTA_MODEL"))
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config, prompt_template="$feature", save=True)
```

The `ClassificationAccuracy` algorithm returns a list of EvalOutput objects that contain the classification accuracy, precision, recall, and balanced accuracy scores for each sample. For more information about these scores, see the Classification section in Foundation model evaluation overview. To run the classification accuracy algorithm, instantiate a `ClassificationAccuracyConfig` and pass in an averaging strategy to `multiclass_average_strategy`. You can choose `micro`, `macro`, `samples`, `weighted`, or `binary`. The default value is `micro`. Then, pass in a list containing the names of the columns that contain the true labels for your classification categories to `valid_labels`. Lastly, call the evaluate method and pass in your desired parameters as shown in the following code example:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.classification_accuracy import ClassificationAccuracy, ClassificationAccuracyConfig

eval_algo = ClassificationAccuracy(ClassificationAccuracyConfig(multiclass_average_strategy="samples",valid_labels=["animal_type","plant_type","fungi_type"]))
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config, prompt_template="$feature", save=True)
```
**Factual knowledge**

You can run the factual knowledge algorithm for open-ended generation. To run the factual knowledge algorithm, instantiate a `FactualKnowledgeConfig` and optionally pass a delimiter string (by default, this is `<OR>`). The factual knowledge algorithm compares the response that your model generates with a known response. The algorithm scores the response as correct if it generates any of the content separated by the delimiter in the answer. If you pass `None` as the `target_output_delimiter`, then the model must generate the same response as the answer to be scored as correct. Lastly, call the `evaluate` method and pass in your desired parameters.

Factual knowledge returns a list of `EvalScore` objects. These contain aggregated scores on how well your model is able to encode factual knowledge as described in the Foundation model evaluation overview section. The scores range between 0 and 1 with the lowest score corresponding to a lower knowledge of real-world facts.

The following code example shows how to evaluate your LLM using the factual knowledge algorithm:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.factual_knowledge import FactualKnowledge,
    FactualKnowledgeConfig

eval_algo = FactualKnowledge(FactualKnowledgeConfig())
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config,
    prompt_template='$feature', save=True)
```

**Prompt stereotyping**

You can run the prompt stereotyping algorithm for open-ended generation. To run the prompt stereotyping algorithm, your `DataConfig` must identify the columns in your input dataset that contain a less stereotypical sentence in `sent_less_input_location` and a more stereotypical sentence in `sent_more_output_location`. For more information about `DataConfig`, see the previous section 2. Configure `ModelRunner`. Next, call the `evaluate` method and pass in your desired parameters.

Prompt stereotyping returns a list of `EvalOutput` objects that contain a score for each input record and overall scores for each type of bias. The scores are calculated by comparing the probability of the more and less stereotypical sentences. The overall score reports how often the model preferred the stereotypical sentence in that the model assigns a higher probability to the
more stereotypical compared to the less stereotypical sentence. A score of 0.5 indicates that your model is unbiased, or that it prefers more and less stereotypical sentences at equal rates. A score of greater than 0.5 indicates that your model is likely to generate a response that is more stereotypical. Scores less than 0.5 indicate that your model is likely to generate a response that is less stereotypical.

The following code example shows how to evaluate your LLM using the prompt stereotyping algorithm:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.prompt_stereotyping import PromptStereotyping

eval_algo = PromptStereotyping()
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config,
prompt_template="$feature", save=True)
```

**Semantic robustness**

You can run a semantic robustness algorithm for any FMEval task, however your model should be deterministic. A deterministic model is one that always generate the same output for the same input. One may typically achieve determinism by setting a random seed in the decoding process. The algorithms are different for each task in order to accommodate the different data input types and problems as follows:

- For open-ended generation, question answering, or task classification run the GeneralSemanticRobustness algorithm with a GeneralSemanticRobustnessConfig file.
- For text summarization, run the SummarizationAccuracySemanticRobustness algorithm with a SummarizationAccuracySemanticRobustnessConfig file.

The GeneralSemanticRobustness algorithm returns a list of EvalScore objects that contain accuracy with values between 0 and 1 quantifying the difference between the perturbed and unperturbed model outputs. To run the general semantic robustness algorithm, instantiate a GeneralSemanticRobustnessConfig and pass in a perturbation_type. You can choose one of the following for perturbation_type:

- **Butterfinger** – A perturbation that mimics spelling mistakes using character swaps based on keyboard distance. Input a probability that a given character is perturbed. Butterfinger is the default value for perturbation_type.
• RandomUpperCase – A perturbation that changes a fraction of characters to uppercase. Input a decimal from 0 to 1.
• WhitespaceAddRemove – The probability that a white space character is added in front of a non-white space character into white.

You can also specify the following parameters:

• num_perturbations – The number of perturbations for each sample to introduce into the generated text. The default is 5.
• butter_finger_perturbation_prob – The probability that a character is be perturbed. Used only when perturbation_type is Butterfinger. The default is 0.1.
• random_uppercase_corrupt_proportion – The fraction of characters to be changed to uppercase. Used only when perturbation_type is RandomUpperCase. The default is 0.1.
• whitespace_add_prob – Given a white space, the probability of removing it from a sample. Used only when perturbation_type is WhitespaceAddRemove. The default is 0.05.
• whitespace_remove_prob – Given a non-white space, the probability of adding a white space in front of it. Used only when perturbation_type is WhitespaceAddRemove. The default is 0.1.

Lastly, call the evaluate method and pass in your desired parameters as shown in the following code example:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.general_semantic_robustness import GeneralSemanticRobustness, GeneralSemanticRobustnessConfig

eval_algo = GeneralSemanticRobustness(GeneralSemanticRobustnessConfig(perturbation_type="RandomUpperCase", num_perturbations=2, random_uppercase_corrupt_proportion=0.3))
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config, prompt_template="$feature", save=True)
```

The SummarizationAccuracySemanticRobustness algorithm returns a list of EvalScore objects that contain the difference (or delta) between the ROUGE-N, Meteor, and BERTScore values between the generated and reference summaries. For more information about these scores, see the Text summarization section in Foundation model evaluation.
To run the text summarization semantic robustness algorithm, instantiate a `SummarizationAccuracySemanticRobustnessConfig` and pass in a `perturbation_type`.

You can choose one of the following for `perturbation_type`:

- **Butterfinger** – A perturbation that mimics spelling mistakes using character swaps based on keyboard distance. Input a probability that a given character is perturbed. Butterfinger is the default value for `perturbation_type`.
- **RandomUpperCase** – A perturbation that changes a fraction of characters to uppercase. Input a decimal from 0 to 1.
- **WhitespaceAddRemove** – Input a probability that a white space character is added in front of a non-white space character into white.

You can also specify the following parameters:

- **num_perturbations** – The number of perturbations for each sample to introduce into the generated text. Default is 5.
- **butter_finger_perturbation_prob** – The probability that a character is perturbed. Used only when `perturbation_type` is Butterfinger. Default is 0.1.
- **random_uppercase_corrupt_proportion** – The fraction of characters to be changed to uppercase. Used only when `perturbation_type` is RandomUpperCase. Default is 0.1.
- **whitespace_add_prob** – Given a white space, the probability of removing it from a sample. Used only when `perturbation_type` is WhitespaceAddRemove. Default is 0.05.
- **whitespace_remove_prob** – Given a non-white space, the probability of adding a white space in front of it. Used only when `perturbation_type` is WhitespaceAddRemove. Default is 0.1.
- **rouge_type** – Metrics that compare generated summaries to reference summaries. Specify the type of ROUGE metric you want to use in your evaluation to `rouge_type`. You can choose `rouge1`, `rouge2`, or `rougeL`. ROUGE-1 compares the generated summaries and reference summaries using overlapping unigrams (sequences of one item such as “the”, “is”). ROUGE-2 compares the generated and reference summaries using bigrams (groups of two sequences such as “the large”, “is home”). ROUGE-L compares the longest matching sequence of words. For more information about ROUGE, see [ROUGE: A Package for Automatic Evaluation of Summaries](https://www.aaai.org/Papers/AAAI/2004/AAAI04-093.pdf).
- Set `user_stemmer_for_rouge` to True or False. A stemmer removes affixes from words before comparing them. For example, a stemmer removes the affixes from “swimming” and “swam” so that they are both “swim” after stemming.
• Set `model_type_for_bertscore` to the model that you want to use to calculate a BERTScore. You can choose `ROBERTA_MODEL` or the more advanced `MICROSOFT_DEBERTA_MODEL`.

Call the `evaluate` method and pass in your desired parameters as shown in the following code example:

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.summarization_accuracy_semantic_robustness import
    SummarizationAccuracySemanticRobustness,
    SummarizationAccuracySemanticRobustnessConfig

eval_algo =
    SummarizationAccuracySemanticRobustness(SummarizationAccuracySemanticRobustnessConfig(perturbation_type="Butterfinger",num_perturbations=3,butter_finger_perturbation_prob=0.2)))
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config,
                                prompt_template="$feature", save=True)
```

**Toxicity**

You can run the a toxicity algorithm for open-ended generation, text summarization, or question answering. There are three distinct classes depending on the task.

• For open-ended generation, run the Toxicity algorithm with a ToxicityConfig file.
• For summarization, use the class Summarization_Toxicity.
• For question answering, use the class QAToxicity.

The toxicity algorithm returns one or more a list of EvalScore objects (depending on the toxicity detector) that contain scores between 0 and 1. To run the toxicity algorithm, instantiate a ToxicityConfig and pass in a toxicity model to use to evaluate your model against in `model_type`. You can choose the following for `model_type`:

• `detoxify` for UnitaryAI Detoxify-unbiased, a multilabel text classifier trained on Toxic Comment Classification Challenge and Jigsaw Unintended Bias in Toxicity Classification. The model provides 7 scores for the following classes: toxicity, severe toxicity, obscenity, threat, insult, sexual explicitity and identity attack.

The following is example output from the toxicity model:

```python
EvalScore(name='toxicity', value=0.01936926692724228),
```
EvalScore(name='severe_toxicity', value=3.375677577573806e-06),
EvalScore(name='obscene', value=0.00022437423467636108),
EvalScore(name='identity_attack', value=0.0006707844440825284),
EvalScore(name='insult', value=0.005559926386922598),
EvalScore(name='threat', value=0.00016682750720065087),
EvalScore(name='sexual_explicit', value=4.828436431125738e-05)

- `'toxigen` for Toxigen-roberta, a binary RoBERTa-based text classifier fine-tuned on the ToxiGen dataset, which contains sentences with subtle and implicit toxicity pertaining to 13 minority groups.

Lastly, call the `evaluate` method and pass in your desired parameters as shown in the following code example.

```python
from fmeval.eval import get_eval_algorithm
from fmeval.eval_algorithms.toxicity import Toxicity, ToxicityConfig

eval_algo = Toxicity(ToxicityConfig(model_type="detoxify"))
eval_output = eval_algo.evaluate(model=model_runner, dataset_config=config,
prompt_template="$feature", save=True)
```

**Customize your workflow using the fmeval library**

⚠️ **Important**
In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see [Migrating from Amazon SageMaker Studio Classic](#). For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).
You can customize your model evaluation to allow for a model that is not a SageMaker JumpStart or Amazon Bedrock model or use a custom workflow for evaluation. If you use your own model, you have to create a custom ModelRunner. If you use your own dataset for evaluation, you must configure a DataConfig object. The following section shows how to format your input dataset, customize a DataConfig object to use your custom dataset, and create a custom ModelRunner.

**Use a custom input dataset**

If you want to use your own dataset to evaluate your model, you must use a DataConfig object to specify the dataset_name and the dataset_uri of the dataset that you want to evaluate. If you use a built-in dataset, the DataConfig object is already configured as the default for evaluation algorithms.

You can use one custom dataset every time you use the evaluate function. You can invoke evaluate any number of times to use any number of datasets that you want.

Configure a custom dataset with your model request specified in the question column, and the target answer specified in the column answer, as follows:

```python
from fmeval.data_loaders.data_config import DataConfig
from fmeval.constants import MIME_TYPE_JSONLINES

config = DataConfig(
    dataset_name="tiny_dataset",
    dataset_uri="tiny_dataset.jsonl",
    dataset_mime_type=MIME_TYPE_JSONLINES,
    model_input_location="question",
    target_output_location="answer",
)
```

The DataConfig class contains the following parameters:

- **dataset_name** – The name of the dataset that you want to use to evaluate your LLM.
- **dataset_uri** – The local path or uniform resource identifier (URI) to the S3 location of your dataset.
- **dataset_mime_type** – The format of the input data that you want to use to evaluate your LLM. The FMEval library can support both MIME_TYPE_JSON and MIME_TYPE_JSONLINES.
- **model_input_location** – (Optional) The name of the column in your dataset that contains the model inputs or prompts that you want to evaluate.
Use a model_input_location that specifies the name of your column. The column must contain the following values corresponding to the following associated tasks:

- For **open-ended generation**, **toxicity**, and **accuracy** evaluations, specify the column that contains the **prompt** that your model should respond to.

- For a **question answering** task, specify the column that contains the **question** that your model should generate a response to.

- For a **text summarization task**, specify the name of the column that contains the **text** that you want your model to summarize.

- For a **classification task**, specify the name of the column that contains the **text** that you want your model to classify.

- For a **factual knowledge** evaluations, specify the name of the column that contains the **question** that you want the model to predict the answer to.

- For **semantic robustness** evaluations, specify the name of the column that contains the **input** that you want your model to perturb.

- For **prompt stereotyping** evaluations, use the sent_more_input_location and sent_less_input_location instead of model_input_location, as shown in the following parameters.

- **model_output_location** – (Optional) The name of the column in your dataset that contains the predicted output that you want to compare against the reference output that is contained in target_output_location. If you provide model_output_location, then FMEval won’t send a request to your model for inference. Instead, it uses the output contained in the specified column to evaluate your model.

- **target_output_location** – The name of the column in the reference dataset that contains the true value to compare against the predicted value that is contained in model_output_location. Required only for factual knowledge, accuracy, and semantic robustness. For factual knowledge, each row in this column should contain all possible answers separated by a delimiter. For example, if the answers for a question are [“UK”, “England”], then the column should contain “UK<OR>England”. The model prediction is correct if it contains any of the answers separated by the delimiter.

- **category_location** – The name of the column that contains the name of a category. If you provide a value for category_location, then scores are aggregated and reported for each category.
• `sent_more_input_location` – The name of the column that contains a prompt with more bias. Required only for prompt stereotyping. Avoid unconscious bias. For bias examples, see the CrowS-Pairs dataset.

• `sent_less_input_location` – The name of the column that contains a prompt with less bias. Required only for prompt stereotyping. Avoid unconscious bias. For bias examples, see the CrowS-Pairs dataset.

• `sent_more_output_location` – (Optional) The name of the column that contains a predicted probability that your model's generated response will contain more bias. This parameter is only used in prompt stereotyping tasks.

• `sent_less_output_location` – (Optional) The name of the column that contains a predicted probability that your model's generated response will contain less bias. This parameter is only used in prompt stereotyping tasks.

If you want to add a new attribute that corresponds to a dataset column to the DataConfig class, you must add the suffix `_location` to the end of the attribute name.

**Use a custom ModelRunner**

To evaluate a custom model, use a base data class to configure your model and create a custom ModelRunner. Then, you can use this ModelRunner to evaluate any language model. Use the following steps to define a model configuration, create a custom ModelRunner, and test it.

The ModelRunner interface has one abstract method as follows:

```python
def predict(self, prompt: str) # Tuple[Optional[str], Optional[float]]
```

This method takes in a prompt as a string input, and returns a Tuple containing a model text response and an input log probability. Every ModelRunner must implement a predict method.

**Create a custom ModelRunner**

1. Define a model configuration.

   The following code example shows how to apply a dataclass decorator to a custom HFModelConfig class so that you can define a model configuration for a Hugging Face model:

```python
from dataclasses import dataclass
from dataclasses import import dataclass
```
@dataclass
class HFModelConfig:
    model_name: str
    max_new_tokens: int
    seed: int = 0
    remove_prompt_from_generated_text: bool = True

In the previous code example, the following applies:

- The parameter `max_new_tokens` is used to limit the length of the response by limiting the number of tokens returned by an LLM. The type of model is set by passing a value for `model_name` when the class is instantiated. In this example, the model name is set to `gpt2`, as shown in the end of this section. The parameter `max_new_tokens` is one option to configure text generation strategies using a gpt2 model configuration for a pre-trained OpenAI GPT model. See `AutoConfig` for other model types.

- If the parameter `remove_prompt_from_generated_text` is set to `True`, then the generated response won’t contain the originating prompt sent in the request.

For other text generation parameters, see the Hugging Face documentation for `GenerationConfig`.

2. Create a custom `ModelRunner` and implement a `predict` method. The following code example shows how to create a custom `ModelRunner` for a Hugging Face model using the `HFModelConfig` class created in the previous code example.

```python
from typing import Tuple, Optional
import torch
from transformers import AutoModelForCausalLM, AutoTokenizer
from fmeval.model_runners.model_runner import ModelRunner

class HuggingFaceCausalLLMModelRunner(ModelRunner):
    def __init__(self, model_config: HFModelConfig):
        self.config = model_config
        self.model = AutoModelForCausalLM.from_pretrained(self.config.model_name)
        self.tokenizer = AutoTokenizer.from_pretrained(self.config.model_name)

    def predict(self, prompt: str) -> Tuple[Optional[str], Optional[float]]:
        input_ids = self.tokenizer(prompt, Optional[float]).to(self.model.device)
```
generations = self.model.generate(
    **input_ids,
    max_new_tokens=self.config.max_new_tokens,
    pad_token_id=self.tokenizer.eos_token_id,
)

generation_contains_input = (input_ids['input_ids'][0] == generations[0][:input_ids['input_ids'].shape[1]]).all()
    if self.config.remove_prompt_from_generated_text and not generation_contains_input:
        warnings.warn('Your model does not return the prompt as part of its generations. '
                      '`remove_prompt_from_generated_text` does nothing.'
    )
    if self.config.remove_prompt_from_generated_text and
    generation_contains_input:
        output = self.tokenizer.batch_decode(generations[:,
    input_ids['input_ids'].shape[1] :])[0]
    else:
        output = self.tokenizer.batch_decode(generations,
    skip_special_tokens=True)[0]

    with torch.inference_mode():
        input_ids = self.tokenizer(self.tokenizer.bos_token + prompt,
    return_tensors="pt")['input_ids']
        model_output = self.model(input_ids, labels=input_ids)
        probability = -model_output[0].item()

    return output, probability

The previous code uses a custom HuggingFaceCausalLLMModelRunner class that inherits properties from the FMEval ModelRunner class. The custom class contains a constructor and a definition for a predict function, which returns a Tuple.

For more ModelRunner examples, see the model_runner section of the fmeval library.

The HuggingFaceCausalLLMModelRunner constructor contains the following definitions:

- The configuration is set to HFModelConfig, defined in the beginning of this section.
• The model is set to a pre-trained model from the Hugging Face Auto Class that is specified using the model_name parameter upon instantiation.

• The tokenizer is set to a class from the Hugging Face tokenizer library that matches the pre-trained model specified by model_name.

The predict method in the HuggingFaceCausalLLMModelRunner class uses the following definitions:

• input_ids – A variable that contains input for your model. The model generates the input as follows.
  • A tokenizer Converts the request contained in prompt into token identifiers (IDs). These token IDs, which are numerical values that represent a specific token (word, sub-word or character), can be used directly by your model as input. The token IDs are returned as a PyTorch tensor objects, as specified by return_tensors="pt". For other types of return tensor types, see the Hugging Face documentation for apply_chat_template.
  • Token IDs are sent to a device where the model is located so that they can be used by the model.

• generations – A variable that contains the response generated by your LLM. The model's generate function uses the following inputs to generate the response:
  • The input_ids from the previous step.
  • The parameter max_new_tokens specified in HFModelConfig.
  • A pad_token_id adds an end of sentence (eos) token to the response. For other tokens that you can use, see the Hugging Face documentation for PreTrainedTokenizer.

• generation_contains_input – A boolean variable that returns True when the generated response includes the input prompt in its response, and False otherwise. The return value is calculated using an element-wise comparison between the following.
  • All of the token IDs in the input prompt that are contained in input_ids["input_ids"][0].
  • The beginning of the generated content that is contained in generations[0][: input_ids["input_ids"].shape[1]].

The predict method returns a warning if you directed the LLM to remove_prompt_from_generated_text in your configuration but the generated response doesn't contain the input prompt.
The output from the `predict` method contains a string returned by the `batch_decode` method, which converts token IDs returned in the response into human readable text. If you specified `remove_prompt_from_generated_text` as `True`, then the input prompt is removed from the generated text. If you specified `remove_prompt_from_generated_text` as `False`, the generated text will be returned without any special tokens that you included in the dictionary `special_token_dict`, as specified by `skip_special_tokens=True`.

3. Test your `ModelRunner`. Send a sample request to your model.

The following example shows how to test a model using the `gpt2` pre-trained model from the Hugging Face `AutoConfig` class:

```python
hf_config = HFModelConfig(model_name='gpt2', max_new_tokens=32)
model = HuggingFaceCausalLMMModelRunner(model_config=hf_config)
```

In the previous code example, `model_name` specifies the name of the pre-trained model. The `HFModelConfig` class is instantiated as `hf_config` with a value for the parameter `max_new_tokens`, and used to initialize `ModelRunner`.

If you want to use another pre-trained model from Hugging Face, choose a `pretrained_model_name_or_path` in `from_pretrained` under `AutoClass`.

Lastly, test your `ModelRunner`. Send a sample request to your model as shown in the following code example:

```python
model_output = model.predict("London is the capital of?")[0]
print(model_output)
eval_algo.evaluate_sample()
```

---

**Notebook tutorials**

⚠️ **Important**

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update
This section provides the following notebook tutorials, which include example code and explanations:

- How to evaluate a SageMaker JumpStart model for prompt stereotyping.
- How to evaluate an Amazon Bedrock model for text summarization accuracy.

**How to evaluate a SageMaker JumpStart model for prompt stereotyping**

You can use a high-level `ModelRunner` wrapper to evaluate an Amazon SageMaker JumpStart model for prompt stereotyping. The prompt stereotyping algorithm measures the probability of your model encoding biases in its response. These biases include those for race, gender, sexual orientation, religion, age, nationality, disability, physical appearance, and socioeconomic status.

This tutorial shows how to load the Falcon 7-B model from the Technology Innovation Institute, available in SageMaker JumpStart, and ask this model to generate responses to prompts. Then, this tutorial shows how to evaluate the responses for prompt stereotyping against the built-in CrowS-Pairs open source challenge dataset.

The sections of this tutorial show how to do the following:

- Set up your environment.
- Run your model evaluation.
- View your analysis results.

**Set up your environment**

**Prerequisites**

- Use a base Python 3.10 kernel environment and an `ml.g4dn.2xlarge` Amazon Elastic Compute Cloud (Amazon EC2) instance before starting this tutorial.

For more information about instance types and their recommended use cases, see Available Studio Classic Instance Types.
Install required libraries

1. Install the SageMaker, fmeval, and other required libraries in your code as follows:

   ```bash
   !pip3 install sagemaker
   !pip3 install -U pyarrow
   !pip3 install -U accelerate
   !pip3 install "ipywidgets>=8"
   !pip3 install jsonlines
   !pip install fmeval
   !pip install boto3==1.28.65
   import sagemaker
   ```

2. Download the sample JSON Lines dataset `crows-pairs_sample.jsonl`, into your current working directory.

3. Check that your environment contains the sample input file using the following code:

   ```python
   import glob

   # Check for fmeval wheel and built-in dataset
   if not glob.glob("crows-pairs_sample.jsonl"):
       print("ERROR - please make sure file exists: crows-pairs_sample.jsonl")
   ```

4. Define a SageMaker JumpStart model as follows:

   ```python
   from sagemaker.jumpstart.model import JumpStartModel

   model_id, model_version, = (
       "huggingface-llm-falcon-7b-instruct-bf16",
       "*",
   )
   ```

5. Deploy the SageMaker JumpStart model and create an endpoint as follows:

   ```python
   my_model = JumpStartModel(model_id=model_id)
   predictor = my_model.deploy()
   endpoint_name = predictor.endpoint_name
   ```

6. Define a prompt and the format of the model request, or payload, as follows:

   ```python
   prompt = "London is the capital of"
   ```
In the previous code example, the following parameters are included in the model request:

- **do_sample** – Instructs the model to sample from the raw model outputs (prior to normalization) during model inference to introduce diversity and creativity into model responses. Defaults to False. If you set `do_sample` to `True`, then you must specify a value for one of the following parameters: `temperature`, `top_k`, `top_p`, or `typical_p`.

- **top_p** – Controls the randomness by limiting the set of tokens to consider when generating the next token. Higher values of `top_p` allow for a set containing a broader vocabulary. Lower values restrict the set of tokens to more probable words. Ranges for `top_p` are greater than 0 and less than 1.

- **temperature** – Controls the randomness of the generated text. Higher values of `temperature` instruct the model to generate more random and diverse responses. Lower values generate responses that are more predictable. Values for `temperature` must be positive.

- **max_new_tokens** – Limits the length of the response by limiting the number of tokens returned by your model. Defaults to 20.

- **decoder_input_details** – Returns information about the log probabilities assigned by the model to each potential next token and the corresponding token IDs. If `decoder_input_details` is set to `True`, you must also set `details` to `True` in order to receive the requested details. Defaults to `False`.

For more information about parameters for this Hugging Face model, see `types.py`.
Send a sample inference request

To test your model, send a sample request to your model and print the model response as follows:

```python
response = predictor.predict(payload)
print(response[0]["generated_text"])
```

In the previous code example, if your model provided the response `[{"response": "this is the output"}]`, then the `print` statement returns `this is the output`.

Set up FMEval

1. Load the required libraries to run FMEval as follows:

```python
import fmeval
from fmeval.data_loaders.data_config import DataConfig
from fmeval.model_runners.sm_jumpstart_model_runner import JumpStartModelRunner
from fmeval.constants import MIME_TYPE_JSONLINES
from fmeval.eval_algorithms.prompt_stereotyping import PromptStereotyping,
    PROMPT_STEREOTYPING
from fmeval.eval_algorithms import EvalAlgorithm
```

2. Set up the data configuration for your input dataset.

   If you don't use a built-in dataset, your data configuration must identify the column that contains more bias in `sent_more_input_location`. You must also identify the column that contains less bias in `sent_less_input_location`. If you are using a built-in dataset from SageMaker JumpStart, these parameters are passed to FMEval automatically through the model metadata.

   Specify the `sent_more_input_location` and `sent_less_input_location` columns for a prompt stereotyping task, the name, uniform resource identifier (URI), and MIME type.

```python
config = DataConfig(
    dataset_name="crows-pairs_sample",
    dataset_uri="crows-pairs_sample.jsonl",
    dataset_mime_type=MIME_TYPE_JSONLINES,
    sent_more_input_location="sent_more",
    sent_less_input_location="sent_less",
    category_location="bias_type",
)
```
For more information about column information that other tasks require, see the Use a custom input dataset section in Use a custom input dataset.

3. Set up a custom ModelRunner as shown in the following code example:

```python
js_model_runner = JumpStartModelRunner(
    endpoint_name=endpoint_name,
    model_id=model_id,
    model_version=model_version,
    output='[0].generated_text',
    log_probability='[0].details.prefill[*].logprob',
    content_template='{"inputs": $prompt, "parameters":
        {"do_sample": true, "top_p": 0.9, "temperature": 0.8, "max_new_tokens": 1024,
        "decoder_input_details": true,"details": true}}',
)
```

The previous code example specifies the following:

- **endpoint_name** – The name of the endpoint that you created in the previous Install required libraries step.

- **model_id** – The id used to specify your model. This parameter was specified when the SageMaker JumpStart model was defined.

- **model_version** – The version of your model used to specify your model. This parameter was specified when the SageMaker JumpStart model was defined.

- **output** – Captures the output from the Falcon 7b model, which returns its response in a generated_text key. If your model provided the response `[{"generated_text": "this is the output"}]`, then `[0].generated_text` returns this is the output.

- **log_probability** – Captures the log probability returned by this SageMaker JumpStart model.

- **content_template** – Specifies how your model interacts with requests. The example configuration template is detailed solely to explain the previous example, and it's not required. The parameters in the content template are the same ones that are declared for payload. For more information about parameters for this Hugging Face model, see types.py.

4. Configure your evaluation report and save it to a directory as shown in the following example code:
import os
eval_dir = "results-eval-prompt-stereotyping"
curr_dir = os.getcwd()
eval_results_path = os.path.join(curr_dir, eval_dir) + ""/"
os.environ["EVAL_RESULTS_PATH"] = eval_results_path
if os.path.exists(eval_results_path):
    print(f"Directory '{eval_results_path}' exists.")
else:
    os.mkdir(eval_results_path)

5. Set up a parallelization factor as follows:

    os.environ["PARALLELIZATION_FACTOR"] = "1"

A PARALLELIZATION_FACTOR is a multiplier for the number of concurrent batches sent to your compute instance. If your hardware allows for parallelization, you can set this number to multiply the number of invocations for your evaluation job. For example, if you have 100 invocations, and PARALLELIZATION_FACTOR is set to 2, then your job will run 200 invocations. You can increase PARALLELIZATION_FACTOR up to 10, or remove the variable entirely. To read a blog about how AWS Lambda uses PARALLELIZATION_FACTOR see New AWS Lambda scaling controls for Kinesis and DynamoDB event sources.

Run your model evaluation

1. Define your evaluation algorithm. The following example shows how to define a PromptStereotyping algorithm:

    eval_algo = PromptStereotyping()

For examples of algorithms that calculate metrics for other evaluation tasks, see Evaluate your model in Use the fmeval library to run an automatic evaluation.

2. Run your evaluation algorithm. The following code example uses the model and data configuration that was previously defined, and a prompt_template that uses feature to pass your prompt to the model as follows:

    eval_output = eval_algo.evaluate(model=js_model_runner, dataset_config=config, prompt_template="$feature", save=True)
Your model output may be different than the previous sample output.

**View your analysis results**

1. Parse an evaluation report from the `eval_output` object returned by the evaluation algorithm as follows:

   ```python
   import json
   print(json.dumps(eval_output, default=vars, indent=4))
   ```

   The previous command returns the following output (condensed for brevity):

   ```json
   [
   {
     "eval_name": "prompt_stereotyping",
     "dataset_name": "crows-pairs_sample",
     "dataset_scores": [
       {
         "name": "prompt_stereotyping",
         "value": 0.6666666666666666
       }
     ],
     "prompt_template": "$feature",
     "category_scores": [
       {
         "name": "disability",
         "scores": [
           {
             "name": "prompt_stereotyping",
             "value": 0.5
           }
         ]
       },
       ...
     ],
     "output_path": "/home/sagemaker-user/results-eval-prompt-stereotyping/
                 prompt_stereotyping_crows-pairs_sample.jsonl",
     "error": null
   }
   ]
   ```
The previous example output displays an overall score for dataset following "name": prompt_stereotyping. This score is the normalized difference in log probabilities between the model response providing more versus less bias. If the score is greater than 0.5, this means that your model response is more likely to return a response containing more bias. If the score is less than 0.5, your model is more likely to return a response containing less bias. If the score is 0.5, the model response does not contain bias as measured by the input dataset. You will use the output_path to create a Pandas DataFrame in the following step.

2. Import your results and read them into a DataFrame, and attach the prompt stereotyping scores to the model input, model output, and target output as follows:

   ```python
   import pandas as pd
   data = []
   with open(os.path.join(eval_results_path, "prompt_stereotyping_crows-pairs_sample.jsonl"), "r") as file:
       for line in file:
           data.append(json.loads(line))
   df = pd.DataFrame(data)
   df['eval_algo'] = df['scores'].apply(lambda x: x[0]['name'])
   df['eval_score'] = df['scores'].apply(lambda x: x[0]['value'])
   df
   
   For a notebook that contains the code examples given in this section, see jumpstart-falcon-stereotyping.ipynb.

How to evaluate an Amazon Bedrock model for text summarization accuracy

You can use a high-level ModelRunner wrapper to create a custom evaluation based on a model that is hosted outside of SageMaker JumpStart.

This tutorial shows how to load the Anthropic Claude 2 model, which is available in Amazon Bedrock, and ask this model to summarize text prompts. Then, this tutorial shows how to evaluate the model response for accuracy using the Rouge-L, Meteor, and BERTScore metrics.

The tutorials show how to do the following:

- Set up your environment.
- Run your model evaluation.
- View your analysis results.
Set up your environment

Prerequisites

- Use a base Python 3.10 kernel environment and an ml.m5.2xlarge Amazon Elastic Compute Cloud (Amazon EC2) instance before starting this tutorial.

For additional information about instance types and their recommended use cases, see Available Studio Classic Instance Types.

Set up Amazon Bedrock

Before you can use an Amazon Bedrock model, you have to request access to it.

1. Sign into your AWS account.
   - If you do not have an AWS account, see Sign up for an AWS account in Set up Amazon Bedrock.

2. Open the Amazon Bedrock console.

3. In the Welcome to Amazon Bedrock! section that opens, choose Manage model access.

4. In the Model access section that appears, choose Manage model access.

5. In the Base models section that appears, check the box next to Claude listed under the Anthropic subsection of Models.

6. Choose Request model access.

7. If your request is successful, a check mark with Access granted should appear under Access status next to your selected model.

8. You may need to log back into your AWS account to be able to access the model.

Install required libraries

1. In your code, install the fmeval and boto3 libraries as follows:

   ```bash
   !pip install fmeval
   !pip3 install boto3==1.28.65
   ```

2. Import libraries, set a parallelization factor, and invoke an Amazon Bedrock client as follows:

   ```python
   import boto3
   ```
import json
import os

# Dependent on available hardware and memory
os.environ["PARALLELIZATION_FACTOR"] = "1"

# Bedrock clients for model inference
bedrock = boto3.client(service_name='bedrock')
bedrock_runtime = boto3.client(service_name='bedrock-runtime')

In the previous code example, the following applies:

- PARALLELIZATION_FACTOR – A multiplier for the number of concurrent batches sent to your compute instance. If your hardware allows for parallelization, you can set this number to multiply the number of invocations for your evaluation job by. For example, if you have 100 invocations, and PARALLELIZATION_FACTOR is set to 2, then your job will run 200 invocations. You can increase PARALLELIZATION_FACTOR up to 10, or remove the variable entirely. To read a blog about how AWS Lambda uses PARALLELIZATION_FACTOR see New AWS Lambda scaling controls for Kinesis and DynamoDB event sources.

3. Download the sample JSON Lines dataset, sample-dataset.jsonl, into your current working directory.

4. Check that your environment contains the sample input file as follows:

```python
import glob

# Check for the built-in dataset
if not glob.glob("sample-dataset.jsonl"):
    print("ERROR - please make sure file exists: sample-dataset.jsonl")
```

**Send a sample inference request to your model**

1. Define the model and the MIME type of your prompt. For an Anthropic Claude 2 model hosted on Amazon Bedrock, your prompt must be structured as follows:

```python
import json
model_id = 'anthropic.claude-v2'
accept = "application/json"
contentType = "application/json"
# Ensure that your prompt has the correct format
```
prompt_data = """Human: Who is Barack Obama? 
Assistant: """

For more information about how to structure the body of your request, see Model invocation request body field. Other models may have different formats.

2. Send a sample request to your model. The body of your request contains the prompt and any additional parameters that you want to set. A sample request with the max_tokens_to_sample set to 500 follows:

```python
body = json.dumps({"prompt": prompt_data, "max_tokens_to_sample": 500})
response = bedrock_runtime.invoke_model(
    body=body, modelId=model_id, accept=accept, contentType=contentType
)
response_body = json.loads(response.get("body").read())
print(response_body.get("completion"))
```

In the previous code example, you can set the following parameters:

- **temperature** – Controls the randomness of the generated text, and accepts positive values. Higher values of temperature instruct the model to generate more random and diverse responses. Lower values generate responses that are more predictable. Ranges for temperature lie between 0 and 1, with a default value of 0.5.

- **topP** – Controls the randomness by limiting the set of tokens to consider when generating the next token. Higher values of topP allow for a set containing a broader vocabulary and lower values restrict the set of tokens to more probable words. Ranges for topP are 0 to 1, with a default value of 1.

- **topK** – Limits the model predictions to the top k most probable tokens. Higher values of topK allow for more inventive responses. Lower values generate responses that are more coherent. Ranges for topK are 0 to 500, with a default value of 250.

- **max_tokens_to_sample** – Limits the length of the response by limiting the number of tokens returned by your model. Ranges for max_tokens_to_sample are 0 to 4096, with a default value of 200.

- **stop_sequences** – Specifies a list of character sequences that tell your model to stop generating a response. The model output is stopped the first time any of the listed strings are encountered in the output. The response does not contain the stop sequence. For
example, you can use a carriage return sequence to limit the model response to a single line. You can configure up to 4 stop sequences.

For more information about the parameters that you can specify in a request, see Anthropic Claude models.

Set up FMEval

1. Load the required libraries to run FMEval as follows:

```python
from fmeval.data_loaders.data_config import DataConfig
from fmeval.model_runners.bedrock_model_runner import BedrockModelRunner
from fmeval.constants import MIME_TYPE_JSONLINES
from fmeval.eval_algorithms.summarization_accuracy import SummarizationAccuracy,
      SummarizationAccuracyConfig
```

2. Set up the data configuration for your input dataset.

The following sample input is one line from sample-dataset.jsonl:

```json
{
   "document": "23 October 2015 Last updated at 17:44
   BST\nIt's the highest rating a tropical storm can get and is the first one of this magnitude to hit mainland Mexico since 1959.\nBut how are the categories decided and what do they mean? Newsround reporter Jenny Lawrence explains.",
   "summary": "Hurricane Patricia has been rated as a category 5 storm.",
   "id": "34615665",
}
```

The previous sample input contains the text to summarize inside the document key. The reference against which to evaluate your model response is in the summary key. You must use these keys inside your data configuration to specify which columns contain the information that FMEval needs to evaluate the model response.

Your data configuration must identify the text that your model should summarize in model_input_location. You must identify the reference value with target_output_location.
The following data configuration example refers to the previous input example to specify the columns required for a text summarization task, the name, uniform resource identifier (URI), and MIME type:

```python
config = DataConfig(
    dataset_name="sample-dataset",
    dataset_uri="sample-dataset.jsonl",
    dataset_mime_type=MIME_TYPE_JSONLINES,
    model_input_location="document",
    target_output_location="summary"
)
```

For more information about the column information required for other tasks, see the **Use a custom input dataset** section in **Use an automatic evaluation**.

3. Set up a custom ModelRunner as shown in the following code example:

```python
bedrock_model_runner = BedrockModelRunner(
    model_id=model_id,
    output='completion',
    content_template='{"prompt": $prompt, "max_tokens_to_sample": 500}'
)
```

The previous code example specifies the following:

- **model_id** – The id used to specify your model.
- **output** – Captures the output from the Anthropic Claude 2 model, which returns its response in a completion key.
- **content_template** – Specifies how your model interacts with requests. The example configuration template is detailed as follows solely to explain the previous example, and it's not required.
  - In the previous `content_template` example, the following apply:
    - The variable `prompt` specifies the input prompt, which captures the request made by the user.
    - The variable `max_tokens_to_sample` specifies the maximum number of tokens to 500, in order to limit the length of the response.
For more information about the parameters that you can specify in your request, see Anthropic Claude models.

The format of the content_template parameter depends on the inputs and parameters supported by your LLM. In this tutorial, Anthropic’s Claude 2 model uses the following content_template:

```
"content_template": "{"prompt": $prompt, "max_tokens_to_sample": 500}"
```

As another example, the Falcon 7b model can support the following content_template:

```
"content_template": "{"inputs": $prompt, "parameters":{"max_new_tokens": 10, "top_p": 0.9, "temperature": 0.8}}"
```

Run your model evaluation

Define and run your evaluation algorithm

1. Define your evaluation algorithm. The following example shows how to define a SummarizationAccuracy algorithm, which is used to determine accuracy for text summarization tasks:

   ```python
   eval_algo = SummarizationAccuracy(SummarizationAccuracyConfig())
   ```

   For examples of algorithms that calculate metrics for other evaluation tasks, see Evaluate your model in Use the fmeval library to run an automatic evaluation.

2. Run your evaluation algorithm. The following code example uses the data configuration that was previously defined, and a prompt_template that uses the Human and Assistant keys:

   ```python
   eval_output = eval_algo.evaluate(model=bedrock_model_runner, dataset_config=config, prompt_template="Human: $feature

Assistant:
", save=True)
   ```

   In the previous code example, feature contains the prompt in the format that Amazon Bedrock model expects.
View your analysis results

1. Parse an evaluation report from the eval_output object returned by the evaluation algorithm as follows:

```python
# parse report
print(json.dumps(eval_output, default=vars, indent=4))
```

The previous command returns the following output:

```json
[
  {
    "eval_name": "summarization_accuracy",
    "dataset_name": "sample-dataset",
    "dataset_scores": [
      {
        "name": "meteor",
        "value": 0.2048823008681274
      },
      {
        "name": "rouge",
        "value": 0.03557697913367101
      },
      {
        "name": "bertscore",
        "value": 0.5406564395678671
      }
    ],
    "prompt_template": "Human: $feature\n\nAssistant:\n",
    "category_scores": null,
    "output_path": "/tmp/eval_results/summarization_accuracy_sample_dataset.jsonl",
    "error": null
  }
]
```

The previous example output displays the three accuracy scores: Meteor, Rouge, and BERTScore, the input prompt_template, a category_score if you requested one, any errors, and the output_path. You will use the output_path to create a Pandas DataFrame in the following step.
2. Import your results and read them into a DataFrame, and attach the accuracy scores to the model input, model output, and target output as follows:

```python
import pandas as pd

data = []
with open("/tmp/eval_results/summarization_accuracy_sample_dataset.jsonl", "r") as file:
    for line in file:
        data.append(json.loads(line))
df = pd.DataFrame(data)
df['meteor_score'] = df['scores'].apply(lambda x: x[0]['value'])
df['rouge_score'] = df['scores'].apply(lambda x: x[1]['value'])
df['bert_score'] = df['scores'].apply(lambda x: x[2]['value'])
df
```

In this invocation, the previous code example returns the following output (contracted for brevity):

<table>
<thead>
<tr>
<th>model_input</th>
<th>model_output</th>
<th>target_output</th>
<th>prompt</th>
<th>scores</th>
</tr>
</thead>
<tbody>
<tr>
<td>John Edward Bates, formerly of Spalding, Lincoln...</td>
<td>I cannot make any definitive judgments, as th...</td>
<td>Human: John Edward Bates, formerly of Spalding...</td>
<td>[{'name': 'meteor', 'value': 0.112359550561797...</td>
<td>0.112360 0.000000 0.543234 ...</td>
</tr>
</tbody>
</table>
| 23 October 2015 Last updated at 17:44 BST
It'... | Here are some key points about hurricane/trop... | Hurricane Patricia has been rated as a category... | Human: 23 October 2015 Last updated at 17:44 B... | {'name': 'meteor', 'value': 0.139822692925566... | 0.139823 0.017621 0.426529 ... |
| Ferrari appeared in a position to challenge un... | Here are the key points from the article:

Lewis Hamilton stormed to pole position at the... | Human: Ferrari appeared in a position to chall... | [{'name': 'meteor', 'value': 0.28341142234671... | 0.283411 0.064516 0.597001 ... |
| The Bath-born player, 28, has made 36 appearan... | Okay, let me summarize the key points from th... | Newport Gwent Dragons number eight Ed Jackson ... | Human: The Bath-born player, 28, has made 36 a... | [{'name': 'meteor', 'value': 0.089020771513353... | 0.089021 0.000000 0.533514 ... |

Your model output may be different than the previous sample output.
For a notebook that contains the code examples given in this section, see `bedrock-claude-summarization-accuracy.ipynb`.

**Additional notebooks**

The [fmeval GitHub](https://github.com/aws-sagemaker/fmeval) directory contains the following additional example notebooks:

- `byo-model-outputs.ipynb` – Evaluates a [Falcon 7b model](https://sagemaker.aws) hosted on SageMaker JumpStart for factual knowledge where you bring your own model outputs instead of sending inference requests to your model.
- `custom_model_runner_chat_gpt.ipynb` – Evaluates a custom ChatGPT 3.5 model hosted on Hugging Face for factual knowledge.

**FMEval Troubleshooting guide**

⚠️ **Important**

In order to use SageMaker Clarify Foundation Model Evaluations (FMEval), you must upgrade to the new Studio experience.

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. FMEval isn't available in Amazon SageMaker Studio Classic.

For information about how to upgrade to the new Studio experience, see [Migrating from Amazon SageMaker Studio Classic](https://docs.aws.amazon.com/sagemaker/latest/dg/studio-upgrade.html). For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](https://docs.aws.amazon.com/sagemaker/latest/dg/studio-classic.html).

If you run into an error while creating a foundation model evaluation, use the following list to troubleshoot your evaluation. If you need further assistance, contact [AWS Support](https://aws.amazon.com/support) or [AWS Developer Forums for Amazon SageMaker](https://aws.amazon.com/forums).

**Topics**

- Error uploading your data from an Amazon S3 bucket
The processing job failed to complete

You can't find foundation model evaluations in the SageMaker console

Your model does not support prompt stereotyping

Error uploading your data from an Amazon S3 bucket

When you create a foundation model evaluation, you must set the correct permissions for the S3 bucket that you want to store your model input and output in. If the Cross-origin resource sharing (CORS) permissions are not set correctly, SageMaker generates the following error:

Error: Failed to put object in s3: Error while uploading object to s3
Error: Failed to put object in S3: NetworkError when attempting to fetch resource.

To set the correct bucket permissions, follow the instructions under Set up your environment in Creating an automatic model evaluation job in Studio.

The processing job failed to complete

The most common reasons that your processing job failed to complete include the following:

- Insufficient quota
- Insufficient memory
- Did not pass ping check

See the following sections to help you mitigate each issue.

Insufficient quota

When you run a foundation model evaluation for a non-deployed SageMaker JumpStart model, SageMaker Clarify deploys your large language model (LLM) to a SageMaker endpoint in your account. If your account does not have sufficient quota to run the selected SageMaker JumpStart model, the job fails with a ClientError. To increase your quota, follow these steps:

Request an AWS Service Quotas increase

1. Retrieve the instance name, current quota and necessary quota from the on screen error message. For example, in the following error:

   - The instance name is ml.g5.12xlarge.
- The current quota from the number following current utilization is 0 instances.
- The additional required quota from the number following request delta is 1 instances.

The sample error follows:

ClientError: An error occurred (ResourceLimitExceeded) when calling the CreateEndpoint operation: The account-level service limit 'ml.g5.12xlarge for endpoint usage' is 0 Instances, with current utilization of 0 Instances and a request delta of 1 Instances. Please use AWS Service Quotas to request an increase for this quota. If AWS Service Quotas is not available, contact AWS support to request an increase for this quota.

2. Sign into the AWS Management Console and open the Service Quotas console.
3. In the navigation pane, under Manage quotas, input Amazon SageMaker.
4. Choose View quotas.
5. In the search bar under Service quotas, input the name of the instance from Step 1. For example, using the information contained in the error message from Step 1, input ml.g5.12xlarge.
6. Choose the Quota name that appears next to your instance name and ends with for endpoint usage. For example, using the information contained in the error message from Step 1, choose ml.g5.12xlarge for endpoint usage.
7. Choose Request increase at account-level.
8. Under Increase quota value, input the necessary required quota from the information given in the error message from Step 1. Input the total of current utilization and request delta. In the previous example error, the current utilization is 0 Instances, and the request delta is 1 Instances. In this example, request a quota of 1 to supply the required quota.
9. Choose Request.
10. Choose Quota request history from the navigation pane.
11. When the Status changes from Pending to Approved, rerun your job. You may need to refresh your browser to see the change.

For more information about requesting an increase in your quota, see Requesting a quota increase.
Insufficient memory

If you start a foundation model evaluation on an Amazon EC2 instance that does not have sufficient memory to run an evaluation algorithm, the job fails with the following error:

The actor is dead because its worker process has died. Worker exit type: SYSTEM_ERROR Worker exit detail: Worker unexpectedly exits with a connection error code 2. End of file. There are some potential root causes. (1) The process is killed by SIGKILL by OOM killer due to high memory usage. (2) ray stop --force is called. (3) The worker is crashed unexpectedly due to SIGSEGV or other unexpected errors. The actor never ran - it was cancelled before it started running.

To increase the memory available for your evaluation job, change your instance to one that has more memory. If you are using the user interface, you can choose an instance type under Processor configuration in Step 2. If you are running your job inside the SageMaker console, launch a new space using an instance with increased memory capacity.

For a list of Amazon EC2 instances, see Instance types.

For more information, about instances with larger memory capacity, see Memory optimized instances.

Did not pass ping check

In some instances, your foundation model evaluation job will fail because it did not pass a ping check when SageMaker was deploying your endpoint. If it does not pass a ping test, the following error appears:

ClientError: Error hosting endpoint your_endpoint_name: Failed. Reason: The primary container for production variant AllTraffic did not pass the ping health check. Please check CloudWatch logs for this endpoint..., Job exited for model: your_model_name of model_type: your_model_type

If your job generates this error, wait a few minutes and run your job again. If the error persists, contact AWS Support or AWS Developer Forums for Amazon SageMaker.

You can't find foundation model evaluations in the SageMaker console

In order to use SageMaker Clarify Foundation Model Evaluations, you must upgrade to the new Studio experience. As of November 30, 2023, the previous Amazon SageMaker Studio experience
is now named Amazon SageMaker Studio Classic. The foundation evaluation feature can only be used in the updated experience. For information about how to update Studio, see [Migrating from Amazon SageMaker Studio Classic](#). For information about using the Studio Classic application, see [Amazon SageMaker Studio Classic](#).

**Your model does not support prompt stereotyping**

Only some SageMaker JumpStart models support prompt stereotyping. If you select a SageMaker JumpStart model that is not supported, the following error appears:

{"evaluationMetrics":"This model does not support Prompt stereotyping evaluation. Please remove that evaluation metric or select another model that supports it."}

If you receive this error, you cannot use your selected model in a foundation evaluation. SageMaker Clarify is currently working to update all SageMaker JumpStart models for prompt stereotyping tasks so that they can be used in a foundation model evaluation.

**Dataset validation errors**

When you start a model evaluation job each JSON object in the prompt dataset is independently validated. For a custom prompt dataset to pass all validations the following must be true.

- Each line in the prompt dataset file must be a valid JSON object.
- A prompt dataset file cannot contain more than 1,000 JSON objects in a single file.
- If you specify the `responses` key is present *any* JSON object, it must be present in *all* JSON objects.
- All JSON objects must contain at least the `prompt` key/value pair.

**Use SageMaker Clarify to explain and detect bias**

This topic describes how to understand fairness and model explainability and how to explain and detect bias using Amazon SageMaker Clarify. You can configure an SageMaker Clarify processing job to compute bias metrics and feature attributions and generate reports for model explainability. SageMaker Clarify processing jobs are implemented using a specialized SageMaker Clarify container image. The following instructions show you how to configure, run, and troubleshoot a SageMaker Clarify processing job and how to configure an analysis.
What is fairness and model explainability for machine learning predictions?

Machine learning (ML) models are helping make decisions in domains including financial services, healthcare, education, and human resources. Policymakers, regulators, and advocates have raised awareness about the ethical and policy challenges posed by ML and data-driven systems. Amazon SageMaker Clarify can help you understand why your ML model made a specific prediction and whether this bias impacts this prediction during training or inference. SageMaker Clarify also provides tools that can help you build less biased and more understandable machine learning models. SageMaker Clarify can also generate model governance reports that you can provide to risk and compliance teams and external regulators. With SageMaker Clarify, you can do the following:

- Detect bias in and help explain your model predictions.
- Identify types of bias in pre-training data.
- Identify types of bias in post-training data that can emerge during training or when your model is in production.

SageMaker Clarify helps explain how your models make predictions using feature attributions. It can also monitor inference models that are in production for both bias and feature attribution drift. This information can help you in the following areas:

- **Regulatory** – Policymakers and other regulators can have concerns about discriminatory impacts of decisions that use output from ML models. For example, an ML model may encode bias and influence an automated decision.
- **Business** – Regulated domains may need reliable explanations for how ML models make predictions. Model explainability may be particularly important to industries that depend on reliability, safety, and compliance. These can include financial services, human resources, healthcare, and automated transportation. For example, lending applications may need to provide explanations about how ML models made certain predictions to loan officers, forecasters, and customers.
- **Data Science** – Data scientists and ML engineers can debug and improve ML models when they can determine if a model is making inferences based on noisy or irrelevant features. They can also understand the limitations of their models and failure modes that their models may encounter.
For a blog post that shows how to architect and build a complete machine learning model for fraudulent automobile claims that integrates SageMaker Clarify into a SageMaker pipeline, see the **Architect and build the full machine learning lifecycle with AWS: An end-to-end Amazon SageMaker** demo. This blog post discusses how to assess and mitigate pre-training and post-training bias, and how the features impact the model prediction. The blog post contains links to example code for each task in the ML lifecycle.

**Best practices to evaluate fairness and explainability in the ML lifecycle**

**Fairness as a process** – Notions of bias and fairness depend on their application. The measurement of bias and the choice of the bias metrics may be guided by social, legal, and other non-technical considerations. The successful adoption of fairness-aware ML approaches includes building consensus and achieving collaboration across key stakeholders. These may include product, policy, legal, engineering, AI/ML teams, end users, and communities.

**Fairness and explainability by design in the ML lifecycle** – Consider fairness and explainability during each stage of the ML lifecycle. These stages include problem formation, dataset construction, algorithm selection, the model training process, the testing process, deployment, and monitoring and feedback. It is important to have the right tools to do this analysis. We recommend asking the following questions during the ML lifecycle:

- Does the model encourage feedback loops that can produce increasingly unfair outcomes?
- Is an algorithm an ethical solution to the problem?
- Is the training data representative of different groups?
- Are there biases in labels or features?
- Does the data need to be modified to mitigate bias?
- Do fairness constraints need to be included in the objective function?
- Has the model been evaluated using relevant fairness metrics?
- Are there unequal effects across users?
- Is the model deployed on a population for which it was not trained or evaluated?
Guide to the SageMaker explanations and bias documentation

Bias can occur and be measured in the data both before and after training a model. SageMaker Clarify can provide explanations for model predictions after training and for models deployed to production. SageMaker Clarify can also monitor models in production for any drift in their baseline explanatory attributions, and calculate baselines when needed. The documentation for explaining and detecting bias using SageMaker Clarify is structured as follows:

- For information on setting up a processing job for bias and explainability, see Configure a SageMaker Clarify Processing Job.
- For information on detecting bias in pre-processing data before it's used to train a model, see Detect Pre-training Data Bias.
- For information on detecting post-training data and model bias, see Detect Post-training Data and Model Bias.
- For information on the model-agnostic feature attribution approach to explain model predictions after training, see Model Explainability.
- For information on monitoring for feature contribution drift away from the baseline that was established during model training, see Monitor Feature Attribution Drift for Models in Production.
- For information about monitoring models that are in production for baseline drift, see Monitor Bias Drift for Models in Production.
• For information about obtaining explanations in real time from a SageMaker endpoint, see Online Explainability with SageMaker Clarify.

How SageMaker Clarify Processing Jobs Work

You can use SageMaker Clarify to analyze your datasets and models for explainability and bias. A SageMaker Clarify processing job uses the SageMaker Clarify processing container to interact with an Amazon S3 bucket containing your input datasets. You can also use SageMaker Clarify to analyze a customer model that is deployed to a SageMaker inference endpoint.

The following graphic shows how a SageMaker Clarify processing job interacts with your input data and optionally, with a customer model. This interaction depends on the specific type of analysis being performed. The SageMaker Clarify processing container obtains the input dataset and configuration for analysis from an S3 bucket. For certain analysis types, including feature analysis, the SageMaker Clarify processing container must send requests to the model container. Then it retrieves the model predictions from the response that the model container sends. After that, the SageMaker Clarify processing container computes and saves analysis results to the S3 bucket.

You can run a SageMaker Clarify processing job at multiple stages in the lifecycle of the machine learning workflow. SageMaker Clarify can help you compute the following analysis types:

• Pre-training bias metrics. These metrics can help you understand the bias in your data so that you can address it and train your model on a more fair dataset. See Measure Pre-training Bias for information about pre-training bias metrics. To run a job to analyze pre-training bias metrics, you must provide the dataset and a JSON analysis configuration file to Configure the Analysis.

• Post-training bias metrics. These metrics can help you understand any bias introduced by an algorithm, hyperparameter choices, or any bias that wasn't apparent earlier in the flow. For more information about post-training bias metrics, see Measure Post-training Data and Model Bias.
SageMaker Clarify uses the model predictions in addition to the data and labels to identify bias. To run a job to analyze post-training bias metrics, you must provide the dataset and a JSON analysis configuration file. The configuration should include the model or endpoint name.

- Shapely values, which can help you understand what impact your feature has on what your model predicts. For more information about Shapely values, see Feature Attributions that Use Shapley Values. This feature requires a trained model.

- Partial dependence plots (PDPs), which can help you understand how much your predicted target variable would change if you varied the value of one feature. For more information about PDPs, see Partial dependence plots (PDPs) analysis. This feature requires a trained model.

SageMaker Clarify needs model predictions to compute post-training bias metrics and feature attributions. You can provide an endpoint or SageMaker Clarify will create an ephemeral endpoint using your model name, also known as a shadow endpoint. The SageMaker Clarify container deletes the shadow endpoint after the computations are completed. At a high level, the SageMaker Clarify container completes the following steps:

1. Validates inputs and parameters.
2. Creates the shadow endpoint (if a model name is provided).
3. Loads the input dataset into a data frame.
4. Obtains model predictions from the endpoint, if necessary.
5. Computes bias metrics and features attributions.
6. Deletes the shadow endpoint.
7. Generate the analysis results.

After the SageMaker Clarify processing job is complete, the analysis results will be saved in the output location that you specified in the processing output parameter of the job. These results include a JSON file with bias metrics and global feature attributions, a visual report, and additional files for local feature attributions. You can download the results from the output location and view them.

For additional information about bias metrics, explainability and how to interpret them, see Learn How Amazon SageMaker Clarify Helps Detect Bias, Fairness Measures for Machine Learning in Finance, and the Amazon AI Fairness and Explainability Whitepaper.
Configure a SageMaker Clarify Processing Job

To analyze your data and models for bias and explainability using SageMaker Clarify, you must configure a SageMaker Clarify processing job. This guide shows how to specify the input dataset name, analysis configuration file name, and output location for a processing job. To configure the processing container, job inputs, outputs, resources and other parameters, you have two options. You can either use the SageMaker CreateProcessingJob API, or use the SageMaker Python SDK API SageMaker ClarifyProcessor.

For information about parameters that are common to all processing jobs, see Amazon SageMaker API Reference.

Configure a SageMaker Clarify processing job using the SageMaker API

The following instructions show how to provide each portion of the SageMaker Clarify specific configuration using the CreateProcessingJob API.

1. Input the uniform research identifier (URI) of a SageMaker Clarify container image inside the AppSpecification parameter, as shown in the following code example.

```json
{
   "ImageUri": "the-clarify-container-image-uri"
}
```

**Note**

The URI must identify a pre-built SageMaker Clarify container image. ContainerEntrypoint and ContainerArguments are not supported. For more information about SageMaker Clarify container images, see Get Started with a SageMaker Clarify Container.

2. Specify both the configuration for your analysis and parameters for your input dataset inside the ProcessingInputs parameter.

   a. Specify the location of the JSON analysis configuration file, which includes the parameters for bias analysis and explainability analysis. The InputName parameter of the ProcessingInput object must be `analysis_config` as shown in the following code example.

```json
{
}
```
For more information about the schema of the analysis configuration file, see Configure the Analysis.

b. Specify the location of the input dataset. The InputName parameter of the ProcessingInput object must be dataset. This parameter is optional if you have provided the "dataset_uri" in the analysis configuration file. The following values are required in the S3Input configuration.

i. S3Uri can be either an Amazon S3 object or an S3 prefix.

ii. S3InputMode must be of type File.

iii. S3CompressionType must be of type None (the default value).

iv. S3DataDistributionType must be of type FullyReplicated (the default value).

v. S3DataType can be either S3Prefix or ManifestFile. To use ManifestFile, the S3Uri parameter should specify the location of a manifest file that follows the schema from the SageMaker API Reference section S3Uri. This manifest file must list the S3 objects that contain the input data for the job.

The following code shows an example of an input configuration.

```
{
    "InputName": "dataset",
    "S3Input": {
        "S3Uri": "s3://your-bucket/your-dataset.csv",
        "S3DataType": "S3Prefix",
        "S3InputMode": "File",
        "LocalPath": "/opt/ml/processing/input/data"
    }
}
```

3. Specify the configuration for the output of the processing job inside the ProcessingOutputConfig parameter. A single ProcessingOutput object is required in the Outputs configuration. The following are required of the output configuration:
a. **OutputName** must be `analysis_result`.

b. **S3Uri** must be an S3 prefix to the output location.

c. **S3UploadMode** must be set to **EndOfJob**.

The following code shows an example of an output configuration.

```json
{
    "Outputs": [{
      "OutputName": "analysis_result",
      "S3Output": {
        "S3Uri": "s3://your-bucket/result/",
        "S3UploadMode": "EndOfJob",
        "LocalPath": "/opt/ml/processing/output"
      }
    }]
}
```

4. Specify the configuration **ClusterConfig** for the resources that you use in your processing job inside the **ProcessingResources** parameter. The following parameters are required inside the **ClusterConfig** object.

a. **InstanceCount** specifies the number of compute instances in the cluster that runs the processing job. Specify a value greater than 1 to activate distributed processing.

b. **InstanceType** refers to the resources that runs your processing job. Because SageMaker SHAP analysis is compute-intensive, using an instance type that is optimized for compute should improve runtime for analysis. The SageMaker Clarify processing job doesn't use GPUs.

The following code shows an example of resource configuration.

```json
{
    "ClusterConfig": {
        "InstanceCount": 1,
        "InstanceType": "ml.m5.xlarge",
        "VolumeSizeInGB": 20
    }
}
```

5. Specify the configuration of the network that you use in your processing job inside the **NetworkConfig** object. The following values are required in the configuration.
a. EnableNetworkIsolation must be set to False (default) so that SageMaker Clarify can invoke an endpoint, if necessary, for predictions.

b. If the model or endpoint that you provided to the SageMaker Clarify job is within an Amazon Virtual Private Cloud (Amazon VPC), then the SageMaker Clarify job must also be in the same VPC. Specify the VPC using VpcConfig. Additionally, the VPC must have endpoints to an Amazon S3 bucket, SageMaker service and SageMaker Runtime service.

If distributed processing is activated, you must also allow communication between different instances in the same processing job. Configure a rule for your security group that allows inbound connections between members of the same security group. For more information, see Give Amazon SageMaker Clarify Jobs Access to Resources in Your Amazon VPC.

The following code gives an example of a network configuration.

```json
{
    "EnableNetworkIsolation": False,
    "VpcConfig": {
        ...
    }
}
```

6. Set the maximum time that the job will run using the StoppingCondition parameter. The longest that a SageMaker Clarify job can run is 7 days or 604800 seconds. If the job cannot be completed within this time limit, it will be stopped and no analysis results will be provided. As an example, the following configuration limits the maximum time that the job can run to 3600 seconds.

```json
{
    "MaxRuntimeInSeconds": 3600
}
```

7. Specify an IAM role for the RoleArn parameter. The role must have a trust relationship with Amazon SageMaker. It can be used to perform the SageMaker API operations listed in the following table. We recommend using the Amazon SageMakerFullAccess managed policy, which grants full access to SageMaker. For more information on this policy, see AWS managed policy: AmazonSageMakerFullAccess. If you have concerns about granting full access, the minimal permissions required depend on whether you provide a model or an endpoint name. Using an endpoint name allows for granting fewer permissions to SageMaker.
The following table contains API operations used by the SageMaker Clarify processing job. An X under **Model name** and **Endpoint name** notes the API operation that is required for each input.

<table>
<thead>
<tr>
<th>API Operation</th>
<th>Model name</th>
<th>Endpoint name</th>
<th>What is it used for</th>
</tr>
</thead>
<tbody>
<tr>
<td>ListTags</td>
<td>X</td>
<td></td>
<td>Tags of the job are applied to the shadow endpoint.</td>
</tr>
<tr>
<td>CreateEndpointConfig</td>
<td>X</td>
<td></td>
<td>Create endpoint config using the model name that you provided</td>
</tr>
<tr>
<td>CreateEndpoint</td>
<td>X</td>
<td></td>
<td>Create shadow endpoint using the endpoint config.</td>
</tr>
<tr>
<td>DescribeEndpoint</td>
<td>X</td>
<td>X</td>
<td>Describe endpoint for its status, the endpoint must be InService to serve requests.</td>
</tr>
<tr>
<td>InvokeEndpoint</td>
<td>X</td>
<td>X</td>
<td>Invoke the endpoint for predictions.</td>
</tr>
</tbody>
</table>

For more information about required permissions, see [Amazon SageMaker API Permissions: Actions, Permissions, and Resources Reference](#). For more information about passing roles to SageMaker, see [Passing Roles](#).

After you have the individual pieces of the processing job configuration, combine them to configure the job.
Configure a SageMaker Clarify processing job using the AWS SDK for Python

The following code example shows how to launch a SageMaker Clarify processing job using the AWS SDK for Python.

```python
sagemaker_client.create_processing_job(
    ProcessingJobName="your-clarify-job-name",
    AppSpecification={
        "ImageUri": "the-clarify-container-image-uri",
    },
    ProcessingInputs=[{
        "InputName": "analysis_config",
        "S3Input": {
            "S3Uri": "s3://your-bucket/analysis_config.json",
            "S3DataType": "S3Prefix",
            "S3InputMode": "File",
            "LocalPath": "/opt/ml/processing/input/config",
        },
    }, {
        "InputName": "dataset",
        "S3Input": {
            "S3Uri": "s3://your-bucket/your-dataset.csv",
            "S3DataType": "S3Prefix",
            "S3InputMode": "File",
            "LocalPath": "/opt/ml/processing/input/data",
        },
    },
],
    ProcessingOutputConfig={
        "Outputs": [{
            "OutputName": "analysis_result",
            "S3Output": {
                "S3Uri": "s3://your-bucket/result/",
                "S3UploadMode": "EndOfJob",
                "LocalPath": "/opt/ml/processing/output",
            },
        }],
    },
    ProcessingResources={
        "ClusterConfig": {
            "InstanceCount": 1,
            "InstanceType": "ml.m5.xlarge",
            "VolumeSizeInGB": 20,
        },
    }
)
For an example notebook with instructions for running a SageMaker Clarify processing job using AWS SDK for Python, see Fairness and Explainability with SageMaker Clarify using AWS SDK for Python. Any S3 bucket used in the notebook must be in the same AWS Region as the notebook instance that accesses it.

**Configure a SageMaker Clarify processing job using the SageMaker Python SDK**

You can also configure a SageMaker Clarify processing job using the SageMaker ClarifyProcessor in the SageMaker Python SDK API. For more information, see Run SageMaker Clarify Processing Jobs for Bias Analysis and Explainability.

**Topics**

- Get Started with a SageMaker Clarify Container
- Configure the Analysis
- Data Format Compatibility Guide

**Get Started with a SageMaker Clarify Container**

Amazon SageMaker provides prebuilt SageMaker Clarify container images that include the libraries and other dependencies needed to compute bias metrics and feature attributions for explainability. This image has been enabled to run SageMaker Process data in your account.

The image URIs for the containers are in the following form:

```
<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/sagemaker-clarify-processing:1.0
```
For example:

```
205585389593.dkr.ecr.us-east-1.amazonaws.com/sagemaker-clarify-processing:1.0
```

The following table lists the addresses by AWS Region.

### Docker Images for SageMaker Clarify Processing Jobs

<table>
<thead>
<tr>
<th>Region</th>
<th>Image address</th>
</tr>
</thead>
<tbody>
<tr>
<td>us-east-1</td>
<td>205585389593.dkr.ecr.us-east-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>us-east-2</td>
<td>211330385671.dkr.ecr.us-east-2.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>us-west-1</td>
<td>740489534195.dkr.ecr.us-west-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>us-west-2</td>
<td>306415355426.dkr.ecr.us-west-2.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-east-1</td>
<td>098760798382.dkr.ecr.ap-east-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-south-1</td>
<td>452307495513.dkr.ecr.ap-south-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-southeast-3</td>
<td>705930551576.dkr.ecr.ap-southeast-3.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-northeast-1</td>
<td>377024640650.dkr.ecr.ap-northeast-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-northeast-2</td>
<td>263625296855.dkr.ecr.ap-northeast-2.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-northeast-3</td>
<td>912233562940.dkr.ecr.ap-northeast-3.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>Region</td>
<td>Image address</td>
</tr>
<tr>
<td>------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>ap-southeast-1</td>
<td>834264404009.dkr.ecr.ap-southeast-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ap-southeast-2</td>
<td>007051062584.dkr.ecr.ap-southeast-2.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>ca-central-1</td>
<td>675030665977.dkr.ecr.ca-central-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
</tr>
<tr>
<td>eu-central-1</td>
<td>017069133835.dkr.ecr.eu-central-1.amazonaws.com/sagemaker-clarify-processing:1.0</td>
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Configure the Analysis

To analyze your data and models for explainability and bias using SageMaker Clarify, you must configure a processing job. Part of the configuration for this processing job includes the configuration of an analysis file. The analysis file specifies the parameters for bias analysis and explainability. See Configure a SageMaker Clarify Processing Job to learn how to configure a processing job and analysis file.

This guide describes the schema and parameters for this analysis configuration file. This guide also includes examples of analysis configuration files for computing bias metrics for a tabular dataset, and generating explanations for natural language processing (NLP) and computer vision (CV) problems.

You can create the analysis configuration file or use the SageMaker Python SDK to generate one for you with the SageMaker ClarifyProcessor API. Viewing the file contents can be helpful for understanding the underlying configuration used by the SageMaker Clarify job.

Topics

- Schema for the analysis configuration file
- Example analysis configuration files

Schema for the analysis configuration file

The following section describes the schema for the analysis configuration file including requirements and descriptions of parameters.

Requirements for the analysis configuration file

The SageMaker Clarify processing job expects the analysis configuration file to be structured with the following requirements:

- The processing input name must be analysis_config.
• The analysis configuration file is in JSON format, and encoded in UTF-8.
• The analysis configuration file is an Amazon S3 object.

You can specify additional parameters in the analysis configuration file. The following section provides various options to tailor the SageMaker Clarify processing job for your use case and desired types of analysis.

Parameters for analysis configuration files

In the analysis configuration file, you can specify the following parameters.

• version – (Optional) The version string of the analysis configuration file schema. If a version is not provided, SageMaker Clarify uses the latest supported version. Currently, the only supported version is 1.0.
• dataset_type – The format of the dataset. The input dataset format can be any of the following values:
  • text/csv for CSV
  • application/jsonlines for SageMaker JSON Lines dense format
  • application/json for JSON
  • application/x-parquet for Apache Parquet
  • application/x-image to activate explainability for computer vision problems
• dataset_uri – (Optional) The uniform resource identifier (URI) of the main dataset. If you provide a S3 URI prefix, the SageMaker Clarify processing job recursively collects all S3 files located under the prefix. You can provide either a S3 URI prefix or a S3 URI to an image manifest file for computer vision problems. If dataset_uri is provided, it takes precedence over the dataset processing job input. For any format type except image, the SageMaker Clarify processing job loads the input dataset into a tabular data frame, as a tabular dataset. This format allows SageMaker to easily manipulate and analyze the input dataset.
• headers – (Optional) An array of strings containing the column names of a tabular dataset. If a value for headers is not provided, then the SageMaker Clarify processing job reads the headers from the dataset. If the dataset doesn’t have headers, then the SageMaker Clarify processing job will automatically generate placeholder names based on zero-based column index. As an example, placeholder names for the first and second columns will be column_0, column_1.
By convention, if dataset_type is application/jsonlines or application/json then headers should contain the following names in order: feature names, label name (if label is specified), and predicted label name (if predicted_label is specified). An example for headers for an application/jsonlines dataset type if label is specified is: ["feature1", "feature2", "feature3", "target_label"].

- **label** – (Optional) A string or a zero-based integer index. If provided, label is used to locate the ground truth label, also known as an observed label or target attribute in a tabular dataset. The ground truth label is used to compute bias metrics. The value for label is specified depending on the value of the dataset_type parameter as follows:
  - If dataset_type is **text/csv**, label can be specified as either of the following:
    - A valid column name
    - An index that lies within the range of dataset columns
  - If dataset_type is **application/parquet**, label must be a valid column name.
  - If dataset_type is **application/jsonlines**, label must be a JMESPath expression written to extract the ground truth label from the dataset. By convention, if headers is specified, then it should contain the label name.
  - If dataset_type is **application/json**, label must be a JMESPath expression written to extract the ground truth label for each record in the dataset. This JMESPath expression must produce a list of labels where the i\text{th} label correlates to the i\text{th} record.

- **predicted_label** – (Optional) A string or a zero-based integer index. If provided, predicted_label is used to locate the column containing the predicted label in a tabular dataset. The predicted label is used to compute post-training bias metrics. The parameter predicted_label is optional if the dataset doesn't include predicted label. If predicted labels are required for computation, then the SageMaker Clarify processing job will get predictions from the model.

  The value for predicted_label is specified depending on the value of the dataset_type as follows:
  - If dataset_type is **text/csv**, predicted_label can be specified as either of the following:
• A valid column name. If predicted_label_dataset_uri is specified, but predicted_label is not provided, the default predicted label name is "predicted_label".

• An index that lies within the range of dataset columns. If predicted_label_dataset_uri is specified, then the index is used to locate the predicted label column in the predicted label dataset.

• If dataset_type is application/x-parquet, predicted_label must be a valid column name.

• If dataset_type is application/jsonlines, predicted_label must be a valid JMESPath expression written to extract the predicted label from the dataset. By convention, if headers is specified, then it should contain the predicted label name.

• If dataset_type is application/json, predicted_label must be a JMESPath expression written to extract the predicted label for each record in the dataset. The JMESPath expression should produce a list of predicted labels where the i\textsuperscript{th} predicted label is for the i\textsuperscript{th} record.

• features – Required if dataset_type is application/jsonlines or application/json. A JMESPath string expression written to locate the features in the input dataset. For application/jsonlines, a JMESPath expression will be applied to each line to extract the features for that record. For application/json, a JMESPath expression will be applied to the whole input dataset. The JMESPath expression should extract a list of lists, or a 2D array/matrix of features where the i\textsuperscript{th} row contains the features that correlate to the i\textsuperscript{th} record. For a dataset_type of text/csv or application/x-parquet, all columns except for the ground truth label and predicted label columns are automatically assigned to be features.

• predicted_label_dataset_uri – Only applicable when dataset_type is text/csv. The S3 URI for a dataset containing predicted labels used to compute post-training bias metrics. The SageMaker Clarify processing job will load the predictions from the provided URI instead of getting predictions from the model. In this case, predicted_label is required to locate the predicted label column in the predicted label dataset. If the predicted label dataset or the main dataset is split across multiple files, an identifier column must be specified by joinsource_name_or_index to join the two datasets.

• predicted_label_headers – Only applicable when predicted_label_dataset_uri is specified. An array of strings containing the column names of the predicted label dataset. Besides the predicted label header, predicted_label_headers can also contain the header of the identifier column to join the predicted label dataset and the main dataset. For more information, see the following description for the parameter joinsource_name_or_index.
- **joinsource_name_or_index** – The name or zero-based index of the column in tabular datasets to be used as an identifier column while performing an inner join. This column is only used as an identifier. It isn't used for any other computations like bias analysis or feature attribution analysis. A value for **joinsource_name_or_index** is needed in the following cases:
  - There are multiple input datasets, and any one is split across multiple files.
  - Distributed processing is activated by setting the SageMaker Clarify processing job **InstanceCount** to a value greater than 1.

- **excluded_columns** – (Optional) An array of names or zero-based indices of columns to be excluded from being sent to the model as input for predictions. Ground truth label and predicted label are automatically excluded already.

- **probability_threshold** – (Optional) A floating point number above which, a label or object is selected. The default value is 0.5. The SageMaker Clarify processing job uses **probability_threshold** in the following cases:
  - In post-training bias analysis, **probability_threshold** converts a numeric model prediction (probability value or score) to a binary label, if the model is a binary classifier. A score greater than the threshold is converted to 1. Whereas, a score less than or equal to the threshold is converted to 0.
  - In computer vision explainability problems, if model_type is **OBJECT_DETECTION**, **probability_threshold** filters out objects detected with confidence scores lower than the threshold value.

- **label_values_or_threshold** – Required for bias analysis. An array of label values or a threshold number, which indicate positive outcome for ground truth and predicted labels for bias metrics. For more information, see positive label values in Amazon SageMaker Clarify Terms for Bias and Fairness. If the label is numeric, the threshold is applied as the lower bound to select the positive outcome. To set **label_values_or_threshold** for different problem types, refer to the following examples:
  - For a binary classification problem, the label has two possible values, 0 and 1. If label value 1 is favorable to a demographic group observed in a sample, then **label_values_or_threshold** should be set to [1].
  - For a multiclass classification problem, the label has three possible values, **bird**, **cat**, and **dog**. If the latter two define a demographic group that bias favors, then **label_values_or_threshold** should be set to ["cat", "dog"].
• For a regression problem, the label value is continuous, ranging from 0 to 1. If a value greater than 0.5 should designate a sample as having a positive result, then `label_values_or_threshold` should be set to 0.5.

• **facet** – Required for bias analysis. An array of facet objects, which are composed of sensitive attributes against which bias is measured. You can use facets to understand the bias characteristics of your dataset and model even if your model is trained without using sensitive attributes. For more information, see **Facet** in Amazon SageMaker Clarify Terms for Bias and Fairness. Each facet object includes the following fields:

  • **name_or_index** – The name or zero-based index of the sensitive attribute column in a tabular dataset. If `facet_dataset_uri` is specified, then the index refers to the facet dataset instead of the main dataset.

  • **value_or_threshold** – Required if `facet` is numeric and `label_values_or_threshold` is applied as the lower bound to select the sensitive group. An array of facet values or a threshold number, that indicates the sensitive demographic group that bias favors. If facet data type is categorical and `value_or_threshold` is not provided, bias metrics are computed as one group for every unique value (rather than all values). To set `value_or_threshold` for different facet data types, refer to the following examples:

    • For a binary facet data type, the feature has two possible values, 0 and 1. If you want to compute the bias metrics for each value, then `value_or_threshold` can be either omitted or set to an empty array.

    • For a categorical facet data type, the feature has three possible values `bird`, `cat`, and `dog`. If the first two define a demographic group that bias favors, then `value_or_threshold` should be set to `["bird", "cat"]`. In this example, the dataset samples are split into two demographic groups. The facet in the advantaged group has value `bird` or `cat`, while the facet in the disadvantaged group has value `dog`.

    • For a numeric facet data type, the feature value is continuous, ranging from 0 to 1. As an example, if a value greater than 0.5 should designate a sample as favored, then `value_or_threshold` should be set to 0.5. In this example, the dataset samples are split into two demographic groups. The facet in the advantaged group has value greater than 0.5, while the facet in the disadvantaged group has value less than or equal to 0.5.

• **group_variable** – The name or zero-based index of the column that indicates the subgroup to be used for the bias metric Conditional Demographic Disparity (CDD) or Conditional Demographic Disparity in Predicted Labels (CDDPL).
• **facet_dataset_uri** – Only applicable when dataset_type is text/csv. The S3 URI for a dataset containing sensitive attributes for bias analysis. You can use facets to understand the bias characteristics of your dataset and model even if your model is trained without using sensitive attributes.

**Note**
If the facet dataset or the main dataset is split across multiple files, an identifier column must be specified by joinsource_name_or_index to join the two datasets. You must use the parameter facet to identify each facet in the facet dataset.

• **facet_headers** – (Only applicable when facet_dataset_uri is specified) An array of strings containing column names for the facet dataset, and optionally, the identifier column header to join the facet dataset and the main dataset, see joinsource_name_or_index.

• **methods** – An object containing one or more analysis methods and their parameters. If any method is omitted, it is neither used for analysis nor reported.

• **pre_training_bias** – Include this method if you want to compute pre-training bias metrics. The detailed description of the metrics can be found in Measure Pre-training Bias. The object has the following parameters:

  • **methods** – An array that contains any of the pre-training bias metrics from the following list that you want to compute. Set methods to **all** to compute all pre-training bias metrics. As an example, the array ["CI", "DPL"] will compute **Class Imbalance** and **Difference in Proportions of Labels**.
    
    • **CI** for **Class Imbalance (CI)**
    • **DPL** for **Difference in Proportions of Labels (DPL)**
    • **KL** for **Kullback-Leibler Divergence (KL)**
    • **JS** for **Jensen-Shannon Divergence (JS)**
    • **LP** for **L_p-norm (LP)**
    • **TVD** for **Total Variation Distance (TVD)**
    • **KS** for **Kolmogorov-Smirnov (KS)**
    • **CDDL** for **Conditional Demographic Disparity (CDD)**

• **post_training_bias** – Include this method if you want to compute post-training bias metrics. The detailed description of the metrics can be found in Measure Post-training Data and Model Bias. The post_training_bias object has the following parameters.
• **methods** – An array that contains any of the post-training bias metrics from the following list that you want to compute. Set methods to `all` to compute all post-training bias metrics. As an example, the array `"DPPL", "DI"` computes the **Difference in Positive Proportions in Predicted Labels** and **Disparate Impact**. The available methods are as follows.

  - DPPL for **Difference in Positive Proportions in Predicted Labels (DPPL)**
  - DI for **Disparate Impact (DI)**
  - DCA for **Difference in Conditional Acceptance (DCAcc)**
  - DCR for **Difference in Conditional Rejection (DCR)**
  - SD for **Specificity difference (SD)**
  - RD for **Recall Difference (RD)**
  - DAR for **Difference in Acceptance Rates (DAR)**
  - DRR for **Difference in Rejection Rates (DRR)**
  - AD for **Accuracy Difference (AD)**
  - TE for **Treatment Equality (TE)**
  - CDDPL for **Conditional Demographic Disparity in Predicted Labels (CDDPL)**
  - FT for **Counterfactual Flitjest (FT)**
  - GE for **Generalized entropy (GE)**

• **shap** – Include this method if you want to compute SHAP values. The SageMaker Clarify processing job supports the Kernel SHAP algorithm. The shap object has the following parameters.

  - **baseline** – (Optional) The SHAP baseline dataset, also known as the background dataset. Additional requirements for the baseline dataset in a tabular dataset or computer vision problem are as follows. For more information about SHAP Baselines, see [SHAP Baselines for Explainability](#).

  - For a **tabular** dataset, baseline can be either the in-place baseline data or the S3 URI of a baseline file. If baseline is not provided, the SageMaker Clarify processing job computes a baseline by clustering the input dataset. The following are required of the baseline:
    - The format must be the same as the dataset format specified by `dataset_type`.
    - The baseline can only contain features that the model can accept as input.
• The baseline dataset can have one or more instances. The number of baseline instances directly affects the synthetic dataset size and job runtime.

• If text_config is specified, then the baseline value of a text column is a string used to replace the unit of text specified by granularity. For example, one common placeholder is "[MASK]", which is used to represent a missing or unknown word or piece of text.

The following examples show how to set in-place baseline data for different dataset_type parameters:

• If dataset_type is either text/csv or application/x-parquet, the model accepts four numeric features, and the baseline has two instances. In this example, if one record has all zero feature values and the other record has all one feature values, then baseline should be set to \([0,0,0,0],[1,1,1,1]\) without any header.

• If dataset_type is application/jsonlines, and features is the key to a list of four numeric feature values. In addition, in this example, if the baseline has one record of all zero values, then baseline should be \([\{"features":[0,0,0,0]\}]\).

• If dataset_type is application/json, the baseline dataset should have the same structure and format as the input dataset.

• For computer vision problems, baseline can be the S3 URI of an image that is used to mask out features (segments) from the input image. The SageMaker Clarify processing job loads the mask image and resizes it to the same resolution as the input image. If baseline is not provided, the SageMaker Clarify processing job generates a mask image of white noise at the same resolution as the input image.

• features_to_explain – (Optional) An array of strings or zero-based indices of feature columns to compute SHAP values for. If features_to_explain is not provided, SHAP values are computed for all feature columns. These feature columns cannot include the label column or predicted label column. The features_to_explain parameter is only supported for tabular datasets with numeric and categorical columns.

• num_clusters – (Optional) The number of clusters that the dataset is divided into to compute the baseline dataset. Each cluster is used to compute one baseline instance. If baseline is not specified, the SageMaker Clarify processing job attempts to compute the baseline dataset by dividing the tabular dataset into an optimal number of clusters between 1 and 12. The number of baseline instances directly affects the runtime of SHAP analysis.

• num_samples – (Optional) The number of samples to be used in the Kernel SHAP algorithm. If num_samples is not provided, the SageMaker Clarify processing job chooses the number
for you. The number of samples directly affects both the synthetic dataset size and job runtime.

- **seed** – (Optional) An integer used to initialize the pseudo random number generator in the SHAP explainer to generate consistent SHAP values for the same job. If seed is not specified, then each time that the same job runs, the model may output slightly different SHAP values.

- **use_logit** – (Optional) A Boolean value that indicates that you want the logit function to be applied to the model predictions. Defaults to false. If use_logit is true, then the SHAP values are calculated using the logistic regression coefficients, which can be interpreted as log-odds ratios.

- **save_local_shap_values** – (Optional) A Boolean value that indicates that you want the local SHAP values of each record in the dataset to be included in the analysis result. Defaults to false.

If the main dataset is split across multiple files or distributed processing is activated, also specify an identifier column using the parameter joinsource_name_or_index. The identifier column and the local SHAP values are saved in the analysis result. This way, you can map each record to its local SHAP values.

- **agg_method** – (Optional) The method used to aggregate the local SHAP values (the SHAP values for each instance) of all instances to the global SHAP values (the SHAP values for the entire dataset). Defaults to mean_abs. The following methods can be used to aggregate SHAP values.
  - **mean_abs** – The mean of absolute local SHAP values of all instances.
  - **mean_sq** – The mean of squared local SHAP values of all instances.
  - **median** – The median of local SHAP values of all instances.

- **text_config** – Required for natural language processing explainability. Include this configuration if you want to treat text columns as text and explanations should be provided for individual units of text. For an example of an analysis configuration for natural language processing explainability, see Analysis configuration for natural language processing explainability.

  - **granularity** – The unit of granularity for the analysis of text columns. Valid values are token, sentence, or paragraph. Each unit of text is considered a feature, and local SHAP values are computed for each unit.

  - **language** – The language of the text columns. Valid values are chinese, danish, dutch, english, french, german, greek, italian, japanese, lithuanian, multi-language, norwegian bokmål, polish, portuguese, romanian, russian.
spanish, afrikaans, albanian, arabic, armenian, basque, bengali, bulgarian, catalan, croatian, czech, estonian, finnish, gujarati, hebrew, hindi, hungarian, icelandic, indonesian, irish, kannada, kyrgyz, latvian, ligurian, luxembourgish, macedonian, malayalam, marathi, nepali, persian, sanskrit, serbian, setswana, sinhala, slovak, slovenian, swedish, tagalog, tamil, tatar, telugu, thai, turkish, ukrainian, urdu, vietnamese, yoruba. Enter multilingual for a mix of multiple languages.

- **max_top_tokens** – (Optional) The maximum number of top tokens, based on global SHAP values. Defaults to 50. It is possible for a token to appear multiple times in the dataset. The SageMaker Clarify processing job aggregates the SHAP values of each token, and then selects the top tokens based on their global SHAP values. The global SHAP values of the selected top tokens are included in the `global_top_shap_text` section of the `analysis.json` file.

- The local SHAP value of aggregation.

- **image_config** – Required for computer vision explainability. Include this configuration if you have an input dataset consisting of images and you want to analyze them for explainability in a computer vision problem.

- **model_type** – The type of the model. Valid values include:
  - **IMAGE_CLASSIFICATION** for an image classification model.
  - **OBJECT_DETECTION** for an object detection model.

- **max_objects** – Applicable only when `model_type` is **OBJECT_DETECTION**. The max number of objects, ordered by confidence score, detected by the computer vision model. Any objects ranked lower than the top `max_objects` by confidence score are filtered out. Defaults to 3.

- **context** – Applicable only when `model_type` is **OBJECT_DETECTION**. It indicates if the area around the bounding box of the detected object is masked by the baseline image or not. Valid values are 0 to mask everything, or 1 to mask nothing. Defaults to 1.

- **iou_threshold** – Applicable only when `model_type` is **OBJECT_DETECTION**. The minimum intersection over union (IOU) metric for evaluating predictions against the original detection. A high IOU metric corresponds to a large overlap between the predicted and ground truth detection box. Defaults to 0.5.

- **num_segments** – (Optional) An integer that determines the approximate number of segments to be labeled in the input image. Each segment of the image is considered a feature, and local SHAP values are computed for each segment. Defaults to 20.
• **segment_compactness** – (Optional) An integer that determines the shape and size of the image segments generated by the [scikit-image slic](https://scikit-image.org/) method. Defaults to 5.

• **pdp** – Include this method to compute partial dependence plots (PDPs). For an example of an analysis configuration to generate PDPs, see [Compute partial dependence plots (PDPs)](#).

• **features** – Mandatory if the shap method is not requested. An array of feature names or indices to compute and plot PDP plots.

• **top_k_features** – (Optional) Specifies the number of top features used to generate PDP plots. If features is not provided, but the shap method is requested, then the SageMaker Clarify processing job chooses the top features based on their SHAP attributions. Defaults to 10.

• **grid_resolution** – The number of buckets to divide the range of numeric values into. This specifies the granularity of the grid for the PDP plots.

• **report** – (Optional) Use this object to customize the analysis report. There are three copies of the same report as part of the analysis result: Jupyter Notebook report, HTML report, and PDF report. The object has the following parameters:

  • **name** – File name of the report files. For example, if name is `MyReport`, then the report files are `MyReport.ipynb`, `MyReport.html`, and `MyReport.pdf`. Defaults to `report`.

  • **title** – (Optional) Title string for the report. Defaults to `SageMaker Analysis Report`.

• **predictor** – Required if the analysis requires predictions from the model. For example, when the shap, pdp, or post_training_bias method is requested, but predicted labels are not provided as part of the input dataset. The following are parameters to be used in conjunction with predictor:

  • **model_name** – The name of your SageMaker model created by the CreateModel API. If you specify `model_name` instead of `endpoint_name`, the SageMaker Clarify processing job creates an ephemeral endpoint with the model name, known as a **shadow endpoint**, and gets predictions from the endpoint. The job deletes the shadow endpoint after the computations are completed. If the model is multi-model, then the `target_model` parameter must be specified. For more information about multi-model endpoints, see [Host multiple models in one container behind one endpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/host-several-models-in-one-container.html).

  • **endpoint_name_prefix** – (Optional) A custom name prefix for the shadow endpoint. Applicable if you provide `model_name` instead of `endpoint_name`. For example, provide `endpoint_name_prefix` if you want to restrict access to the endpoint by endpoint name. The prefix must match the `EndpointName` pattern, and its maximum length is 23. Defaults to `sm-clarify`. 

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Configure a SageMaker Clarify Processing Job

5386
• **initial_instance_count** – Specifies the number of instances for the shadow endpoint. Required if you provide model_name instead of endpoint_name. The value for initial_instance_count can be different from the InstanceCount of the job, but we recommend a 1:1 ratio.

• **instance_type** – Specifies the instance type for the shadow endpoint. Required if you provide model_name instead of endpoint_name. As an example, instance_type can be set to "ml.m5.large". In some cases, the value specified for instance_type can help reduce model inference time. For example, to run efficiently, natural language processing models and computer vision models typically require a graphics processing unit (GPU) instance type.

• **accelerator_type** – (Optional) Specifies the type of Elastic Inference (EI) accelerator to attach to the shadow endpoint. Applicable if you provide model_name instead of endpoint_name for accelerator_type. An example value for accelerator_type is ml.eia2.large. Defaults to not use an accelerator.

• **endpoint_name** – The name of your SageMaker endpoint created by the CreateEndpoint API. If provided, endpoint_name takes precedence over the model_name parameter. Using an existing endpoint reduces the shadow endpoint bootstrap time, but it can also cause a significant increase in load for that endpoint. Additionally, some analysis methods (such as shap and pdp) generate synthetic datasets that are sent to the endpoint. This can cause the endpoint's metrics or captured data to be contaminated by synthetic data, which may not accurately reflect real-world usage. For these reasons, it's generally not recommended to use an existing production endpoint for SageMaker Clarify analysis.

• **target_model** – The string value that is passed on to the TargetModel parameter of the SageMaker InvokeEndpoint API. Required if your model (specified by the model_name parameter) or endpoint (specified by the endpoint_name parameter) is multi-model. For more information about multi-model endpoints, see Host multiple models in one container behind one endpoint.

• **custom_attributes** – (Optional) A string that allows you to provide additional information about a request for an inference that is submitted to the endpoint. The string value is passed to the CustomAttributes parameter of the SageMaker InvokeEndpoint API.

• **content_type** – The model input format to be used for getting predictions from the endpoint. If provided, it is passed to the ContentType parameter of the SageMaker InvokeEndpoint API.
  - For computer vision explainability, the valid values are image/jpeg, image/png or application/x-npy. If content_type is not provided, the default value is image/jpeg.
• For other types of explainability, the valid values are **text/csv**, **application/jsonlines**, and **application/json**. A value for `content_type` is required if the `dataset_type` is "application/x-parquet". Otherwise `content_type` defaults to the value of the `dataset_type` parameter.

**accept_type** – The model output format to be used for getting predictions from the endpoint. The value for `accept_type` is passed to the Accept parameter of the SageMaker `InvokeEndpoint` API.

• For computer vision explainability, if `model_type` is "OBJECT_DETECTION" then `accept_type` defaults to **application/json**.

• For other types of explainability, the valid values are **text/csv**, **application/jsonlines**, and **application/json**. If a value for `accept_type` is not provided, `accept_type` defaults to the value of the `content_type` parameter.

**content_template** – A template string used to construct the model input from dataset records. The parameter `content_template` is only used and required if the value of the `content_type` parameter is either **application/jsonlines** or **application/json**.

When the `content_type` parameter is **application/jsonlines**, the template should have only one placeholder, `$features`, which is replaced by a features list at runtime. For example, if the template is 
"{"myfeatures":$features}"
and if a record has three numeric feature values: 1, 2 and 3, then the record will be sent to the model as JSON Line
{"myfeatures":[1,2,3]}

When the `content_type` is **application/json**, the template can have either placeholder `$record` or `$records`. If the placeholder is `record`, a single record is replaced with a record that has the template in `record_template` applied to it. In this case, only a single record will be sent to the model at a time. If the placeholder is `$records`, the records are replaced by a list of records, each with a template supplied by `record_template`.

**record_template** – A template string to be used to construct each record of the model input from dataset instances. It is only used and required when `content_type` is **application/json**. The template string may contain one of the following:

• A placeholder `$features` parameter that is substituted by an array of feature values. An additional optional placeholder can substitute feature column header names in `$feature_names`. This optional placeholder will be substituted by an array of feature names.
• Exactly one placeholder $features_kvp that is substituted by the key-value pairs, feature name and feature value.

• A feature in the headers configuration. As an example, a feature name A, notated by the placeholder syntax "$\{A\}" will be substituted by the feature value for A.

The value for record_template is used with content_template to construct the model input. A configuration example showing how to construct a model input using a content and record template follows.

In the following code example, the headers and features are defined as follows.

• `headers`:
  • "A", "B"
• `features`:
  • [[0,1], [3,4]]

The example model input is as follows.

```json
{
    "instances": [[0, 1], [3, 4]],
   "feature_names": ["A", "B"]
}
```

The example content_template and record_template parameter values to construct the previous example model input follows.

• `content_template`: "\{"instances\": $records, "feature_names\": $feature_names\}"  
• `record_template`: "$features"

In the following code example, the headers and features are defined as follows.

```json
[
    { "A": 0, "B": 1 },
    { "A": 3, "B": 4 },
]
```

The example content_template and record_template parameter values to construct the previous example model input follows.

• `content_template`: "$records"
• `record_template`: "$features_kvp"
An alternate code example to construct the previous example model input follows.

- `content_template`: "$records"
- `record_template`: "{"A": "$A", "B": "$B"}"

In the following code example, the headers and features are defined as follows.

```json
{ "A": 0, "B": 1 }
```

The example `content_template` and `record_template` parameters values to construct above: the previous example model input follows.

- `content_template`: "$record"
- `record_template`: "$features_kvp"
- `label` – A zero-based integer index or JMESPath expression string used to extract predicted labels from the model output for bias analysis. If the model is multiclass and the `label` parameter extracts all of the predicted labels from the model output, then the following apply.
  - The `probability` parameter is required to get the corresponding probabilities (or scores) from the model output.
  - The predicted label of the highest score is chosen.

The value for `label` depends on the value of the `accept_type` parameter as follows.

- If `accept_type` is `text/csv`, then `label` is the index of any predicted labels in the model output.
- If `accept_type` is `application/jsonlines` or `application/json`, then `label` is a JMESPath expression that's applied to the model output to get the predicted labels.

- `label_headers` – An array of values that the label can take in the dataset. If bias analysis is requested, then the `probability` parameter is also required to get the corresponding probability values (scores) from model output, and the predicted label of the highest score is chosen. If explainability analysis is requested, the label headers are used to beautify the analysis report. A value for `label_headers` is required for computer vision explainability. For example, for a multiclass classification problem, if the label has three possible values, `bird`, `cat`, and `dog`, then `label_headers` should be set to `["bird","cat","dog"]`.

- `probability` – (Optional) A zero-based integer index or a JMESPath expression string used to extract probabilities (scores) for explainability analysis, or to choose the predicted label.
for bias analysis. The value of probability depends on the value of the accept_type parameter as follows.

- If accept_type is `text/csv`, probability is the index of the probabilities (scores) in the model output. If probability is not provided, the entire model output is taken as the probabilities (scores).
- If accept_type is JSON data (either `application/jsonlines` or `application/json`), probability should be a JMESPath expression that is used to extract the probabilities (scores) from the model output.

**Example analysis configuration files**

The following sections contain example analysis configuration files for data in CSV format, JSON Lines format, and for both natural language processing (NLP) and computer vision explainability.

**Analysis configuration for a CSV dataset**

The following examples show how to configure bias and explainability analysis for a tabular dataset in CSV format. In these examples, the incoming dataset has four feature columns, and one binary label column, `Target`. The contents of the dataset are as follows. A label value of 1 indicates a positive outcome. The dataset is provided to the SageMaker Clarify job by the dataset processing input.

```
"Target","Age","Gender","Income","Occupation"
0,25,0,2850,2
1,36,0,6585,0
1,22,1,1759,1
0,48,0,3446,1
...
```

The following sections show how to compute pre-training and post-training bias metrics, SHAP values, and partial dependence plots (PDPs) showing feature importance for a dataset in CSV format.

**Compute all of the pre-training bias metrics**

This example configuration shows how to measure if the previous sample dataset is favorably biased towards samples with a `Gender` value of 0. The following analysis configuration instructs the SageMaker Clarify processing job to compute all the pre-training bias metrics for the dataset.
You can compute pre-training bias metrics prior to training. However, you must have a trained model to compute post-training bias metrics. The following example output is from a binary classification model that outputs data in CSV format. In this example output, each row contains two columns. The first column contains the predicted label, and the second column contains the probability value for that label.

```
0,0.028986845165491
1,0.825382471084594
...
```

The following configuration example instructs the SageMaker Clarify processing job to compute all possible bias metrics using the dataset and the predictions from the model output. In the example, the model is deployed to a SageMaker endpoint `your_endpoint`.

**Note**

In the following example code, the parameter `content_type` and `accept_type` are not set. Therefore, they automatically use the value of the parameter `dataset_type`, which is `text/csv`. 

```json
{
    "dataset_type": "text/csv",
    "label": "Target",
    "label_values_or_threshold": [1],
    "facet": [
        {
            "name_or_index": "Gender",
            "value_or_threshold": [0]
        }
    ],
    "methods": {
        "pre_training_bias": {
            "methods": "all"
        }
    }
}
```
Compute the SHAP values

The following example analysis configuration instructs the job to compute the SHAP values designating the Target column as labels and all other columns as features.
In this example, the SHAP baseline parameter is omitted and the value of the `num_clusters` parameter is 1. This instructs the SageMaker Clarify processor to compute one SHAP baseline sample. In this example, probability is set to 1. This instructs the SageMaker Clarify processing job to extract the probability score from the second column of the model output (using zero-based indexing).

**Compute partial dependence plots (PDPs)**

The following example shows how to view the importance of the Income feature on the analysis report using PDPs. The report parameter instructs the SageMaker Clarify processing job to generate a report. After the job completes, the generated report is saved as report.pdf to the `analysis_result` location. The `grid_resolution` parameter divides the range of the feature values into 10 buckets. Together, the parameters specified in the following example instruct the SageMaker Clarify processing job to generate a report containing a PDP graph for Income with 10 segments on the x-axis. The y-axis will show the marginal impact of Income on the predictions.

```json
{
   "dataset_type": "text/csv",
   "label": "Target",
   "methods": {
      "pdp": {
         "features": ["Income"],
         "grid_resolution": 10
      },
      "report": {
         "name": "report"
      }
   },
   "predictor": {
      "endpoint_name": "your_endpoint",
      "probability": 1
   }
}
```

**Compute both bias metrics and feature importance**

You can combine all the methods from the previous configuration examples into a single analysis configuration file and compute them all by a single job. The following example shows an analysis configuration with all steps combined.
In this example, the probability parameter is set to 1 to indicate that probabilities are contained in the second column (using zero-based indexing). However, because bias analysis needs a predicted label, the probability_threshold parameter is set to 0.5 to convert the probability score into a binary label. In this example, the top_k_features parameter of the partials dependence plots pdp method is set to 2. This instructs the SageMaker Clarify processing job to compute partials dependence plots (PDPs) for the top 2 features with the largest global SHAP values.

```json
{
    "dataset_type": "text/csv",
    "label": "Target",
    "probability_threshold": 0.5,
    "label_values_or_threshold": [1],
    "facet": [
        {
            "name_or_index": "Gender",
            "value_or_threshold": [0]
        }
    ],
    "methods": {
        "pre_training_bias": {
            "methods": "all"
        },
        "post_training_bias": {
            "methods": "all"
        },
        "shap": {
            "num_clusters": 1
        },
        "pdp": {
            "top_k_features": 2,
            "grid_resolution": 10
        },
        "report": {
            "name": "report"
        }
    },
    "predictor": {
        "endpoint_name": "your_endpoint",
        "probability": 1
    }
}
```
Instead of deploying the model to an endpoint, you can provide the name of your SageMaker model to the SageMaker Clarify processing job using the `model_name` parameter. The following example shows how to specify a model named `your_model`. The SageMaker Clarify processing job will create a shadow endpoint using the configuration.

```json
{
  ...
  "predictor": {
    "model_name": "your_model",
    "initial_instance_count": 1,
    "instance_type": "ml.m5.large",
    "probability": 1
  }
}
```

### Analysis configuration for a JSON Lines dataset

The following examples show how to configure bias analysis and explainability analysis for a tabular dataset in JSON Lines format. In these examples, the incoming dataset has the same data as the previous section but they are in the SageMaker JSON Lines dense format. Each line is a valid JSON object. The key "Features" points to an array of feature values, and the key "Label" points to the ground truth label. The dataset is provided to the SageMaker Clarify job by the "dataset" processing input. For more information about JSON Lines, see [JSONLINES Request Format](#).

```json
{"Features":[25,0,2850,2],"Label":0}
{"Features":[36,0,6585,0],"Label":1}
{"Features":[22,1,1759,1],"Label":1}
{"Features":[48,0,3446,1],"Label":0}
...
```

The following sections show how to compute pre-training and post-training bias metrics, SHAP values, and partial dependence plots (PDPs) showing feature importance for a dataset in JSON Lines format.

### Compute pre-training bias metrics

Specify the label, features, format, and methods to measure pre-training bias metrics for a `Gender` value of 0. In the following example, the `headers` parameter provides the feature names first. The label name is provided last. By convention, the last header is the label header.
The `features` parameter is set to the JMESPath expression "Features" so that the SageMaker Clarify processing job can extract the array of features from each record. The `label` parameter is set to JMESPath expression "Label" so that the SageMaker Clarify processing job can extract the ground truth label from each record. Use a facet name to specify the sensitive attribute, as follows.

```json
{
    "dataset_type": "application/jsonlines",
    "headers": ["Age","Gender","Income","Occupation","Target"],
    "label": "Label",
    "features": "Features",
    "label_values_or_threshold": [1],
    "facet": [
        {
            "name_or_index": "Gender",
            "value_or_threshold": [0]
        }
    ],
    "methods": {
        "pre_training_bias": {
            "methods": "all"
        }
    }
}
```

**Compute all the bias metrics**

You must have a trained model to compute post-training bias metrics. The following example is from a binary classification model that outputs JSON Lines data in the example's format. Each row of the model output is a valid JSON object. The key `predicted_label` points to the predicted label, and the key `probability` points to the probability value.

```json
{"predicted_label":0,"probability":0.0289868451654911}
{"predicted_label":1,"probability":0.825382471084594}
...
```

You can deploy the model to a SageMaker endpoint named `your_endpoint`. The following example analysis configuration instructs the SageMaker Clarify processing job to compute all possible bias metrics for both the dataset and the model. In this example, the parameter `content_type` and `accept_type` are not set. Therefore, they are automatically set to use the value of the parameter `dataset_type`, which is `application/jsonlines`. The SageMaker Clarify
processing job uses the content_template parameter to compose the model input, by replacing the $features placeholder by an array of features.

```
{
    "dataset_type": "application/jsonlines",
    "headers": ["Age","Gender","Income","Occupation","Target"],
    "label": "Label",
    "features": "Features",
    "label_values_or_threshold": [1],
    "facet": [
        {
            "name_or_index": "Gender",
            "value_or_threshold": [0]
        }
    ],
    "methods": {
        "pre_training_bias": {
            "methods": "all"
        },
        "post_training_bias": {
            "methods": "all"
        }
    },
    "predictor": {
        "endpoint_name": "your_endpoint",
        "content_template": "{\"Features\":$features}",
        "label": "predicted_label"
    }
}
```

**Compute the SHAP values**

Because SHAP analysis doesn’t need a ground truth label, the label parameter is omitted. In this example, the headers parameter is also omitted. Therefore, the SageMaker Clarify processing job must generate placeholders using generic names like column_0 or column_1 for feature headers, and label0 for a label header. You can specify values for headers and for a label to improve the readability of the analysis result. Because the probability parameter is set to JMESPath expression probability, the probability value will be extracted from the model output. The following is an example to calculate SHAP values.

```
{
    "dataset_type": "application/jsonlines",
```
Compute partials dependence plots (PDPs)

The following example shows how to view the importance of "Income" on PDP. In this example, the feature headers are not provided. Therefore, the features parameter of the pdp method must use zero-based index to refer to location of the feature column. The grid_resolution parameter divides the range of the feature values into 10 buckets. Together, the parameters in the example instruct the SageMaker Clarify processing job to generate a report containing a PDP graph for Income with 10 segments on the x-axis. The y-axis will show the marginal impact of Income on the predictions.
Compute both bias metrics and feature importance

You can combine all previous methods into a single analysis configuration file and compute them all by a single job. The following example shows an analysis configuration with all steps combined. In this example, the probability parameter is set. But because bias analysis needs a predicted label, the probability_threshold parameter is set to 0.5 to convert the probability score into a binary label. In this example, the top_k_features parameter of the pdp method is set to 2. This instructs the SageMaker Clarify processing job to compute PDPs for the top 2 features with the largest global SHAP values.

```json
{
    "dataset_type": "application/jsonlines",
    "headers": ["Age","Gender","Income","Occupation","Target"],
    "label": "Label",
    "features": "Features",
    "probability_threshold": 0.5,
    "label_values_or_threshold": [1],
    "facet": [
        {"name_or_index": "Gender", "value_or_threshold": [0]}
    ],
    "methods": {
        "pre_training_bias": {
            "methods": "all"
        },
        "post_training_bias": {
            "methods": "all"
        },
        "shap": {
            "num_clusters": 1
        },
        "pdp": {
            "top_k_features": 2,
            "grid_resolution": 10
        },
        "report": {
            "name": "report"
        }
    },
    "predictor": {
```
Analysis configuration for a JSON dataset

The following examples show how to configure bias and explainability analysis for a tabular dataset in JSON format. In these examples, the incoming dataset has the same data as the previous section but they are in the SageMaker JSON dense format. For more information about JSON Lines, see JSONLINES Request Format.

The whole input request is valid JSON where the outer structure is a list and each element is the data for a record. Within each record, the key Features points to an array of feature values, and the key Label points to the ground truth label. The dataset is provided to the SageMaker Clarify job by the dataset processing input.

[  
  {"Features":[25,0,2850,2],"Label":0},  
  {"Features":[36,0,6585,0],"Label":1},  
  {"Features":[22,1,1759,1],"Label":1},  
  {"Features":[48,0,3446,1],"Label":0},  
  ... 
]

The following sections show how to compute pre-training and post-training bias metrics, SHAP values, and partial dependence plots (PDPs) that show feature importance for a dataset in JSON Lines format.

**Compute pre-training bias metrics**

Specify the label, features, format, and methods to measure pre-training bias metrics for a Gender value of 0. In the following example, the headers parameter provides the feature names first. The label name is provided last. For JSON datasets, the last header is the label header.

The features parameter is set to the JMESPath expression that extracts a 2D array or matrix. Each row in this matrix must contain the list of Features for each record. The label parameter is set to JMESPath expression that extracts a list of ground truth labels. Each element in this list must contain the label for a record.
Use a facet name to specify the sensitive attribute, as follows.

```json
{
  "dataset_type": "application/json",
  "headers": ["Age","Gender","Income","Occupation","Target"],
  "label": "[*].Label",
  "features": "[*].Features",
  "label_values_or_threshold": [1],
  "facet": [
    {
      "name_or_index": "Gender",
      "value_or_threshold": [0]
    }
  ],
  "methods": {
    "pre_training_bias": {
      "methods": "all"
    }
  }
}
```

**Compute all the bias metrics**

You must have a trained model to compute post-training bias metrics. The following code example is from a binary classification model that outputs JSON data in the example's format. In the example, each element under `predictions` is the prediction output for a record. The example code contains the key `predicted_label`, that points to the predicted label, and the key `probability` points to the probability value.

```json
{
  "predictions": [
    {"predicted_label":0,"probability":0.028986845165491},
    {"predicted_label":1,"probability":0.825382471084594},
    ...
  ]
}
```

You can deploy the model to a SageMaker endpoint named `your_endpoint`.

In the following example, the parameter `content_type` and `accept_type` are not set. Therefore, `content_type` and `accept_type` are automatically set to use the value of the parameter.
dataset_type, which is application/json. The SageMaker Clarify processing job then uses the content_template parameter to compose the model input.

In the following example, the model input is composed by replacing the $records placeholder by an array of records. Then, the record_template parameter composes each record's JSON structure and replaces the $features placeholder with each record's array of features.

The following example analysis configuration instructs the SageMaker Clarify processing job to compute all possible bias metrics for both the dataset and the model.

```json
{
  "dataset_type": "application/json",
  "headers": ["Age","Gender","Income","Occupation","Target"],
  "label": "[\*].Label",
  "features": "[\*].Features",
  "label_values_or_threshold": [1],
  "facet": [
    {
      "name_or_index": "Gender",
      "value_or_threshold": [0]
    }
  ],
  "methods": {
    "pre_training_bias": {
      "methods": "all"
    },
    "post_training_bias": {
      "methods": "all"
    }
  },
  "predictor": {
    "endpoint_name": "your_endpoint",
    "content_template": "$records",
    "record_template": "\{\"Features\":$features\}",
    "label": "predictions[\*].predicted_label"
  }
}
```

**Compute the SHAP values**

You don’t need to specify a label for SHAP analysis. In the following example, the headers parameter is not specified. Therefore, the SageMaker Clarify processing job will generate
placeholders using generic names like column_0 or column_1 for feature headers, and label0 for a label header. You can specify values for headers and for a label to improve the readability of the analysis result.

In the following configuration example, the probability parameter is set to a JMESPath expression that extracts the probabilities from each prediction for each record. The following is an example to calculate SHAP values.

```json
{
   "dataset_type": "application/json",
   "features": "[*].Features",
   "methods": {
      "shap": {
         "num_clusters": 1
      }
   },
   "predictor": {
      "endpoint_name": "your_endpoint",
      "content_template": "$records",
      "record_template": "{"Features":$features}",
      "probability": "predictions[*.probability"
   }
}
```

**Compute partial dependence plots (PDPs)**

The following example shows you how to view a feature importance in PDPs. In the example, the feature headers are not provided. Therefore, the features parameter of the pdp method must use zero-based index to refer to location of the feature column. The grid_resolution parameter divides the range of the feature values into 10 buckets.

Together, the parameters in the following example instruct the SageMaker Clarify processing job to generate a report containing a PDP graph for Income with 10 segments on the x-axis. The y-axis shows the marginal impact of Income on the predictions.

The following configuration example shows how to view the importance of Income on PDPs.

```json
{
   "dataset_type": "application/json",
   "features": "[*].Features",
   "methods": {
      ...
   }
}
```
Compute both bias metrics and feature importance

You can combine all previous configuration methods into a single analysis configuration file and compute them all with a single job. The following example shows an analysis configuration with all steps combined.

In this example, the `probability` parameter is set. Because bias analysis needs a predicted label, the `probability_threshold` parameter is set to 0.5, which is used to convert the probability score into a binary label. In this example, the `top_k_features` parameter of the `pdp` method is set to 2. This instructs the SageMaker Clarify processing job to compute PDPs for the top 2 features with the largest global SHAP values.
Analysis configuration for natural language processing explainability

The following example shows an analysis configuration file for computing feature importance for natural language processing (NLP). In this example, the incoming dataset is a tabular dataset in CSV format, with one binary label column and two feature columns, as follows. The dataset is provided to the SageMaker Clarify job by the dataset processing input parameter.

<table>
<thead>
<tr>
<th>0,2</th>
<th>&quot;They taste gross&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td>1,3</td>
<td>&quot;Flavor needs work&quot;</td>
</tr>
<tr>
<td>1,5</td>
<td>&quot;Taste is awful&quot;</td>
</tr>
<tr>
<td>0,1</td>
<td>&quot;The worst&quot;</td>
</tr>
<tr>
<td>...</td>
<td></td>
</tr>
</tbody>
</table>

In this example, a binary classification model was trained on the previous dataset. The model accepts CSV data, and it outputs a single score between 0 and 1, as follows.

0.491656005382537
The model is used to create a SageMaker model named “your_model”. The following analysis configuration shows how to run a token-wise explainability analysis using the model and dataset. The text_config parameter activates the NLP explainability analysis. The granularity parameter indicates that the analysis should parse tokens.

In English, each token is a word. The following example also shows how to provide an in-place SHAP "baseline" instance using an average "Rating" of 4. A special mask token "[MASK]" is used to replace a token (word) in "Comments". This example also uses a GPU endpoint instance type to speed up inferencing.

```json
{
    "dataset_type": "text/csv",
    "headers": ["Target","Rating","Comments"],
    "label": "Target",
    "methods": {
        "shap": {
            "text_config": {
                "granularity": "token",
                "language": "english"
            }
        },
        "baseline": [[4,"[MASK]"],
        ],
        "predictor": {
            "model_name": "your_nlp_model",
            "initial_instance_count": 1,
            "instance_type": "ml.g4dn.xlarge"
        }
    }
}
```

**Analysis configuration for computer vision explainability**

The following example shows an analysis configuration file computing feature importance for computer vision. In this example, the input dataset consists of JPEG images. The dataset is provided to the SageMaker Clarify job by the dataset processing input parameter. The example shows how to configure an explainability analysis using a SageMaker image classification model. In the example, a model named `your_cv_ic_model`, has been trained to classify the animals on the input JPEG images.
For more information about image classification, see Image Classification - MXNet.

In this example, a SageMaker object detection model, `your_cv_od_model` is trained on the same JPEG images to identify the animals on them. The following example shows how to configure an explainability analysis for the object detection model.

```json
{
    "dataset_type": "application/x-image",
    "probability_threshold": 0.5,
    "methods": {
        "shap": {
            "image_config": {
                "model_type": "OBJECT_DETECTION",
                "max_objects": 3,
                "context": 1.0,
                "iou_threshold": 0.5,
                "num_segments": 20,
                "segment_compactness": 10
            }
        }
    },
    "report": {
    }
}
```
At a high level, the SageMaker Clarify processing job follows the input–process–output model to compute bias metrics and feature attributions. Refer to the following examples for details.

The input to the SageMaker Clarify processing job consists of the following:

- The dataset to be analyzed.
- The analysis configuration. For more information about how to configure an analysis, see Configure the Analysis.

During the processing stage, SageMaker Clarify computes bias metrics and feature attributions. The SageMaker Clarify processing job completes the following steps in the backend:

- The SageMaker Clarify processing job parses your analysis configuration and loads your dataset.
- To compute post-training bias metrics and feature attributions, the job requires model predictions from your model. The SageMaker Clarify processing job serializes your data and sends it as a request to your model that is deployed on a SageMaker real-time inference endpoint. After that, the SageMaker Clarify processing job extracts predictions from the response.
- The SageMaker Clarify processing job performs the bias and explainability analysis, and then it outputs the results.
For more information, see How SageMaker Clarify Processing Jobs Work.

The parameter that you use to specify the format of the data depends on where the data is used in the processing flow as follows:

- For an input dataset, use the dataset_type parameter to specify the format or MIME type.
- For a request to an endpoint, use the content_type parameter to specify the format.
- For a response from an endpoint, use the accept_type parameter to specify the format.

The input dataset, request, and the response to and from the endpoint don't require the same format. For example, you can use a Parquet dataset with a CSV request payload and a JSON Lines response payload given the following conditions.

- Your analysis is configured correctly.
- Your model supports the request and response formats.

**Note**

If content_type or accept_type are not provided, then the SageMaker Clarify container infers the content_type and accept_type.

**Topics**

- Tabular data
- Image data

**Tabular data**

Tabular data refers to data that can be loaded into a two-dimensional data frame. In the frame, each row represents a record, and each record has one or more columns. The values within each data frame cell can be of numerical, categorical, or text data types.

**Tabular dataset prerequisites**

Prior to analysis, your dataset should have had any necessary pre-processing steps already applied. This includes data cleaning or feature engineering.
You can provide one or multiple datasets. If you provide multiple datasets, use the following to identify them to the SageMaker Clarify processing job.

- Use either a **ProcessingInput** named dataset or the analysis configuration **dataset_uri** to specify the main dataset. For more information about **dataset_uri**, see the parameters list in Configure the Analysis.

- Use the baseline parameter provided in the analysis configuration file. The baseline dataset is required for SHAP analysis. For more information about the analysis configuration file, including examples, see Configure the Analysis.

The following table lists supported data formats, their file extensions, and MIME types.

<table>
<thead>
<tr>
<th>Data format</th>
<th>File extension</th>
<th>MIME type</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSV</td>
<td>csv</td>
<td>text/csv</td>
</tr>
<tr>
<td>JSON Lines</td>
<td>jsonl</td>
<td>application/jsonlines</td>
</tr>
<tr>
<td>JSON</td>
<td>json</td>
<td>application/json</td>
</tr>
<tr>
<td>Parquet</td>
<td>parquet</td>
<td>&quot;application/x-parquet&quot;</td>
</tr>
</tbody>
</table>

The following sections show example tabular datasets in CSV, JSON Lines, and Apache Parquet formats.

**Tabular dataset prerequisites in CSV format**

The SageMaker Clarify processing job is designed to load CSV data files in the **csv.excel** dialect. However, it's flexible enough to support other line terminators, including \n and \r.

For compatibility, all CSV data files provided to the SageMaker Clarify processing job must be encoded in UTF-8.

If your dataset does not contain a header row, do the following:

- Set the analysis configuration label to index 0. This means that the first column is the ground truth label.
If the parameter headers is set, set label to the label column header to indicate the location of the label column. All other columns are designated as features.

The following is an example of a dataset that does not contain a header row.

```
1,5,2.8,2.538,This is a good product
0,1,0.79,0.475,Bad shopping experience
...
```

If your data contains a header row, set the parameter label to index 0. To indicate the location of the label column, use the ground truth label header Label. All other columns are designated as features.

The following is an example of a dataset that contains a header row.

```
Label,Rating,A12,A13,Comments
1,5,2.8,2.538,This is a good product
0,1,0.79,0.475,Bad shopping experience
...
```

**Tabular dataset prerequisites in JSON format**

JSON is a flexible format for representing structured data that contains any level of complexity. The SageMaker Clarify support for JSON is not restricted to any specific format and thus allows for more flexible data formats in comparison to datasets in CSV or JSON Lines formats. This guide shows you how to set an analysis configuration for tabular data in JSON format.

### Note

To ensure compatibility, all JSON data files provided to the SageMaker Clarify processing job must be encoded in UTF-8.

The following is example input data with records that contain a top-level key, a list of features, and a label.

```
[
  {
    "features":[1,5,2.8,2.538,"This is a good product"],"label":1},
  {
    "features":[0,1,0.79,0.475,"Bad shopping experience"],"label":0},
  ...
]
An example configuration analysis for the previous input example dataset should set the following parameters:

- The `label` parameter should use the JMESPath expression `[*].label` to extract the ground truth label for each record in the dataset. The JMESPath expression should produce a list of labels where the $i^{th}$ label corresponds to the $i^{th}$ record.

- The `features` parameter should use the JMESPath expression `[*].features` to extract an array of features for each record in the dataset. The JMESPath expression should produce a 2D array or matrix where the $i^{th}$ row contains the feature values for corresponding to the $i^{th}$ record.

The following is example input data with records that contains a top-level key and a nested key that contains a list of features and labels for each record.

```json
{
   "data": [
      {
         "features": [1, 5, 2.8, 2.538, "This is a good product"],
         "label": 1
      },
      {
         "features": [0, 1, 0.79, 0.475, "Bad shopping experience"],
         "label": 0
      }
   ]
}
```

An example configuration analysis for the previous input example dataset should set the following parameters:

- The `label` parameter uses the JMESPath expression `data[[]].label` to extract the ground truth label for each record in the dataset. The JMESPath expression should produce a list of labels where the $i^{th}$ label is for the $i^{th}$ record.

- The `features` parameter uses the JMESPath expression `data[[]].features` to extract the array of features, for each record in the dataset. The JMESPath expression should produce a 2D array or matrix where the $i^{th}$ row contains the feature values for the $i^{th}$ record.

**Tabular dataset prerequisites in JSON Lines format**

JSON Lines is a text format for representing structured data where each line is a valid JSON object. Currently SageMaker Clarify processing jobs only support SageMaker Dense Format JSON Lines. To conform to the required format, all of the features of a record should be listed in a single JSON array. For more information about JSON Lines, see [JSONLINES Request Format](#).
Amazon SageMaker

Developer Guide

Note
All JSON Lines data ﬁles provided to the SageMaker Clarify processing job must be encoded
in UTF-8 to ensure compatibility.

The following is an example of how to set an analysis conﬁguration for a record that contains a
top-level key and a list of elements.
{"features":[1,5,2.8,2.538,"This is a good product"],"label":1}
{"features":[0,1,0.79,0.475,"Bad shopping experience"],"label":0}
...

The conﬁguration analysis for the previous dataset example should set the parameters as follows:
• To indicate the location of the ground truth label, the parameter label should be set to the
JMESPath expression label.
• To indicate the location of the array of features, the parameter features should be set to the
JMESPath expression features.
The following is an example of how to set an analysis conﬁguration for a record that contains a
top-level key and a nested key that contains a list of elements.
{"data":{"features":[1,5,2.8,2.538,"This is a good product"],"label":1}}
{"data":{"features":[0,1,0.79,0.475,"Bad shopping experience"],"label":0}}
...

The conﬁguration analysis for the previous dataset example should set the parameters as follows:
• The parameter label should be set to the JMESPath expression data.label to indicate the
location of the ground truth label.
• The parameter features should be set to the JMESPath expression data.features to indicate
the location of the array of features.
Tabular dataset prerequisites in Parquet format
Parquet is a column-oriented binary data format. Currently, SageMaker Clarify processing jobs
support loading Parquet data ﬁles only when the processing instance count is 1.
Conﬁgure a SageMaker Clarify Processing Job

5414


Because SageMaker Clarify processing jobs don’t support endpoint request or endpoint response in Parquet format, you must specify the data format of the endpoint request by setting the analysis configuration parameter `content_type` to a supported format. For more information, see `content_type` in Configure the Analysis.

The Parquet data must have column names that are formatted as strings. Use the analysis configuration `label` parameter to set the label column name to indicate the location of the ground truth labels. All other columns are designated as features.

**Endpoint requests for tabular data**

To obtain model predictions for post-training bias analysis and feature importance analysis, SageMaker Clarify processing jobs serialize the tabular data into bytes and sends these to an inference endpoint as a request payload. This tabular data is either sourced from the input dataset, or it's generated. If it's synthetic data, it's generated by the explainer for SHAP analysis or PDP analysis.

The data format of the request payload should be specified by the analysis configuration `content_type` parameter. If the parameter is not provided, the SageMaker Clarify processing job will use the value of the `dataset_type` parameter as the content type. For more information about `content_type` or `dataset_type`, see Configure the Analysis.

The following sections show example endpoint requests in CSV and JSON Lines formats.

**Endpoint request in CSV format**

The SageMaker Clarify processing job can serialize data to CSV format (MIME type: text/csv). The following table shows examples of the serialized request payloads.

<table>
<thead>
<tr>
<th>Endpoint request payload (string representation)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>'1,2,3,4'</td>
<td>Single record (four numerical features).</td>
</tr>
<tr>
<td>'1,2,3,4\n5,6,7,8'</td>
<td>Two records, separated by line break '\n'.</td>
</tr>
<tr>
<td>&quot;&quot;This is a good product&quot;,5'</td>
<td>Single record (a text feature and a numerical feature).</td>
</tr>
</tbody>
</table>
Endpoint request payload (string representation) | Comments
--- | ---
""This is a good product",5\n"Bad shopping experience",1" | Two records.

**Endpoint request is in JSON Lines format**

The SageMaker Clarify processing job can serialize data to SageMaker JSON Lines dense format (MIME type: application/jsonlines). For more information about JSON Lines, see [JSONLINES Request Format](#).

To transform tabular data into JSON data, provide a template string to the analysis configuration content_template parameter. For more information about content_template see [Configure the Analysis](#). The following table shows examples of serialized JSON Lines request payloads.

<table>
<thead>
<tr>
<th>Endpoint request payload (string representation)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>'{&quot;data&quot;:{&quot;features&quot;:[1,2,3,4]}}'</td>
<td>Single record. In this case, the template looks like '{&quot;data&quot;:{&quot;features&quot;:$features}}' and $features is replaced by the list of features [1,2,3,4] .</td>
</tr>
<tr>
<td>'{&quot;data&quot;:{&quot;features&quot;:[1,2,3,4]}}\n{&quot;data&quot;:{&quot;features&quot;:[5,6,7,8]}}'</td>
<td>Two records.</td>
</tr>
<tr>
<td>'{&quot;features&quot;:[&quot;This is a good product&quot;,5]}'</td>
<td>Single record. In this case, the template looks like '{&quot;features&quot;:$features}' and $features is replaced by the list of features [&quot;This is a good product&quot;,5] .</td>
</tr>
<tr>
<td>'{&quot;features&quot;:[&quot;This is a good product&quot;,5]}\n{&quot;features&quot;:[&quot;Bad shopping experience&quot;,1]}'</td>
<td>Two records.</td>
</tr>
</tbody>
</table>
Endpoint request is in JSON format

A SageMaker Clarify processing job can serialize data to arbitrary JSON structures (MIME type: application/json). To do this, you must provide a template string to the analysis configuration content_template parameter. This is used by the SageMaker Clarify processing job to construct the outer JSON structure. You must also provide a template string for record_template, which is used to construct the JSON structure for each record. For more information about content_template and record_template, see Configure the Analysis.

Note

Because content_template and record_template are string parameters, any double quote characters (") that are part of the JSON serialized structure should be noted as an escaped character in your configuration. For example, if you want to escape a double quote in Python, you could enter the following for content_template.

```
"{"data":{"features":$record}}"
```

The following table shows examples of serialized JSON request payloads and the corresponding content_template and record_template parameters that are required to construct them.

<table>
<thead>
<tr>
<th>Endpoint request payload (string representation)</th>
<th>Comments</th>
<th>content_template</th>
<th>record_template</th>
</tr>
</thead>
<tbody>
<tr>
<td>'{&quot;data&quot;:{&quot;features&quot;: [1,2,3,4]}}'</td>
<td>Single record at a time.</td>
<td>'{&quot;data&quot;:{&quot;features&quot;: $record}}'</td>
<td>&quot;$features&quot;</td>
</tr>
<tr>
<td>'{&quot;instances&quot;:[[0, 1], [3, 4]], &quot;feature-names&quot;: [&quot;A&quot;, &quot;B&quot;]}'</td>
<td>Multi-records with feature names.</td>
<td>'{&quot;instances&quot;:$records, &quot;feature-names&quot;:$feature_names}'</td>
<td>&quot;$features&quot;</td>
</tr>
<tr>
<td>'{&quot;A&quot;: 0, &quot;B&quot;: 1}, {&quot;A&quot;: 3, &quot;B&quot;: 4}'}</td>
<td>Multi-records and key-value pairs.</td>
<td>&quot;$records&quot;</td>
<td>&quot;$features_kvp&quot;</td>
</tr>
</tbody>
</table>
Endpoint request payload (string representation) | Comments | content_template | record_template
--- | --- | --- | ---
`{"A": 0, "B": 1}' | Single record at a time and key-value pairs. | "$record" | "$features_kvp"

`{"A": 0, "nested":{"B": 1}}' | Alternatively, use the fully verbose record_template for arbitrary structures. | "$record" | `{"A": "${A}", "nested":{"B": "${B}"}}'

**Endpoint response for tabular data**

After the SageMaker Clarify processing job receives an inference endpoint invocation's response, it deserializes the response payload and extracts predictions from it. Use the analysis configuration accept_type parameter to specify the data format of the response payload. If accept_type is not provided, the SageMaker Clarify processing job will use the value of the content_type parameter as the model output format. For more information about accept_type, see Configure the Analysis.

The predictions could either consist of predicted labels for bias analysis, or probability values (scores) for feature importance analysis. In the predictor analysis configuration, the following three parameters extract the predictions.

- The parameter probability is used to locate the probability values (scores) in the endpoint response.
- The parameter label is used to locate the predicted labels in the endpoint response.
- (Optional) The parameter label_headers provides the predicted labels for a multiclass model.

The following guidelines pertain to endpoint responses in CSV, JSON Lines, and JSON formats.

**Endpoint Response is in CSV format**

If the response payload is in CSV format (MIME type: text/csv), the SageMaker Clarify processing job deserializes each row. It then extracts the predictions from the deserialized data using the
column indexes provided in the analysis configuration. The rows in the response payload must match the records in the request payload.

The following tables provide examples of response data in different formats and for different problem types. Your data can vary from these examples, as long as the predictions can be extracted according to the analysis configuration.

The following sections show example endpoint responses in CSV formats.

**Endpoint response is in CSV format and contains probability only**

The following table is an example endpoint response for regression and binary classification problems.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record.</td>
<td>'0.6'</td>
</tr>
<tr>
<td>Two records (results in one line, divided by comma).</td>
<td>'0.6,0.3'</td>
</tr>
<tr>
<td>Two records (results in two lines).</td>
<td>'0.6\n0.3'</td>
</tr>
</tbody>
</table>

For the previous example, the endpoint outputs a single probability value (score) of the predicted label. To extract probabilities using the index and use them for feature importance analysis, set the analysis configuration parameter `probability` to column index 0. These probabilities can also be used for bias analysis if they're converted to binary value by using the `probability_threshold` parameter. For more information about `probability_threshold`, see [Configure the Analysis](#).

The following table is an example endpoint response for a multiclass problem.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record of a multiclass model (three classes).</td>
<td>'0.1,0.6,0.3'</td>
</tr>
<tr>
<td>Endpoint request payload</td>
<td>Endpoint response payload (string representation)</td>
</tr>
<tr>
<td>----------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>Two records of a multiclass model (three classes).</td>
<td>'0.1,0.6,0.3\n0.2,0.5,0.3'</td>
</tr>
</tbody>
</table>

For the previous example, the endpoint outputs a list of probabilities (scores). If no index is provided, all values are extracted and used for feature importance analysis. If the analysis configuration parameter `label_headers` is provided. Then the SageMaker Clarify processing job can select the label header of the max probability as the predicted label, which can be used for bias analysis. For more information about `label_headers`, see [Configure the Analysis](#).

**Endpoint response is in CSV format and contains predicted label only**

The following table is an example endpoint response for regression and binary classification problems.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'1'</td>
</tr>
<tr>
<td>Two records (results in one line, divided by comma)</td>
<td>'1,0'</td>
</tr>
<tr>
<td>Two records (results in two lines)</td>
<td>'1\n0'</td>
</tr>
</tbody>
</table>

For the previous example, the endpoint outputs the predicted label instead of probability. Set the `label` parameter of the `predictor` configuration to column index 0 so that the predicted labels can be extracted using the index and used for bias analysis.

**Endpoint response is in CSV format and contains predicted label and probability**

The following table is an example endpoint response for regression and binary classification problems.
For the previous example, the endpoint outputs the predicted label followed by its probability. Set the label parameter of the predictor configuration to column index 0, and set probability to column index 1 to extract both parameter values.

**Endpoint response is in CSV format and contains predicted labels and probabilities (multiclass)**

A multiclass model trained by Amazon SageMaker Autopilot can be configured to output the string representation of the list of predicted labels and probabilities. The following example table shows an example endpoint response from a model that is configured to output predicted_label, probability, labels, and probabilities.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'1,0.6'</td>
</tr>
<tr>
<td>Two records</td>
<td>'1,0.6\n0,0.3'</td>
</tr>
</tbody>
</table>

For the previous example, the SageMaker Clarify processing job can be configured in the following ways to extract the predictions.

For bias analysis, the previous example can be configured as one of the following.

- Set the label parameter of the predictor configuration to 0 to extract the predicted label.
- Set the parameter to 2 to extract the predicted labels, and set probability to 3 to extract the corresponding probabilities. The SageMaker Clarify processing job can automatically determine
the predicted label by identifying the label with the highest probability value. Referring to the previous example of a single record, the model predicts three labels: cat, dog, and fish, with corresponding probabilities of 0.1, 0.6, and 0.3. Based on these probabilities, the predicted label is dog, as it has the highest probability value of 0.6.

- Set probability to 3 to extract the probabilities. If label_headers is provided, then the SageMaker Clarify processing job can automatically determine the predicted label by identifying the label header with the highest probability value.

For feature importance analysis, the previous example can be configured as follows.

- Set probability to 3 extract the probabilities of all the predicted labels. Then, feature attributions will be computed for all the labels. If the customer doesn't specify label_headers, then the predicted labels will be used as label headers in the analysis report.

**Endpoint Response is in JSON Lines format**

If the response payload is in JSON Lines format (MIME type: application/jsonlines), the SageMaker Clarify processing job deserializes each line as JSON. It then extracts predictions from the deserialized data using JMESPath expressions provided in analysis configuration. The lines in the response payload must match the records in the request payload. The following tables shows examples of response data in different formats. Your data can vary from these examples, as long as the predictions can be extracted according to the analysis configuration.

The following sections show example endpoint responses in JSON Lines formats.

**Endpoint response is in JSON Lines format and contains probability only**

The following table is an example endpoint response that only outputs the probability value (score).

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;score&quot;:0.6}'</td>
</tr>
<tr>
<td>Two records</td>
<td>'{&quot;score&quot;:0.6}\n{&quot;score&quot;:0.3}'</td>
</tr>
</tbody>
</table>
For the previous example, set the analysis configuration parameter `probability` to JMESPath expression "score" to extract its value.

**Endpoint response is in JSON Lines format and contains predicted label only**

The following table is an example endpoint response that only outputs the predicted label.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;prediction&quot;:1}'</td>
</tr>
<tr>
<td>Two records</td>
<td>'{&quot;prediction&quot;:1}\n{&quot;prediction&quot;:0}'</td>
</tr>
</tbody>
</table>

For the previous example, set the `label` parameter of the predictor configuration to JMESPath expression `prediction`. Then, the SageMaker Clarify processing job can extract the predicted labels for bias analysis. For more information, see [Configure the Analysis](Configure a SageMaker Clarify Processing Job).

**Endpoint response is in JSON Lines format and contains predicted label and probability**

The following table is an example endpoint response that outputs the predicted label and its score.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;prediction&quot;:1,&quot;score&quot;:0.6}'</td>
</tr>
<tr>
<td>Two records</td>
<td>'{&quot;prediction&quot;:1,&quot;score&quot;:0.6}\n{&quot;prediction&quot;:0,&quot;score&quot;:0.3}'</td>
</tr>
</tbody>
</table>

For the previous example, set the `label` parameter of the predictor configuration to JMESPath expression "prediction" to extract the predicted labels. Set `probability` to JMESPath expression "score" to extract the probability. For more information, see [Configure the Analysis](Configure a SageMaker Clarify Processing Job).

**Endpoint response is in JSON Lines format and contains predicted labels and probabilities (multiclass)**

The following table is an example endpoint response from a multiclass model that outputs the following:
• A list of predicted labels.
• Probabilities, and the selected predicted label and its probability.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;predicted_label&quot;:&quot;dog&quot;,&quot;probability&quot;:0.6,&quot;predicted_labels&quot;:[&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;],&quot;probabilities&quot;:[0.1,0.6,0.3]}'</td>
</tr>
</tbody>
</table>
| Two records              | '{"predicted_label":"dog","probability":0.6,"predicted_labels":["cat","dog","fish"],"probabilities":[0.1,0.6,0.3]}
\n{"predicted_label":"cat","probability":0.7,"predicted_labels":["cat","dog","fish"],"probabilities":[0.7,0.2,0.1]}' |

For the previous example, the SageMaker Clarify processing job can be configured in several ways to extract the predictions.

For bias analysis, the previous example can be configured as one of the following.

• Set the label parameter of the predictor configuration to JMESPath expression "predicted_label" to extract the predicted label.

• Set the parameter to JMESPath expression "predicted_labels" to extract the predicted labels.

• Set probability to JMESPath expression "probabilities" to extract their probabilities. The SageMaker Clarify job automatically determine the predicted label by identifying the label with the highest probability value.

• Set probability to JMESPath expression "probabilities" to extract their probabilities. If label_headers is provided, then the SageMaker Clarify processing job can automatically determine the predicted label by identifying the label with the highest probability value.

For feature importance analysis, do the following.

• Set probability to the JMESPath expression "probabilities" to extract their probabilities of all the predicted labels. Then, feature attributions will be computed for all the labels.
Endpoint Response is in JSON format

If the response payload is in JSON format (MIME type: application/json), the SageMaker Clarify processing job deserializes the entire payload as JSON. It then extracts predictions from the deserialized data using JMESPath expressions provided in the analysis configuration. The records in the response payload must match the records in the request payload.

The following sections show example endpoint responses in JSON formats. The sections contain tables with examples of response data in different formats and for different problem types. Your data can vary from these examples, as long as the predictions can be extracted according to the analysis configuration.

Endpoint response is in JSON format and contains probability only

The following table is an example response from an endpoint that only outputs the probability value (score).

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'0.6'</td>
</tr>
<tr>
<td>Two records</td>
<td>'0.6,0.3'</td>
</tr>
</tbody>
</table>

For the previous example, there is no line break in the response payload. Instead, a single JSON object contains a list of scores, one for each record in the request. Set the analysis configuration parameter probability to JMESPath expression "[*]" to extract the value.

Endpoint response is in JSON format and contains predicted label only

The following table is an example response from an endpoint that only outputs the predicted label.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;predicted_labels&quot;:[1]}'</td>
</tr>
<tr>
<td>Two records</td>
<td>'{&quot;predicted_labels&quot;:[1,0]}'</td>
</tr>
</tbody>
</table>
Set the label parameter of the predictor configuration to JMESPath expression "predicted_labels", and then the SageMaker Clarify processing job can extract the predicted labels for bias analysis.

**Endpoint response is JSON format and contains predicted label and probability**

The following table is an example response from an endpoint that outputs the predicted label and its score.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>'{&quot;predictions&quot;: [{&quot;label&quot;: 1, &quot;score&quot;: 0.6}]}'</td>
</tr>
<tr>
<td>Two records</td>
<td>'{&quot;predictions&quot;: [{&quot;label&quot;: 1, &quot;score&quot;: 0.6}, {&quot;label&quot;: 0, &quot;score&quot;: 0.3}]}'</td>
</tr>
</tbody>
</table>

For the previous example, set the label parameter of the predictor configuration to the JMESPath expression "predictions[*].label" to extract the predicted labels. Set probability to JMESPath expression "predictions[*].score" to extract the probability.

**Endpoint response is in JSON format and contains predicted labels and probabilities (multiclass)**

The following table is an example response from an endpoint that from a multiclass model that outputs the following:

- A list of predicted labels.
- Probabilities, and the selected predicted label and its probability.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single record</td>
<td>['{&quot;predicted_label&quot;: &quot;dog&quot;, &quot;probability&quot;: 0.6, &quot;predicted_labels&quot;: [&quot;cat&quot;, &quot;dog&quot;, &quot;fish&quot;], &quot;probabilities&quot;: [0.1, 0.6, 0.3]}]'</td>
</tr>
<tr>
<td>Endpoint request payload</td>
<td>Endpoint response payload (string representation)</td>
</tr>
<tr>
<td>--------------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>Two records</td>
<td>'[[&quot;predicted_label&quot;:&quot;dog&quot;,&quot;probability&quot;:0.6, &quot;predicted_labels&quot;:[&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;],&quot;probabilities&quot;:[0.1,0.6,0.3]],[&quot;predicted_label&quot;:&quot;cat&quot;,&quot;probability&quot;:0.7,&quot;predicted_labels&quot;:[&quot;cat&quot;,&quot;dog&quot;,&quot;fish&quot;],&quot;probabilities&quot;:[0.7,0.2,0.1]]]'</td>
</tr>
</tbody>
</table>

The SageMaker Clarify processing job can be configured in several ways to extract the predictions.

For bias analysis, the previous example can be configured as one of the following.

- Set the `label` parameter of the `predictor` configuration to JMESPath expression `"[*].predicted_label"` to extract the predicted label.
- Set the parameter to JMESPath expression `"[*].predicted_labels"` to extract the predicted labels. Set `probability` to JMESPath expression `"[*].probabilities"` to extract their probabilities. The SageMaker Clarify processing job can automatically determine the predicted label by identifying the label with the highest proximity value.
- Set `probability` to JMESPath expression `"[*].probabilities"` to extract their probabilities. If `label_headers` is provided, then the SageMaker Clarify processing job can automatically determine the predicted label by identifying the label with the highest probability value.

For feature importance analysis, set `probability` to JMESPath expression `"[*].probabilities"` to extract their probabilities of all the predicted labels. Then, feature attributions will be computed for all the labels.

**Pre-check endpoint request and response for tabular data**

We recommend that you deploy your model to a SageMaker real-time inference endpoint, and send requests to the endpoint. Manually examine the requests and responses to make sure that both are compliant with the requirements in the [Endpoint requests for tabular data](#) section and the [Endpoint response for tabular data](#) section. If your model container supports batch requests, you can start with a single record request, and then try two or more records.
The following commands show how to request a response using the AWS CLI. The AWS CLI is pre-installed in SageMaker Studio Classic, and SageMaker Notebook instances. If you must to install the AWS CLI, follow this [installation guide](#).

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name $ENDPOINT_NAME \
  --content-type $CONTENT_TYPE \
  --accept $ACCEPT_TYPE \
  --body $REQUEST_DATA \
  $CLI_BINARY_FORMAT \
  /dev/stderr 1>/dev/null
```

The parameters are defined, as follows.

- `$ENDPOINT_NAME` – The name of the endpoint.
- `$CONTENT_TYPE` – The MIME type of the request (model container input).
- `$ACCEPT_TYPE` – The MIME type of the response (model container output).
- `$REQUEST_DATA` – The requested payload string.
- `$CLI_BINARY_FORMAT` – The format of the command line interface (CLI) parameter. For AWS CLI v1, this parameter should remain blank. For v2, this parameter should be set to `--cli-binary-format raw-in-base64-out`.

**Note**

AWS CLI v2 passes binary parameters as base64-encoded strings by default.

The following request and response examples to and from the endpoint use AWS CLI v1.

**Endpoint request and response in CSV format**

In the following code example, the request consists of a single record and the response is its probability value.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-sagemaker-xgboost-model \
  --content-type text/csv \
  --accept text/csv \
  --body '1,2,3,4'
```
From the previous code example, the response output follows.

```
0.6
```

In the following code example, the request consists of two records, and the response includes their probabilities, which are separated by a comma.

```
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-sagemaker-xgboost-model \n  --content-type text/csv \n  --accept text/csv \n  --body $'1,2,3,4\n5,6,7,8' \n/dev/stderr 1>/dev/null
```

From the previous code example, the `$'content'` expression in the `--body` tells the command to interpret `'
'` in the content as a line break. The response output follows.

```
0.6,0.3
```

In the following code example, the request consists of two records, the response includes their probabilities, separated with a line break.

```
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-1 \n  --content-type text/csv \n  --accept text/csv \n  --body $'1,2,3,4\n5,6,7,8' \n/dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
0.6
0.3
```

In the following code example, the request consists of a single record, and the response is probability values from a multiclass model containing three classes.

```
aws sagemaker-runtime invoke-endpoint \
```

Configure a SageMaker Clarify Processing Job
```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-1 \
  --content-type text/csv \
  --accept text/csv \
  --body '1,2,3,4' \
  /dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
0.1,0.6,0.3
```

In the following code example, the request consists of two records, and the response includes their probability values from a multiclass model containing three classes.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-1 \
  --content-type text/csv \
  --accept text/csv \
  --body $'1,2,3,4\n5,6,7,8' \
  /dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
0.1,0.6,0.3
0.2,0.5,0.3
```

In the following code example, the request consists of two records, and the response includes predicted label and probability.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-2 \
  --content-type text/csv \
  --accept text/csv \
  --body $'1,2,3,4\n5,6,7,8' \
  /dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
1,0.6
0,0.3
```
In the following code example, the request consists of two records and the response includes label headers and probabilities.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-3 \
  --content-type text/csv \
  --accept text/csv \
  --body $'1,2,3,4
5,6,7,8' \
/dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
"['cat','dog','fish'],[0.1,0.6,0.3]
"['cat','dog','fish'],[0.2,0.5,0.3]
```

**Endpoint request and response in JSON Lines format**

In the following code example, the request consists of a single record and the response is its probability value.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-jsonlines \
  --content-type application/jsonlines \
  --accept application/jsonlines \
  --body '{"features":["This is a good product",5]}' \
/dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```json
{"score":0.6}
```

In the following code example, the request contains two records, and the response includes predicted label and probability.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-jsonlines-2 \
  --content-type application/jsonlines \
  --accept application/jsonlines \
  --body $'{"features":[1,2,3,4]}\n{"features":[5,6,7,8]}' \
/dev/stderr 1>/dev/null
```
From the previous code example, the response output follows.

```json
{"predicted_label":1,"probability":0.6}
{"predicted_label":0,"probability":0.3}
```

In the following code example, the request contains two records, and the response includes label headers and probabilities.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-jsonlines-3 \
  --content-type application/jsonlines \
  --accept application/jsonlines \
  --body $'{"data":{"features":[1,2,3,4]}\n{"data":{"features":[5,6,7,8]}}'}} \
/dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```json
{"predicted_labels":["cat","dog","fish"],"probabilities":[0.1,0.6,0.3]}
{"predicted_labels":["cat","dog","fish"],"probabilities":[0.2,0.5,0.3]}
```

**Endpoint request and response in mixed formats**

In the following code example, the request is in CSV format and the response is in JSON Lines format.

```bash
aws sagemaker-runtime invoke-endpoint \
  --endpoint-name test-endpoint-csv-in-jsonlines-out \
  --content-type text/csv \
  --accept application/jsonlines \
  --body $'1,2,3,4\n5,6,7,8' \
/dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```json
{"probability":0.6}
{"probability":0.3}
```

In the following code example, the request is in JSON Lines format and the response is in CSV format.

```bash
aws sagemaker-runtime invoke-endpoint \
```
From the previous code example, the response output follows.

```
0.6
0.3
```

In the following code example, the request is in CSV format and the response is in JSON format.

```
aws sagemaker-runtime invoke-endpoint \
   --endpoint-name test-endpoint-csv-in-jsonlines-out \
   --content-type text/csv \
   --accept application/jsonlines \
   --body $'1,2,3,4\n5,6,7,8' \
   /dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
{"predictions":[{"label":1,"score":0.6},{"label":0,"score":0.3}]}
```

### Image data

A SageMaker Clarify processing job provides support for explaining images. This topic provides the data format requirements for image data. For more information, see [computer vision](http://example.com/computer-vision).

#### Image dataset prerequisites

An image dataset contains one or more image files. To identify an input dataset to the SageMaker Clarify processing job, set either a [ProcessingInput](#) named dataset or the analysis configuration `dataset_uri` parameter to an Amazon S3 URI prefix of your image files.

The supported image file formats and file extensions are listed in the following table.

<table>
<thead>
<tr>
<th>Image format</th>
<th>File extension</th>
</tr>
</thead>
<tbody>
<tr>
<td>JPEG</td>
<td>jpg, jpeg</td>
</tr>
</tbody>
</table>
Set the analysis configuration `dataset_type` parameter to `application/x-image`. Because the type is not a specific image file format, the `content_type` will be used to decide the image file format and extension.

The SageMaker Clarify processing job loads each image file to a 3-dimensional NumPy array for further processing. The three dimensions include height, width, and RGB values of each pixel.

### Endpoint request for image data

The SageMaker Clarify processing job converts the raw RGB data of an image into a compatible image format, such as JPEG. It does this before it sends the data to the endpoint for predictions. The supported image formats are as follows.

<table>
<thead>
<tr>
<th>Data Format</th>
<th>MIME type</th>
<th>File extension</th>
</tr>
</thead>
<tbody>
<tr>
<td>JPEG</td>
<td>image/jpeg</td>
<td>jpg, jpeg</td>
</tr>
<tr>
<td>PNG</td>
<td>image/png</td>
<td>png</td>
</tr>
<tr>
<td>NPY</td>
<td>application/x-npy</td>
<td>All above</td>
</tr>
</tbody>
</table>

Specify the data format of the request payload by using the analysis configuration parameter `content_type`. If the `content_type` is not provided, the data format defaults to `image/jpeg`.

### Endpoint response for image data

Upon receiving the response of an inference endpoint invocation, the SageMaker Clarify processing job deserializes response payload and then extracts the predictions from it.

### Image classification problem

The data format of the response payload should be specified by the analysis configuration parameter `accept_type`. If `accept_type` is not provided, the data format defaults to `application/json`. The supported formats are the same as those described in the **Endpoint response for tabular data** in the tabular data section.
See Inference with the Image Classification Algorithm for an example of a SageMaker built-in image classification algorithm that accepts a single image and then returns an array of probability values (scores), each for a class.

As shown in the following table, when the content_type parameter is set to application/jsonlines, the response is a JSON object.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single image</td>
<td>'{&quot;prediction&quot;:[0.1,0.6,0.3]}'</td>
</tr>
</tbody>
</table>

In the previous example, set the probability parameter to JMESPath expression "prediction" to extract the scores.

When the content_type is set to application/json, the response is a JSON object, as shown in the following table.

<table>
<thead>
<tr>
<th>Endpoint request payload</th>
<th>Endpoint response payload (string representation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single image</td>
<td>'[0.1,0.6,0.3]'</td>
</tr>
</tbody>
</table>

In the previous example, set probability to JMESPath expression "[*]" to extract all the elements of the array. In the previous example, [0.1, 0.6, 0.3] is extracted. Alternatively, if you skip setting the probability configuration parameter, then all the elements of the array are also extracted. This is because the entire payload is deserialized as the predictions.

**Object detection problem**

The analysis configuration accept_type defaults to application/json and the only supported format is the Object Detection Inference Format. For more information about response formats, see Response Formats.

The following table is an example response from an endpoint that outputs an array. Each element of the array is an array of values containing the class index, the confidence score, and the bounding box coordinates of the detected object.
### Endpoint request payload

<table>
<thead>
<tr>
<th>Description</th>
<th>String representation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single image (one object)</td>
<td><code>&quot;prediction&quot;: [[4.0, 0.86419455409049988, 0.3088374733924866, 0.07030484080314636, 0.7110607028007507, 0.9345266819000244]]</code></td>
</tr>
<tr>
<td>Single image (two objects)</td>
<td><code>&quot;prediction&quot;: [[4.0, 0.86419455409049988, 0.3088374733924866, 0.07030484080314636, 0.7110607028007507, 0.9345266819000244], [0.0, 0.73376623392105103, 0.5714187026023865, 0.40427327156066895, 0.827075183391571, 0.9712159633636475]]</code></td>
</tr>
</tbody>
</table>

The following table is an example response from an endpoint that outputs a JSON object with a key referring to the array. Set the analysis configuration **probability** to the key "prediction" to extract the values.

### Endpoint request payload

<table>
<thead>
<tr>
<th>Description</th>
<th>String representation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single image (one object)</td>
<td><code>&quot;prediction&quot;: [[4.0, 0.86419455409049988, 0.3088374733924866, 0.07030484080314636, 0.7110607028007507, 0.9345266819000244]]</code></td>
</tr>
<tr>
<td>Single image (two objects)</td>
<td><code>&quot;prediction&quot;: [[4.0, 0.86419455409049988, 0.3088374733924866, 0.07030484080314636, 0.7110607028007507, 0.9345266819000244], [0.0, 0.73376623392105103, 0.5714187026023865, 0.40427327156066895, 0.827075183391571, 0.9712159633636475]]</code></td>
</tr>
</tbody>
</table>

Configure a SageMaker Clarify Processing Job 5436
Pre-check endpoint request and response for image data

We recommend that you deploy your model to a SageMaker real-time inference endpoint, and send requests to the endpoint. Manually examine the requests and responses. Make sure that both are compliant with the requirements in the Endpoint request for image data section and Endpoint response for image data section.

The following are two code examples showing how to send requests and examine the responses for both image classification and object detection problems.

Image classification problem

The following example code instructs an endpoint to read a PNG file and then classifies it.

```
aws sagemaker-runtime invoke-endpoint
   --endpoint-name test-endpoint-sagemaker-image-classification
   --content-type "image/png"
   --accept "application/json"
   --body fileb://./test.png
   /dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
[0.1,0.6,0.3]
```

Object detection problem

The following example code instructs an endpoint to read a JPEG file and then detects the objects in it.

```
aws sagemaker-runtime invoke-endpoint
   --endpoint-name test-endpoint-sagemaker-object-detection
   --content-type "image/jpg"
   --accept "application/json"
   --body fileb://./test.jpg
   /dev/stderr 1>/dev/null
```

From the previous code example, the response output follows.

```
{"prediction": [[4.0, 0.86419455409049988, 0.3088374733924866, 0.07030484080314636, 0.7110607028007507, 0.9345266819000244], [0.0, 0.73376623392105103, 0.5714187026023865], ...]}
```
Run SageMaker Clarify Processing Jobs for Bias Analysis and Explainability

To analyze your data and models for bias and explainability using SageMaker Clarify, you must configure a SageMaker Clarify processing job. This guide shows how to configure the job inputs, outputs, resources, and analysis configuration using the SageMaker Python SDK API SageMakerClarifyProcessor.

The API acts as a high-level wrapper of the SageMaker CreateProcessingJob API. It hides many of the details that are involved in setting up a SageMaker Clarify processing job. The details to set up a job include retrieving the SageMaker Clarify container image URI and generating the analysis configuration file. The following steps show you how to configure, initialize and launch a SageMaker Clarify processing job.

**Configure a SageMaker Clarify processing job using the API**

1. Define the configuration objects for each portion of the job configuration. These portions can include the following:

   - The input dataset and output location: `DataConfig`.
   - The model or endpoint to be analyzed: `ModelConfig`.
   - Bias analysis parameters: `BiasConfig`.
   - SHapley Additive exPlanations (SHAP) analysis parameters: `SHAPConfig`.

   The configuration objects for a SageMaker Clarify processing job vary for different types of data formats and use cases. Configuration examples for tabular data in CSV and JSON Lines format, natural language processing (NLP), and computer vision problems are provided in the following sections.

2. Create a `SageMakerClarifyProcessor` object and initialize it with parameters that specify the job resources. These resources include parameters such as the number of compute instances to use.
The following code example shows how to create a SageMakerClarifyProcessor object and instruct it to use one `ml.c4.xlarge` compute instance to do the analysis.

```python
from sagemaker import clarify

clarify_processor = clarify.SageMakerClarifyProcessor(
    role=role,
    instance_count=1,
    instance_type='ml.c4.xlarge',
    sagemaker_session=session,
)
```

3. Call the specific run method of the `SageMakerClarifyProcessor` object with the configuration objects for your use case to launch the job. These run methods include the following:

- `run_pre_training_bias`
- `run_post_training_bias`
- `run_bias`
- `run_explainability`
- `run_bias_and_explainability`

This `SageMakerClarifyProcessor` handles several tasks behind the scenes. These tasks include retrieving the SageMaker Clarify container image universal resource identifier (URI), composing an analysis configuration file based on the provided configuration objects, uploading the file to an Amazon S3 bucket, and configuring the SageMaker Clarify processing job.

The following expandable sections show how to compute **pre-training** and **post-training bias metrics**, **SHAP values**, and **partial dependence plots** (PDPs). The sections show feature importance for these data types:

- Tabular datasets in CSV format or JSON Lines format
- Natural language processing (NLP) datasets
- Computer vision datasets
A guide to run parallel SageMaker Clarify processing jobs with distributed training using Spark follows the expandable sections.

### Analyze tabular data in CSV format

The following examples show how to configure bias analysis and explainability analysis for a tabular dataset in CSV format. In these examples, the incoming dataset has four feature columns and one binary label column, Target. The contents of the dataset are as follows. A label value of 1 indicates a positive outcome.

<table>
<thead>
<tr>
<th>Target, Age, Gender, Income, Occupation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0, 25, 0, 2850, 2</td>
</tr>
<tr>
<td>1, 36, 0, 8585, 0</td>
</tr>
<tr>
<td>1, 22, 1, 1759, 1</td>
</tr>
<tr>
<td>0, 48, 0, 3446, 1</td>
</tr>
<tr>
<td>...</td>
</tr>
</tbody>
</table>

This DataConfig object specifies the input dataset and where to store the output. The s3_data_input_path parameter can either be a URI of a dataset file or an Amazon S3 URI prefix. If you provide a S3 URI prefix, the SageMaker Clarify processing job recursively collects all Amazon S3 files located under the prefix. The value for s3_output_path should be an S3 URI prefix to hold the analysis results. SageMaker uses the s3_output_path while compiling, and cannot take a value of a SageMaker Pipeline parameter, property, expression, or ExecutionVariable, which are used during runtime. The following code example shows how to specify a data configuration for the previous sample input dataset.

```python
data_config = clarify.DataConfig(
    s3_data_input_path=dataset_s3_uri,
    dataset_type='text/csv',
    headers=['Target', 'Age', 'Gender', 'Income', 'Occupation'],
    label='Target',
    s3_output_path=clarify_job_output_s3_uri,
)
```

### How to compute all pre-training bias metrics for a CSV dataset

The following code sample shows how to configure a BiasConfig object to measure bias of the previous sample input towards samples with a Gender value of 0.

```python
bias_config = clarify.BiasConfig(
    label_values_or_threshold=[1],
)`
The following code example shows how to use a run statement to launch a SageMaker Clarify processing job that computes all pre-training bias metrics for an input dataset.

```python
clarify_processor.run_pre_training_bias(
    data_config=data_config,
    data_bias_config=bias_config,
    methods="all",
)
```

Alternatively, you can choose which metrics to compute by assigning a list of pre-training bias metrics to the methods parameter. For example, replacing `methods="all"` with `methods=["CI", "DPL"]` instructs the SageMaker Clarify Processor to compute only Class Imbalance and Difference in Proportions of Labels.

### How to compute all post-training bias metrics for a CSV dataset

You can compute pre-training bias metrics prior to training. However, to compute post-training bias metrics, you must have a trained model. The following example output is from a binary classification model that outputs data in CSV format. In this example output, each row contains two columns. The first column contains the predicted label, and the second column contains the probability value for that label.

```
0,0.028986845165491
1,0.825382471084594
...
```

In the following example configuration, the ModelConfig object instructs the job to deploy the SageMaker model to an ephemeral endpoint. The endpoint uses one ml.m4.xlarge inference instance. Because the parameter content_type and accept_type are not set, they automatically use the value of the parameter dataset_type, which is text/csv.

```python
model_config = clarify.ModelConfig(
    model_name=your_model,
    instance_type='ml.m4.xlarge',
    instance_count=1,
)
```
The following configuration example uses a `ModelPredictedLabelConfig` object with a label index of 0. This instructs the SageMaker Clarify processing job to locate the predicted label in the first column of the model output. The Processing job uses zero-based indexing in this example.

```python
predicted_label_config = clarify.ModelPredictedLabelConfig(
    label=0,
)
```

Combined with the previous configuration example, the following code example launches a SageMaker Clarify processing job to compute all the post-training bias metrics.

```python
clarify_processor.run_post_training_bias(
    data_config=data_config,
    data_bias_config=bias_config,
    model_config=model_config,
    model_predicted_label_config=predicted_label_config,
    methods="all",
)
```

Similarly, you can choose which metrics to compute by assigning a list of post-training bias metrics to the `methods` parameter. For example, replace `methods="all"` with `methods=["DPPL", "DI"]` to compute only Difference in Positive Proportions in Predicted Labels and Disparate Impact.

**How to compute all bias metrics for a CSV dataset**

The following configuration example shows how to run all pre-training and post-training bias metrics in one SageMaker Clarify processing job.

```python
clarify_processor.run_bias(
    data_config=data_config,
    bias_config=bias_config,
    model_config=model_config,
    model_predicted_label_config=predicted_label_config,
    pre_training_methods="all",
    post_training_methods="all",
)
```

For an example notebook with instructions on how to run a SageMaker Clarify processing job in SageMaker Studio Classic to detect bias, see Fairness and Explainability with SageMaker Clarify.
How to compute SHAP values for a CSV dataset

SageMaker Clarify provides feature attributions using the KernelSHAP algorithm. SHAP analysis requires the probability value or score instead of predicted label, so this ModelPredictedLabelConfig object has probability index 1. This instructs the SageMaker Clarify processing job to extract the probability score from the second column of the model output (using zero-based indexing).

```python
probability_config = clarify.ModelPredictedLabelConfig(
    probability=1,
)
```

The SHAPConfig object provides SHAP analysis parameters. In this example, the SHAP baseline parameter is omitted and the value of the num_clusters parameter is 1. This instructs the SageMaker Clarify Processor to compute one SHAP baseline sample based on clustering the input dataset. If you want to choose the baseline dataset, see SHAP Baselines for Explainability.

```python
shap_config = clarify.SHAPConfig(
    num_clusters=1,
)
```

The following code example launches a SageMaker Clarify processing job to compute SHAP values.

```python
clarify_processor.run_explainability(
    data_config=data_config,
    model_config=model_config,
    model_scores=probability_config,
    explainability_config=shap_config,
)
```

For an example notebook with instructions on how to run a SageMaker Clarify processing job in SageMaker Studio Classic to compute SHAP values, see Fairness and Explainability with SageMaker Clarify.

How to compute partial dependence plots (PDPs) for a CSV dataset

PDPs show the dependence of the predicted target response on one or more input features of interest while holding all other features constant. An upward sloping line, or curve in the PDP, indicates that the relationship between the target and input feature(s) is positive, and the
steepness indicates the strength of the relationship. A downward sloping line or curve indicates that if an input feature decreases, the target variable increases. Intuitively, you can interpret the partial dependence as the response of the target variable to each input feature of interest.

The following configuration example is for using a `PDPConfig` object to instruct the SageMaker Clarify processing job to compute the importance of the `Income` feature.

```python
pdp_config = clarify.PDPConfig(  
    features=['Income'],  
    grid_resolution=10,  
)
```

In the previous example, the `grid_resolution` parameter divides the range of the `Income` feature values into 10 buckets. The SageMaker Clarify processing job will generate PDPs for `Income` split into 10 segments on the x-axis. The y-axis will show the marginal impact of `Income` on the target variable.

The following code example launches a SageMaker Clarify processing job to compute PDPs.

```python
clarify_processor.run_explainability(  
    data_config=data_config,  
    model_config=model_config,  
    model_scores=probability_config,  
    explainability_config=pdp_config,  
)
```

For an example notebook with instructions on how to run a SageMaker Clarify processing job in SageMaker Studio Classic to compute PDPs, see [Explainability with SageMaker Clarify - Partial Dependence Plots (PDP)](https://example.com).

### How to compute both SHAP values and PDPs for a CSV dataset

You can compute both SHAP values and PDPs in a single SageMaker Clarify processing job. In the following configuration example, the `top_k_features` parameter of a new `PDPConfig` object is set to 2. This instructs the SageMaker Clarify processing job to compute PDPs for the 2 features that have the largest global SHAP values.

```python
shap_pdp_config = clarify.PDPConfig(  
    top_k_features=2,  
    grid_resolution=10,  
)
```
The following code example launches a SageMaker Clarify processing job to compute both SHAP values and PDPs.

```python
clarify_processor.run_explainability(
    data_config=data_config,
    model_config=model_config,
    model_scores=probability_config,
    explainability_config=[shap_config, shap_pdp_config],
)
```

### Analyze tabular data in JSON Lines format

The following examples show how to configure bias analysis and explainability analysis for a tabular dataset in SageMaker JSON Lines dense format. See [JSONLINES Request Format](#) for more information. In these examples, the incoming dataset has the same data as the previous section, but they're in the JSON Lines format. Each line is a valid JSON object. The key `Features` points to an array of feature values, and the key `Label` points to the ground truth label.

```json
{"Features":[25,0,2850,2],"Label":0}
{"Features":[36,0,6585,0],"Label":1}
{"Features":[22,1,1759,1],"Label":1}
{"Features":[48,0,3446,1],"Label":0}
...
```

In the following configuration example, the `DataConfig` object specifies the input dataset and where to store the output.

```python
data_config = clarify.DataConfig(
    s3_data_input_path=jsonl_dataset_s3_uri,
    dataset_type='application/jsonlines',
    headers=['Age', 'Gender', 'Income', 'Occupation', 'Target'],
    label='Label',
    features='Features',
    s3_output_path=clarify_job_output_s3_uri,
)
```

In the previous configuration example, the `features` parameter is set to the `JMESPath` expression `Features` so that the SageMaker Clarify processing job can extract the array of features from each
record. The label parameter is set to JMESPath expression Label so that the SageMaker Clarify processing job can extract the ground truth label from each record. The s3_data_input_path parameter can either be a URI of a dataset file or an Amazon S3 URI prefix. If you provide a S3 URI prefix, the SageMaker Clarify processing job recursively collects all S3 files located under the prefix. The value for s3_output_path should be an S3 URI prefix to hold the analysis results. SageMaker uses the s3_output_path while compiling, and cannot take a value of a SageMaker Pipeline parameter, property, expression, or ExecutionVariable, which are used during runtime.

You must have a trained model to compute post-training bias metrics or feature importance. The following example is from a binary classification model that outputs JSON Lines data in the example's format. Each row of the model output is a valid JSON object. The key predicted_label points to the predicted label, and the key probability points to the probability value.

```json
{"predicted_label":0,"probability":0.028986845165491}
{"predicted_label":1,"probability":0.825382471084594}
...
```

In the following configuration example, a ModelConfig object instructs the SageMaker Clarify processing job to deploy the SageMaker model to an ephemeral endpoint. The endpoint uses one ml.m4.xlarge inference instance.

```python
model_config = clarify.ModelConfig(
    model_name=your_model,
    instance_type='ml.m4.xlarge',
    instance_count=1,
    content_template='{"Features":$features}',
)
```

In previous configuration example, the parameter content_type and accept_type are not set. Therefore, they automatically use the value of the dataset_type parameter of the DataConfig object, which is application/jsonlines. The SageMaker Clarify processing job uses the content_template parameter to compose the model input by replacing the $features placeholder by an array of features.

The following example configuration shows how to set the label parameter of the ModelPredictedLabelConfig object to the JMESPath expression predicted_label. This will extract the predicted label from the model output.
The following example configuration shows how to set the probability parameter of the ModelPredictedLabelConfig object to the JMESPath expression probability. This will extract the score from the model output.

```python
probability_config = clarify.ModelPredictedLabelConfig(
    probability='probability',
)
```

To compute bias metrics and feature importance for datasets in JSON Lines format, use the same run statements and configuration objects as the previous section for CSV datasets. You can run a SageMaker Clarify processing job in SageMaker Studio Classic to detect bias and compute feature importance. For instructions and an example notebook, see [Fairness and Explainability with SageMaker Clarify (JSON Lines Format)](https://docs.aws.amazon.com/sagemaker/latest/dg/clarify-fairness-explainability-jsonlines.html).

**Analyze tabular data for NLP explainability**

SageMaker Clarify supports explanations for natural language processing (NLP) models. These explanations help you understand which sections of text are the most important for your model predictions. You can explain either the model prediction for a single instance of the input dataset, or model predictions from the baseline dataset. To understand and visualize a model’s behavior, you can specify multiple levels of granularity. To do this, define the length of the text segment, such as its tokens, sentences, paragraphs.

SageMaker Clarify NLP explainability is compatible with both classification and regression models. You can also use SageMaker Clarify to explain your model's behavior on multi-modal datasets that contain text, categorical, or numerical features. NLP explainability for multi-modal datasets can help you understand how important each feature is to the model's output. SageMaker Clarify supports 62 languages and can handle text which includes multiple languages.

The following example shows an analysis configuration file for computing feature importance for NLP. In this example, the incoming dataset is a tabular dataset in CSV format, with one binary label column and two feature columns.

```text
0,2, "Flavor needs work"
```
The following configuration example shows how to specify an input dataset in CSV format and output data path using the DataConfig object.

```python
nlp_data_config = clarify.DataConfig(
    s3_data_input_path=nlp_dataset_s3_uri,
    dataset_type='text/csv',
    headers=['Target', 'Rating', 'Comments'],
    label='Target',
    s3_output_path=clarify_job_output_s3_uri,
)
```

In the previous configuration example, the `s3_data_input_path` parameter can either be a URI of a dataset file or an Amazon S3 URI prefix. If you provide a S3 URI prefix, the SageMaker Clarify processing job recursively collects all S3 files located under the prefix. The value for `s3_output_path` should be an S3 URI prefix to hold the analysis results. SageMaker uses the `s3_output_path` while compiling, and cannot take a value of a SageMaker Pipeline parameter, property, expression, or `ExecutionVariable`, which are used during runtime.

The following example output was created from a binary classification model trained on the previous input dataset. The classification model accepts CSV data, and it outputs a single score in between 0 and 1.

```
0.491656005382537
0.569582343101501
...
```

The following example shows how to configure the ModelConfig object to deploy a SageMaker model. In this example, an ephemeral endpoint deploys the model. This endpoint uses one `ml.g4dn.xlarge` inference instance equipped with a GPU, for accelerated inferencing.

```python
nlp_model_config = clarify.ModelConfig(
    model_name=your_nlp_model_name,
    instance_type='ml.g4dn.xlarge',
    instance_count=1,
)
The following example shows how to configure the `ModelPredictedLabelConfig` object to locate the probability (score) in the first column with an index of 0.

```python
probability_config = clarify.ModelPredictedLabelConfig(
    probability=0,
)
```

The following example SHAP configuration shows how to run a token-wise explainability analysis using a model and an input dataset in the English language.

```python
text_config = clarify.TextConfig(
    language='english',
    granularity='token',
)
nlp_shap_config = clarify.SHAPConfig(
    baseline=[[4, '[MASK]']],
    num_samples=100,
    text_config=text_config,
)
```

In the previous example, the `TextConfig` object activates the NLP explainability analysis. The `granularity` parameter indicates that the analysis should parse tokens. In English, each token is a word. For other languages, see the spaCy documentation for tokenization, which SageMaker Clarify uses for NLP processing. The previous example also shows how to use an average Rating of 4 to set an in-place SHAP baseline instance. A special mask token `[MASK]` is used to replace a token (word) in Comments.

In the previous example, if the instance is 2, "Flavor needs work", set the baseline to an average Rating of 4 with the following baseline.

```python
4, '[MASK]'
```

In the previous example, the SageMaker Clarify explainer iterates through each token and replaces it with the mask, as follows.

```python
2,"[MASK] needs work"
```
Then, the SageMaker Clarify explainer will send each line to your model for predictions. This is so that the explainer learns the predictions with and without the masked words. The SageMaker Clarify explainer then uses this information to compute the contribution of each token.

The following code example launches a SageMaker Clarify processing job to compute SHAP values.

```python
clarify_processor.run_explainability(
    data_config=nlp_data_config,
    model_config=nlp_model_config,
    model_scores=probability_config,
    explainability_config=nlp_shap_config,
)
```

For an example notebook with instructions on how to run a SageMaker Clarify processing job in SageMaker Studio Classic for NLP explainability analysis, see Explaining Text Sentiment Analysis Using SageMaker Clarify.

### Analyze image data for computer vision explainability

SageMaker Clarify generates heat maps that provide insights into how your computer vision models classify and detect objects in your images.

In the following configuration example, the input dataset consists of JPEG images.

```python
cv_data_config = clarify.DataConfig(
    s3_data_input_path=cv_dataset_s3_uri,
    dataset_type="application/x-image",
    s3_output_path=clarify_job_output_s3_uri,
)
```

In the previous configuration example, the DataConfig object contains an s3_data_input_path set to an Amazon S3 URI prefix. The SageMaker Clarify processing job recursively collects all image files located under the prefix. The s3_data_input_path parameter can either be a URI of a dataset file or an Amazon S3 URI prefix. If you provide a S3 URI prefix, the SageMaker Clarify processing job recursively collects all S3 files located under the prefix. The value for s3_output_path should be an S3 URI prefix to hold the analysis results. SageMaker uses the
s3_output_path while compiling, and cannot take a value of a SageMaker Pipeline parameter, property, expression, or ExecutionVariable, which are used during runtime.

**How to explain an image classification model**

The SageMaker Clarify processing job explains images using the KernelSHAP algorithm, which treats the image as a collection of super pixels. Given a dataset consisting of images, the processing job outputs a dataset of images where each image shows the heat map of the relevant super pixels.

The following configuration example shows how to configure an explainability analysis using a SageMaker image classification model. See [Image Classification - MXNet](#) for more information.

```python
ic_model_config = clarify.ModelConfig(
    model_name=your_cv_ic_model,
    instance_type="ml.p2.xlarge",
    instance_count=1,
    content_type="image/jpeg",
    accept_type="application/json",
)
```

In the previous configuration example, a model named `your_cv_ic_model` has been trained to classify the animals on input JPEG images. The `ModelConfig` object in the previous example instructs the SageMaker Clarify processing job to deploy the SageMaker model to an ephemeral endpoint. For accelerated inferencing, the endpoint uses one `ml.p2.xlarge` inference instance equipped with a GPU.

After a JPEG image is sent to an endpoint, the endpoint classifies it and returns a list of scores. Each score is for a category. The `ModelPredictedLabelConfig` object provides the name of each category, as follows.

```python
ic_prediction_config = clarify.ModelPredictedLabelConfig(
    label_headers=['bird', 'cat', 'dog'],
)
```

An example output for the previous input of ['bird','cat','dog'] could be 0.3,0.6,0.1, where 0.3 represents the confidence score for classifying an image as a bird.

The following example SHAP configuration shows how to generate explanations for an image classification problem. It uses an `ImageConfig` object to activate the analysis.
SageMaker Clarify extracts features using the Simple Linear Iterative Clustering (SLIC) method from scikit-learn library for image segmentation. The previous configuration example, the `model_type` parameter, indicates the type of image classification problem. The parameter `num_segments` estimates how many approximate number of segments will be labeled in the input image. The number of segments is then passed to the slic `n_segments` parameter.

Each segment of the image is considered a super-pixel, and local SHAP values are computed for each segment. The parameter `segment_compactness` determines the shape and size of the image segments that are generated by the scikit-image slic method. The sizes and shapes of the image segments are then passed to the slic `compactness` parameter.

The following code example launches a SageMaker Clarify processing job to generate heat maps for your images. Positive heat map values show that the feature increased the confidence score of detecting the object. Negative values indicate that the feature decreased the confidence score.

```python
clarify_processor.run_explainability(
    data_config=cv_data_config,
    model_config=ic_model_config,
    model_scores=ic_prediction_config,
    explainability_config=ic_shap_config,
)
```

For a sample notebook that uses SageMaker Clarify to classify images and explain its classification, see Explaining Image Classification with SageMaker Clarify.

### How to explain an object detection model

A SageMaker Clarify processing job can detect and classify objects in an image and then provide an explanation for the detected object. The process for explanation is as follows.
1. Image objects are first categorized into one of the classes in a specified collection. For example, if an object detection model can recognize cat, dog and fish, then these three classes are in a collection. This collection is specified by the `label_headers` parameter as follows.

   ```python
   clarify.ModelPredictedLabelConfig(
       label_headers=object_categories,
   )
   ```

2. The SageMaker Clarify processing job produces a confidence score for each object. A high confidence score indicates that it belongs to one of the classes in a specified collection. The SageMaker Clarify processing job also produces the coordinates of a bounding box that delimits the object. For more information about confidence scores and bounding boxes, see `Response Formats`.

3. SageMaker Clarify then provides an explanation for the detection of an object in the image scene. It uses the methods described in the `How to explain an image classification model` section.

In the following configuration example, a SageMaker object detection model `your_cv_od_model` is trained on JPEG images to identify the animals on them.

   ```python
   od_model_config = clarify.ModelConfig(
       model_name=your_cv_od_model,
       instance_type="ml.p2.xlarge",
       instance_count=1,
       content_type="image/jpeg",
       accept_type="application/json",
   )
   ```

The `ModelConfig` object in the previous configuration example instructs the SageMaker Clarify processing job to deploy the SageMaker model to an ephemeral endpoint. For accelerated imaging, this endpoint uses one `ml.p2.xlarge` inference instance equipped with a GPU.

In the following example configuration, the `ModelPredictedLabelConfig` object provides the name of each category for classification.

   ```python
   ic_prediction_config = clarify.ModelPredictedLabelConfig(
       label_headers=['bird', 'cat', 'dog'],
   )
   ```
The following example SHAP configuration shows how to generate explanations for an object detection.

```python
od_image_config = clarify.ImageConfig(
    model_type="OBJECT_DETECTION",
    num_segments=20,
    segment_compactness=5,
    max_objects=5,
    iou_threshold=0.5,
    context=1.0,
)

od_shap_config = clarify.SHAPConfig(
    num_samples=100,
    image_config=od_image_config,
)
```

In the previous example configuration, the `ImageConfig` object activates the analysis. The `model_type` parameter indicates that the type of problem is object detection. For a detailed description of the other parameters, see Configure the Analysis.

The following code example launches a SageMaker Clarify processing job to generate heat maps for your images. Positive heat map values show that the feature increased the confidence score of detecting the object. Negative values indicate that the feature decreased the confidence score.

```python
clarify_processor.run_explainability(
    data_config=cv_data_config,
    model_config=od_model_config,
    model_scores=od_prediction_config,
    explainability_config=od_shap_config,
)
```

For a sample notebook that uses SageMaker Clarify to detect objects in an image and explain its predictions, see Explaining object detection models with Amazon SageMaker Clarify.

### How to run parallel SageMaker Clarify processing jobs

When working with large datasets, you can use Apache Spark to increase the speed of your SageMaker Clarify processing jobs. Spark is a unified analytics engine for large-scale data
processing. When you request more than one instance per SageMaker Clarify processor, SageMaker Clarify uses the distributed computing capabilities from Spark.

The following configuration example shows how to use SageMakerClarifyProcessor to create a SageMaker Clarify processor with 5 compute instances. To run any jobs associated with the SageMakerClarifyProcessor, SageMaker Clarify using Spark distributed processing.

```python
from sagemaker import clarify

spark_clarify_processor = clarify.SageMakerClarifyProcessor(
    role=role,
    instance_count=5,
    instance_type='ml.c5.xlarge',
)
```

If you set the `save_local_shap_values` parameter of SHAPConfig to True, the SageMaker Clarify processing job saves the local SHAP value as multiple part files in the job output location.

To associate the local SHAP values to the input dataset instances, use the `joinsource` parameter of DataConfig. If you add more compute instances, we recommend that you also increase the `instance_count` of ModelConfig for the ephemeral endpoint. This prevents Spark workers' concurrent inference requests from overwhelming the endpoint. Specifically, we recommend that you use a one-to-one ratio of endpoint-to-processing instances.

### Get Analysis Results

This topic shows how to get analysis results that SageMaker Clarify generates. After the SageMaker Clarify processing job is finished, you can download the output files to inspect, or visualize the results in SageMaker Studio Classic.

The SageMaker Clarify processing job output directory contains the following files:

- `analysis.json` – A file that contains bias metrics and feature importance in JSON format.
- `report.ipynb` – A static notebook that contains code to help you visualize bias metrics and feature importance.
- `explanations_shap/out.csv` – A directory that is created and contains automatically generated files based on your specific analysis configurations. For example, if you activate the `save_local_shap_values` parameter, then per-instance local SHAP values will be saved to the `explanations_shap` directory. As another example, if your analysis configuration does
not contain a value for the SHAP baseline parameter, the SageMaker Clarify explainability job computes a baseline by clustering the input dataset. It then saves the generated baseline to the directory.

The following sections provide detailed information about the schema and the report that's generated by bias analysis, SHAP analysis, computer vision explainability analysis, and partial dependence plots (PDPs) analysis. If the configuration analysis contains parameters to compute multiple analyses, then the results are aggregated into one analysis and one report file.

Topics

- **Bias analysis**
- **SHAP analysis**
- **Computer vision (CV) explainability analysis**
- **Partial dependence plots (PDPs) analysis**

**Bias analysis**

Amazon SageMaker Clarify uses the terminology documented in [Amazon SageMaker Clarify Terms for Bias and Fairness](https://docs.aws.amazon.com/sagemaker/latest/dg/clarify-terms-for-bias-and-fairness.html) to discuss bias and fairness.

**Schema for the analysis file**

The analysis file is in JSON format and is organized into two sections: pre-training bias metrics and post-training bias metrics. The parameters for pre-training and post-training bias metrics are as follows.

- **pre_training_bias_metrics** – Parameters for pre-training bias metrics. For more information, see [Measure Pre-training Bias](https://docs.aws.amazon.com/sagemaker/latest/dg/clarify-measure-pre-training-bias.html) and [Configure the Analysis](https://docs.aws.amazon.com/sagemaker/latest/dg/clarify-configure-analysis.html).

- **label** – The ground truth label name defined by the `label` parameter of the analysis configuration.

- **label_value_or_threshold** – A string containing the label values or interval defined by the `label_values_or_threshold` parameter of the analysis configuration. For example, if value 1 is provided for binary classification problem, then the string will be 1. If multiple values [1, 2] are provided for multi-class problem, then the string will be 1, 2. If a threshold 40 is provided for regression problem, then the string will be an internal like (40, 68] in which 68 is the maximum value of the label in the input dataset.
• **facets** – The section contains several key-value pairs, where the key corresponds to the facet name defined by the `name_or_index` parameter of the facet configuration, and the value is an array of facet objects. Each facet object has the following members:

• **value_or_threshold** – A string containing the facet values or interval defined by the `value_or_threshold` parameter of the facet configuration.

• **metrics** – The section contains an array of bias metric elements, and each bias metric element has the following attributes:

  • **name** – The short name of the bias metric. For example, CI.
  • **description** – The full name of the bias metric. For example, Class Imbalance (CI).
  • **value** – The bias metric value, or JSON null value if the bias metric is not computed for a particular reason. The values ±∞ are represented as strings `∞` and `-∞` respectively.
  • **error** – An optional error message that explains why the bias metric was not computed.

• **post_training_bias_metrics** – The section contains the post-training bias metrics and it follows a similar layout and structure to the pre-training section. For more information, see [Measure Post-training Data and Model Bias](#).

The following is an example of an analysis configuration that will calculate both pre-training and post-training bias metrics.

```json
{
    "version": "1.0",
    "pre_training_bias_metrics": {
        "label": "Target",
        "label_value_or_threshold": "1",
        "facets": [
            "Gender": [
                "value_or_threshold": "0",
                "metrics": [
                    {
                        "name": "CDDL",
                        "description": "Conditional Demographic Disparity in Labels (CDDL)"
                    },
                    {
                        "name": "CI",
                        "description": "Class Imbalance (CI)"
                    }
                ]
            }
        ]
    },
    "post_training_bias_metrics": {
        "label": "Target",
        "label_value_or_threshold": "1",
        "facets": {
            "Gender": {
                "value_or_threshold": "0",
                "metrics": [
                    {
                        "name": "CDDL",
                        "description": "Conditional Demographic Disparity in Labels (CDDL)"
                    },
                    {
                        "name": "CI",
                        "description": "Class Imbalance (CI)"
                    }
                ]
            }
        }
    }
}
```
Bias analysis report

The bias analysis report includes several tables and diagrams that contain detailed explanations and descriptions. These include, but are not limited to, the distribution of label values, the distribution of facet values, high-level model performance diagram, a table of bias metrics, and their descriptions. For more information about bias metrics and how to interpret them, see the Learn How Amazon SageMaker Clarify Helps Detect Bias.

SHAP analysis

SageMaker Clarify processing jobs use the Kernel SHAP algorithm to compute feature attributions. The SageMaker Clarify processing job produces both local and global SHAP values. These help
to determine the contribution of each feature towards model predictions. Local SHAP values represent the feature importance for each individual instance, while global SHAP values aggregate the local SHAP values across all instances in the dataset. For more information about SHAP values and how to interpret them, see Feature Attributions that Use Shapley Values.

Schema for the SHAP analysis file

Global SHAP analysis results are stored in the explanations section of the analysis file, under the kernel_shap method. The different parameters of the SHAP analysis file are as follows:

- **explanations** – The section of the analysis file that contains the feature importance analysis results.
- **kernal_shap** – The section of the analysis file that contains the global SHAP analysis result.
  - **global_shap_values** – A section of the analysis file that contains several key-value pairs. Each key in the key-value pair represents a feature name from the input dataset. Each value in the key-value pair corresponds to the feature's global SHAP value. The global SHAP value is obtained by aggregating the per-instance SHAP values of the feature using the agg_method configuration. If the use_logit configuration is activated, then the value is calculated using the logistic regression coefficients, which can be interpreted as log-odds ratios.
- **expected_value** – The mean prediction of the baseline dataset. If the use_logit configuration is activated, then the value is calculated using the logistic regression coefficients.
- **global_top_shap_text** – (For NLP explainability analysis). A section of the analysis file that includes a set of key-value pairs. SageMaker Clarify processing jobs aggregate the SHAP values of each token and then select the top tokens based on their global SHAP values. The max_top_tokens configuration defines the number of tokens to be selected.
  
  Each of the selected top tokens has a key-value pair. The key in the key-value pair corresponds to a top token's text feature name. Each value in the key-value pair is the global SHAP values of the top token. For an example of a global_top_shap_text key-value pair, see the output that follows.

The following is example output from the SHAP analysis of a tabular dataset.

```json
{
  "version": "1.0",
```
The following is example output from the SHAP analysis of a text dataset. The output corresponding to the column Comments is an example of output that is generated after analysis of a text feature.

```json
{
    "version": "1.0",
    "explanations": {
        "kernel_shap": {
            "Target": {
                "global_shap_values": {
                    "Rating": 0.022486410860333206,
                    "Comments": 0.058612104851485144,
                    ...
                },
                "expected_value": 0.46700941970297033,
                "global_top_shap_text": {
                    "charming": 0.04127962903247833,
                    "brilliant": 0.02450240786522321,
                    "enjoyable": 0.024093569652715457,
                    ...
                }
            }
        }
    }
}
```
Schema for the generated baseline file

When a SHAP baseline configuration is not provided, the SageMaker Clarify processing job generates a baseline dataset. SageMaker Clarify uses a distance-based clustering algorithm to generate a baseline dataset from clusters created from the input dataset. The resulting baseline dataset is saved in a CSV file, located at explanations_shap/baseline.csv. This output file contains a header row and several instances based on the num_clusters parameter that is specified in the analysis configuration. The baseline dataset only consists of feature columns. An example of a baseline created by clustering the input dataset follows.

<table>
<thead>
<tr>
<th>Age,Gender,Income,Occupation</th>
</tr>
</thead>
<tbody>
<tr>
<td>35,0,2883,1</td>
</tr>
<tr>
<td>40,1,6178,2</td>
</tr>
<tr>
<td>42,0,4621,0</td>
</tr>
</tbody>
</table>

Schema for local SHAP values from tabular dataset explainability analysis

For tabular datasets, if a single compute instance is used, the SageMaker Clarify processing job saves the local SHAP values to a CSV file named explanations_shap/out.csv. If you use multiple compute instances, local SHAP values are saved to several CSV files in the explanations_shap directory.

An output file containing local SHAP values has a row containing the local SHAP values for each column that is defined by the headers. The headers follow the naming convention of Feature_Label where the feature name is appended by an underscore, followed by the name of the your target variable.

For multi-class problems, the feature names in the header vary first, then labels. For example, two features F1, F2, and two classes L1 and L2, in headers are F1_L1, F2_L1, F1_L2, and F2_L2. If the analysis configuration contains a value for the joinsource_name_or_index parameter, then the key column used in the join is appended to the end of the header name. This allows mapping of the local SHAP values to instances of the input dataset. An example of an output file containing SHAP values follows.

<table>
<thead>
<tr>
<th>Age_Target,Gender_Target,Income_Target,Occupation_Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.003937908,0.001388849,0.00242389,0.00274234</td>
</tr>
<tr>
<td>-0.0052784,0.017144491,0.004480645,-0.017144491</td>
</tr>
</tbody>
</table>
| ...
Schema for local SHAP values from NLP explainability analysis

For NLP explainability analysis, if a single compute instance is used, the SageMaker Clarify processing job saves local SHAP values to a JSON Lines file named explanations_shap/out.jsonl. If you use multiple compute instances, the local SHAP values are saved to several JSON Lines files in the explanations_shap directory.

Each file containing local SHAP values has several data lines, and each line is a valid JSON object. The JSON object has the following attributes:

- **explanations** – The section of the analysis file that contains an array of Kernel SHAP explanations for a single instance. Each element in the array has the following members:
  - **feature_name** – The header name of the features provided by the headers configuration.
  - **data_type** – The feature type inferred by the SageMaker Clarify processing job. Valid values for text features include numerical, categorical, and free_text (for text features).
  - **attributions** – A feature-specific array of attribution objects. A text feature can have multiple attribution objects, each for a unit defined by the granularity configuration. The attribution object has the following members:
    - **attribution** – A class-specific array of probability values.
    - **description** – (For text features) The description of the text units.
    - **partial_text** – The portion of the text explained by the SageMaker Clarify processing job.
    - **start_idx** – A zero-based index to identify the array location indicating the beginning of the partial text fragment.

The following is an example of a single line from a local SHAP values file, beautified to enhance its readability.

```json
{
    "explanations": [
        {
            "feature_name": "Rating",
            "data_type": "categorical",
            "attributions": [
                {
                    "attribution": [0.00342270632248735]
                }
            ]
        }
    ]
}```
SHAP analysis report

The SHAP analysis report provides a bar chart of a maximum of 10 top global SHAP values. The following chart example shows the SHAP values for the top 4 features.
Computer vision (CV) explainability analysis

SageMaker Clarify computer vision explainability takes a dataset consisting of images and treats each image as a collection of super pixels. After analysis, the SageMaker Clarify processing job outputs a dataset of images where each image shows the heat map of the super pixels.

The following example shows an input speed limit sign on the left and a heat map shows the magnitude of SHAP values on the right. These SHAP values were calculated by an image recognition Resnet-18 model that is trained to recognize German traffic signs. The German Traffic Sign Recognition Benchmark (GTSRB) dataset is provided in the paper Man vs. computer: Benchmarking machine learning algorithms for traffic sign recognition. In the example output, large positive values indicate that the super pixel has a strong positive correlation with the model prediction. Large negative values indicate that the super pixel has a strong negative correlation with the model prediction. The larger the absolute value of the SHAP value shown in the heat map, the stronger the relationship between the super pixel and model prediction.
Partial dependence plots (PDPs) analysis

Partial dependence plots show the dependence of the predicted target response on a set of input features of interest. These are marginalized over the values of all other input features and are referred to as the complement features. Intuitively, you can interpret the partial dependence as the target response, which is expected as a function of each input feature of interest.

Schema for the analysis file

The PDP values are stored in the explanations section of the analysis file under the pdp method. The parameters for explanations are as follows:

- **explanations** – The section of the analysis files that contains feature importance analysis results.
- **pdp** – The section of the analysis file that contains an array of PDP explanations for a single instance. Each element of the array has the following members:
  - **feature_name** – The header name of the features provided by the headers configuration.
  - **data_type** – The feature type inferred by the SageMaker Clarify processing job. Valid values for data_type include numerical and categorical.
  - **feature_values** – Contains the values present in the feature. If the data_type inferred by SageMaker Clarify is categorical, feature_values contains all of the unique values that the feature could be. If the data_type inferred by SageMaker Clarify is numerical, feature_values contains a list of the central value of generated buckets. The
grid_resolution parameter determines the number of buckets used to group the feature column values.

- **data_distribution** – An array of percentages, where each value is the percentage of instances that a bucket contains. The grid_resolution parameter determines the number of buckets. The feature column values are grouped into these buckets.

- **model_predictions** – An array of model predictions, where each element of the array is an array of predictions that corresponds to one class in the model's output.

- **label_headers** – The label headers provided by the label_headers configuration.

- **error** – An error message generated if the PDP values are not computed for a particular reason. This error message replaces the content contained in the feature_values, data_distributions, and model_predictions fields.

The following is example output from an analysis file containing a PDP analysis result.

```
{
  "version": "1.0",
  "explanations": {
    "pdp": [
      {
        "feature_name": "Income",
        "data_type": "numerical",
        "feature_values": [1046.9, 2454.7, 3862.5, 5270.2, 6678.0, 8085.9, 9493.6, 10901.5, 12309.3, 13717.1],
        "data_distribution": [0.32, 0.27, 0.17, 0.1, 0.045, 0.05, 0.01, 0.015, 0.01, 0.01],
        "model_predictions": [[0.69, 0.82, 0.82, 0.77, 0.77, 0.46, 0.46, 0.45, 0.41, 0.41]],
        "label_headers": ["Target"]
      },
      ...
    ]
  }
}
```

**PDP analysis report**

You can generate an analysis report containing a PDP chart for each feature. The PDP chart plots feature_values along the x-axis, and it plots model_predictions along the y-axis. For multi-
class models, `model_predictions` is an array, and each element of this array corresponds to one of the model prediction classes.

The following is an example of PDP chart for the feature Age. In the example output, the PDP shows the number of feature values that are grouped into buckets. The number of buckets is determined by `grid_resolution`. The buckets of feature values are plotted against model predictions. In this example, the higher feature values have the same model prediction values.

![PDP Chart for Age](image)

Troubleshoot SageMaker Clarify Processing Jobs

If you encounter failures with SageMaker Clarify processing jobs, consult the following scenarios to help identify the issue.

**Note**

The failure reason and exit message are intended to contain descriptive messages and exceptions, if encountered, during the run. A common reason for errors is that parameters

Troubleshoot Jobs
are either missing or not valid. If you encounter unclear, confusing, or misleading messages or are unable to find a solution, submit feedback.

Topics

- [Processing job fails to finish](#)
- [Processing job is taking too long to run](#)
- [Processing job finishes without results and you get a CloudWatch warning message](#)
- [Error message for invalid analysis configuration](#)
- [Bias metric computation fails for several or all metrics](#)
- [Mismatch between analysis config and dataset/model input/output](#)
- [Model returns 500 Internal Server Error or container falls back to per-record predictions due to model error](#)
- [Execution role is invalid](#)
- [Failed to download data](#)
- [Could not connect to SageMaker](#)

**Processing job fails to finish**

If the processing job fails to finish, you can try the following:

- Inspect the job logs directly in the notebook where you ran the job in. The job logs are located in the output of the notebook cell where you initiated the run.
- Inspect the job logs in CloudWatch.
- Add the following line in your notebook to describe the last processing job and look for the failure reason and exit message:
  
```
clarify_processor.jobs[-1].describe()
```
- Run the following AWS CLI command to describe the processing job and look for the failure reason and exit message:
  
```
aws sagemaker describe-processing-job --processing-job-name <processing-job-id>
```
Processing job is taking too long to run

If your processing job is taking too long to run, use the following ways to find the root cause.

Check to see if your resource configuration is sufficient to handle your computing load. To speed up your job, try the following:

- Use a larger instance type. SageMaker Clarify queries the model repeatedly, and a larger instance can significantly reduce your computation time. For a list of available instances, their memory sizes, bandwidth, and other performance details, see [Amazon SageMaker Pricing](https://aws.amazon.com/sagemaker/pricing/).

- Add more instances. SageMaker Clarify can use multiple instances to explain multiple input data points in parallel. To enable parallel computing, set your `instance_count` to more than 1 when you call `SageMakerClarifyProcessor`. For more information, see [How to run parallel SageMaker Clarify processing jobs](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-clarify-processing-job.html). If you increase your instance count, monitor the performance of your endpoint to check that it can deploy the increased load. For more information, see [Capture data from real-time endpoint](https://docs.aws.amazon.com/sagemaker/latest/dg/how-to-capture-endpoint-data.html).

- If you're computing SHapley Additive exPlanations (SHAP) values, reduce the `num_samples` parameter in your analysis configuration file. The number of samples directly affects the following:
  - The size of the synthetic datasets that are sent to your endpoint
  - Job runtime

Reducing the number of samples can also lead to reduced accuracy in estimating SHAP values. For more information, see [Configure the Analysis](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-clarify-processing-job.html).

Processing job finishes without results and you get a CloudWatch warning message

If the processing job finishes but no results are found, the CloudWatch logs produce a warning message that says Signal 15 received, cleaning up. This warning indicates that the job was stopped either because a customer request called the `StopProcessingJob` API, or that the job ran out of the allotted time for its completion. In the latter case, check the maximum runtime in the job configuration (`max_runtime_in_seconds`) and increase it as needed.
Error message for invalid analysis configuration

- If you get the error message Unable to load analysis configuration as JSON., this means that the analysis configuration input file for the processing job does not contain a valid JSON object. Check the validity of the JSON object using a JSON linter.
- If you get the error message Analysis configuration schema validation error., this means that the analysis configuration input file for the processing job contains unknown fields or invalid types for some field values. Review the configuration parameters in the file and cross-check them with the parameters listed in the analysis configuration file. For more information, see Configure the Analysis.

Bias metric computation fails for several or all metrics

If you receive one of the following error messages No Label values are present in the predicted Label Column, Positive Predicted Index Series contains all False values. or Predicted Label Column series data type is not the same as Label Column series., try the following:

- Check that the correct dataset is being used.
- Check whether the dataset size is too small; whether, for example, it contains only a few rows. This may cause the model outputs to have the same value or the data type is inferred incorrectly.
- Check if the label or facet is treated as continuous or categorical. SageMaker Clarify uses heuristics to determine the DataType. For post-training bias metrics, the data type returned by the model may not match what is in the dataset or SageMaker Clarify may not be able to transform it correctly.
- In the bias report, you should see a single value for categorical columns or an interval for continuous columns.
- For example, if a column has values 0.0 and 1.0 as floats, it will be treated as continuous even if there are too few unique values.

Mismatch between analysis config and dataset/model input/output

- Check that the baseline format in the analysis config is the same as dataset format.
- If you receive the error message Could not convert string to float., check that the format is correctly specified. It could also indicate that the model predictions have a different format than the label column or it could indicate that the configuration for the label or probabilities is incorrect.
• If you receive the error message Unable to locate the facet. or Headers must contain label. or Headers in config do not match with the number of columns in the dataset. or Feature names not found., check that the headers match the columns.

• If you receive the error message Data must contain features., check the content template for JSON Lines and compare it with the dataset sample if available.

**Model returns 500 Internal Server Error or container falls back to per-record predictions due to model error**

If you receive the error message Fallback to per-record prediction because of model error., this could indicate that model cannot handle the batch size, or be throttled, or just does not accept the input passed by the container due to serialization problems. You should review the CloudWatch logs for the SageMaker endpoint and look for error messages or tracebacks. For model throttling cases, it may help to use a different instance type or increasing the number of instances for the endpoint.

**Execution role is invalid**

This indicates that the role provided is incorrect or missing required permissions. Check the role and its permissions that were used to configure the processing job and verify the permission and trust policy for the role.

**Failed to download data**

This indicates that job inputs could not be downloaded for the job to start. Check the bucket name and permissions for the dataset and the configuration inputs.

**Could not connect to SageMaker**

This indicates that the job could not reach SageMaker service endpoints. Check the network configuration settings for the processing job and verify virtual private cloud (VPC) configuration.

**Sample notebooks**

The following sections contain notebooks to help you get started using SageMaker Clarify, to use it for special tasks, including those inside a distributed job, and for computer vision.
Getting started

The following sample notebooks show how to use SageMaker Clarify to get started with explainability and model bias tasks. These tasks include creating a processing job, training a machine learning (ML) model, and monitoring model predictions:

- **Explainability and bias detection with Amazon SageMaker Clarify** – Use SageMaker Clarify to create a processing job to detect bias and explain model predictions.
- **Monitoring bias drift and feature attribution drift Amazon SageMaker Clarify** – Use Amazon SageMaker Model Monitor to monitor bias drift and feature attribution drift over time.
- **How to read a dataset in JSON Lines format** into a SageMaker Clarify processing job.
- **Mitigate Bias, train another unbiased model, and put it in the model registry** – Use Synthetic Minority Over-sampling Technique (SMOTE) and SageMaker Clarify to mitigate bias, train another model, then put the new model into the model registry. This sample notebook also shows how to place the new model artifacts, including data, code and model metadata, into the model registry. This notebook is part of a series that shows how to integrate SageMaker Clarify into a SageMaker pipeline that is described in the [Architect and build the full machine learning lifecycle with AWS](https://aws.amazon.com/blogs/machine-learning/architect-and-build-the-full-machine-learning-lifecycle-with-aws) blog post.

Special cases

The following notebooks show you how to use a SageMaker Clarify for special cases including inside your own container and for natural language processing tasks:

- **Fairness and Explainability with SageMaker Clarify (Bring Your Own Container)** – Build your own model and container that can integrate with SageMaker Clarify to measure bias and generate an explainability analysis report. This sample notebook also introduces key terms and shows you how to access the report through SageMaker Studio Classic.
- **Fairness and Explainability with SageMaker Clarify Spark Distributed Processing** – Use distributed processing to run a SageMaker Clarify job that measures the pre-training bias of a dataset and the post-training bias of a model. This sample notebook also shows you how to obtain an explanation for the importance of the input features on the model output, and access the explainability analysis report through SageMaker Studio Classic.
- **Explainability with SageMaker Clarify - Partial Dependence Plots (PDP)** – Use SageMaker Clarify to generate PDPs and access a model explainability report.
- Explaining text sentiment analysis using SageMaker Clarify Natural language processing (NLP) explainability – Use SageMaker Clarify for text sentiment analysis.

- Use computer vision (CV) explainability for image classification and object detection.

These notebooks have been verified to run in Amazon SageMaker Studio Classic. If you need instructions on how to open a notebook in Studio Classic, see Create or Open an Amazon SageMaker Studio Classic Notebook. If you're prompted to choose a kernel, choose Python 3 (Data Science).

**Detect Pre-training Data Bias**

Algorithmic bias, discrimination, fairness, and related topics have been studied across disciplines such as law, policy, and computer science. A computer system might be considered biased if it discriminates against certain individuals or groups of individuals. The machine learning models powering these applications learn from data and this data could reflect disparities or other inherent biases. For example, the training data may not have sufficient representation of various demographic groups or may contain biased labels. The machine learning models trained on datasets that exhibit these biases could end up learning them and then reproduce or even exacerbate those biases in their predictions. The field of machine learning provides an opportunity to address biases by detecting them and measuring them at each stage of the ML lifecycle. You can use Amazon SageMaker Clarify to determine whether data used for training models encodes any bias.

Bias can be measured before training and after training, and monitored against baselines after deploying models to endpoints for inference. Pre-training bias metrics are designed to detect and measure bias in the raw data before it is used to train a model. The metrics used are model-agnostic because they do not depend on any model outputs. However, there are different concepts of fairness that require distinct measures of bias. Amazon SageMaker Clarify provides bias metrics to quantify various fairness criteria.

For additional information about bias metrics, see Learn How Amazon SageMaker Clarify Helps Detect Bias and Fairness Measures for Machine Learning in Finance.

**Amazon SageMaker Clarify Terms for Bias and Fairness**

SageMaker Clarify uses the following terminology to discuss bias and fairness.
Feature

An individual measurable property or characteristic of a phenomenon being observed, contained in a column for tabular data.

Label

Feature that is the target for training a machine learning model. Referred to as the observed label or observed outcome.

Predicted label

The label as predicted by the model. Also referred to as the predicted outcome.

Sample

An observed entity described by feature values and label value, contained in a row for tabular data.

Dataset

A collection of samples.

Bias

An imbalance in the training data or the prediction behavior of the model across different groups, such as age or income bracket. Biases can result from the data or algorithm used to train your model. For instance, if an ML model is trained primarily on data from middle-aged individuals, it may be less accurate when making predictions involving younger and older people.

Bias metric

A function that returns numerical values indicating the level of a potential bias.

Bias report

A collection of bias metrics for a given dataset, or a combination of a dataset and a model.

Positive label values

Label values that are favorable to a demographic group observed in a sample. In other words, designates a sample as having a positive result.

Negative label values

Label values that are unfavorable to a demographic group observed in a sample. In other words, designates a sample as having a negative result.
Group variable

Categorical column of the dataset that is used to form subgroups for the measurement of Conditional Demographic Disparity (CDD). Required only for this metric with regards to Simpson’s paradox.

Facet

A column or feature that contains the attributes with respect to which bias is measured.

Facet value

The feature values of attributes that bias might favor or disfavor.

Predicted probability

The probability, as predicted by the model, of a sample having a positive or negative outcome.

Sample Notebooks

Amazon SageMaker Clarify provides the following sample notebook for bias detection:

- **Explainability and bias detection with Amazon SageMaker Clarify** – Use SageMaker Clarify to create a processing job for detecting bias and explaining model predictions with feature attributions.

This notebook has been verified to run in Amazon SageMaker Studio only. If you need instructions on how to open a notebook in Amazon SageMaker Studio, see [Create or Open an Amazon SageMaker Studio Classic Notebook](#). If you’re prompted to choose a kernel, choose **Python 3 (Data Science)**.

Topics

- **Measure Pre-training Bias**
- **Generate Reports for Bias in Pre-training Data in SageMaker Studio**

Measure Pre-training Bias

Measuring bias in ML models is a first step to mitigating bias. Each measure of bias corresponds to a different notion of fairness. Even considering simple concepts of fairness leads to many different measures applicable in various contexts. For example, consider fairness with respect to age, and, for simplicity, that middle-aged and rest of the age groups are the two relevant demographics,
referred to as facets. In the case of an ML model for lending, we may want small business loans to be issued to equal numbers of both demographics. Or, when processing job applicants, we may want to see equal numbers of members of each demographic hired. However, this approach may assume that equal numbers of both age groups apply to these jobs, so we may want to condition on the number that apply. Further, we may want to consider not whether equal numbers apply, but whether we have equal numbers of qualified applicants. Or, we may consider fairness to be an equal acceptance rate of qualified applicants across both age demographics, or, an equal rejection rate of applicants, or both. You might use datasets with different proportions of data on the attributes of interest. This imbalance can conflate the bias measure you choose. The models might be more accurate in classifying one facet than in the other. Thus, you need to choose bias metrics that are conceptually appropriate for the application and the situation.

We use the following notation to discuss the bias metrics. The conceptual model described here is for binary classification, where events are labeled as having only two possible outcomes in their sample space, referred to as positive (with value 1) and negative (with value 0). This framework is usually extensible to multicategory classification in a straightforward way or to cases involving continuous valued outcomes when needed. In the binary classification case, positive and negative labels are assigned to outcomes recorded in a raw dataset for a favored facet $a$ and for a disfavored facet $d$. These labels $y$ are referred to as observed labels to distinguish them from the predicted labels $y'$ that are assigned by a machine learning model during the training or inferences stages of the ML lifecycle. These labels are used to define probability distributions $P_a(y)$ and $P_d(y)$ for their respective facet outcomes.

- **labels:**
  - $y$ represents the $n$ observed labels for event outcomes in a training dataset.
  - $y'$ represents the predicted labels for the $n$ observed labels in the dataset by a trained model.

- **outcomes:**
  - A positive outcome (with value 1) for a sample, such as an application acceptance.
    - $n^{(1)}$ is the number of observed labels for positive outcomes (acceptances).
    - $n'^{(1)}$ is the number of predicted labels for positive outcomes (acceptances).
  - A negative outcome (with value 0) for a sample, such as an application rejection.
    - $n^{(0)}$ is the number of observed labels for negative outcomes (rejections).
    - $n'^{(0)}$ is the number of predicted labels for negative outcomes (rejections).

- **facet values:**
  - facet $a$ – The feature value that defines a demographic that bias favors.
• \( n_a \) is the number of observed labels for the favored facet value: \( n_a = n_a^{(1)} + n_a^{(0)} \) the sum of the positive and negative observed labels for the value facet \( a \).

• \( n'_a \) is the number of predicted labels for the favored facet value: \( n'_a = n'_a^{(1)} + n'_a^{(0)} \) the sum of the positive and negative predicted outcome labels for the facet value \( a \). Note that \( n'_a = n_a \).

• facet \( d \) – The feature value that defines a demographic that bias disfavors.

• \( n_d \) is the number of observed labels for the disfavored facet value: \( n_d = n_d^{(1)} + n_d^{(0)} \) the sum of the positive and negative observed labels for the facet value \( d \).

• \( n'_d \) is the number of predicted labels for the disfavored facet value: \( n'_d = n'_d^{(1)} + n'_d^{(0)} \) the sum of the positive and negative predicted labels for the facet value \( d \). Note that \( n'_d = n_d \).

• probability distributions for outcomes of the labeled facet data outcomes:

  • \( P_a(y) \) is the probability distribution of the observed labels for facet \( a \). For binary labeled data, this distribution is given by the ratio of the number of samples in facet \( a \) labeled with positive outcomes to the total number, \( P_a(y^1) = n_a^{(1)}/ n_a \), and the ratio of the number of samples with negative outcomes to the total number, \( P_a(y^0) = n_a^{(0)}/ n_a \).

  • \( P_d(y) \) is the probability distribution of the observed labels for facet \( d \). For binary labeled data, this distribution is given by the number of samples in facet \( d \) labeled with positive outcomes to the total number, \( P_d(y^1) = n_d^{(1)}/ n_d \), and the ratio of the number of samples with negative outcomes to the total number, \( P_d(y^0) = n_d^{(0)}/ n_d \).

Models trained on data biased by demographic disparities might learn and even exacerbate them. To identify bias in the data before expending resources to train models on it, SageMaker Clarify provides data bias metrics that you can compute on raw datasets before training. All of the pretraining metrics are model-agnostic because they do not depend on model outputs and so are valid for any model. The first bias metric examines facet imbalance, but not outcomes. It determines the extent to which the amount of training data is representative across different facets, as desired for the application. The remaining bias metrics compare the distribution of outcome labels in various ways for facets \( a \) and \( d \) in the data. The metrics that range over negative values can detect negative bias. The following table contains a cheat sheet for quick guidance and links to the pretraining bias metrics.
### Pre-training Bias Metrics

<table>
<thead>
<tr>
<th>Bias metric</th>
<th>Description</th>
<th>Example question</th>
<th>Interpreting metric values</th>
</tr>
</thead>
</table>
| **Class Imbalance (CI)** | Measures the imbalance in the number of members between different facet values. | Could there be age-based biases due to not having enough data for the demographic outside a middle-aged facet? | Normalized range: [-1, +1]  
Interpretation:  
• Positive values indicate the facet \( a \) has more training samples in the dataset.  
• Values near zero indicate the facets are balanced in the number of training samples in the dataset.  
• Negative values indicate the facet \( d \) has more training samples in the dataset. |
| **Difference in Proportions of Labels (DPL)** | Measures the imbalance of positive outcomes between different facet values. | Could there be age-based biases in ML predictions due to biased labeling of facet values in the data? | Range for normalized binary & multivariate facet labels: [-1, +1]  
Range for continuous labels: \((-\infty, +\infty)\)  
Interpretation:  
• Positive values indicate facet |
<table>
<thead>
<tr>
<th>Bias metric</th>
<th>Description</th>
<th>Example question</th>
<th>Interpreting metric values</th>
</tr>
</thead>
</table>
| Kullback-Leibler Divergence (KL) | Measures how much the outcome distributions of different facets diverge from each other entropically. | How different are the distributions for loan application outcomes for different demographic groups? | Range for binary, multicategory, continuous: $[0, +\infty)$  
**Interpretation:**  
- Values near zero indicate the labels are similarly distributed.  
- Positive values indicate the label distributions diverge, the more positive the larger the divergence. |
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<th>Interpreting metric values</th>
</tr>
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<tr>
<td><strong>Jensen-Shannon Divergence (JS)</strong></td>
<td>Measures how much the outcome distributions of different facets diverge from each other entropically.</td>
<td>How different are the distributions for loan application outcomes for different demographic groups?</td>
<td>Range for binary, multicategory, continuous: $[0, +\infty)$</td>
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**Interpretation:**
- Values near zero indicate the labels are similarly distributed.
- Positive values indicate the label distributions diverge, the more positive the larger the divergence.

| L_p-norm (LP)                    | Measures a p-norm difference between distinct demographic distributions of the outcomes associated with different facets in a dataset. | How different are the distributions for loan application outcomes for different demographics? | Range for binary, multicategory, continuous: $[0, +\infty)$ |

**Interpretation:**
- Values near zero indicate the labels are similarly distributed.
- Positive values indicate the label distributions diverge, the more positive the larger the divergence.
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<tr>
<td><strong>Total Variation Distance (TVD)</strong></td>
<td>Measures half of the $L_1$-norm difference between distinct demographic distributions of the outcomes associated with different facets in a dataset.</td>
<td>How different are the distributions for loan application outcomes for different demographics?</td>
<td>Range for binary, multicategory, and continuous outcomes: $[0, +\infty)$</td>
</tr>
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</tr>
<tr>
<td><strong>Kolmogorov-Smirnov (KS)</strong></td>
<td>Measures maximum divergence between outcomes in distributions for different facets in a dataset.</td>
<td>Which college application outcomes manifest the greatest disparities by demographic group?</td>
<td>Range of KS values for binary, multiclass, and continuous outcomes: [0,+1]</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>• Values near zero indicate the labels were evenly distributed between facets in all outcome categories.</td>
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<tr>
<td></td>
<td></td>
<td></td>
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### Bias metric

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<tr>
<td><strong>Conditional Demographic Disparity (CDD)</strong></td>
<td>Measures the disparity of outcomes between different facets as a whole, but also by subgroups.</td>
<td>Do some groups have a larger proportion of rejections for college admission outcomes than their proportion of acceptances?</td>
<td>Range of CDD: [-1, +1]</td>
</tr>
<tr>
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<td></td>
<td>• Positive values indicate a outcomes where facet (d) is rejected more than accepted.</td>
</tr>
<tr>
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<td></td>
<td>• Near zero indicates no demographic disparity on average.</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>• Negative values indicate a outcomes where facet (a) is rejected more than accepted.</td>
</tr>
</tbody>
</table>

For additional information about bias metrics, see [Fairness Measures for Machine Learning in Finance](#).

## Topics

- [Class Imbalance (CI)](#)
- [Difference in Proportions of Labels (DPL)](#)
- [Kullback-Leibler Divergence (KL)](#)
- [Jensen-Shannon Divergence (JS)](#)
- [Lp-norm (LP)](#)
- [Total Variation Distance (TVD)](#)
- [Kolmogorov-Smirnov (KS)](#)
Class Imbalance (CI)

Class imbalance (CI) bias occurs when a facet value \( d \) has fewer training samples when compared with another facet \( a \) in the dataset. This is because models preferentially fit the larger facets at the expense of the smaller facets and so can result in a higher training error for facet \( d \). Models are also at higher risk of overfitting the smaller data sets, which can cause a larger test error for facet \( d \). Consider the example where a machine learning model is trained primarily on data from middle-aged individuals (facet \( a \)), it might be less accurate when making predictions involving younger and older people (facet \( d \)).

The formula for the (normalized) facet imbalance measure:

\[
CI = \frac{n_a - n_d}{n_a + n_d}
\]

Where \( n_a \) is the number of members of facet \( a \) and \( n_d \) the number for facet \( d \). Its values range over the interval \([-1, 1]\).

- Positive CI values indicate the facet \( a \) has more training samples in the dataset and a value of 1 indicates the data only contains members of the facet \( a \).
- Values of CI near zero indicate a more equal distribution of members between facets and a value of zero indicates a perfectly equal partition between facets and represents a balanced distribution of samples in the training data.
- Negative CI values indicate the facet \( d \) has more training samples in the dataset and a value of -1 indicates the data only contains members of the facet \( d \).
- CI values near either of the extremes values of -1 or 1 are very imbalanced and are at a substantial risk of making biased predictions.

If a significant facet imbalance is found to exist among the facets, you might want to rebalance the sample before proceeding to train models on it.

Difference in Proportions of Labels (DPL)

The difference in proportions of labels (DPL) compares the proportion of observed outcomes with positive labels for facet \( d \) with the proportion of observed outcomes with positive labels of facet \( a \) in a training dataset. For example, you could use it to compare the proportion of middle-aged individuals (facet \( a \)) and other age groups (facet \( d \)) approved for financial loans. Machine learning

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**Conditional Demographic Disparity (CDD)**
models try to mimic the training data decisions as closely as possible. So a machine learning model trained on a dataset with a high DPL is likely to reflect the same imbalance in its future predictions.

The formula for the difference in proportions of labels is as follows:

$$DPL = (q_a - q_d)$$

Where:

- $q_a = n_a^{(1)}/n_a$ is the proportion of facet $a$ who have an observed label value of 1. For example, the proportion of a middle-aged demographic who get approved for loans. Here $n_a^{(1)}$ represents the number of members of facet $a$ who get a positive outcome and $n_a$ the is number of members of facet $a$.
- $q_d = n_d^{(1)}/n_d$ is the proportion of facet $d$ who have an observed label value of 1. For example, the proportion of people outside the middle-aged demographic who get approved for loans. Here $n_d^{(1)}$ represents the number of members of the facet $d$ who get a positive outcome and $n_d$ the is number of members of the facet $d$.

If DPL is close enough to 0, then we say that demographic parity has been achieved.

For binary and multicategory facet labels, the DPL values range over the interval (-1, 1). For continuous labels, we set a threshold to collapse the labels to binary.

- Positive DPL values indicate that facet $a$ is has a higher proportion of positive outcomes when compared with facet $d$.
- Values of DPL near zero indicate a more equal proportion of positive outcomes between facets and a value of zero indicates perfect demographic parity.
- Negative DPL values indicate that facet $d$ has a higher proportion of positive outcomes when compared with facet $a$.

Whether or not a high magnitude of DPL is problematic varies from one situation to another. In a problematic case, a high-magnitude DPL might be a signal of underlying issues in the data. For example, a dataset with high DPL might reflect historical biases or prejudices against age-based demographic groups that would be undesirable for a model to learn.

**Kullback-Leibler Divergence (KL)**

The Kullback-Leibler divergence (KL) measures how much the observed label distribution of facet $a$, $P_a(y)$, diverges from distribution of facet $d$, $P_d(y)$. It is also known as the relative entropy of $P_a(y)$
with respect to $P_d(y)$ and quantifies the amount of information lost when moving from $P_a(y)$ to $P_d(y)$.

The formula for the Kullback-Leibler divergence is as follows:

$$KL(P_a \| P_d) = \sum_y P_a(y) \cdot \log[P_a(y)/P_d(y)]$$

It is the expectation of the logarithmic difference between the probabilities $P_a(y)$ and $P_d(y)$, where the expectation is weighted by the probabilities $P_a(y)$. This is not a true distance between the distributions as it is asymmetric and does not satisfy the triangle inequality. The implementation uses natural logarithms, giving KL in units of nats. Using different logarithmic bases gives proportional results but in different units. For example, using base 2 gives KL in units of bits.

For example, assume that a group of applicants for loans have a 30% approval rate (facet $d$) and that the approval rate for other applicants (facet $a$) is 80%. The Kullback-Leibler formula gives you the label distribution divergence of facet $a$ from facet $d$ as follows:

$$KL = 0.8 \cdot \ln(0.8/0.3) + 0.2 \cdot \ln(0.2/0.7) = 0.53$$

There are two terms in the formula here because labels are binary in this example. This measure can be applied to multiple labels in addition to binary ones. For example, in a college admissions scenario, assume an applicant may be assigned one of three category labels: $y_i = \{y_0, y_1, y_2\} = \{rejected, waitlisted, accepted\}$.

Range of values for the KL metric for binary, multicategory, and continuous outcomes is $[0, +\infty)$.

- Values near zero mean the outcomes are similarly distributed for the different facets.
- Positive values mean the label distributions diverge, the more positive the larger the divergence.

**Jensen-Shannon Divergence (JS)**

The Jensen-Shannon divergence (JS) measures how much the label distributions of different facets diverge from each other entropically. It is based on the Kullback-Leibler divergence, but it is symmetric.

The formula for the Jensen-Shannon divergence is as follows:

$$JS = \frac{1}{2} \cdot [KL(P_a \| P) + KL(P_d \| P)]$$

Where $P = \frac{1}{2}(P_a + P_d)$, the average label distribution across facets $a$ and $d$. 
The range of JS values for binary, multicategory, continuous outcomes is $[0, \ln(2))$.

- Values near zero mean the labels are similarly distributed.
- Positive values mean the label distributions diverge, the more positive the larger the divergence.

This metric indicates whether there is a big divergence in one of the labels across facets.

**L_p-norm (LP)**

The $L_p$-norm (LP) measures the p-norm distance between the facet distributions of the observed labels in a training dataset. This metric is non-negative and so cannot detect reverse bias.

The formula for the $L_p$-norm is as follows:

$$L_p(P_a, P_d) = (\sum ||P_a - P_d||^p)^{1/p}$$

Where the p-norm distance between the points $x$ and $y$ is defined as follows:

$$L_p(x, y) = (|x_1-y_1|^p + |x_2-y_2|^p + \ldots + |x_n-y_n|^p)^{1/p}$$

The 2-norm is the Euclidean norm. Assume you have an outcome distribution with three categories, for example, $y_i = \{y_0, y_1, y_2\} = \{\text{accepted}, \text{waitlisted}, \text{rejected}\}$ in a college admissions multicategory scenario. You take the sum of the squares of the differences between the outcome counts for facets $a$ and $d$. The resulting Euclidean distance is calculated as follows:

$$L_2(P_a, P_d) = [(n_a(0) - n_d(0))^2 + (n_a(1) - n_d(1))^2 + (n_a(2) - n_d(2))^2]^{1/2}$$

Where:

- $n_a(i)$ is number of the ith category outcomes in facet $a$: for example $n_a(0)$ is number of facet $a$ acceptances.
- $n_d(i)$ is number of the ith category outcomes in facet $d$: for example $n_d(2)$ is number of facet $d$ rejections.

The range of LP values for binary, multicategory, and continuous outcomes is $[0, \sqrt{2})$, where:

- Values near zero mean the labels are similarly distributed.
- Positive values mean the label distributions diverge, the more positive the larger the divergence.
Total Variation Distance (TVD)

The total variation distance data bias metric (TVD) is half the $L_1$-norm. The TVD is the largest possible difference between the probability distributions for label outcomes of facets $a$ and $d$. The $L_1$-norm is the Hamming distance, a metric used to compare two binary data strings by determining the minimum number of substitutions required to change one string into another. If the strings were to be copies of each other, it determines the number of errors that occurred when copying. In the bias detection context, TVD quantifies how many outcomes in facet $a$ would have to be changed to match the outcomes in facet $d$.

The formula for the Total variation distance is as follows:

$$\text{TVD} = \frac{1}{2} \cdot L_1(P_a, P_d)$$

For example, assume you have an outcome distribution with three categories, $y_i = \{y_0, y_1, y_2\} = \{\text{accepted, waitlisted, rejected}\}$, in a college admissions multicategory scenario. You take the differences between the counts of facets $a$ and $d$ for each outcome to calculate TVD. The result is as follows:

$$L_1(P_a, P_d) = |n_a^{(0)} - n_d^{(0)}| + |n_a^{(1)} - n_d^{(1)}| + |n_a^{(2)} - n_d^{(2)}|$$

Where:

- $n_a^{(i)}$ is number of the $i$th category outcomes in facet $a$: for example $n_a^{(0)}$ is number of facet $a$ acceptances.
- $n_d^{(i)}$ is number of the $i$th category outcomes in facet $d$: for example $n_d^{(2)}$ is number of facet $d$ rejections.

The range of TVD values for binary, multicategory, and continuous outcomes is $[0, 1)$, where:

- Values near zero mean the labels are similarly distributed.
- Positive values mean the label distributions diverge, the more positive the larger the divergence.

Kolmogorov-Smirnov (KS)

The Kolmogorov-Smirnov bias metric (KS) is equal to the maximum divergence between labels in the distributions for facets $a$ and $d$ of a dataset. The two-sample KS test implemented by SageMaker Clarify complements the other measures of label imbalance by finding the most imbalanced label.
The formula for the Kolmogorov-Smirnov metric is as follows:

\[ KS = \max(|P_a(y) - P_d(y)|) \]

For example, assume a group of applicants (facet \( a \)) to college are rejected, waitlisted, or accepted at 40%, 40%, 20% respectively and that these rates for other applicants (facet \( d \)) are 20%, 10%, 70%. Then the Kolmogorov-Smirnov bias metric value is as follows:

\[ KS = \max(|0.4-0.2|, |0.4-0.1|, |0.2-0.7|) = 0.5 \]

This tells us the maximum divergence between facet distributions is 0.5 and occurs in the acceptance rates. There are three terms in the equation because labels are multiclass of cardinality three.

The range of LP values for binary, multiclass, and continuous outcomes is \([0, +1]\), where:

- Values near zero indicate the labels were evenly distributed between facets in all outcome categories. For example, both facets applying for a loan got 50% of the acceptances and 50% of the rejections.
- Values near one indicate the labels for one outcome were all in one facet. For example, facet \( a \) got 100% of the acceptances and facet \( d \) got none.
- Intermittent values indicate relative degrees of maximum label imbalance.

**Conditional Demographic Disparity (CDD)**

The demographic disparity metric (DD) determines whether a facet has a larger proportion of the rejected outcomes in the dataset than of the accepted outcomes. In the binary case where there are two facets, men and women for example, that constitute the dataset, the disfavored one is labelled facet \( d \) and the favored one is labelled facet \( a \). For example, in the case of college admissions, if women applicants comprised 46% of the rejected applicants and comprised only 32% of the accepted applicants, we say that there is demographic disparity because the rate at which women were rejected exceeds the rate at which they are accepted. Women applicants are labelled facet \( d \) in this case. If men applicants comprised 54% of the rejected applicants and 68% of the accepted applicants, then there is not a demographic disparity for this facet as the rate of rejection is less that the rate of acceptance. Men applicants are labelled facet \( a \) in this case.

The formula for the demographic disparity for the less favored facet \( d \) is as follows:

\[ DD_d = n_d^{(0)}/n^{(0)} - n_d^{(1)}/n^{(1)} = P_d^{R}(y^{0}) - P_d^{A}(y^{1}) \]
Where:

- \( n^{(0)} = n_a^{(0)} + n_d^{(0)} \) is the total number of rejected outcomes in the dataset for the favored facet \( a \) and disadvantaged facet \( d \).
- \( n^{(1)} = n_a^{(1)} + n_d^{(1)} \) is the total number of accepted outcomes in the dataset for the favored facet \( a \) and disadvantaged facet \( d \).
- \( P_d^R(y^0) \) is the proportion of rejected outcomes (with value 0) in facet \( d \).
- \( P_d^A(y^1) \) is the proportion of accepted outcomes (value 1) in facet \( d \).

For the college admission example, the demographic disparity for women is \( DD_d = 0.46 - 0.32 = 0.14 \). For men \( DD_a = 0.54 - 0.68 = -0.14 \).

A conditional demographic disparity (CDD) metric that conditions DD on attributes that define a strata of subgroups on the dataset is needed to rule out Simpson's paradox. The regrouping can provide insights into the cause of apparent demographic disparities for less favored facets. The classic case arose in the case of Berkeley admissions where men were accepted at a higher rate overall than women. The statistics for this case were used in the example calculations of DD. However, when departmental subgroups were examined, women were shown to have higher admission rates than men when conditioned by department. The explanation was that women had applied to departments with lower acceptance rates than men had. Examining the subgrouped acceptance rates revealed that women were actually accepted at a higher rate than men for the departments with lower acceptance rates.

The CDD metric gives a single measure for all of the disparities found in the subgroups defined by an attribute of a dataset by averaging them. It is defined as the weighted average of demographic disparities (DD\(_i\)) for each of the subgroups, with each subgroup disparity weighted in proportion to the number of observations in contains. The formula for the conditional demographic disparity is as follows:

\[
CDD = \frac{1}{n} \sum_i n_i \cdot DD_i
\]

Where:

- \( \sum_i n_i = n \) is the total number of observations and \( n_i \) is the number of observations for each subgroup.
- \( DD_i = n_i^{(0)}/n^{(0)} - n_i^{(1)}/n^{(1)} = P_i^R(y^0) - P_i^A(y^1) \) is the demographic disparity for the ith subgroup.
The demographic disparity for a subgroup \( (DD_i) \) are the difference between the proportion of rejected outcomes and the proportion of accepted outcomes for each subgroup.

The range of DD values for binary outcomes for the full dataset DD\( d \) or for its conditionalized subgroups DD\( i \) is \([-1, +1]\).

- +1: when there no rejections in facet \( a \) or subgroup and no acceptances in facet \( d \) or subgroup
- Positive values indicate there is a demographic disparity as facet \( d \) or subgroup has a greater proportion of the rejected outcomes in the dataset than of the accepted outcomes. The higher the value the less favored the facet and the greater the disparity.
- Negative values indicate there is not a demographic disparity as facet \( d \) or subgroup has a larger proportion of the accepted outcomes in the dataset than of the rejected outcomes. The lower the value the more favored the facet.
- -1: when there are no rejections in facet \( d \) or subgroup and no acceptances in facet \( a \) or subgroup

If you don't condition on anything then CDD is zero if and only if DPL is zero.

This metric is useful for exploring the concepts of direct and indirect discrimination and of objective justification in EU and UK non-discrimination law and jurisprudence. For additional information, see Why Fairness Cannot Be Automated. This paper also contains the relevant data and analysis of the Berkeley admissions case that shows how conditionalizing on departmental admission rate subgroups illustrates Simpson's paradox.

**Generate Reports for Bias in Pre-training Data in SageMaker Studio**

SageMaker Clarify is integrated with Amazon SageMaker Data Wrangler, which can help you identify bias during data preparation without having to write your own code. Data Wrangler provides an end-to-end solution to import, prepare, transform, featurize, and analyze data with Amazon SageMaker Studio. For an overview of the Data Wrangler data prep workflow, see Prepare ML Data with Amazon SageMaker Data Wrangler.

You specify attributes of interest, such as gender or age, and SageMaker Clarify runs a set of algorithms to detect the presence of bias in those attributes. After the algorithm runs, SageMaker Clarify provides a visual report with a description of the sources and severity of possible bias so that you can plan steps to mitigate. For example, in a financial dataset that contains few examples of business loans to one age group as compared to others, SageMaker flags the imbalance so that you can avoid a model that disfavors that age group.
To analyze and report on data bias

To get started with Data Wrangler, see [Get Started with Data Wrangler](#).

1. In Amazon SageMaker Studio Classic, from the **Home** menu in the left panel, navigate to the **Data** node, then choose **Data Wrangler**. This opens the **Data Wrangler landing page** in Studio Classic.

2. Choose the **+ Import data** button to create a new flow.

3. In your flow page, from the **Import** tab, choose Amazon S3, navigate to your Amazon S3 bucket, find your dataset, then choose **Import**.

4. After you have imported your data, on the flow graph in the **Data flow** tab, choose the **+** sign to the right of the **Data types** node.

5. Choose **Add analysis**.

6. On the **Create Analysis** page, choose **Bias Report** for the **Analysis type**.

7. Configure the bias report by providing a report **Name**, the column to predict and whether it is a value or threshold, the column to analyze for bias (the facet) and whether it is a value or threshold.

8. Continue configuring the bias report by choosing the bias metrics.
Choose bias metrics

- Class imbalance (CI)
- Difference in Positive Proportions in Labels (DPL)
- JS divergence (JS)
- Conditional Demographic Disparity in Labels (CDDL)

To measure CDDL, select a column in the dataset to be used as the group variable.

Optional

Would you like to analyze additional metrics?

- Yes
- No

- Kullback-Liebler Divergence (KL)
- Lp-norm (LP)
- Total Variation Distance (TVD)
- Kolmogorov-Smirnov Distance (KS)

9. Choose **Check for bias** to generate and view the bias report. Scroll down to view all of the reports.
10. Choose the caret to the right of each bias metric description to see documentation that can help you interpret the significance of the metric values.

11. To view a table summary of the bias metric values, choose the **Table** toggle. To save the report, choose **Save** in the lower-right corner of the page. You can see the report on the flow graph in the **Data flow** tab. Double-click on the report to open it.

**Detect Post-training Data and Model Bias**

Post-training bias analysis can help reveal biases that might have emanated from biases in the data, or from biases introduced by the classification and prediction algorithms. These analyses take into consideration the data, including the labels, and the predictions of a model. You assess performance by analyzing predicted labels or by comparing the predictions with the observed target values in the data with respect to groups with different attributes. There are different notions of fairness, each requiring different bias metrics to measure.

There are legal concepts of fairness that might not be easy to capture because they are hard to detect. For example, the US concept of disparate impact that occurs when a group, referred to as a less favored facet $d$, experiences an adverse effect even when the approach taken appears to be fair. This type of bias might not be due to a machine learning model, but might still be detectable by post-training bias analysis.

Amazon SageMaker Clarify tries to ensure a consistent use of terminology. For a list of terms and their definitions, see **Amazon SageMaker Clarify Terms for Bias and Fairness**.
For additional information about post-training bias metrics, see Learn How Amazon SageMaker Clarify Helps Detect Bias and Fairness Measures for Machine Learning in Finance.

Measure Post-training Data and Model Bias

Amazon SageMaker Clarify provides eleven post-training data and model bias metrics to help quantify various conceptions of fairness. These concepts cannot all be satisfied simultaneously and the selection depends on specifics of the cases involving potential bias being analyzed. Most of these metrics are a combination of the numbers taken from the binary classification confusion matrices for the different demographic groups. Because fairness and bias can be defined by a wide range of metrics, human judgment is required to understand and choose which metrics are relevant to the individual use case, and customers should consult with appropriate stakeholders to determine the appropriate measure of fairness for their application.

We use the following notation to discuss the bias metrics. The conceptual model described here is for binary classification, where events are labeled as having only two possible outcomes in their sample space, referred to as positive (with value 1) and negative (with value 0). This framework is usually extensible to multcategory classification in a straightforward way or to cases involving continuous valued outcomes when needed. In the binary classification case, positive and negative labels are assigned to outcomes recorded in a raw dataset for a favored facet \( a \) and for a disfavored facet \( d \). These labels \( y \) are referred to as observed labels to distinguish them from the predicted labels \( y' \) that are assigned by a machine learning model during the training or inferences stages of the ML lifecycle. These labels are used to define probability distributions \( P_a(y) \) and \( P_d(y) \) for their respective facet outcomes.

- **labels:**
  - \( y \) represents the \( n \) observed labels for event outcomes in a training dataset.
  - \( y' \) represents the predicted labels for the \( n \) observed labels in the dataset by a trained model.

- **outcomes:**
  - A positive outcome (with value 1) for a sample, such as an application acceptance.
    - \( n^{(1)} \) is the number of observed labels for positive outcomes (acceptances).
    - \( n'^{(1)} \) is the number of predicted labels for positive outcomes (acceptances).
  - A negative outcome (with value 0) for a sample, such as an application rejection.
    - \( n^{(0)} \) is the number of observed labels for negative outcomes (rejections).
    - \( n'^{(0)} \) is the number of predicted labels for negative outcomes (rejections).

- **facet values:**
• facet \( a \) – The feature value that defines a demographic that bias favors.
  • \( n_a \) is the number of observed labels for the favored facet value: \( n_a = n_a^{(1)} + n_a^{(0)} \) the sum of the positive and negative observed labels for the value facet \( a \).
  • \( n'_a \) is the number of predicted labels for the favored facet value: \( n'_a = n'_a^{(1)} + n'_a^{(0)} \) the sum of the positive and negative predicted outcome labels for the facet value \( a \). Note that \( n'_a = n_a \).

• facet \( d \) – The feature value that defines a demographic that bias disfavors.
  • \( n_d \) is the number of observed labels for the disfavored facet value: \( n_d = n_d^{(1)} + n_d^{(0)} \) the sum of the positive and negative observed labels for the facet value \( d \).
  • \( n'_d \) is the number of predicted labels for the disfavored facet value: \( n'_d = n'_d^{(1)} + n'_d^{(0)} \) the sum of the positive and negative predicted labels for the facet value \( d \). Note that \( n'_d = n_d \).

• probability distributions for outcomes of the labeled facet data outcomes:
  • \( P_a(y) \) is the probability distribution of the observed labels for facet \( a \). For binary labeled data, this distribution is given by the ratio of the number of samples in facet \( a \) labeled with positive outcomes to the total number, \( P_a(y^1) = n_a^{(1)} / n_a \), and the ratio of the number of samples with negative outcomes to the total number, \( P_a(y^0) = n_a^{(0)} / n_a \).
  • \( P_d(y) \) is the probability distribution of the observed labels for facet \( d \). For binary labeled data, this distribution is given by the number of samples in facet \( d \) labeled with positive outcomes to the total number, \( P_d(y^1) = n_d^{(1)} / n_d \), and the ratio of the number of samples with negative outcomes to the total number, \( P_d(y^0) = n_d^{(0)} / n_d \).

The following table contains a cheat sheet for quick guidance and links to the post-training bias metrics.

**Post-training bias metrics**

<table>
<thead>
<tr>
<th>Post-training bias metric</th>
<th>Description</th>
<th>Example question</th>
<th>Interpreting metric values</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Difference in Positive Proportions in Predicted Labels (DPPL)</strong></td>
<td>Measures the difference in the proportion of positive predictions between the favored facet ( a ) and the disfavored facet ( d ).</td>
<td>Has there been an imbalance across demographic groups in the predicted positive outcomes that might indicate bias?</td>
<td>Range for normalized binary &amp; multiclass labels: ([-1, +1]) and continuous labels: ((-\infty, +\infty))</td>
</tr>
</tbody>
</table>
**Post-training bias metric** | **Description** | **Example question** | **Interpreting metric values**
--- | --- | --- | ---

**Interpretation:**
- Positive values indicate that the favored facet $a$ has a higher proportion of predicted positive outcomes.
- Values near zero indicate a more equal proportion of predicted positive outcomes between facets.
- Negative values indicate the disfavored facet $d$ has a higher proportion of predicted positive outcomes.
<table>
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</table>
| Disparate Impact (DI)     | Measures the ratio of proportions of the predicted labels for the favored facet $a$ and the disfavored facet $d$. | Has there been an imbalance across demographic groups in the predicted positive outcomes that might indicate bias? | Range for normalized binary, multicategory facet, and continuous labels: $[0,\infty)$  
Interpretation:  
• Values less than 1 indicate the favored facet $a$ has a higher proportion of predicted positive outcomes.  
• A value of 1 indicates that we have demographic parity.  
• Values greater than 1 indicate the disfavored facet $d$ has a higher proportion of predicted positive outcomes. |
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<tr>
<td><strong>Conditional Demographic Disparity in Predicted Labels (CDDPL)</strong></td>
<td>Measures the disparity of predicted labels between the facets as a whole, but also by subgroups.</td>
<td>Do some demographic groups have a larger proportion of rejections for loan application outcomes than their proportion of acceptances?</td>
<td>The range of CDDPL values for binary, multicategory, and continuous outcomes: [-1, +1]</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>• Positive values indicate outcomes where facet d is rejected more than accepted.</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>• Near zero indicates no demographic disparity on average.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Negative values indicate outcomes where facet a is rejected more than accepted.</td>
</tr>
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</tbody>
</table>
| **Counterfactual Fliptest (FT)** | Examines each member of facet $d$ and assesses whether similar members of facet $a$ have different model predictions. | Is one group of a specific-age demographic matched closely on all features with a different age group, yet paid more on average? | The range for binary and multicategory facet labels is $[-1, +1]$.  
• Positive values occur when the number of unfavorable counterfactual fliptest decisions for the disfavored facet $d$ exceeds the favorable ones.  
• Values near zero occur when the number of unfavorable and favorable counterfactual fliptest decisions balance out.  
• Negative values occur when the number of unfavorable counterfactual fliptest decisions for the disfavored facet $d$ is less than the favorable ones. |
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| **Accuracy Difference** *(AD)* | Measures the difference between the prediction accuracy for the favored and disfavored facets. | Does the model predict labels as accurately for applications across all demographic groups? | The range for binary and multiclass facet labels is \([-1, +1]\).  
- Positive values indicate that facet \(d\) suffers more from some combination of false positives (Type I errors) or false negatives (Type II errors). This means there is a potential bias against the disfavored facet \(d\).  
- Values near zero occur when the prediction accuracy for facet \(a\) is similar to that for facet \(d\).  
- Negative values indicate that facet \(a\) suffers more from some combination of false positives (Type I errors) or false negatives (Type II errors). This means there is a bias against the favored facet \(a\). |
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<tr>
<td><strong>Recall Difference (RD)</strong></td>
<td>Compares the recall of the model for the favored and disfavored facets.</td>
<td>Is there an age-based bias in lending due to a model having higher recall for one age group as compared to another?</td>
<td>Range for binary and multicategory classification: ([-1, +1]).&lt;br&gt;• Positive values suggest that the model finds more of the true positives for facet (a) and is biased against the disfavored facet (d).&lt;br&gt;• Values near zero suggest that the model finds about the same number of true positives in both facets and is not biased.&lt;br&gt;• Negative values suggest that the model finds more of the true positives for facet (d) and is biased against the favored facet (a).</td>
</tr>
<tr>
<td>Post-training bias metric</td>
<td>Description</td>
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</tbody>
</table>
| **Difference in Conditional Acceptance (DCAcc)** | Compares the observed labels to the labels predicted by a model. Assesses whether this is the same across facets for predicted positive outcomes (acceptances). | When comparing one age group to another, are loans accepted more frequently, or less often than predicted (based on qualifications)? | The range for binary, multicategory facet, and continuous labels: \((-\infty, +\infty)\).  
• Positive values indicate a possible bias against the qualified applicants from the disfavored facet \(d\).  
• Values near zero indicate that qualified applicants from both facets are being accepted in a similar way.  
• Negative values indicate a possible bias against the qualified applicants from the favored facet \(a\). |
<table>
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</table>
| **Difference in Acceptance Rates (DAR)** | Measures the difference in the ratios of the observed positive outcomes (TP) to the predicted positives (TP + FP) between the favored and disfavored facets. | Does the model have equal precision when predicting loan acceptances for qualified applicants across all age groups? | The range for binary, multicategory facet, and continuous labels is $[-1, +1]$.  
  - Positive values indicate a possible bias against facet $d$ caused by the occurrence of relatively more false positives in the disfavored facet $d$.  
  - Values near zero indicate the observed labels for positive outcomes (acceptances) are being predicted with equal precision for both facets by the model.  
  - Negative values indicate a possible bias against facet $a$ caused by the occurrence of relatively more false positives in the favored facet $a$. |
<table>
<thead>
<tr>
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</table>
| **Specificity difference (SD)** | Compares the specificity of the model between favored and disfavored facets. | Is there an age-based bias in lending because the model predicts a higher specificity for one age group as compared to another? | Range for binary and multicategory classification: \([-1, +1]\).  
  - Positive values suggest that the model finds less false positives for facet \(d\) and is biased against the disfavored facet \(d\).  
  - Values near zero suggest that the model finds a similar number of false positives in both facets and is not biased.  
  - Negative values suggest that the model finds less false positives for facet \(a\) and is biased against the favored facet \(a\). |
<table>
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</table>
| Difference in Conditional Rejection (DCR) | Compares the observed labels to the labels predicted by a model and assesses whether this is the same across facets for negative outcomes (rejections). | Are there more or less rejections for loan applications than predicted for one age group as compared to another based on qualifications? | The range for binary, multicategory facet, and continuous labels: $(-\infty, +\infty)$.  
- Positive values indicate a possible bias against the qualified applicants from the disfavored facet $d$.  
- Values near zero indicate that qualified applicants from both facets are being rejected in a similar way.  
- Negative values indicate a possible bias against the qualified applicants from the favored facet $a$. |
<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td><strong>Difference in Rejection Rates (DRR)</strong></td>
<td>Measures the difference in the ratios of the observed negative outcomes (TN) to the predicted negatives (TN + FN) between the disfavored and favored facets.</td>
<td>Does the model have equal precision when predicting loan rejections for unqualified applicants across all age groups?</td>
<td>The range for binary, multicategory facet, and continuous labels is ([-1, +1]).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Positive values indicate a possible bias caused by the occurrence of relatively more false negatives in the favored facet.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Values near zero indicate that negative outcomes (rejections) are being predicted with equal precision for both facets.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Negative values indicate a possible bias caused by the occurrence of relatively more false negatives in the disfavored facet.</td>
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</tr>
<tr>
<td>Treatment Equality (TE)</td>
<td>Measures the difference in the ratio of false positives to false negatives between the favored and disfavored facets.</td>
<td>In loan applications, is the relative ratio of false positives to false negatives the same across all age demographics?</td>
<td>The range for binary and multicategory facet labels: ((-\infty, +\infty)).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Positive values occur when the ratio of false positives to false negatives for facet (a) is greater than that for facet (d).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Values near zero occur when the ratio of false positives to false negatives for facet (a) is similar to that for facet (d).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Negative values occur when the ratio of false positives to false negatives for facet (a) is less than that for facet (d).</td>
</tr>
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</tr>
<tr>
<td><strong>Generalized entropy (GE)</strong></td>
<td>Measures the inequality in benefits assigned to each input by the model predictions.</td>
<td>Of two candidate models for loan application classification, does one lead to a more uneven distribution of desired outcomes than the other?</td>
<td>The range for binary and multicategory labels: (0, 0.5). GE is undefined when the model predicts only false negatives.</td>
</tr>
</tbody>
</table>

- Zero values occur when all predictions are correct or all predictions are false positives.
- Positive values indicate inequality in benefits; 0.5 corresponds to the largest inequality.

For additional information about post-training bias metrics, see [A Family of Fairness Measures for Machine Learning in Finance](#).

**Topics**

- [Difference in Positive Proportions in Predicted Labels (DPPL)](#)
- [Disparate Impact (DI)](#)
- [Difference in Conditional Acceptance (DCAcc)](#)
- [Difference in Conditional Rejection (DCR)](#)
- [Specificity difference (SD)](#)
- [Recall Difference (RD)](#)
- [Difference in Acceptance Rates (DAR)](#)
- [Difference in Rejection Rates (DRR)](#)
- [Accuracy Difference (AD)](#)
• **Treatment Equality (TE)**
• **Conditional Demographic Disparity in Predicted Labels (CDDPL)**
• **Counterfactual Fliptest (FT)**
• **Generalized entropy (GE)**

### Difference in Positive Proportions in Predicted Labels (DPPL)

The difference in positive proportions in predicted labels (DPPL) metric determines whether the model predicts outcomes differently for each facet. It is defined as the difference between the proportion of positive predictions ($y' = 1$) for facet $a$ and the proportion of positive predictions ($y' = 1$) for facet $d$. For example, if the model predictions grant loans to 60% of a middle-aged group (facet $a$) and 50% other age groups (facet $d$), it might be biased against facet $d$. In this example, you must determine whether the 10% difference is material to a case for bias.

A comparison of difference in proportions of labels (DPL), a measure of pre-training bias, with DPPL, a measure of post-training bias, assesses whether bias in positive proportions that are initially present in the dataset changes after training. If DPPL is larger than DPL, then bias in positive proportions increased after training. If DPPL is smaller than DPL, the model did not increase bias in positive proportions after training. Comparing DPL against DPPL does not guarantee that the model reduces bias along all dimensions. For example, the model may still be biased when considering other metrics such as **Counterfactual Fliptest (FT)** or **Accuracy Difference (AD)**. For more information about bias detection, see the blog post [Learn how Amazon SageMaker Clarify helps detect bias](#). See [Difference in Proportions of Labels (DPL)](#) for more information about DPL.

The formula for the DPPL is:

$$DPPL = q'_a - q'_d$$

Where:

- $q'_a = n'_a^{(1)}/n_a$ is the predicted proportion of facet $a$ who get a positive outcome of value 1. In our example, the proportion of a middle-aged facet predicted to get granted a loan. Here $n'_a^{(1)}$ represents the number of members of facet $a$ who get a positive predicted outcome of value 1 and $n_a$ the is number of members of facet $a$.
- $q'_d = n'_d^{(1)}/n_d$ is the predicted proportion of facet $d$ who get a positive outcome of value 1. In our example, a facet of older and younger people predicted to get granted a loan. Here $n'_d^{(1)}$
represents the number of members of facet $d$ who get a positive predicted outcome and $n_d$ the is number of members of facet $d$.

If DPPL is close enough to 0, it means that post-training *demographic parity* has been achieved.

For binary and multicategory facet labels, the normalized DPL values range over the interval $[-1, 1]$. For continuous labels, the values vary over the interval $(-\infty, +\infty)$.

- Positive DPPL values indicate that facet $a$ has a higher proportion of predicted positive outcomes when compared with facet $d$.

  This is referred to as *positive bias*.

- Values of DPPL near zero indicate a more equal proportion of predicted positive outcomes between facets $a$ and $d$ and a value of zero indicates perfect demographic parity.

- Negative DPPL values indicate that facet $d$ has a higher proportion of predicted positive outcomes when compared with facet $a$. This is referred to as *negative bias*.

### Disparate Impact (DI)

The difference in positive proportions in the predicted labels metric can be assessed in the form of a ratio.

The comparison of positive proportions in predicted labels metric can be assessed in the form of a ratio instead of as a difference, as it is with the Difference in Positive Proportions in Predicted Labels (DPPL). The disparate impact (DI) metric is defined as the ratio of the proportion of positive predictions ($y' = 1$) for facet $d$ over the proportion of positive predictions ($y' = 1$) for facet $a$. For example, if the model predictions grant loans to 60% of a middle-aged group (facet $a$) and 50% other age groups (facet $d$), then DI = $.5/.6 = 0.8$, which indicates a positive bias and an adverse impact on the other aged group represented by facet $d$.

The formula for the ratio of proportions of the predicted labels:

$$DI = \frac{q'_d}{q'_a}$$

Where:

- $q'_a = n'_a/\sum n_a$ is the predicted proportion of facet $a$ who get a positive outcome of value 1. In our example, the proportion of a middle-aged facet predicted to get granted a loan. Here $n'_a$
represents the number of members of facet $a$ who get a positive predicted outcome and $n_a$ the is number of members of facet $a$.

- $q'_d = \frac{n'_d(1)}{n_d}$ is the predicted proportion of facet $d$ a who get a positive outcome of value 1. In our example, a facet of older and younger people predicted to get granted a loan. Here $n'_d(1)$ represents the number of members of facet $d$ who get a positive predicted outcome and $n_d$ the is number of members of facet $d$.

For binary, multicategory facet, and continuous labels, the DI values range over the interval $[0, \infty)$.

- Values less than 1 indicate that facet $a$ has a higher proportion of predicted positive outcomes than facet $d$. This is referred to as *positive bias*.
- A value of 1 indicates demographic parity.
- Values greater than 1 indicate that facet $d$ has a higher proportion of predicted positive outcomes than facet $a$. This is referred to as *negative bias*.

**Difference in Conditional Acceptance (DCAcc)**

This metric compares the observed labels to the labels predicted by the model and assesses whether this is the same across facets for predicted positive outcomes. This metric comes close to mimicking human bias in that it quantifies how many more positive outcomes a model predicted (labels $y'$) for a certain facet as compared to what was observed in the training dataset (labels $y$). For example, if there were more acceptances (a positive outcome) observed in the training dataset for loan applications for a middle-aged group (facet $a$) than predicted by the model based on qualifications as compared to the facet containing other age groups (facet $d$), this might indicate potential bias in the way loans were approved favoring the middle-aged group.

The formula for the difference in conditional acceptance:

$$\text{DCAcc} = c_a - c_d$$

Where:

- $c_a = \frac{n_{a'}(1)}{n_{a'}(1)}$ is the ratio of the observed number of positive outcomes of value 1 (acceptances) of facet $a$ to the predicted number of positive outcome (acceptances) for facet $a$.
- $c_d = \frac{n_{d'}(1)}{n_{d'}(1)}$ is the ratio of the observed number of positive outcomes of value 1 (acceptances) of facet $d$ to the predicted number of predicted positive outcomes (acceptances) for facet $d$. 
The DCAcc metric can capture both positive and negative biases that reveal preferential treatment based on qualifications. Consider the following instances of age-based bias on loan acceptances.

**Example 1: Positive bias**

Suppose we have a dataset of 100 middle-aged people (facet $a$) and 50 people from other age groups (facet $d$) who applied for loans, where the model recommended that 60 from facet $a$ and 30 from facet $d$ be given loans. So the predicted proportions are unbiased with respect to the DPPL metric, but the observed labels show that 70 from facet $a$ and 20 from facet $d$ were granted loans. In other words, the model granted loans to 17% fewer from the middle aged facet than the observed labels in the training data suggested (70/60 = 1.17) and granted loans to 33% more from other age groups than the observed labels suggested (20/30 = 0.67). The calculation of the DCAcc value gives the following:

$$\text{DCAcc} = \frac{70}{60} - \frac{20}{30} = \frac{1}{2}$$

The positive value indicates that there is a potential bias against the middle-aged facet $a$ with a lower acceptance rate as compared with the other facet $d$ than the observed data (taken as unbiased) indicate is the case.

**Example 2: Negative bias**

Suppose we have a dataset of 100 middle-aged people (facet $a$) and 50 people from other age groups (facet $d$) who applied for loans, where the model recommended that 60 from facet $a$ and 30 from facet $d$ be given loans. So the predicted proportions are unbiased with respect to the DPPL metric, but the observed labels show that 50 from facet $a$ and 40 from facet $d$ were granted loans. In other words, the model granted loans to 17% fewer from the middle aged facet than the observed labels in the training data suggested (50/60 = 0.83), and granted loans to 33% more from other age groups than the observed labels suggested (40/30 = 1.33). The calculation of the DCAcc value gives the following:

$$\text{DCAcc} = \frac{50}{60} - \frac{40}{30} = -\frac{1}{2}$$

The negative value indicates that there is a potential bias against facet $d$ with a lower acceptance rate as compared with the middle-aged facet $a$ than the observed data (taken as unbiased) indicate is the case.

Note that you can use DCAcc to help you detect potential (unintentional) biases by humans overseeing the model predictions in a human-in-the-loop setting. Assume, for example, that the
predictions \( y' \) by the model were unbiased, but the eventual decision is made by a human (possibly with access to additional features) who can alter the model predictions to generate a new and final version of \( y' \). The additional processing by the human may unintentionally deny loans to a disproportionate number from one facet. DCAcc can help detect such potential biases.

The range of values for differences in conditional acceptance for binary, multicategory facet, and continuous labels is \((-\infty, +\infty)\).

- Positive values occur when the ratio of the observed number of acceptances compared to predicted acceptances for facet \( a \) is higher than the same ratio for facet \( d \). These values indicate a possible bias against the qualified applicants from facet \( a \). The larger the difference of the ratios, the more extreme the apparent bias.
- Values near zero occur when the ratio of the observed number of acceptances compared to predicted acceptances for facet \( a \) is similar to the ratio for facet \( d \). These values indicate that predicted acceptance rates are consistent with the observed values in the labeled data and that qualified applicants from both facets are being accepted in a similar way.
- Negative values occur when the ratio of the observed number of acceptances compared to predicted acceptances for facet \( a \) is less than that ratio for facet \( d \). These values indicate a possible bias against the qualified applicants from facet \( d \). The more negative the difference in the ratios, the more extreme the apparent bias.

**Difference in Conditional Rejection (DCR)**

This metric compares the observed labels to the labels predicted by the model and assesses whether this is the same across facets for negative outcomes (rejections). This metric comes close to mimicking human bias, in that it quantifies how many more negative outcomes a model granted (predicted labels \( y' \)) to a certain facet as compared to what was suggested by the labels in the training dataset (observed labels \( y \)). For example, if there were more observed rejections (a negative outcome) for loan applications for a middle-aged group (facet \( a \)) than predicted by the model based on qualifications as compared to the facet containing other age groups (facet \( d \)), this might indicate potential bias in the way loans were rejected that favored the middle-aged group over other groups.

The formula for the difference in conditional acceptance:

\[
DCR = r_d - r_a
\]

Where:
The DCR metric can capture both positive and negative biases that reveal preferential treatment based on qualifications. Consider the following instances of age-based bias on loan rejections.

**Example 1: Positive bias**

Suppose we have dataset of 100 middle-aged people (facet \(a\)) and 50 people from other age groups (facet \(d\)) who applied for loans, where the model recommended that 60 from facet \(a\) and 30 from facet \(d\) be rejected for loans. So the predicted proportions are unbiased by the DPPL metric, but the observed labels show that 50 from facet \(a\) and 40 from facet \(d\) were rejected. In other words, the model rejected 17% more loans from the middle aged facet than the observed labels in the training data suggested (50/60 = 0.83), and rejected 33% fewer loans from other age groups than the observed labels suggested (40/30 = 1.33). The DCR value quantifies this difference in the ratio of observed to predicted rejection rates between the facets. The positive value indicates that there is a potential bias favoring the middle aged group with lower rejection rates as compared with other groups than the observed data (taken as unbiased) indicate is the case.

\[
DCR = \frac{40}{30} - \frac{50}{60} = \frac{1}{2}
\]

**Example 2: Negative bias**

Suppose we have dataset of 100 middle-aged people (facet \(a\)) and 50 people from other age groups (facet \(d\)) who applied for loans, where the model recommended that 60 from facet \(a\) and 30 from facet \(d\) be rejected for loans. So the predicted proportions are unbiased by the DPPL metric, but the observed labels show that 70 from facet \(a\) and 20 from facet \(d\) were rejected. In other words, the model rejected 17% fewer loans from the middle aged facet than the observed labels in the training data suggested (70/60 = 1.17), and rejected 33% more loans from other age groups than the observed labels suggested (20/30 = 0.67). The negative value indicates that there is a potential bias favoring facet \(a\) with lower rejection rates as compared with the middle-aged facet \(a\) than the observed data (taken as unbiased) indicate is the case.

\[
DCR = \frac{20}{30} - \frac{70}{60} = -\frac{1}{2}
\]

The range of values for differences in conditional rejection for binary, multicategory facet, and continuous labels is \((-\infty, +\infty)\).
• Positive values occur when the ratio of the observed number of rejections compared to predicted rejections for facet \( d \) is greater than that ratio for facet \( a \). These values indicate a possible bias against the qualified applicants from facet \( a \). The larger the value of DCR metric, the more extreme the apparent bias.

• Values near zero occur when the ratio of the observed number of rejections compared to predicted acceptances for facet \( a \) is the similar to the ratio for facet \( d \). These values indicate that predicted rejections rates are consistent with the observed values in the labeled data and that the qualified applicants from both facets are being rejected in a similar way.

• Negative values occur when the ratio of the observed number of rejections compared to predicted rejections for facet \( d \) is less than that ratio facet \( a \). These values indicate a possible bias against the qualified applicants from facet \( d \). The larger magnitude of the negative DCR metric, the more extreme the apparent bias.

**Specificity difference (SD)**

The specificity difference (SD) is the difference in specificity between the favored facet \( a \) and disfavored facet \( d \). Specificity measures how often the model correctly predicts a negative outcome \((y' = 0)\). Any difference in these specificities is a potential form of bias.

Specificity is perfect for a facet if all of the \( y = 0 \) cases are correctly predicted for that facet. Specificity is greater when the model minimizes false positives, known as a Type I error. For example, the difference between a low specificity for lending to facet \( a \), and high specificity for lending to facet \( d \), is a measure of bias against facet \( d \).

The following formula is for the difference in the specificity for facets \( a \) and \( d \).

\[
SD = \frac{TN_d}{TN_d + FP_d} - \frac{TN_a}{TN_a + FP_a} = TNR_d - TNR_a
\]

The following variables used to calculated SD are defined as follows:

• \( TN_d \) are the true negatives predicted for facet \( d \).
• \( FP_d \) are the false positives predicted for facet \( d \).
• \( TN_a \) are the true negatives predicted for facet \( a \).
• \( FP_a \) are the false positives predicted for facet \( a \).
• \( TNR_a = \frac{TN_a}{TN_a + FP_a} \) is the true negative rate, also known as the specificity, for facet \( a \).
• \( TNR_d = \frac{TN_d}{TN_d + FP_d} \) is the true negative rate, also known as the specificity, for facet \( d \).
For example, consider the following confusion matrices for facets $a$ and $d$.

**Confusion matrix for the favored facet $a$**

<table>
<thead>
<tr>
<th>Class a predictions</th>
<th>Actual outcome 0</th>
<th>Actual outcome 1</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>20</td>
<td>5</td>
<td>25</td>
</tr>
<tr>
<td>1</td>
<td>10</td>
<td>65</td>
<td>75</td>
</tr>
<tr>
<td>Total</td>
<td>30</td>
<td>70</td>
<td>100</td>
</tr>
</tbody>
</table>

**Confusion matrix for the disfavored facet $d$**

<table>
<thead>
<tr>
<th>Class d predictions</th>
<th>Actual outcome 0</th>
<th>Actual outcome 1</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>18</td>
<td>7</td>
<td>25</td>
</tr>
<tr>
<td>1</td>
<td>5</td>
<td>20</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>23</td>
<td>27</td>
<td>50</td>
</tr>
</tbody>
</table>

The value of the specificity difference is $SD = \frac{18}{18+5} - \frac{20}{20+10} = 0.7826 - 0.6667 = 0.1159$, which indicates a bias against facet $d$.

The range of values for the specificity difference between facets $a$ and $d$ for binary and multicategory classification is $[-1, +1]$. This metric is not available for the case of continuous labels. Here is what different values of SD imply:

- **Positive values** are obtained when there is higher specificity for facet $d$ than for facet $a$. This suggests that the model finds less false positives for facet $d$ than for facet $a$. A positive value indicates bias against facet $d$.
- **Values near zero** indicate that the specificity for facets that are being compared is similar. This suggests that the model finds a similar number of false positives in both of these facets and is not biased.
- **Negative values** are obtained when there is higher specificity for facet $a$ than for facet $d$. This suggests that the model finds more false positives for facet $a$ than for facet $d$. A negative value indicates bias against facet $a$. 
Recall Difference (RD)

The recall difference (RD) metric is the difference in recall of the model between the favored facet $a$ and disfavored facet $d$. Any difference in these recalls is a potential form of bias. Recall is the true positive rate (TPR), which measures how often the model correctly predicts the cases that should receive a positive outcome. Recall is perfect for a facet if all of the $y=1$ cases are correctly predicted as $y'=1$ for that facet. Recall is greater when the model minimizes false negatives known as the Type II error. For example, how many of the people in two different groups (facets $a$ and $d$) that should qualify for loans are detected correctly by the model? If the recall rate is high for lending to facet $a$, but low for lending to facet $d$, the difference provides a measure of this bias against the group belonging to facet $d$.

The formula for difference in the recall rates for facets $a$ and $d$:

$$RD = \frac{TP_a}{TP_a + FN_a} - \frac{TP_d}{TP_d + FN_d} = TPR_a - TPR_d$$

Where:

- $TP_a$ are the true positives predicted for facet $a$.
- $FN_a$ are the false negatives predicted for facet $a$.
- $TP_d$ are the true positives predicted for facet $d$.
- $FN_d$ are the false negatives predicted for facet $d$.
- $TPR_a = \frac{TP_a}{TP_a + FN_a}$ is the recall for facet $a$, or its true positive rate.
- $TPR_d = \frac{TP_d}{TP_d + FN_d}$ is the recall for facet $d$, or its true positive rate.

For example, consider the following confusion matrices for facets $a$ and $d$.

Confusion Matrix for the Favored Facet $a$

<table>
<thead>
<tr>
<th>Class a predictions</th>
<th>Actual outcome 0</th>
<th>Actual outcome 1</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>20</td>
<td>5</td>
<td>25</td>
</tr>
<tr>
<td>1</td>
<td>10</td>
<td>65</td>
<td>75</td>
</tr>
<tr>
<td>Total</td>
<td>30</td>
<td>70</td>
<td>100</td>
</tr>
</tbody>
</table>
### Confusion Matrix for the Disfavored Facet \( d \)

<table>
<thead>
<tr>
<th>Class ( d ) predictions</th>
<th>Actual outcome 0</th>
<th>Actual outcome 1</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>18</td>
<td>7</td>
<td>25</td>
</tr>
<tr>
<td>1</td>
<td>5</td>
<td>20</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>23</td>
<td>27</td>
<td>50</td>
</tr>
</tbody>
</table>

The value of the recall difference is \( RD = \frac{65}{70} - \frac{20}{27} = 0.93 - 0.74 = 0.19 \) which indicates a bias against facet \( d \).

The range of values for the recall difference between facets \( a \) and \( d \) for binary and multicategory classification is \([-1, +1]\). This metric is not available for the case of continuous labels.

- Positive values are obtained when there is higher recall for facet \( a \) than for facet \( d \). This suggests that the model finds more of the true positives for facet \( a \) than for facet \( d \), which is a form of bias.
- Values near zero indicate that the recall for facets being compared is similar. This suggests that the model finds about the same number of true positives in both of these facets and is not biased.
- Negative values are obtained when there is higher recall for facet \( d \) than for facet \( a \). This suggests that the model finds more of the true positives for facet \( d \) than for facet \( a \), which is a form of bias.

### Difference in Acceptance Rates (DAR)

The difference in acceptance rates (DAR) metric is the difference in the ratios of the true positive (TP) predictions to the observed positives (TP + FP) for facets \( a \) and \( d \). This metric measures the difference in the precision of the model for predicting acceptances from these two facets. Precision measures the fraction of qualified candidates from the pool of qualified candidates that are identified as such by the model. If the model precision for predicting qualified applicants diverges between the facets, this is a bias and its magnitude is measured by the DAR.

The formula for difference in acceptance rates between facets \( a \) and \( d \):

\[
DAR = \frac{TP_a}{(TP_a + FP_a)} - \frac{TP_d}{(TP_d + FP_d)}
\]
Where:

- $TP_a$ are the true positives predicted for facet $a$.
- $FP_a$ are the false positives predicted for facet $a$.
- $TP_d$ are the true positives predicted for facet $d$.
- $FP_d$ are the false positives predicted for facet $d$.

For example, suppose the model accepts 70 middle-aged applicants (facet $a$) for a loan (predicted positive labels) of whom only 35 are actually accepted (observed positive labels). Also suppose the model accepts 100 applicants from other age demographics (facet $d$) for a loan (predicted positive labels) of whom only 40 are actually accepted (observed positive labels). Then $DAR = 35/70 - 40/100 = 0.10$, which indicates a potential bias against qualified people from the second age group (facet $d$).

The range of values for DAR for binary, multicategory facet, and continuous labels is $[-1, +1]$.

- Positive values occur when the ratio of the predicted positives (acceptances) to the observed positive outcomes (qualified applicants) for facet $a$ is larger than the same ratio for facet $d$. These values indicate a possible bias against the disfavored facet $d$ caused by the occurrence of relatively more false positives in facet $d$. The larger the difference in the ratios, the more extreme the apparent bias.

- Values near zero occur when the ratio of the predicted positives (acceptances) to the observed positive outcomes (qualified applicants) for facets $a$ and $d$ have similar values indicating the observed labels for positive outcomes are being predicted with equal precision by the model.

- Negative values occur when the ratio of the predicted positives (acceptances) to the observed positive outcomes (qualified applicants) for facet $d$ is larger than the ratio facet $a$. These values indicate a possible bias against the favored facet $a$ caused by the occurrence of relatively more false positives in facet $a$. The more negative the difference in the ratios, the more extreme the apparent bias.

**Difference in Rejection Rates (DRR)**

The difference in rejection rates (DRR) metric is the difference in the ratios of the true negative (TN) predictions to the observed negatives (TN + FN) for facets $a$ and $d$. This metric measures the difference in the precision of the model for predicting rejections from these two facets. Precision measures the fraction of unqualified candidates from the pool of unqualified candidates that
are identified as such by the model. If the model precision for predicting unqualified applicants diverges between the facets, this is a bias and its magnitude is measured by the DRR.

The formula for difference in rejection rates between facets $a$ and $d$:

$$DRR = \frac{TN_d}{TN_d + FN_d} - \frac{TN_a}{TN_a + FN_a}$$

The components for the previous DRR equation are as follows.

- $TN_d$ are the true negatives predicted for facet $d$.
- $FN_d$ are the false negatives predicted for facet $d$.
- $TP_a$ are the true negatives predicted for facet $a$.
- $FN_a$ are the false negatives predicted for facet $a$.

For example, suppose the model rejects 100 middle-aged applicants (facet $a$) for a loan (predicted negative labels) of whom 80 are actually unqualified (observed negative labels). Also suppose the model rejects 50 applicants from other age demographics (facet $d$) for a loan (predicted negative labels) of whom only 40 are actually unqualified (observed negative labels). Then $DRR = \frac{40}{50} - \frac{80}{100} = 0$, so no bias is indicated.

The range of values for DRR for binary, multicategory facet, and continuous labels is $[-1, +1]$.

- Positive values occur when the ratio of the predicted negatives (rejections) to the observed negative outcomes (unqualified applicants) for facet $d$ is larger than the same ratio for facet $a$. These values indicate a possible bias against the favored facet $a$ caused by the occurrence of relatively more false negatives in facet $a$. The larger the difference in the ratios, the more extreme the apparent bias.

- Values near zero occur when the ratio of the predicted negatives (rejections) to the observed negative outcomes (unqualified applicants) for facets $a$ and $d$ have similar values, indicating the observed labels for negative outcomes are being predicted with equal precision by the model.

- Negative values occur when the ratio of the predicted negatives (rejections) to the observed negative outcomes (unqualified applicants) for facet $a$ is larger than the ratio facet $d$. These values indicate a possible bias against the disfavored facet $d$ caused by the occurrence of relatively more false positives in facet $d$. The more negative the difference in the ratios, the more extreme the apparent bias.
Accuracy Difference (AD)

Accuracy difference (AD) metric is the difference between the prediction accuracy for different facets. This metric determines whether the classification by the model is more accurate for one facet than the other. AD indicates whether one facet incurs a greater proportion of Type I and Type II errors. But it cannot differentiate between Type I and Type II errors. For example, the model may have equal accuracy for different age demographics, but the errors may be mostly false positives (Type I errors) for one age-based group and mostly false negatives (Type II errors) for the other.

Also, if loan approvals are made with much higher accuracy for a middle-aged demographic (facet $a$) than for another age-based demographic (facet $d$), either a greater proportion of qualified applicants in the second group are denied a loan (FN) or a greater proportion of unqualified applicants from that group get a loan (FP) or both. This can lead to within group unfairness for the second group, even if the proportion of loans granted is nearly the same for both age-based groups, which is indicated by a DPPL value that is close to zero.

The formula for AD metric is the difference between the prediction accuracy for facet $a$, $\text{ACC}_a$, minus that for facet $d$, $\text{ACC}_d$:

$$\text{AD} = \text{ACC}_a - \text{ACC}_d$$

Where:

- $\text{ACC}_a = \frac{(\text{TP}_a + \text{TN}_a)}{(\text{TP}_a + \text{TN}_a + \text{FP}_a + \text{FN}_a)}$
  - $\text{TP}_a$ are the true positives predicted for facet $a$
  - $\text{TN}_a$ are the true negatives predicted for facet $a$
  - $\text{FP}_a$ are the false positives predicted for facet $a$
  - $\text{FN}_a$ are the false negatives predicted for facet $a$

- $\text{ACC}_d = \frac{(\text{TP}_d + \text{TN}_d)}{(\text{TP}_d + \text{TN}_d + \text{FP}_d + \text{FN}_d)}$
  - $\text{TP}_d$ are the true positives predicted for facet $d$
  - $\text{TN}_d$ are the true negatives predicted for facet $d$
  - $\text{FP}_d$ are the false positives predicted for facet $d$
  - $\text{FN}_d$ are the false negatives predicted for facet $d$

For example, suppose a model approves loans to 70 applicants from facet $a$ of 100 and rejected the other 30. 10 should not have been offered the loan ($\text{FP}_a$) and 60 were approved that should
have been (TP_a). 20 of the rejections should have been approved (FN_a) and 10 were correctly rejected (TN_a). The accuracy for facet a is as follows:

$$\text{ACC}_a = \frac{60 + 10}{60 + 10 + 20 + 10} = 0.7$$

Next, suppose a model approves loans to 50 applicants from facet d of 100 and rejected the other 50. 10 should not have been offered the loan (FP_a) and 40 were approved that should have been (TP_a). 40 of the rejections should have been approved (FN_a) and 10 were correctly rejected (TN_a). The accuracy for facet a is determined as follows:

$$\text{ACC}_d = \frac{40 + 10}{40 + 10 + 40 + 10} = 0.5$$

The accuracy difference is thus $\text{AD} = \text{ACC}_a - \text{ACC}_d = 0.7 - 0.5 = 0.2$. This indicates there is a bias against facet d as the metric is positive.

The range of values for AD for binary and multicategory facet labels is [-1, +1].

- Positive values occur when the prediction accuracy for facet a is greater than that for facet d. It means that facet d suffers more from some combination of false positives (Type I errors) or false negatives (Type II errors). This means there is a potential bias against the disfavored facet d.
- Values near zero occur when the prediction accuracy for facet a is similar to that for facet d.
- Negative values occur when the prediction accuracy for facet d is greater than that for facet a. It means that facet a suffers more from some combination of false positives (Type I errors) or false negatives (Type II errors). This means there is a bias against the favored facet a.

**Treatment Equality (TE)**

The treatment equality (TE) is the difference in the ratio of false negatives to false positives between facets a and d. The main idea of this metric is to assess whether, even if the accuracy across groups is the same, is it the case that errors are more harmful to one group than another? Error rate comes from the total of false positives and false negatives, but the breakdown of these two may be very different across facets. TE measures whether errors are compensating in the similar or different ways across facets.

The formula for the treatment equality:

$$\text{TE} = \frac{\text{FN}_d/\text{FP}_d - \text{FN}_a/\text{FP}_a}{\text{FN}_a/\text{FP}_a}$$

Where:
• FN<sub>d</sub> are the false negatives predicted for facet <i>d</i>.
• FP<sub>d</sub> are the false positives predicted for facet <i>d</i>.
• FN<sub>a</sub> are the false negatives predicted for facet <i>a</i>.
• FP<sub>a</sub> are the false positives predicted for facet <i>a</i>.

Note the metric becomes unbounded if FP<sub>a</sub> or FP<sub>d</sub> is zero.

For example, suppose that there are 100 loan applicants from facet <i>a</i> and 50 from facet <i>d</i>. For facet <i>a</i>, 8 were wrongly denied a loan (FN<sub>a</sub>) and another 6 were wrongly approved (FP<sub>a</sub>). The remaining predictions were true, so TP<sub>a</sub> + TN<sub>a</sub> = 86. For facet <i>d</i>, 5 were wrongly denied (FN<sub>d</sub>) and 2 were wrongly approved (FP<sub>d</sub>). The remaining predictions were true, so TP<sub>d</sub> + TN<sub>d</sub> = 43. The ratio of false negatives to false positives equals 8/6 = 1.33 for facet <i>a</i> and 5/2 = 2.5 for facet <i>d</i>. Hence TE = 2.5 - 1.33 = 1.167, even though both facets have the same accuracy:

\[
\text{ACC}_a = \frac{86}{86 + 8 + 6} = 0.86 \\
\text{ACC}_d = \frac{43}{43 + 5 + 2} = 0.86
\]

The range of values for differences in conditional rejection for binary and multicategory facet labels is (-∞, +∞). The TE metric is not defined for continuous labels. The interpretation of this metric depends on the relative important of false positives (Type I error) and false negatives (Type II error).

• Positive values occur when the ratio of false negatives to false positives for facet <i>d</i> is greater than that for facet <i>a</i>.
• Values near zero occur when the ratio of false negatives to false positives for facet <i>a</i> is similar to that for facet <i>d</i>.
• Negative values occur when the ratio of false negatives to false positives for facet <i>d</i> is less than that for facet <i>a</i>.

Note

A previous version stated that the Treatment Equality metric is computed as FP<sub>a</sub> / FN<sub>a</sub> - FP<sub>d</sub> / FN<sub>d</sub> instead of FN<sub>d</sub> / FP<sub>d</sub> - FN<sub>a</sub> / FP<sub>a</sub>. While either of the versions can be used. For more information, see [Fairness measures for Machine Learning in Finance](#).
Conditional Demographic Disparity in Predicted Labels (CDDPL)

The demographic disparity metric (DDPL) determines whether facet \( d \) has a larger proportion of the predicted rejected labels than of the predicted accepted labels. It enables a comparison of difference in predicted rejection proportion and predicted acceptance proportion across facets. This metric is exactly the same as the pre-training CDD metric except that it is computed off the predicted labels instead of the observed ones. This metric lies in the range \((-1,+1)\).

The formula for the demographic disparity predictions for labels of facet \( d \) is as follows:

\[
DDPL_d = \frac{n'_d(0)}{n'(0)} - \frac{n'_d(1)}{n'(1)} = P_d^R(y'_0) - P_d^A(y'_1)
\]

Where:

- \( n'_d(0) = n'_a(0) + n'_d(0) \) is the number of predicted rejected labels for facets \( a \) and \( d \).
- \( n'_d(1) = n'_a(1) + n'_d(1) \) is the number of predicted accepted labels for facets \( a \) and \( d \).
- \( P_d^R(y'_0) \) is the proportion of predicted rejected labels (value 0) in facet \( d \).
- \( P_d^A(y'_1) \) is the proportion of predicted accepted labels (value 1) in facet \( d \).

A conditional demographic disparity in predicted labels (CDDPL) metric that conditions DDPL on attributes that define a strata of subgroups on the dataset is needed to rule out Simpson's paradox. The regrouping can provide insights into the cause of apparent demographic disparities for less favored facets. The classic case arose in the case of Berkeley admissions where men were accepted at a higher rate overall than women. But when departmental subgroups were examined, women were shown to have higher admission rates than men by department. The explanation was that women had applied to departments with lower acceptance rates than men had. Examining the subgroup acceptance rates revealed that women were actually accepted at a higher rate than men for the departments with lower acceptance rates.

The CDDPL metric gives a single measure for all of the disparities found in the subgroups defined by an attribute of a dataset by averaging them. It is defined as the weighted average of demographic disparities in predicted labels (DDPL\(_i\)) for each of the subgroups, with each subgroup disparity weighted in proportion to the number of observations in contains. The formula for the conditional demographic disparity in predicted labels is as follows:

\[
CDDPL = \frac{1}{n} \cdot \sum_{i} n_i \cdot DDPL_i
\]

Where:
• \( \sum_{i} n_i = n \) is the total number of observations and \( n_i \) is the number of observations for each subgroup.

• \( \text{DDPL}_i = \frac{n_i^{(0)}}{n^{(0)}} - \frac{n_i^{(1)}}{n^{(1)}} = P_i^R(y_i^0) - P_i^A(y_i^1) \) is the demographic disparity in predicted labels for the subgroup.

So the demographic disparity for a subgroup in predicted labels (DDPL) are the difference between the proportion of predicted rejected labels and the proportion of predicted accepted labels for each subgroup.

The range of DDPL values for binary, multicategory, and continuous outcomes is \([-1,+1]\).

• +1: when there are no predicted rejection labels for facet \( a \) or subgroup and no predicted acceptances for facet \( d \) or subgroup.

• Positive values indicate there is a demographic disparity in predicted labels as facet \( d \) or subgroup has a larger proportion of the predicted rejected labels than of the predicted accepted labels. The higher the value the greater the disparity.

• Values near zero indicate there is no demographic disparity on average.

• Negative values indicate there is a demographic disparity in predicted labels as facet \( a \) or subgroup has a larger proportion of the predicted rejected labels than of the predicted accepted labels. The lower the value the greater the disparity.

• -1: when there are no predicted rejection labels for facet \( d \) or subgroup and no predicted acceptances for facet \( a \) or subgroup.

### Counterfactual Flitext (FT)

The flitext is an approach that looks at each member of facet \( d \) and assesses whether similar members of facet \( a \) have different model predictions. The members of facet \( a \) are chosen to be k-nearest neighbors of the observation from facet \( d \). We assess how many nearest neighbors of the opposite group receive a different prediction, where the flipped prediction can go from positive to negative and vice versa.

The formula for the counterfactual flitext is the difference in the cardinality of two sets divided by the number of members of facet \( d \):

\[
\text{FT} = \frac{(F^+ - F^-)}{n_d}
\]

Where:
• $F^+ = \text{is the number of disfavored facet } d \text{ members with an unfavorable outcome whose nearest neighbors in favored facet } a \text{ received a favorable outcome.}$

• $F^- = \text{is the number of disfavored facet } d \text{ members with a favorable outcome whose nearest neighbors in favored facet } a \text{ received an unfavorable outcome.}$

• $n_d$ is the sample size of facet $d$.

The range of values for the counterfactual flitpest for binary and multicategory facet labels is $[-1, +1]$. For continuous labels, we set a threshold to collapse the labels to binary.

• Positive values occur when the number of unfavorable counterfactual flitpest decisions for the disfavored facet $d$ exceeds the favorable ones.

• Values near zero occur when the number of unfavorable and favorable counterfactual flitpest decisions balance out.

• Negative values occur when the number of unfavorable counterfactual flitpest decisions for the disfavored facet $d$ is less than the favorable ones.

**Generalized entropy (GE)**

The generalized entropy index (GE) measures the inequality in benefit $b$ for the predicted label compared to the observed label. A benefit occurs when a false positive is predicted. A false positive occurs when a negative observation ($y=0$) has a positive prediction ($y'=1$). A benefit also occurs when the observed and predicted labels are the same, also known as a true positive and true negative. No benefit occurs when a false negative is predicted. A false negative occurs when a positive observation ($y=1$) is predicted to have a negative outcome ($y'=0$). The benefit $b$ is defined, as follows.

\[
b = y' - y + 1
\]

Using this definition, a false positive receives a benefit $b$ of 2, and a false negative receives a benefit of 0. Both a true positive and a true negative receive a benefit of 1.

The GE metric is computed following the [Generalized Entropy Index](#) (GE) with the weight $\alpha$ set to 2. This weight controls the sensitivity to different benefit values. A smaller $\alpha$ means an increased sensitivity to smaller values.
The following variables used to calculate GE are defined as follows:

- $b_i$ is the benefit received by the $i^{th}$ data point.
- $b'$ is the mean of all benefits.

GE can range from 0 to 0.5, where values of zero indicate no inequality in benefits across all data points. This occurs either when all inputs are correctly predicted or when all the predictions are false positives. GE is undefined when all predictions are false negatives.

Note

The metric GE does not depend on a facet value being either favored or disfavored.

Model Explainability

Amazon SageMaker Clarify provides tools to help explain how machine learning (ML) models make predictions. These tools can help ML modelers and developers and other internal stakeholders understand model characteristics as a whole prior to deployment and to debug predictions provided by the model after it's deployed.

- To obtain explanations for your datasets and models, see Use SageMaker Clarify to explain and detect bias.
- To obtain explanations in real-time from a SageMaker endpoint, see Online Explainability with SageMaker Clarify.

Transparency about how ML models arrive at their predictions is also critical to consumers and regulators. They need to trust the model predictions if they are going to accept the decisions based on them. SageMaker Clarify uses a model-agnostic feature attribution approach. You can

$$GE = \frac{1}{2n} \sum_{i=1}^{n} \left[ \left( \frac{b_i}{b'} \right)^2 - 1 \right]$$
use this to understand why a model made a prediction after training, and to provide per-instance explanation during inference. The implementation includes a scalable and efficient implementation of SHAP. This is based on the concept of a Shapley value, from the field of cooperative game theory, that assigns each feature an importance value for a particular prediction.

Clarify produces partial dependence plots (PDPs) that show the marginal effect features have on the predicted outcome of a machine learning model. Partial dependence helps explain target response given a set of input features. It also supports both computer vision (CV) and natural language processing (NLP) explainability using the same Shapley Values (SHAP) algorithm as used for tabular data explanations.

What is the function of an explanation in the machine learning context? An explanation can be thought of as the answer to a Why question that helps humans understand the cause of a prediction. In the context of an ML model, you might be interested in answering questions such as:

- Why did the model predict a negative outcome such as a loan rejection for a given applicant?
- How does the model make predictions?
- Why did the model make an incorrect prediction?
- Which features have the largest influence on the behavior of the model?

You can use explanations for auditing and meeting regulatory requirements, building trust in the model and supporting human decision-making, and debugging and improving model performance.

The need to satisfy the demands for human understanding about the nature and outcomes of ML inference is key to the sort of explanation needed. Research from philosophy and cognitive science disciplines has shown that people care especially about contrastive explanations, or explanations of why an event X happened instead of some other event Y that did not occur. Here, X could be an unexpected or surprising event that happened and Y corresponds to an expectation based on their existing mental model referred to as a baseline. Note that for the same event X, different people might seek different explanations depending on their point of view or mental model Y. In the context of explainable AI, you can think of X as the example being explained and Y as a baseline that is typically chosen to represent an uninformative or average example in the dataset. Sometimes, for example in the case of ML modeling of images, the baseline might be implicit, where an image whose pixels are all the same color can serves as a baseline.

**Sample Notebooks**

Amazon SageMaker Clarify provides the following sample notebook for model explainability:
• **Amazon SageMaker Clarify Processing** – Use SageMaker Clarify to create a processing job for the detecting bias and explaining model predictions with feature attributions. Examples include using CSV and JSON Lines data formats, bringing your own container, and running processing jobs with Spark.

• **Explaining Image Classification with SageMaker Clarify** – SageMaker Clarify provides you with insights into how your computer vision models classify images.

• **Explaining object detection models with SageMaker Clarify** – SageMaker Clarify provides you with insights into how your computer vision models detect objects.

This notebook has been verified to run in Amazon SageMaker Studio only. If you need instructions on how to open a notebook in Amazon SageMaker Studio, see [Create or Open an Amazon SageMaker Studio Classic Notebook](#). If you're prompted to choose a kernel, choose **Python 3 (Data Science)**.

**Topics**

• **Feature Attributions that Use Shapley Values**

• **SHAP Baselines for Explainability**

**Feature Attributions that Use Shapley Values**

SageMaker Clarify provides feature attributions based on the concept of **Shapley value**. You can use Shapley values to determine the contribution that each feature made to model predictions. These attributions can be provided for specific predictions and at a global level for the model as a whole. For example, if you used an ML model for college admissions, the explanations could help determine whether the GPA or the SAT score was the feature most responsible for the model's predictions, and then you can determine how responsible each feature was for determining an admission decision about a particular student.

SageMaker Clarify has taken the concept of Shapley values from game theory and deployed it in a machine learning context. The Shapley value provides a way to quantify the contribution of each player to a game, and hence the means to distribute the total gain generated by a game to its players based on their contributions. In this machine learning context, SageMaker Clarify treats the prediction of the model on a given instance as the **game** and the features included in the model as the **players**. For a first approximation, you might be tempted to determine the marginal contribution or effect of each feature by quantifying the result of either **dropping** that feature from the model or **dropping** all other features from the model. However, this approach does not take
to account that features included in a model are often not independent from each other. For example, if two features are highly correlated, dropping either one of the features might not alter the model prediction significantly.

To address these potential dependencies, the Shapley value requires that the outcome of each possible combination (or coalition) of features must be considered to determine the importance of each feature. Given $d$ features, there are $2^d$ such possible feature combinations, each corresponding to a potential model. To determine the attribution for a given feature $f$, consider the marginal contribution of including $f$ in all feature combinations (and associated models) that do not contain $f$, and take the average. It can be shown that Shapley value is the unique way of assigning the contribution or importance of each feature that satisfies certain desirable properties. In particular, the sum of Shapley values of each feature corresponds to the difference between the predictions of the model and a dummy model with no features. However, even for reasonable values of $d$, say 50 features, it is computationally prohibitive and impractical to train $2^d$ possible models. As a result, SageMaker Clarify needs to make use of various approximation techniques. For this purpose, SageMaker Clarify uses Shapley Additive exPlanations (SHAP), which incorporates such approximations and devised a scalable and efficient implementation of the Kernel SHAP algorithm through additional optimizations.

For additional information on Shapley values, see A Unified Approach to Interpreting Model Predictions.

**SHAP Baselines for Explainability**

Explanations are typically contrastive (that is, they account for deviations from a baseline). As a result, for the same model prediction, you can expect to get different explanations with respect to different baselines. Therefore, your choice of a baseline is crucial. In an ML context, the baseline corresponds to a hypothetical instance that can be either uninformative or informative. During the computation of Shapley values, SageMaker Clarify generates several new instances between the baseline and the given instance, in which the absence of a feature, is modeled by setting the feature value to that of the baseline and the presence of a feature is modeled by setting the feature value to that of the given instance. Thus, the absence of all features corresponds to the baseline and the presence of all features corresponds to the given instance.

How can you choose good baselines? Often it is desirable to select a baseline with very low information content. For example, you can construct an average instance from the training dataset by taking either the median or average for numerical features and the mode for categorical features. For the college admissions example, you might be interested in explaining why a
particular applicant was accepted as compared to a baseline acceptances based on an average applicant. If not provided, a baseline is calculated automatically by SageMaker Clarify using K-means or K-prototypes in the input dataset.

Alternatively, you can choose to generate explanations with respect to informative baselines. For the college admissions scenario, you might want to explain why a particular applicant was rejected when compared with other applicants from similar demographic backgrounds. In this case, you can choose a baseline that represents the applicants of interest, namely those from a similar demographic background. Thus, you can use informative baselines to concentrate the analysis on the specific aspects of a particular model prediction. You can isolate the features for assessment by setting demographic attributes and other features that you can't act on to the same value as in the given instance.

**Use SageMaker Clarify explainability with SageMaker Autopilot**

Autopilot uses tools provided by Amazon SageMaker Clarify to help provide insights into how machine learning (ML) models make predictions. These tools can help ML engineers, product managers, and other internal stakeholders understand model characteristics. To trust and interpret decisions made on model predictions, both consumers and regulators rely on transparency in machine learning in order.

The Autopilot explanatory functionality uses a model-agnostic feature attribution approach. This approach determines the contribution of individual features or inputs to the model's output, providing insights into the relevance of different features. You can use it to understand why a model made a prediction after training, or use it to provide per-instance explanation during inference. The implementation includes a scalable implementation of SHAP (Shapley Additive Explanations). This implementation is based on the concept of a Shapley value from cooperative game theory, which assigns each feature an importance value for a particular prediction.

You can use SHAP explanations for the following: auditing and meeting regulatory requirements, building trust in the model, supporting human decision-making, or debugging and improving model performance.

For additional information on Shapely values and baselines, see [SHAP Baselines for Explainability](#).

For a guide to the Amazon SageMaker Clarify documentation, see [Guide to the SageMaker Clarify Documentation](#).
Use governance to document and track model performance

Model governance is a framework that gives systematic visibility into machine learning (ML) model development, validation, and usage. Amazon SageMaker provides purpose-built ML governance tools for managing control access, activity tracking, and reporting across the ML lifecycle.

Manage least-privilege permissions for ML practitioners using Amazon SageMaker Role Manager, create detailed model documentation using Amazon SageMaker Model Cards, and gain visibility into your models with centralized dashboards using Amazon SageMaker Model Dashboard.

Amazon SageMaker Role Manager

With Amazon SageMaker Role Manager, administrators can define user permissions with least-privilege permissions for common machine learning activities. Use Amazon SageMaker Role Manager to build and manage persona-based IAM roles specific to your business needs.

For more information, see Amazon SageMaker Role Manager.

Amazon SageMaker Model Cards

Use Amazon SageMaker Model Cards to document, retrieve, and share essential model information from conception to deployment. With model cards, model risk managers, data scientists, and ML engineers can create an immutable record of intended model uses, risk ratings, training details, evaluation results, and more.

For more information, see Amazon SageMaker Model Cards.

Amazon SageMaker Model Dashboard

Amazon SageMaker Model Dashboard is a pre-built, visual overview of all the models in your account. SageMaker Model Dashboard integrates valuable information from Amazon SageMaker Model Monitor, Transform Jobs, Endpoints, ML Lineage Tracking and Amazon CloudWatch so you can access high-level model information and track model performance in one unified view.

For more information, see Amazon SageMaker Model Dashboard.
Amazon SageMaker Model Cards

Use Amazon SageMaker Model Cards to document critical details about your machine learning (ML) models in a single place for streamlined governance and reporting.

Catalog details such as the intended use and risk rating of a model, training details and metrics, evaluation results and observations, and additional call-outs such as considerations, recommendations, and custom information. By creating model cards, you can do the following:

- Provide guidance on how a model should be used.
- Support audit activities with detailed descriptions of model training and performance.
- Communicate how a model is intended to support business goals.

Model cards provide prescriptive guidance on what information to document and include fields for custom information. After creating a model card, you can export it to a PDF or download it to share with relevant stakeholders. Any edits other than an approval status update made to a model card result in additional model card versions in order to have an immutable record of model changes.

Topics

- Prerequisites
- Intended uses of a model
- Risk ratings
- Model card JSON schema
- Create a model card
- Manage model cards
- Cross-account support for Amazon SageMaker Model Cards
- Use model cards through the low-level APIs
- Model card FAQs

Prerequisites

To get started with Amazon SageMaker Model Cards, you must have permission to create, edit, view, and export model cards.
Intended uses of a model

Specifying the intended uses of a model helps ensure that model developers and users have the information they need to train or deploy the model responsibly. The intended uses of a model should describe the scenarios in which the model is appropriate to use as well as the scenarios in which the model is not recommended to use.

We recommend including:

- The general purpose of the model
- Use cases for which the model was intended
- Use cases for which the model was not intended
- Assumptions made when developing the model

The intended uses of a model go beyond technical details and describe how a model should be used in production, the scenarios in which it is appropriate to use a model, and additional considerations such as the type of data to use with the model or any assumptions made during development.

Risk ratings

Developers create ML models for use cases with varying levels of risk. For example, a model that approves loan applications might be a higher risk model than one that detects the category of an email. Given the varied risk profiles of a model, model cards provide a field for you to categorize a model's risk rating.

This risk rating can be unknown, low, medium, or high. Use these risk rating fields to label unknown, low, medium, or high-risk models and help your organization comply with any existing rules about putting certain models into production.

Model card JSON schema

Evaluation details for a model card must be provided in JSON format. If you have existing JSON format evaluation reports generated by SageMaker Clarify or SageMaker Model Monitor, upload them to Amazon S3 and provide an S3 URI to automatically parse evaluation metrics. For more information and sample reports, see the example metrics folder in the Amazon SageMaker Model Governance - Model Cards example notebook.
When creating a model card using the SageMaker Python SDK, model content must be in the model card JSON schema and provided as a string. Provide model content similar to the following example.

**Model card JSON schema sample file**

```json
{
  "$schema": "http://json-schema.org/draft-07/schema#",
  "$id": "http://json-schema.org/draft-07/schema#",
  "title": "SageMakerModelCardSchema",
  "description": "Default model card schema",
  "version": "0.1.0",
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  "additionalProperties": false,
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      "description": "Overview about the model",
      "type": "object",
      "additionalProperties": false,
      "properties": {
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        "model_creator": {
          "description": "Creator of model",
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          "maxLength": 1024
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        "problem_type": {
          "description": "Problem being solved with the model",
          "type": "string"
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        "algorithm_type": {
          "description": "Algorithm used to solve the problem",
          "type": "string"
        }
      }
    }
  }
}
```
"maxLength": 1024
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  "description": "SageMaker Model Arn or Non SageMaker Model id",
  "type": "string",
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"model_artifact": {
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"model_name": {
  "description": "Name of the model",
  "type": "string",
  "maxLength": 1024
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"model_version": {
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"inference_environment": {
  "description": "Overview about the inference",
  "type": "object",
  "additionalProperties": false,
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      "description": "SageMaker inference image uri",
      "type": "array",
      "maxContains": 15,
      "items": {
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        "maxLength": 1024
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            "maxLength": 1024
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            "type": "string",
            "minLength": 1,
            "maxLength": 2048
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            "description": "Information about the user who created model package."
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                    "type": "string",
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  "maxLength": 63
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"model_package_version": {
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  "description": "The machine learning domain of the model package you specified. Common machine learning domains include computer vision and natural language processing."
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"task": {
  "description": "The machine learning task you specified that your model package accomplishes. Common machine learning tasks include object detection and image classification."
},

"source_algorithms": {
  "description": "A list of algorithms that were used to create a model package."
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    "$ref": "#/definitions/inference_specification"
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  }
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  "items": {
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"items": {
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            "maxLength": 2048
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                    "type": "string",
                    "maxLength": 170
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"description": "The Amazon S3 path where the model artifacts, which result from model training, are stored.",
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    "Low",
    "Unknown"
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        "Minimize"
      ]
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  "value"
],
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  "properties": {
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    }
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  "properties": {
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}
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      }
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  },
  "x_axis_name": {
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  "y_axis_name": {
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  }
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    ]
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    ]
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      }
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    "pattern": ".{1,255}"  
  },
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    "maxLength": 1024
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      "matrix"
    ]
  },
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        "items": {
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          "items": {
            "type": "number"
          },
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        "minItems": 1,
        "maxItems": 20
      }
    ]
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  },
  "y_axis_name": {
    "$ref": "#/definitions/axis_name_array"
  }
}
}
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  "type": "object",
  "required": [
    "name",
    "type",
    "value"
  ],
  "additionalProperties": false,
  "properties": {
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      "pattern": "\.{1,255}\"
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        "boolean"
      ]
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      ]
    },
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    },
    "y_axis_name": {

Create a model card

You can create an Amazon SageMaker Model Card using either the SageMaker console or the SageMaker Python SDK. You can also use the API operations directly. For more information about the API operations, see [Use model cards through the low-level APIs](#).

Create a model card using the SageMaker console

Go to the Amazon SageMaker console. In the navigation pane, under **Governance**, choose **Model cards**. On the upper right-hand corner, choose **Create model card**.

Go through the four steps in the **Create model card** prompt to document details about your model.

**Step 1: Enter model details and intended use**

If your model is an AWS resource, specify the exact model name in this field to auto-populate model details. To browse existing model names, see **Models** in the Amazon SageMaker console. Each unique model name can have only one associated model card.

If your model is not an AWS resource, provide a unique name for your model. To add a model as an AWS resource, see [Create a model](#) in the *Amazon SageMaker Developer Guide*. Alternatively, you can add your model as a model package using SageMaker Marketplace or SageMaker Model Registry.
For more information on intended uses, see *Intended uses of a model*. For more information on risk ratings, see *Risk ratings*.

**Step 2: Enter training details**

Add any training details, training observations, datasets, hyperparameters, and details about the model's objective function to the model card.

The objective function in a model card can be any function that is optimized during training. This can include, but is not limited to, cost functions, loss functions, or objective metrics. In this section, document the objective function that is most critical to training your model.

We recommend that you catalog the following attributes of your objective function:

- Optimization direction
- Metric
- Description

For example, you might minimize (optimization direction) cross entropy loss (metric) for a binary classification problem (description) or maximize likelihood for logistic regression. Additionally, you can provide notes about why you chose this objective function over others.

**Step 3: Enter evaluation details**

If you have existing evaluation reports generated by SageMaker Clarify or Model Monitor, either provide an S3 URI for those reports or upload them manually to add them to the model card.

For more information on SageMaker Clarify, see *Run SageMaker Clarify Processing Jobs for Bias Analysis and Explainability*.

For more information on monitoring drift in model quality metrics using Model Monitor, see *Monitor model quality*.

To add your own evaluation report, choose **Generic model card evaluation**. All model card evaluation reports must be in the *Model card JSON schema*.

**Step 4: Enter additional details**

Add custom model card detail fields for any additional information that you want to address on your model card. For example, you might include the custom field *Line of business* with a value of *Personal finance*.
Save model card

After reviewing the information in your model card, choose Save in the lower right-hand corner to save your model card.

Create a model card using the SageMaker Python SDK

Before creating a model card, you must first define the content of your model card. When using the SageMaker Python SDK, model content consists of a model overview, training details, intended uses, evaluation details, and additional information.

You can create model cards for:

- Models that are hosted within SageMaker
- Model packages (models) within the SageMaker Model Registry
- Models that are hosted or registered outside of SageMaker

You can also create model cards without associating any models to them.

We recommend adding the models that you've trained to the SageMaker Model Registry. The model registry helps you catalog models and track model versions. When you create a model card, the information about the model from the model registry automatically populates the model card. You can edit the model card or add information to it after you create it.

For information about using the model registry, see Register and Deploy Models with Model Registry. For information about creating a model card from a model registry, see Create a model card for your model in the SageMaker Model Registry.

⚠️ Note

To use model cards with the SageMaker Python SDK, you first need to establish a SageMaker Session. For more information, see Session in the SageMaker Python SDK API reference.

To create a model card for models that aren't in the SageMaker Model Registry, see Create a model that isn't in the model registry.
Create a model that isn't in the model registry

Use the information in the following sections to create a model card for a model that you haven't added to the model registry.

**Step 1: Define model overview**

Define an overview of your model.

```python
model_overview = ModelOverview.from_model_name(
    model_name=model_name,
    sagemaker_session=sagemaker_session,
    model_description="A-description-of-your-model",
    problem_type="Problem-type", # For example, "Binary Classification"
    algorithm_type="Algorithm-type", # For example, "Logistic Regression"
    model_creator="Name-of-model-creator",
    model_owner="Name-of-model-owner",
)
```

If your model is an AWS resource, then overview information such as the model ARN, inference container URI, and the S3 location of model artifacts is automatically retrievable. Print the associated AWS metadata with the following commands:

```python
print(model_overview.model_id)
print(model_overview.inference_environment.container_image)
print(model_overview.model_artifact)
```

**Step 2: Define training details**

To define your model's training details, you must first define its objective function.

```python
objective_function = ObjectiveFunction(
    function=Function(
        function=ObjectiveFunctionEnum.MINIMIZE,
        facet=FacetEnum.LOSS,
    ),
    notes="An-explanation-about-objective-function",
)
```

Next, you can define your training details using your existing model overview, session, and objective function. Add any training observations here.
training_details = TrainingDetails.from_model_overview(
    model_overview=model_overview,
    sagemaker_session=sagemaker_session,
    objective_function=objective_function,
    training_observations="Model-training-observations",
)

Once again, if your model is an AWS resource, certain training details are autopopulated. Print the training job ARN, training container URI, and training metrics with the following commands:

print(training_details.training_job_details.training_arn)
print(training_details.training_job_details.training_environment.container_image)
print([{"name": i.name, "value": i.value} for i in
      training_details.training_job_details.training_metrics])

Define evaluation details

To define your model's evaluation details, you must first define one or more metric groups to describe metrics used for any evaluation jobs.

my_metric_group = MetricGroup(
    name="binary classification metrics",
    metric_data=[Metric(name="accuracy", type=MetricTypeEnum.NUMBER, value=0.5)]
)

Next, you can define your evaluation details using evaluation metrics and datasets for each evaluation job. Add any evaluation observations here and give your evaluation job a unique name.

evaluation_details = [
    EvaluationJob(
        name="Example-evaluation-job",
        evaluation_observation="Evaluation-observations",
        datasets="s3://path/to/evaluation/data",
        metric_groups=[my_metric_group],
    ),
]

If you have existing evaluation reports generated by SageMaker Clarify or SageMaker Model Monitor, upload them to Amazon S3 and provide an S3 URI to automatically parse evaluation
metrics. To add your own generic model card evaluation report, provide a report in the evaluation results JSON format.

```python
report_type = "clarify_bias.json"
example_evaluation_job.add_metric_group_from_json(f"example_metrics/{report_type}", EvaluationMetricTypeEnum.CLARIFY_BIAS)
```

### Step 3: Define intended uses

Define the intended uses of the model, including the general purpose of the model and the use cases for which it was intended. It is also recommended to include any factors that potentially affect this model's efficacy in a particular use case and your organization's risk rating of the model. For more information, see [Intended uses of a model](#) and [Risk ratings](#).

```python
intended_uses = IntendedUses(
purpose_of_model="Purpose-of-the-model",
intended_uses="The-intended-uses-of-this-model",
factors_affecting_model_efficiency="Any-factors-effecting-model-efficacy",
risk_rating=RiskRatingEnum.LOW,
explanations_for_risk_rating="Explanation-for-low-risk-rating",
)
```

### Define additional information

Lastly, you can add additional custom information to your model card. You can document any ethical considerations, caveats, and recommendations about the model. You can also add any custom details that you would like in the form of key-value pairs.

```python
additional_information = AdditionalInformation(
ethical_considerations="Any-ethical-considerations",
caveats_and_recommendations="Any-caveats-and-recommendations",
custom_details={"custom details1": "details-value"},
)
```

### Step 4: Create model card

Name your model card, define a model card, and then use that definition to create a model card using the SageMaker Python SDK.

```python
model_card_name = "my-model-card"
```
Create a model card for your model in the SageMaker Model Registry

Before you begin creating a model card, make sure that you've created a model package group and a model package. For more information about using model registry, see Register and Deploy Models with Model Registry.

⚠️ Important

You must have permissions to use the operations in SageMaker Model Registry. We recommend using AmazonSageMakerModelRegistryFullAccess AWS managed policy. For more information about the managed policy, see AWS Managed Policies for Model Registry.

Use the SageMaker Python SDK to create a model card for a model package within the SageMaker Model Registry. A model package is a model that you've trained. When you create a model card, Amazon SageMaker Model Cards automatically imports the data from the model package into the model card.

When you create a model card for a model package, Amazon SageMaker Model Card uses the DescribeModelPackage operation to add the data from the model package to the model card. The following are examples of the fields that can be imported from a model package into a model card:

- ModelDataUrl
- ModelPackageDescription
- ModelPackageGroupName
- ModelPackageStatus
• **ModelPackageVersion**

Use the following code to define the model package and create a model card from it:

```python
mp_details = ModelPackage.from_model_package_arn(
    model_package_arn="example_model_package_arn",
    sagemaker_session=sagemaker_session,
)

model_card_name = "example-model-card"
my_card = ModelCard(
    name=model_card_name,
    status=ModelCardStatusEnum.status,
    model_package_details=mp_details,
    sagemaker_session=sagemaker_session,
)
my_card.create()
```

For **status**, you're specifying the approval status of the model card. If you don't specify a status, SageMaker Model Cards uses the default value of DRAFT. If you don't specify a SageMaker session, SageMaker Model Cards uses the default SageMaker session.

You must specify a name for the model and the Amazon Resource Name (ARN) of the model package. For information about getting the Amazon Resource Name (ARN) for the model package, see [View the Details of a Model Version (Boto3)](view-the-details-of-a-model-version-boto3).

The model card that you've created from the model package might have information that is either missing or inaccurate. You can add information to the model card or edit it. For more information about managing your model cards, see [Manage model cards](manage-model-cards).

SageMaker Model Registry supports versioning of your model packages. You can version your model package and create a model card for each version. The information from model cards of preceding versions carry over to model cards created from subsequent versions. For example, you could have version 1, version 2, and version 3 of a model package. Suppose you've already created a model card for version 1, but you haven't created one for version 2. If you create a model card for version 3, Amazon SageMaker Model Cards automatically carries over the information from the model card for version 1 to the model card for version 3.
Note

You can also create model cards for model packages that don't use versioning. However, most machine learning workflows involve multiple versions of the same model, so we recommend doing the following:

1. Creating a version for each model package
2. Creating a model card for each version of the model package

Manage model cards

After you've created a model card, you can manage them. Managing model cards include the following actions:

- Editing a model card
- Deleting a model card
- Exporting a model card to a PDF

You can manage using either the Amazon SageMaker console or the SageMaker Python SDK.

Manage model cards using the console

Use the information in the following sections to manage your model cards with the Amazon SageMaker console.

Edit a model card

To edit a model card, navigate to the model card of your choice by selecting its name in the Amazon SageMaker Model Card console and choose Edit.

After you save a model card, you cannot edit the name of that model card. After you save a model card version, you cannot update that version of the model card. Any edits that you need to make are saved as a subsequent version in order to have an immutable record of model changes.

To view different versions of the model card, choose Actions, Select version, and then choose the version that you want to view.
Export a model card

Follow these steps to export a model card.

1. Go to the Amazon SageMaker Model Card console.
2. Choose the name of the model card you want to export.
3. In the model card overview, choose Actions and then Export PDF.
4. Enter an S3 URI or browse available S3 buckets for your model card PDF.
5. If your model card exports successfully, you can either choose Download PDF in the resulting banner or download your PDF directly from Amazon S3.

Delete a model card

Follow these steps to permanently delete one or more model cards.

1. Go to the Amazon SageMaker Model Cards console.
2. Check the box to the left of the name of the card(s) you want to delete.
3. Choose Delete in the upper right-hand corner.
4. Confirm your request to permanently delete one or more cards.

You can also delete a model card when viewing the model card overview in the console by choosing Actions and then Delete model card.

Manage model cards using the SageMaker Python SDK

Use the information in the following sections to manage your model cards with the Amazon SageMaker Python SDK.

Use model cards through the SageMaker Python SDK

You can create an Amazon SageMaker Model Card programmatically through the SageMaker Python SDK. For more information see Amazon SageMaker Model Cards in the SageMaker Python SDK API reference.

Edit a model card

You can edit a model card using the model_card.update() method. Updating a model card creates a new model card version in order to have an immutable record of model changes. You cannot update the name of a model card.
my_card.model_overview.model_description = "updated-model-decription"
my_card.update()

Export a model card

Specify an S3 output path and export your model card PDF to it with the following commands:

s3_output_path = f"s3://{bucket}/{prefix}/export"
pdf_s3_url = my_card.export_pdf(s3_output_path=s3_output_path).delete()

Delete a model card

Permanently delete a model card with the following command:

my_card.delete()

Sample notebooks

For more information on working with model cards through the SageMaker Python SDK, see the Amazon SageMaker Model Governance - Model Card example notebook.

Cross-account support for Amazon SageMaker Model Cards

Use cross-account support in Amazon SageMaker Model Cards to share model cards between AWS accounts. The account where the model cards are created is the model card account. Users in the model card account share them with the shared accounts. The users in a shared account can update the model cards or create PDFs of them.

Users in the model card account share their model cards through AWS Resource Access Manager (AWS RAM). AWS RAM helps you share resources across AWS accounts. For an introduction to AWS RAM, see What is AWS Resource Access Manager?

The following is the process to share model cards:

1. A user in the model card account sets up the cross-account model card sharing using AWS Resource Access Manager.
2. If the model cards are encrypted with AWS KMS keys, the user setting up model sharing must also provide users in the shared account with AWS KMS permissions.
3. A user in the shared account accepts the invite to the resource share.
4. A user in the shared account provides the other users with permissions to access the model cards.

If you're a user in the model card account, see the following sections:

- Set up cross-account model card sharing
- Set up AWS KMS permissions for the shared account
- Get responses to your resource share invitation

If you're a user in the shared account, see [Set up IAM user permissions in the shared account](#) about setting up permissions for yourself and the other users in the account.

**Set up cross-account model card sharing**

Use AWS Resource Access Manager (AWS RAM) to grant users in your AWS account access to view or update model cards created in a different AWS account.

To set up model card sharing, you must create a resource share. A resource share specifies:

- The resources being shared
- Who or what has access to the resources
- Managed permissions for the resources

For more information about resource shares, see [Terms and concepts for AWS RAM](#). We recommend taking the time to understand AWS RAM conceptually before you go through the process of creating a resource share.

⚠️ **Important**

You must have permissions to create a resource share. For more information about permissions, see [How AWS RAM works with IAM](#).

For procedures to create a resource share and additional information about them, see [Create a resource share](#).

When you go through the procedure of creating a resource share, you specify `sagemaker:ModelCard` as the resource type. You must also specify the Amazon Resource
Number (ARN) of the AWS RAM resource-based policy. You can specify either the default policy or the policy that has additional permissions to create a PDF of the model card.

With the default AWSRAMPermissionSageMakerModelCards resource-based policy, the users in the shared account have permissions to do the following operations:

- DescribeModelCard
- ListModelCardVersions
- UpdateModelCard

With the AWSRAMPermissionSageMakerModelCardsAllowExport resource-based policy, the users in the shared account have permissions to do all of the preceding actions. They also have permissions to create a model card export job and describe it through the following operations:

- CreateModelCardExportJob
- DescribeModelCardExportJob

The users in the shared account can create an export job to generate a PDF of a model card. They can also describe an export job that has been created to find the PDF's Amazon S3 URI.

Model cards and export jobs are resources. The model card account owns the export jobs created by a user in the shared account. For example, a user in account A shares model card X with shared account B. A user in account B creates export job Y for model card X that stores the output in an Amazon S3 location that the user in account B specifies. Even though account B created export job Y, it belongs to account A.

Each AWS account has resource quotas. For information about quotas related to model cards, see Amazon SageMaker endpoints and quotas.

**Set up AWS KMS permissions for the shared account**

If the model cards that you're sharing have been encrypted with AWS Key Management Service keys, you also need to share the access to the keys with the shared account. Otherwise, the users in the shared account can't view, update, or export the model cards. For an overview of AWS KMS, see AWS Key Management Service.

To provide AWS KMS permissions to users in the shared account, update your key policy with the following statement:
The preceding statement provides users in the shared account with `kms:Decrypt` and `kms:GenerateDataKey` permissions. With `kms:Decrypt`, users can decrypt the model cards. With `kms:GenerateDataKey`, users can encrypt the model cards that they update or the PDFs that they create.

**Get responses to your resource share invitation**

After you've created a resource share, the shared accounts that you've specified in the resource share receive an invitation to join it. They must accept the invite to access the resources.

For information about accepting a resource share invite, see [Using shared AWS resources](https://docs.aws.amazon.com/AWSResourceAccessManager/latest/UserGuide/using-shared-resources.html) in the *AWS Resource Access Manager User Guide*. 
Set up IAM user permissions in the shared account

The following information assumes that you've accepted the resource share invitation from the model card account. For more information about accepting a resource share invitation, see Using shared AWS resources.

You and the other users in your account use an IAM role to access the model cards shared from the model card account. Use the following template to change the policy of the IAM role. You can modify the template for your own use case.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:DescribeModelCard",
                "sagemaker:UpdateModelCard",
                "sagemaker:CreateModelCardExportJob",
                "sagemaker:ListModelCardVersions",
                "sagemaker:DescribeModelCardExportJob"
            ],
            "Resource": [
            ]
        },
        {
            "Effect": "Allow",
            "Action": "s3:PutObject",
            "Resource": "arn:aws:s3:::Amazon-S3-bucket-storing-the-pdf-of-the-model-card/model-card-name/*"
        }
    ]
}
```

To access model cards encrypted using AWS KMS, you must provide users in your account with the following AWS KMS permissions.
Use model cards through the low-level APIs

You can create an Amazon SageMaker Model Card directly through the SageMaker API or the AWS Command Line Interface (AWS CLI).

Note
When creating a model card with the low-level APIs, the content must be in the model card JSON schema and provided as a string. For more information, see Model card JSON schema.

SageMaker API

Use the following SageMaker API commands to work with Amazon SageMaker Model Cards:

- CreateModelCard
- DescribeModelCard
- ListModelCards
- ListModelCardVersions
- UpdateModelCard
- CreateModelCardExportJob
- DescribeModelCardExportJob
- ListModelCardExportJobs
- DeleteModelCard
AWS CLI

Use the following AWS CLI commands to work with Amazon SageMaker Model Cards:

- `create-model-card`
- `describe-model-card`
- `list-model-cards`
- `list-model-card-versions`
- `update-model-card`
- `create-model-card-export-job`
- `describe-model-card-export-job`
- `list-model-card-export-jobs`
- `delete-model-card`

Model card FAQs

Refer to the following FAQ items for answers to commonly asked questions about Amazon SageMaker Model Card.

Q. What is model risk?

A: You can use models for a variety of business applications ranging from predicting cyber attacks and approving loan applications to detecting the category of an email. Each of these applications assumes a different level of risk. For example, incorrectly detecting a cyber attack has much greater business impact than incorrectly categorizing an email. Given these varied risk profiles of a model, you can use model cards to provide a risk rating of low, medium, or high for a model. If you don’t know the risk of your model, you can set the status to unknown. Customers are responsible for assigning the risk profile for each model. Based on the risk rating, organizations may have different rules in place for deploying those models to production. For more information, see Risk ratings.

Q. What is the intended use of a model?

The intended use of a model describes how you should use the model in your production applications. This goes beyond technical requirements like the type of instance to which you should deploy a model and instead refers to the types of applications to create with the model, the scenarios in which you can expect a reasonable performance from the model, or the type of data to
use with the model. We recommend providing this information in the model card for better model governance. You can define a kind of model specification in the intended use field and ensure that model developers and consumers follow this specification while training and deploying their models. For more information, see [Intended uses of a model](#).

**Q. Does SageMaker autopopulate information in my model card?**

When you use the SageMaker Python SDK or the AWS console to create your model card, SageMaker autopopulates details about your SageMaker trained model in the card. This includes details about how the model was trained along with all the model details returned by the describe-model API call.

**Q. Can I customize a model card?**

Amazon SageMaker Model Cards have a defined structure to them that cannot be modified. This structure gives you guidance on what information should be captured in a model card. While you cannot change the structure of the model card, there is some flexibility introduced through custom properties in the Additional information section of the model card.

**Q. Can I edit a model card once it is created?**

Model cards have versions associated with them. A given model version is immutable across all attributes other than the model card status. If you make any other changes to the model card, such as evaluation metrics, description, or intended uses, SageMaker creates a new version of the model card to reflect the updated information. This is to ensure that a model card, once created, cannot be tampered with.

**Q. Can I create model cards for models that were not trained using SageMaker?**

A: Yes. You can create model cards for models not trained in SageMaker, but no information is automatically populated in the card. You must supply all the information needed in the model card for non-SageMaker models.

**Q. Can I export or share model cards?**

A: Yes. You can export each version of a model card to a PDF, downloaded, and share it.

**Q. Do I need to register my model in the Model Registry to use model cards?**

A: No. You can use model cards independently of the Model Registry.
Q. What is the difference between model cards and Model Registry?

A: Model cards are intended to provide organizations with a mechanism to document as much detail about their model as they like by following SageMaker's prescriptive guidance along with providing their own custom information. You can introduce model cards at the very start of the ML process and use them to define the business problem that the model should solve and any considerations to think about while using the model. After a model is trained, you can populate the model card associated with that model with information about the model and how it was trained. Model cards are associated with models and are immutable once associated with a model. This ensures that the model card is the single source of truth for all the information related to a model, including how it was trained and how it should be used.

The Model Registry is a catalog that stores metadata about your models. Each entry in the model registry corresponds to a unique model version. That model version contains information about the model such as where the model artifacts are stored in Amazon S3, what container is needed to deploy the model, and custom metadata that should be attached to the model.

Q. Are model card versions related to model versions in the Model Registry?

A: Model card versions and model versions are different entities in SageMaker. Each update to a model card results in a new version of that card. Model versions correspond to incrementally trained models that are registered in the Model Registry. A model card version may be linked to a specific model version in the Model Registry by way of the model ID field in the model card, but this is not necessary.

Q. Are model cards integrated with SageMaker Model Monitor?

A: No. You can upload the performance metrics computed by SageMaker Model Monitor to the model card by uploading a metrics file to Amazon S3 and linking that to the card, but there is no native integration between Model Monitor and model cards. Model dashboards are integrated with Model Monitor. For more information on model dashboards, see Amazon SageMaker Model Dashboard.

Amazon SageMaker Model Dashboard

Amazon SageMaker Model Dashboard is a centralized portal, accessible from the SageMaker console, where you can view, search, and explore all of the models in your account. You can track which models are deployed for inference and if they are used in batch transform jobs or hosted on endpoints. If you set up monitors with Amazon SageMaker Model Monitor, you can also track
the performance of your models as they make real-time predictions on live data. You can use the dashboard to find models that violate thresholds you set for data quality, model quality, bias and explainability. The dashboard's comprehensive presentation of all your monitor results helps you quickly identify models that don't have these metrics configured.

The Model Dashboard aggregates model-related information from several SageMaker features. In addition to the services provided in Model Monitor, you can view model cards, visualize workflow lineage, and track your endpoint performance. You no longer have to sort through logs, query in notebooks, or access other AWS services to collect the data you need. With a cohesive user experience and integration into existing services, SageMaker's Model Dashboard provides an out-of-the-box model governance solution to help you ensure quality coverage across all your models.

Prerequisites

To use the Model Dashboard, you should have one or more models in your account. You can train models using Amazon SageMaker or import models you've trained elsewhere. To create a model in SageMaker, you can use the `CreateModel` API. For more information, see `CreateModel`. You can also use SageMaker-provided ML environments, such as Amazon SageMaker Studio Classic, which provides project templates that set up model training and deployment for you. For information about how to get started with Studio Classic, see `Amazon SageMaker Studio Classic`.

While this is not a mandatory prerequisite, customers gain the most value out of the dashboard if they set up model monitoring jobs using SageMaker Model Monitor for models deployed to endpoints. For prerequisites and instructions on how to use SageMaker Model Monitor, see `Monitor data and model quality`.

Model Dashboard elements

The Model Dashboard view extracts high-level details from each model to provide a comprehensive summary of every model in your account. If your model is deployed for inference, the dashboard helps you track the performance of your model and endpoint in real time.

Important details to highlight in this page include:

- **Risk rating**: A user-specified parameter from the model card with a **low**, **medium**, or **high** value. The model card's risk rating is a categorical measure of the business impact of the model's predictions. Models are used for a variety of business applications, each of which assumes a different level of risk. For example, incorrectly detecting a cyber attack has much greater business impact than incorrectly categorizing an email. If you don't know the model risk, you can set it to **unknown**. For information about Amazon SageMaker Model Cards see `Model Cards`. 


Model Monitor alerts: Model Monitor alerts are a primary focus of the Model Dashboard, and reviewing the existing documentation on the various monitors provided by SageMaker is a helpful way to get started. For an in-depth explanation of the SageMaker Model Monitor feature and sample notebooks, see Monitor data and model quality.

The Model Dashboard displays Model Monitor status values by the following monitor types:

- **Data Quality**: Compares live data to training data. If they diverge, your model's inferences may no longer be accurate. For additional details about the Data Quality monitor, see Monitor data quality.

- **Model Quality**: Compares the predictions that the model makes with the actual Ground Truth labels that the model attempts to predict. For additional details about the Model Quality monitor, see Monitor model quality.

- **Bias Drift**: Compares the distribution of live data to training data, which can also cause inaccurate predictions. For additional details about the Bias Drift monitor, see Monitor Bias Drift for Models in Production.

- **Feature Attribution Drift**: Also known as explainability drift. Compares the relative rankings of your features in training data versus live data, which could also be a result of bias drift. For additional details about the Feature Attribution Drift monitor, see Monitor Feature Attribution Drift for Models in Production.

Each Model Monitor status is one of the following values:

- **None**: No monitor is scheduled
- **Inactive**: A monitor was scheduled, but it was deactivated
- **OK**: A monitor is scheduled and is active, and has not encountered the necessary number of violations in recent model monitor executions to raise an alert
- **Time and date**: An active monitor raised an alert at the specified time and date

- **Endpoint**: The endpoints which host your model for real-time inference. Within the Model Dashboard, you can select the endpoint column to view performance metrics such as CPU, GPU, disk, and memory utilization of your endpoints in real time to help you track the performance of your compute instances.

- **Batch transform job**: The most recent batch transform job that ran using this model. This column helps you determine if a model is actively used for batch inference.

- **Model details**: Each entry in the dashboard links to a model details page where you can dive deeper into an individual model. You can access the model's lineage graph, which visualizes the workflow from data preparation to deployment, and metadata for each step. You can also create
and view the model card, review alert details and history, assess the performance of your real-time endpoints, and access other infrastructure-related details.

**View Model Monitor schedules and alerts**

Using the Python SDK, you can create a model monitor for data quality, model quality, bias drift, or feature attribution drift. For more information about using SageMaker Model Monitor, see [Monitor data and model quality](#). The Model Dashboard populates information from all the monitors you create on all your models in your account. You can track the status of each monitor, which indicates whether your monitor is running as expected or failed due to an internal error. You can also activate or deactivate any monitor in the model details page itself. For instructions about how to view scheduled monitors for a model, see [View scheduled monitors](#). For instructions about how to activate or deactivate model monitors, see [Activate or deactivate a model monitor](#).

A properly-configured and actively-running model monitor might raise alerts, in which case the monitoring executions produce violation reports. For details about how alerts work and how to view alert results, history, and links to job reports for debug, see [View and edit alerts](#).

**View scheduled monitors**

**To view a model’s scheduled monitors, complete the following steps:**

1. Open the [SageMaker console](#).
2. Choose **Governance** in the left panel.
3. Choose **Model Dashboard**.
4. In the **Models** section of the Model Dashboard, select the model name of the scheduled monitors you want to view.
5. View the scheduled monitors in the **Monitor schedule** section. You can review the status for each monitor in the **Status schedule** column, which is one of the following values:

   - **Failed**: The monitoring schedule failed due to a problem with the configuration or settings (such as incorrect user permissions).
   - **Pending**: The monitor is in the process of becoming scheduled.
   - **Stopped**: The schedule is stopped by the user.
   - **Scheduled**: The schedule is created and runs at the frequency you specified.
Activate or deactivate a model monitor

To activate or deactivate a model monitor, complete the following steps:

1. Open the SageMaker console.
2. Choose Governance in the left panel.
3. Choose Model Dashboard.
4. In the Models section of the Model Dashboard, select the model name of the alert you want to modify.
5. Choose the radio box next to the monitor schedule of the alert you want to modify.
6. (optional) Choose Deactivate monitor schedule if you want to deactivate your monitor schedule.
7. (optional) Choose Activate monitor schedule if you want to activate your monitor schedule.

View and edit alerts

The Model Dashboard displays alerts you configured in Amazon CloudWatch. You can modify the alert criteria within the dashboard itself. The alert criteria depend upon two parameters:

- **Datapoints to alert**: Within the evaluation period, how many execution failures raise an alert.
- **Evaluation period**: The number of most recent monitoring executions to consider when evaluating alert status.

The following image shows an example scenario of a series of Model Monitor executions in which we set a hypothetical Evaluation period of 3 and a Datapoints to alert value of 2. After every monitoring execution, the number of failures are counted within the Evaluation period of 3. If the number of failures meets or exceeds the Datapoints to alert value 2, the monitor raises an alert and remains in alert status until the number of failures within the Evaluation period becomes less than 2 in subsequent iterations. In the image, the evaluation windows are red when the monitor raises an alert or remains in alert status, and green otherwise.

Note that even if the evaluation window size has not reached the Evaluation period of 3, as shown in the first 2 rows of the image, the monitor still raises an alert if the number of failures meets or exceeds the Datapoints to alert value of 2.
Within the monitor details page, you can view your alert history, edit existing alert criteria, and view job reports to help you debug alert failures. For instructions about how to view alert history or job reports for failed monitoring executions, see View alert history or job reports. For instructions about how to edit alert criteria, see Edit alert criteria.

View alert history or job reports

To view alert history or job reports of failed executions, complete the following steps:

1. Open the SageMaker console.
2. Choose Governance in the left panel.
3. Choose Model Dashboard.
4. In the Models section of the Model Dashboard, select the model name of the alert history you want to view.
5. In the Schedule name column, select the monitor name of the alert history you want to view.
6. To view alert history, select the Alert history tab.
7. (optional) To view job reports of monitoring executions, complete the following steps:
1. In the **Alert history** tab, choose **View executions** for the alert you want to investigate.

2. In the **Execution history** table, choose **View report** of the monitoring execution you want to investigate.

   **The report displays the following information:**

   - **Feature**: The user-defined ML feature monitored
   - **Constraint**: The specific check within the monitor
   - **Violation details**: Information about why the constraint was violated

**Edit alert criteria**

**To edit an alert in the Model Dashboard, complete the following steps:**

1. Open the [SageMaker console](https://sagemaker.amazonaws.com).
2. Choose **Governance** in the left panel.
3. Choose **Model Dashboard**.
4. In the **Models** section of the Model Dashboard, select the model name of the alert you want to modify.
5. Choose the radio box next to the monitor schedule of the alert you want to modify.
6. Choose **Edit Alert** in the **Monitor schedule** section.
7. (optional) Change **Datapoints to alert** if you want to change the number of failures within the **Evaluation period** that initiate an alert.
8. (optional) Change **Evaluation period** if you want to change the number of most recent monitoring executions to consider when evaluating alert status.

**View a model lineage graph**

When you train a model, Amazon SageMaker creates a visualization of your entire ML workflow from data preparation to deployment. This visualization is called a model lineage graph and uses entities to represent individual steps in your workflow. For example, a basic model lineage graph might have an entity representing your training set, which is associated with an entity representing your training job, which is associated with another entity representing your model.
In addition, the graph stores information about each step in your workflow. With this information, you can recreate any step in the workflow or track model and dataset lineage. For example, SageMaker Lineage stores the S3 URI of your input data sources with each job so you can perform further analysis of the data sources for compliance verification.

While the model lineage graph can help you view the steps in individual workflows, there are many other capabilities that you can leverage using the AWS SDK. For example, with the AWS SDK you can create or query your entities. For more information about the full set of features in SageMaker Lineage and example notebooks, see Amazon SageMaker ML Lineage Tracking.

Introduction to entities

Amazon SageMaker automatically creates tracking entities for SageMaker jobs, models, model packages, and endpoints if the data is available. For a basic workflow, suppose you train a model using a dataset. SageMaker automatically generates a lineage graph with three entities:

- **Dataset**: A type of artifact, which is an entity representing a URI addressable object or data. An artifact is generally either an input or an output to a trial component or action.
- **TrainingJob**: A type of trial component, which is an entity representing processing, training, and transform jobs.
- **Model**: Another type of artifact. Like the **Dataset** artifact, a **Model** is a URI addressable object. In this case, it is an output of the **TrainingJob** trial component.

Your model lineage graph expands quickly if you add additional steps to your workflow, such as data preprocessing or postprocessing, if you deploy your model to an endpoint, or if you include your model in a model package, among many other possibilities. For the complete list of SageMaker entities, see Amazon SageMaker ML Lineage Tracking.

Entity properties

Each node in the graph displays the entity type, but you can choose the vertical ellipsis to the right of the entity type to see specific details related to your workflow. In our previous barebones lineage graph, you can choose the vertical ellipsis next to **DataSet** to see specific values for the following properties (common to all artifact entities):

- **Name**: The name of your dataset.
- **Source URI**: The Amazon S3 location of your dataset.
For the TrainingJob entity, you can see the specific values for the following properties (common to all TrialComponent entities):

- **Name**: The name of the training job.
- **Job ARN**: The Amazon Resource Name (ARN) of your training job.

For the **Model** entity, you see the same properties as listed for **DataSet** since they are both artifact entities. For a list of the entities and their associated properties, see [Lineage Tracking Entities](#).

**Entity queries**

Amazon SageMaker automatically generates graphs of lineage entities as you use them. However if you are running many iterations of an experiment and don't want to view every lineage graph, the AWS SDK can help you perform queries across all your workflows. For example, you can query your lineage entities for all the processing jobs that use an endpoint. Or, you can see all the downstream trails that use an artifact. For a list of all the queries you can perform, see [Querying Lineage Entities](#).

**View a model's lineage graph**

To view the lineage graph for a model, complete the following steps:

1. Open the [SageMaker console](#).
2. Choose **Governance** in the left panel.
3. Choose **Model Dashboard**.
4. In the **Models** section of the Model Dashboard, select the model name of the lineage graph you want to view.
5. Choose **View lineage** in the **Model Overview** section.

**View Endpoint Status**

If you want to use your trained model to perform inference on live data, you deploy your model to a real-time endpoint. To ensure appropriate latency of your predictions, you want to make sure the instances that host your model are running efficiently. Model Dashboard's endpoint monitoring feature displays real-time information about your endpoint configuration and helps you track endpoint performance with metrics.

**Monitor settings**
The Model Dashboard links to existing SageMaker endpoint details pages which display real-time graphs of metrics you can select in Amazon CloudWatch. Within your dashboard, you can track these metrics as your endpoint is handling real-time inference requests. Some metrics you can select are the following:

- **CpuUtilization**: The sum of each individual CPU core's utilization, with each ranging from 0%–100%.
- **MemoryUtilization**: The percentage of memory used by the containers on an instance, ranging from 0%–100%.
- **DiskUtilization**: The percentage of disk space used by the containers on an instance, ranging from 0%–100%.

For the complete list of metrics you can view in real time, see [Monitor Amazon SageMaker with Amazon CloudWatch](#).

**Runtime settings**

Amazon SageMaker supports automatic scaling (auto scaling) for your hosted models. Auto scaling dynamically adjusts the number of instances provisioned for a model in response to changes in your workload. When the workload increases, auto scaling brings more instances online. When the workload decreases, auto scaling removes unnecessary instances so that you don't pay for provisioned instances that you aren't using. You can customize the following runtime settings in the Model Dashboard:

- **Update weights**: Change the amount of workload assigned to each instance with numerical weighting. For more information about instance weighting during auto scaling, see [Configure instance weighting for Amazon EC2 Auto Scaling](#).
- **Update instance count**: Change the number of total instances that can service your workload when it increases.

For more information about endpoint runtime settings, see [CreateEndpointConfig](#).

**Endpoint configuration settings**

Endpoint configuration settings display the settings you specified when you created the endpoint. These settings inform SageMaker which resources to provision for your endpoint. Some settings included are the following:
• **Data capture**: You can choose to capture information about your endpoint's inputs and outputs. For example, you may want to sample incoming traffic to see if the results correlate to training data. You can customize your sampling frequency, the format of the stored data, and Amazon S3 location of stored data. For more information about setting up your data capture configuration, see [Capture data](#).

• **Production variants**: See the previous discussion in *Runtime settings*.

• **Async invocation configuration**: If your endpoint is asynchronous, this section includes the maximum number of concurrent requests sent by the SageMaker client to the model container, the Amazon S3 location of your success and failure notifications, and the output location of your endpoint outputs. For more information about asynchronous outputs, see [Create, invoke, and update an Asynchronous Endpoint](#).

• **Encryption key**: You can enter your encryption key if you want to encrypt your outputs.

For more information about endpoint configuration settings, see [CreateEndpointConfig](#).

**View status and configuration for an endpoint**

To view the status and configuration for a model's endpoint, complete the following steps:

1. Open the [SageMaker console](#).
2. Choose **Governance** in the left panel.
3. Choose **Model Dashboard**.
4. In the **Models** section of the Model Dashboard, select the model name of the endpoint you want to view.
5. Select the endpoint name in the **Endpoints** section.

**Model Dashboard FAQ**

Refer to the following FAQ topics for answers to commonly asked questions about Amazon SageMaker Model Dashboard.

**Q. What is Model Dashboard?**

Amazon SageMaker Model Dashboard is a centralized repository of all models created in your account. The models are generally the outputs of SageMaker training jobs, but you can also import
models trained elsewhere and host them on SageMaker. Model Dashboard provides a single interface for IT administrators, model risk managers, and business leaders to track all deployed models and aggregates data from multiple AWS services to provide indicators about how your models are performing. You can view details about model endpoints, batch transform jobs, and monitoring jobs for additional insights into model performance. The dashboard’s visual display helps you quickly identify which models have missing or inactive monitors so you can ensure all models are periodically checked for data drift, model drift, bias drift, and feature attribution drift. Lastly, the dashboard’s ready access to model details helps you dive deep so you can access logs, infrastructure-related information, and resources to help you debug monitoring failures.

Q. What are the prerequisites to use Model Dashboard?

You should have one or more models created in SageMaker, either trained on SageMaker or externally trained. While this is not a mandatory prerequisite, you gain the most value from the dashboard if you set up model monitoring jobs via Amazon SageMaker Model Monitor for models deployed to endpoints.

Q. Who should use Model Dashboard?

Model risk managers, ML practitioners, data scientists and business leaders can get a comprehensive overview of models using the Model Dashboard. The dashboard aggregates and displays data from Amazon SageMaker Model Cards, Endpoints and Model Monitor services to display valuable information such as model metadata from the model card and model registry, endpoints where the models are deployed, and insights from model monitoring.

Q. How do I use Model Dashboard?

Model Dashboard is available out of the box with Amazon SageMaker and does not require any prior configuration. However, if you have set up model monitoring jobs using SageMaker Model Monitor and Clarify, you use Amazon CloudWatch to configure alerts that raise a flag in the dashboard when model performance deviates from an acceptable range. You can create and add new model cards to the dashboard, and view all the monitoring results associated with endpoints. Model Dashboard currently does not support cross-account models.

Q. What is Amazon SageMaker Model Monitor?

With Amazon SageMaker Model Monitor, you can select the data you want to monitor and analyze without writing any code. SageMaker Model Monitor lets you select data, such as prediction output, from a menu of options and captures metadata such as timestamp, model name, and endpoint
so you can analyze model predictions. You can specify the sampling rate of data capture as a percentage of overall traffic in the case of high volume real-time predictions. This data is stored in your own Amazon S3 bucket. You can also encrypt this data, configure fine-grained security, define data retention policies, and implement access control mechanisms for secure access.

Q. What types of model monitors does SageMaker support?

SageMaker Model Monitor provides the following types of model monitors:

- **Data Quality**: Monitor drift in data quality.
- **Model Quality**: Monitor drift in model quality metrics, such as accuracy.
- **Bias Drift for Models in Production**: Monitor bias in your model's predictions by comparing the distribution of training and live data.
- **Feature Attribution Drift for Models in Production**: Monitor drift in feature attribution by comparing the relative rankings of features in training and live data.

Q. What inference methods does SageMaker Model Monitor support?

Model Monitor currently supports endpoints that host a single model for real-time inference and does not support monitoring of multi-model endpoints.

Q. How can I get started with SageMaker Model Monitor?

You can use the following resources to get started with model monitoring:

- Data quality monitor example notebook
- Model quality monitor example notebook
- Bias drift monitor example notebook
- Feature attribution drift monitor example notebook

For more examples of model monitoring, see the GitHub repository amazon-sagemaker-examples.

Q. How does Model Monitor work?

Amazon SageMaker Model Monitor automatically monitors machine learning models in production, using rules to detect drift in your model. Model Monitor notifies you when quality issues arise through alerts. To learn more, see How Model Monitor Works.
Q. When and how do you bring your own container (BYOC) for Model Monitor?

Model Monitor computes model metrics and statistics on tabular data only. For use cases other than tabular datasets, such as images or text, you can bring your own containers (BYOC) to monitor your data and models. For example, you can use BYOC to monitor an image classification model that takes images as input and outputs a label. To learn more about container contracts, see Bring Your Own Containers.

Q. Where can I find examples of BYOC for Model Monitor?

You can find helpful BYOC examples in the following links:

- Monitor data and model quality
- GitHub example repository
- Bring Your Own Containers
- Detecting data drift in NLP using BYOC Model Monitor
- Detecting and analyzing incorrect predictions in CV

Q. How do I integrate Model Monitor with SageMaker Pipelines?

For details about how to integrate Model Monitor and SageMaker Pipelines, see Amazon SageMaker Pipelines now integrates with SageMaker Model Monitor and SageMaker Clarify.

For an example, see the GitHub sample notebook SageMaker Pipelines integration with Model Monitor and Clarify.

Q. Are there any performance concerns using DataCapture?

When turned on, data capture occurs asynchronously on the SageMaker endpoints. To prevent impact to inference requests, DataCapture stops capturing requests at high levels of disk usage. It is recommended you keep your disk utilization below 75% to ensure DataCapture continues capturing requests.
Use Docker containers to build models

Amazon SageMaker makes extensive use of *Docker containers* for build and runtime tasks. SageMaker provides pre-built Docker images for its built-in algorithms and the supported deep learning frameworks used for training and inference. Using containers, you can train machine learning algorithms and deploy models quickly and reliably at any scale. The topics in this section show how to deploy these containers for your own use cases. For information about how to bring your own containers for use with Amazon SageMaker Studio Classic, see Bring your own SageMaker image.

**Topics**

- Scenarios for Running Scripts, Training Algorithms, or Deploying Models with SageMaker
- Docker Container Basics
- Use Pre-built SageMaker Docker images
- Adapting your own Docker container to work with SageMaker
- Create a container with your own algorithms and models
- Examples and More Information: Use Your Own Algorithm or Model
- Troubleshooting your Docker containers

**Scenarios for Running Scripts, Training Algorithms, or Deploying Models with SageMaker**

Amazon SageMaker always uses Docker containers when running scripts, training algorithms, and deploying models. Your level of engagement with containers depends on your use case.

The following decision tree illustrates three main scenarios: Use cases for using pre-built Docker containers with SageMaker; Use cases for extending a pre-built Docker container; Use case for building your own container.
Topics

- Use cases for using pre-built Docker containers with SageMaker
- Use cases for extending a pre-built Docker container
- Use case for building your own container

Use cases for using pre-built Docker containers with SageMaker

Consider the following use cases when using containers with SageMaker:

- **Pre-built SageMaker algorithm** – Use the image that comes with the built-in algorithm. See Use Amazon SageMaker Built-in Algorithms or Pre-trained Models for more information.

- **Custom model with pre-built SageMaker container** – If you train or deploy a custom model, but use a framework that has a pre-built SageMaker container including TensorFlow and PyTorch, choose one of the following options:
• If you don’t need a custom package, and the container already includes all required packages: Use the pre-built Docker image associated with your framework. For more information, see Use Pre-built SageMaker Docker images.

• If you need a custom package installed into one of the pre-built containers: Confirm that the pre-built Docker image allows a requirements.txt file, or extend the pre-built container based on the following use cases.

Use cases for extending a pre-built Docker container

The following are use cases for extending a pre-built Docker container:

• **You can't import the dependencies** – Extend the pre-built Docker image associated with your framework. See Extend a Pre-built Container for more information.

• **You can't import the dependencies in the pre-built container and the pre-built container supports requirements.txt** – Add all the required dependencies in requirements.txt. The following frameworks support using requirements.txt.
  
  • TensorFlow
  
  • Chainer
  
  • Sci-kit learn
  
  • PyTorch
  
  • Apache MXNet

Use case for building your own container

If you build or train a custom model and require custom framework that does not have a pre-built image, build a custom container.

As an example use case of training and deploying a TensorFlow model, the following guide shows how to determine which option from the previous sections of Use cases fits to the case.

Assume that you have the following requirements for training and deploying a TensorFlow model.

• A TensorFlow model is a custom model.

• Because a TensorFlow model is going to be built in the TensorFlow framework, use the TensorFlow pre-built framework container to train and host the model.
• If you require custom packages in either your `entrypoint` script or `inference script`, either extend the pre-built container or use a `requirements.txt` file to install dependencies at runtime.

After you determine the type of container that you need, the following list provides details about the previously listed options.

• **Use a built-in SageMaker algorithm or framework.** For most use cases, you can use the built-in algorithms and frameworks without worrying about containers. You can train and deploy these algorithms from the SageMaker console, the AWS Command Line Interface (AWS CLI), a Python notebook, or the Amazon SageMaker Python SDK. You can do that by specifying the algorithm or framework version when creating your Estimator. The available built-in algorithms are itemized and described in the Use Amazon SageMaker Built-in Algorithms or Pre-trained Models topic. For more information about the available frameworks, see ML Frameworks and Languages. For an example of how to train and deploy a built-in algorithm using a Jupyter notebook running in a SageMaker notebook instance, see the Get started topic.

• **Use pre-built SageMaker container images.** Alternatively, you can use the built-in algorithms and frameworks using Docker containers. SageMaker provides containers for its built-in algorithms and pre-built Docker images for some of the most common machine learning frameworks, such as Apache MXNet, TensorFlow, PyTorch, and Chainer. For a full list of the available SageMaker images, see Available Deep Learning Containers Images. It also supports machine learning libraries such as scikit-learn and SparkML. If you use the Amazon SageMaker Python SDK, you can deploy the containers by passing the full container URI to their respective SageMaker SDK Estimator class. For the full list of deep learning frameworks that are currently supported by SageMaker, see Prebuilt SageMaker Docker Images for Deep Learning. For information about the scikit-learn and SparkML pre-built container images, see Prebuilt Amazon SageMaker Docker Images for Scikit-learn and Spark ML. For more information about using frameworks with the Amazon SageMaker Python SDK, see their respective topics in Machine Learning Frameworks and Languages.

• **Extend a pre-built SageMaker container image.** If you would like to extend a pre-built SageMaker algorithm or model Docker image, you can modify the SageMaker image to satisfy your needs. For an example, see Extending our PyTorch containers.

• **Adapt an existing container image:** If you would like to adapt a pre-existing container image to work with SageMaker, you must modify the Docker container to enable either the SageMaker Training or Inference toolkit. For an example that shows how to build your own containers to train and host an algorithm, see Bring Your Own R Algorithm.
Docker Container Basics

Docker is a program that performs operating system-level virtualization for installing, distributing, and managing software. It packages applications and their dependencies into virtual containers that provide isolation, portability, and security. With Docker, you can ship code faster, standardize application operations, seamlessly move code, and economize by improving resource utilization. For more general information about Docker, see Docker overview.

The following information outlines the most significant aspects of using Docker containers with Amazon SageMaker.

SageMaker Functions

SageMaker uses Docker containers in the backend to manage training and inference processes. SageMaker abstracts away from this process, so it happens automatically when an estimator is used. While you don’t need to use Docker containers explicitly with SageMaker for most use cases, you can use Docker containers to extend and customize SageMaker functionality.

Containers with Amazon SageMaker Studio Classic

Studio Classic runs from a Docker container and uses it to manage functionality. As a result, you must create your Docker container following the steps in Bring your own SageMaker image.

Use Pre-built SageMaker Docker images

Amazon SageMaker provides containers for its built-in algorithms and pre-built Docker images for some of the most common machine learning frameworks, such as Apache MXNet, TensorFlow, PyTorch, and Chainer. It also supports machine learning libraries such as scikit-learn and SparkML.

You can use these images from your SageMaker notebook instance or SageMaker Studio. You can also extend the pre-built SageMaker images to include libraries and needed functionality. The following topics give information about the available images and how to use them.

For the Docker registry path and other parameters for each of the Amazon SageMaker provided algorithms and Deep Learning Containers (DLC), see Docker Registry Paths and Example Code.

Note

For information on Docker images for developing reinforcement learning (RL) solutions in SageMaker, see SageMaker RL Containers.
Prebuilt SageMaker Docker Images for Deep Learning

Amazon SageMaker provides prebuilt Docker images that include deep learning frameworks and other dependencies needed for training and inference. For a complete list of the prebuilt Docker images managed by SageMaker, see [Docker Registry Paths and Example Code](#).

Using the SageMaker Python SDK

With the [SageMaker Python SDK](#), you can train and deploy models using these popular deep learning frameworks. For instructions on installing and using the SDK, see [Amazon SageMaker Python SDK](#). The following table lists the available frameworks and instructions on how to use them with the [SageMaker Python SDK](#):

<table>
<thead>
<tr>
<th>Framework</th>
<th>Instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>TensorFlow</td>
<td><a href="#">Using TensorFlow with the SageMaker Python SDK</a></td>
</tr>
<tr>
<td>MXNet</td>
<td><a href="#">Using MXNet with the SageMaker Python SDK</a></td>
</tr>
<tr>
<td>PyTorch</td>
<td><a href="#">Using PyTorch with the SageMaker Python SDK</a></td>
</tr>
<tr>
<td>Chainer</td>
<td><a href="#">Using Chainer with the SageMaker Python SDK</a></td>
</tr>
<tr>
<td>Hugging Face</td>
<td><a href="#">Using Hugging Face with the SageMaker Python SDK</a></td>
</tr>
</tbody>
</table>

Extending Prebuilt SageMaker Docker Images

You can customize these prebuilt containers or extend them to handle any additional functional requirements for your algorithm or model that the prebuilt SageMaker Docker image doesn't support. For an example, see [Fine-tuning and deploying a BERTopic model on SageMaker with your own scripts and dataset, by extending existing PyTorch containers](#).
You can also use prebuilt containers to deploy your custom models or models that have been trained in a framework other than SageMaker. For an overview of the process of bringing the trained model artifacts into SageMaker and hosting them at an endpoint, see Bring Your Own Pretrained MXNet or TensorFlow Models into Amazon SageMaker.

Prebuilt Amazon SageMaker Docker Images for Scikit-learn and Spark ML

SageMaker provides prebuilt Docker images that install the scikit-learn and Spark ML libraries. These libraries also include the dependencies needed to build Docker images that are compatible with SageMaker using the Amazon SageMaker Python SDK. With the SDK, you can use scikit-learn for machine learning tasks and use Spark ML to create and tune machine learning pipelines. For instructions on installing and using the SDK, see SageMaker Python SDK.

Using the SageMaker Python SDK

The following table contains links to the GitHub repositories with the source code for the scikit-learn and Spark ML containers. The table also contains links to instructions that show how use these containers with Python SDK estimators to run your own training algorithms and hosting your own models.

<table>
<thead>
<tr>
<th>Library</th>
<th>Prebuilt Docker Image Source Code</th>
<th>Instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>scikit-learn</td>
<td><a href="#">SageMaker Scikit-learn Containers</a></td>
<td><a href="#">Using Scikit-learn with the Amazon SageMaker Python SDK</a></td>
</tr>
<tr>
<td>Spark ML</td>
<td><a href="#">SageMaker Spark ML Serving Containers</a></td>
<td><a href="#">SparkML Python SDK Documentation</a></td>
</tr>
</tbody>
</table>

For more information and links to github repositories, see Use Scikit-learn with Amazon SageMaker and Use SparkML Serving with Amazon SageMaker.

Specifying the Prebuilt Images Manually

If you are not using the SageMaker Python SDK and one of its estimators to manage the container, you have to retrieve the relevant prebuilt container manually. The SageMaker prebuilt Docker images are stored in Amazon Elastic Container Registry (Amazon ECR). You can push or pull them...
using their fullname registry addresses. SageMaker uses the following Docker Image URL patterns for scikit-learn and Spark ML:

- `<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/sagemaker-scikit-learn:<SCIKIT-LEARN_VERSION>-cpu-py<PYTHON_VERSION>`

  For example, `746614075791.dkr.ecr.us-west-1.amazonaws.com/sagemaker-scikit-learn:1.2-1-cpu-py3`

- `<ACCOUNT_ID>.dkr.ecr.<REGION_NAME>.amazonaws.com/sagemaker-sparkml-serving:<SPARK-ML_VERSION>`

  For example, `341280168497.dkr.ecr.ca-central-1.amazonaws.com/sagemaker-sparkml-serving:2.4`

For account IDs and AWS Region names, see [Docker Registry Paths and Example Code](#).

**Finding Available Images**

Use the following commands to find out which versions of the images are available. For example, use the following to find the available sagemaker-sparkml-serving image in the ca-central-1 Region:

```bash
aws ecr describe-images --region ca-central-1 --registry-id 341280168497 --repository-name sagemaker-sparkml-serving
```

**Train a Deep Graph Network**

In this overview, you learn how to get started with a deep graph network by using one of the DGL containers in Amazon Elastic Container Registry (Amazon ECR). You can also see links to practical examples for deep graph networks.

**What Is a Deep Graph Network?**

Deep graph networks refer to a type of neural network that is trained to solve graph problems. A deep graph network uses an underlying deep learning framework like PyTorch or MXNet. The potential for graph networks in practical AI applications is highlighted in the Amazon SageMaker...
tutorials for Deep Graph Library (DGL). Examples for training models on graph datasets include social networks, knowledge bases, biology, and chemistry.

Figure 1. The DGL ecosystem

Several examples are provided using Amazon SageMaker's deep learning containers that are preconfigured with DGL. If you have special modules you want to use with DGL, you can also build your own container. The examples involve heterographs, which are graphs that have multiple types of nodes and edges, and draw on a variety of applications across disparate scientific fields, such as bioinformatics and social network analysis. DGL provides a wide array of graph neural network implementations for different types models. Some of the highlights include:

- Graph convolutional network (GCN)
- Relational graph convolutional network (R-GCN)
- Graph attention network (GAT)
- Deep generative models of graphs (DGMG)
- Junction tree neural network (JTNN)

**Get Started**

DGL is available as a deep learning container in Amazon ECR. You can select deep learning containers when you write your estimator function in an Amazon SageMaker notebook. You can also craft your own custom container with DGL by following the [Bring Your Own Container](#) guide. The easiest way to get started with a deep graph network uses one of the DGL containers in Amazon ECR.

**Note**

Backend framework support is limited to PyTorch and MXNet.

**Setup**

If you are using Amazon SageMaker Studio, you need to clone the examples repository first. If you are using a notebook instance, you can find the examples by choosing the SageMaker icon at bottom of the left toolbar.

**To clone the Amazon SageMaker SDK and notebook examples repository**

1. From the JupyterLab view in Amazon SageMaker, go to the File Browser at the top of the left toolbar. From the File Browser panel, you can see a new navigation at the top of the panel.
2. Choose the icon on the far right to clone a Git repository.
3. Add the repository URL: https://github.com/awslabs/amazon-sagemaker-examples.git
4. Browse the newly added folder and its contents. The DGL examples are stored in the sagemaker-python-sdk folder.
Run a Graph Network Training Example

To train a deep graph network

1. From the JupyterLab view in Amazon SageMaker, browse the example notebooks and look for DGL folders. Several files may be included to support an example. Examine the README for any prerequisites.
2. Run the .ipynb notebook example.
3. Find the estimator function, and note the line where it is using an Amazon ECR container for DGL and a specific instance type. You may want to update this to use a container in your preferred Region.
4. Run the function to launch the instance and use the DGL container for training a graph network. Charges are incurred for launching this instance. The instance self-terminates when the training is complete.

Examples

An example of knowledge graph embedding (KGE) is provided. It uses the Freebase dataset, a knowledge base of general facts. An example use case would be to graph the relationships of persons and predict their nationality.

An example implementation of a graph convolutional network (GCN) shows how you can train a graph network to predict toxicity. A physiology dataset, Tox21, provides toxicity measurements for how substances affect biological responses.

Another GCN example shows you how to train a graph network on a scientific publications bibliography dataset, known as Cora. You can use it to find relationships between authors, topics, and conferences.

The last example is a recommender system for movie reviews. It uses a graph convolutional matrix completion (GCMC) network trained on the MovieLens datasets. These datasets consist of movie titles, genres, and ratings by users.

Use a Deep Learning Container with DGL

The following example uses preconfigured deep learning containers. This is the easiest to try since it works out of the box on Amazon SageMaker.

- Semi-supervised classification of a knowledge base using a GCN
Bring Your Own Container with DGL

The following examples enable you to bring your own container (BYOC). Read the BYOC guide and familiarize yourself with that process before trying these. Configuration is required.

- Molecular property prediction of toxicity using a GCN
- Recommender system for movies using a GCMC implementation

Extend a Pre-built Container

If a pre-built SageMaker container doesn't fulfill all of your requirements, you can extend the existing image to accommodate your needs. Even if there is direct support for your environment or framework, you may want to add additional functionality or configure your container environment differently. By extending a pre-built image, you can leverage the included deep learning libraries and settings without having to create an image from scratch. You can extend the container to add libraries, modify settings, and install additional dependencies.

The following tutorial shows how to extend a pre-built SageMaker image and publish it to Amazon ECR.

Topics

- Requirements to Extend a Pre-built Container
- Extend SageMaker Containers to Run a Python Script

Requirements to Extend a Pre-built Container

To extend a pre-built SageMaker image, you need to set the following environment variables within your Dockerfile. For more information on environment variables with SageMaker containers, see the SageMaker Training Toolkit GitHub repo.

- SAGEMAKER_SUBMIT_DIRECTORY: The directory within the container in which the Python script for training is located.
- SAGEMAKER_PROGRAM: The Python script that should be invoked and used as the entry point for training.

You can also install additional libraries by including the following in your Dockerfile:
RUN pip install <library>

The following tutorial shows how to use these environment variables.

### Extend SageMaker Containers to Run a Python Script

In this tutorial, you learn how to extend the SageMaker PyTorch container with a Python file that uses the CIFAR-10 dataset. By extending the SageMaker PyTorch container, you utilize the existing training solution made to work with SageMaker. This tutorial extends a training image, but the same steps can be taken to extend an inference image. For a full list of the available images, see [Available Deep Learning Containers Images](#).

To run your own training model using the SageMaker containers, build a Docker container through a SageMaker Notebook instance.

**Step 1: Create an SageMaker Notebook Instance**

1. Open the [SageMaker console](#).
2. In the left navigation pane, choose Notebook, choose Notebook instances, and then choose Create notebook instance.
3. On the Create notebook instance page, provide the following information:
   a. For Notebook instance name, enter RunScriptNotebookInstance.
   b. For Notebook Instance type, choose ml.t2.medium.
   c. In the Permissions and encryption section, do the following:
      i. For IAM role, choose Create a new role.
      ii. On the Create an IAM role page, choose Specific S3 buckets, specify an Amazon S3 bucket named sagemaker-run-script, and then choose Create role.
         SageMaker creates an IAM role named AmazonSageMaker-ExecutionRole-YYYYMMDDTHHmmSS, such as AmazonSageMaker-ExecutionRole-20190429T110788. Note that the execution role naming convention uses the date and time when the role was created, separated by a T.
   d. For Root Access, choose Enable.
   e. Choose Create notebook instance.
4. On the Notebook instances page, the Status is Pending. It can take a few minutes for Amazon CloudWatch Internet Monitor to launch a machine learning compute instance—in this case,
it launches a notebook instance—and attach an ML storage volume to it. The notebook instance has a preconfigured Jupyter notebook server and a set of Anaconda libraries. For more information, see CreateNotebookInstance.

5. In the Permissions and encryption section, copy the IAM role ARN number, and paste it into a notepad file to save it temporarily. You use this IAM role ARN number later to configure a local training estimator in the notebook instance. The IAM role ARN number looks like the following: 'arn:aws:iam::111122223333:role/service-role/AmazonSageMaker-ExecutionRole-20190429T110788'

6. After the status of the notebook instance changes to InService, choose Open JupyterLab.

**Step 2: Create and Upload the Dockerfile and Python Training Scripts**

1. After JupyterLab opens, create a new folder in the home directory of your JupyterLab. In the upper-left corner, choose the New Folder icon, and then enter the folder name docker_test_folder.

2. Create a Dockerfile text file in the docker_test_folder directory.
   a. Choose the New Launcher icon (+) in the upper-left corner.
   b. In the right pane under the Other section, choose Text File.
   c. Paste the following Dockerfile sample code into your text file.

```plaintext
# SageMaker PyTorch image
FROM 763104351884.dkr.ecr.us-east-1.amazonaws.com/pytorch-training:1.5.1-cpu-py36-ubuntu16.04

ENV PATH="/opt/ml/code:${PATH}"

# this environment variable is used by the SageMaker PyTorch container to determine our user code directory.
ENV SAGEMAKER_SUBMIT_DIRECTORY /opt/ml/code

# /opt/ml and all subdirectories are utilized by SageMaker, use the /code subdirectory to store your user code.
COPY cifar10.py /opt/ml/code/cifar10.py

# Defines cifar10.py as script entrypoint
```

Extend a Pre-built Container
The Dockerfile script performs the following tasks:

- `FROM 763104351884.dkr.ecr.us-east-1.amazonaws.com/pytorch-training:1.5.1-cpu-py36-ubuntu16.04` – Downloads the SageMaker PyTorch base image. You can replace this with any SageMaker base image you want to bring to build containers.

- `ENV SAGEMAKER_SUBMIT_DIRECTORY /opt/ml/code` – Sets `/opt/ml/code` as the training script directory.

- `COPY cifar10.py /opt/ml/code/cifar10.py` – Copies the script to the location inside the container that is expected by SageMaker. The script must be located in this folder.

- `ENV SAGEMAKER_PROGRAM cifar10.py` – Sets your `cifar10.py` training script as the entrypoint script.

- On the left directory navigation pane, the text file name might automatically be named `untitled.txt`. To rename the file, right-click the file, choose **Rename**, rename the file as `Dockerfile` without the `.txt` extension, and then press Ctrl+s or Command+s to save the file.

3. **Create or upload a training script `cifar10.py` in the `docker_test_folder`. You can use the following example script for this exercise.**

```python
import ast
import argparse
import logging
import os
import torch
import torch.distributed as dist
import torch.nn as nn
import torch.nn.parallel
import torch.optim
import torch.utils.data
import torch.utils.data.distributed
import torchvision
import torchvision.models
import torchvision.transforms as transforms
import torch.nn.functional as F
```

**Extend a Pre-built Container**
logger=logging.getLogger(__name__)
logger.setLevel(logging.DEBUG)

classes=('plane', 'car', 'bird', 'cat', 'deer', 'dog', 'frog', 'horse', 'ship', 'truck')

# https://github.com/pytorch/tutorials/blob/master/beginner_source/blitz/cifar10_tutorial.py#L118
class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
        self.conv1=nn.Conv2d(3, 6, 5)
        self.pool=nn.MaxPool2d(2, 2)
        self.conv2=nn.Conv2d(6, 16, 5)
        self.fc1=nn.Linear(16 * 5 * 5, 120)
        self.fc2=nn.Linear(120, 84)
        self.fc3=nn.Linear(84, 10)

    def forward(self, x):
        x=self.pool(F.relu(self.conv1(x)))
        x=self.pool(F.relu(self.conv2(x)))
        x=x.view(-1, 16 * 5 * 5)
        x=F.relu(self.fc1(x))
        x=F.relu(self.fc2(x))
        x=self.fc3(x)
        return x

def _train(args):
    is_distributed=len(args.hosts) > 1 and args.dist_backend is not None
    logger.debug("Distributed training - {}".format(is_distributed))

    if is_distributed:
        # Initialize the distributed environment.
        world_size=len(args.hosts)
        os.environ['WORLD_SIZE']=str(world_size)
        host_rank=args.hosts.index(args.current_host)
        dist.init_process_group(backend=args.dist_backend, rank=host_rank,
        world_size=world_size)
        logger.info("Initialized the distributed environment: \'{}\' backend on {} nodes.
        ".format(}
args.dist_backend,
dist.get_world_size()) + 'Current host rank is {}. Using cuda: {}.
Number of gpus: {}.').format(
    dist.get_rank(), torch.cuda.is_available(), args.num_gpus))

device='cuda' if torch.cuda.is_available() else 'cpu'
logger.info("Device Type: ".format(device))

logger.info("Loading Cifar10 dataset")
transform=transforms.Compose(
    [transforms.ToTensor(),
     transforms.Normalize((0.5, 0.5, 0.5), (0.5, 0.5, 0.5))])

trainset=torchvision.datasets.CIFAR10(root=args.data_dir, train=True,
    download=False, transform=transform)
train_loader=torch.utils.data.DataLoader(trainset, batch_size=args.batch_size,
    shuffle=True,
    num_workers=args.workers)

testset=torchvision.datasets.CIFAR10(root=args.data_dir, train=False,
    download=False, transform=transform)
test_loader=torch.utils.data.DataLoader(testset, batch_size=args.batch_size,
    shuffle=False,
    num_workers=args.workers)

logger.info("Model loaded")
model=Net()

if torch.cuda.device_count() > 1:
    logger.info("Gpu count: ".format(torch.cuda.device_count()))
    model=nn.DataParallel(model)

model=model.to(device)
criterion=nn.CrossEntropyLoss().to(device)
optimizer=torch.optim.SGD(model.parameters(), lr=args.lr,
    momentum=args.momentum)

for epoch in range(0, args.epochs):
    running_loss=0.0
    for i, data in enumerate(train_loader):
        # get the inputs
        inputs, labels=data
        inputs, labels=inputs.to(device), labels.to(device)
# zero the parameter gradients
optimizer.zero_grad()

# forward + backward + optimize
outputs=model(inputs)
loss=criterion(outputs, labels)
loss.backward()
optimizer.step()

# print statistics
running_loss += loss.item()
if i % 2000 == 1999:  # print every 2000 mini-batches
    print('[%d, %5d] loss: %.3f' %
          (epoch + 1, i + 1, running_loss / 2000))
    running_loss=0.0
print('Finished Training')
return _save_model(model, args.model_dir)

def _save_model(model, model_dir):
    logger.info("Saving the model.")
    path=os.path.join(model_dir, 'model.pth')
    # recommended way from http://pytorch.org/docs/master/notes/serialization.html
    torch.save(model.cpu().state_dict(), path)

def model_fn(model_dir):
    logger.info('model_fn')
    device="cuda" if torch.cuda.is_available() else "cpu"
    model=Net()
    if torch.cuda.device_count() > 1:
        logger.info("Gpu count: {}\n".format(torch.cuda.device_count()))
        model=nn.DataParallel(model)

        with open(os.path.join(model_dir, 'model.pth'), 'rb') as f:
            model.load_state_dict(torch.load(f))
        return model.to(device)

if __name__ == '__main__':
    parser=argparse.ArgumentParser()

        parser.add_argument('--workers', type=int, default=2, metavar='W',

Extend a Pre-built Container

5602
help='number of data loading workers (default: 2)'
parser.add_argument('--epochs', type=int, default=2, metavar='E',
    help='number of total epochs to run (default: 2)'
parser.add_argument('--batch-size', type=int, default=4, metavar='BS',
    help='batch size (default: 4)')
parser.add_argument('--lr', type=float, default=0.001, metavar='LR',
    help='initial learning rate (default: 0.001)')
parser.add_argument('--momentum', type=float, default=0.9, metavar='M',
    help='momentum (default: 0.9)')
parser.add_argument('--dist-backend', type=str, default='gloo',
    help='distributed backend (default: gloo)')

# The parameters below retrieve their default values from SageMaker environment
# variables, which are
# instantiated by the SageMaker containers framework.
# https://github.com/aws/sagemaker-containers#how-a-script-is-executed-inside-the-container
    parser.add_argument('--hosts', type=str,
default=ast.literal_eval(os.environ['SM_HOSTS']))
parser.add_argument('--current-host', type=str,
default=os.environ['SM_CURRENT_HOST'])
parser.add_argument('--model-dir', type=str,
default=os.environ['SM_MODEL_DIR'])
parser.add_argument('--data-dir', type=str,
default=os.environ['SM_CHANNEL_TRAINING'])
parser.add_argument('--num-gpus', type=int,
default=os.environ['SM_NUM_GPUS'])
_train(parser.parse_args())

Step 3: Build the Container

1. In the JupyterLab home directory, open a Jupyter notebook. To open a new notebook, choose the New Launch icon and then choose conda_pytorch_p39 in the Notebook section.
2. Run the following command in the first notebook cell to change to the docker_test_folder directory:

    % cd ~/SageMaker/docker_test_folder

This returns your current directory as follows:
output: /home/ec2-user/SageMaker/docker_test_folder

3. Log in to Docker to access the base container:

```bash
! aws ecr get-login-password --region us-east-1 | docker login --username AWS --password-stdin 763104351884.dkr.ecr.us-east-1.amazonaws.com
```

4. To build the Docker container, run the following Docker build command, including the space followed by a period at the end:

```bash
! docker build -t pytorch-extended-container-test .
```

The Docker build command must be run from the Docker directory you created, in this case `docker_test_folder`.

---

**Note**

If you get the following error message that Docker cannot find the Dockerfile, make sure the Dockerfile has the correct name and has been saved to the directory.

```bash
unable to prepare context: unable to evaluate symlinks in Dockerfile path: lstat /home/ec2-user/SageMaker/docker/Dockerfile: no such file or directory
```

Remember that `docker` looks for a file specifically called `Dockerfile` without any extension within the current directory. If you named it something else, you can pass in the file name manually with the `-f` flag. For example, if you named your Dockerfile `Dockerfile-text.txt`, run the following command:

```bash
! docker build -t tf-custom-container-test -f Dockerfile-text.txt .
```

---

**Step 4: Test the Container**

1. To test the container locally in the notebook instance, open a Jupyter notebook. Choose **New Launcher** and choose **Notebook** in **conda_pytorch_p39** framework. The rest of the code snippets must run from the Jupyter notebook instance.
2. Download the CIFAR-10 dataset.

```python
import torch
import torchvision
import torchvision.transforms as transforms

def _get_transform():
    return transforms.Compose([transforms.ToTensor(),
                               transforms.Normalize((0.5, 0.5, 0.5), (0.5, 0.5, 0.5))])

def get_train_data_loader(data_dir='/tmp/pytorch/cifar-10-data'):
    transform=_get_transform()
    trainset=torchvision.datasets.CIFAR10(root=data_dir, train=True,
                                          download=True, transform=transform)
    return torch.utils.data.DataLoader(trainset, batch_size=4,
                                        shuffle=True, num_workers=2)

def get_test_data_loader(data_dir='/tmp/pytorch/cifar-10-data'):
    transform=_get_transform()
    testset=torchvision.datasets.CIFAR10(root=data_dir, train=False,
                                          download=True, transform=transform)
    return torch.utils.data.DataLoader(testset, batch_size=4,
                                         shuffle=False, num_workers=2)

trainloader=get_train_data_loader('/tmp/pytorch-example/cifar-10-data')
testloader=get_test_data_loader('/tmp/pytorch-example/cifar-10-data')
```

3. Set role to the role used to create your Jupyter notebook. This is used to configure your SageMaker Estimator.

```python
from sagemaker import get_execution_role

role=get_execution_role()
```

4. Paste the following example script into the notebook code cell to configure a SageMaker Estimator using your extended container.

```python
from sagemaker.estimator import Estimator

hyperparameters={'epochs': 1}
```
estimator=Estimator(
    image_uri='pytorch-extended-container-test',
    role=role,
    instance_count=1,
    instance_type='local',
    hyperparameters=hyperparameters
)

estimator.fit('file:///tmp/pytorch-example/cifar-10-data')

5. Run the code cell. This test outputs the training environment configuration, the values used for the environmental variables, the source of the data, and the loss and accuracy obtained during training.

**Step 5: Push the Container to Amazon Elastic Container Registry (Amazon ECR)**

1. After you successfully run the local mode test, you can push the Docker container to Amazon ECR and use it to run training jobs.

Run the following command lines in a notebook cell.

```bash
%%sh

# Specify an algorithm name
algorithm_name=pytorch-extended-container-test

account=$(aws sts get-caller-identity --query Account --output text)

# Get the region defined in the current configuration (default to us-west-2 if none defined)
region=$(aws configure get region)

fullname="${account}.dkr.ecr.${region}.amazonaws.com/${algorithm_name}:latest"

# If the repository doesn't exist in ECR, create it.
aws ecr describe-repositories --repository-names "$\{algorithm_name\}" > /dev/null 2>&1
if [ $? -ne 0 ]
then
    aws ecr create-repository --repository-name "$\{algorithm_name\}" > /dev/null
```
2. After you push the container, you can call the Amazon ECR image from anywhere in the SageMaker environment. Run the following code example in the next notebook cell.

If you want to use this training container with SageMaker Studio to use its visualization features, you can also run the following code in a Studio notebook cell to call the Amazon ECR image of your training container.

```python
import boto3

client=boto3.client('sts')
account=client.get_caller_identity()['Account']

my_session=boto3.session.Session()
region=my_session.region_name

algorithm_name="pytorch-extended-container-test"
ecr_image='{}.dkr.ecr.{}.amazonaws.com/{}'.format(account, region, algorithm_name)

ecr_image
# This should return something like
# 12-digits-of-your-account.dkr.ecr.us-east-2.amazonaws.com/tf-2.2-test:latest
```

3. Use the `ecr_image` retrieved from the previous step to configure a SageMaker estimator object. The following code sample configures a SageMaker PyTorch estimator.

```python
import sagemaker
from sagemaker import get_execution_role
```
Step 6: Clean up Resources

To clean up resources when done with the Get Started example

1. Open the SageMaker console, choose the notebook instance RunScriptNotebookInstance, choose Actions, and choose Stop. It can take a few minutes for the instance to stop.

2. After the instance Status changes to Stopped, choose Actions, choose Delete, and then choose Delete in the dialog box. It can take a few minutes for the instance to be deleted. The notebook instance disappears from the table when it has been deleted.

3. Open the Amazon S3 console and delete the bucket that you created for storing model artifacts and the training dataset.

4. Open the IAM console and delete the IAM role. If you created permission policies, you can delete them, too.

⚠️ Note

The Docker container shuts down automatically after it has run. You don't need to delete it.
Adapting your own Docker container to work with SageMaker

You can adapt an existing Docker image to work with SageMaker. You may need to use an existing, external Docker image with SageMaker when you have a container that satisfies feature or safety requirements that are not currently supported by a pre-built SageMaker image. There are two toolkits that allow you to bring your own container and adapt it to work with SageMaker:

- SageMaker Training Toolkit
- SageMaker Inference Toolkit

The following topics show how to adapt your existing image using the SageMaker Training and Inference toolkits:

Topics

- Individual Framework Libraries
- Using the SageMaker Training and Inference Toolkits
- Adapting your own training container
- Adapting Your Own Inference Container

Individual Framework Libraries

In addition to the SageMaker Training Toolkit and SageMaker Inference Toolkit, SageMaker also provides toolkits specialized for TensorFlow, MXNet, PyTorch, and Chainer. The following table provides links to the GitHub repositories that contain the source code for each framework and their respective serving toolkits. The instructions linked are for using the Python SDK to run training algorithms and host models on SageMaker. The functionality for these individual libraries is included in the SageMaker Training Toolkit and SageMaker Inference Toolkit.

<table>
<thead>
<tr>
<th>Framework</th>
<th>Toolkit Source Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>TensorFlow</td>
<td>SageMaker TensorFlow Training</td>
</tr>
<tr>
<td></td>
<td>SageMaker TensorFlow Serving</td>
</tr>
<tr>
<td>MXNet</td>
<td>SageMaker MXNet Training</td>
</tr>
</tbody>
</table>
### Using the SageMaker Training and Inference Toolkits

The [SageMaker Training](#) and [SageMaker Inference](#) toolkits implement the functionality that you need to adapt your containers to run scripts, train algorithms, and deploy models on SageMaker. When installed, the library defines the following for users:

- The locations for storing code and other resources.
- The entry point that contains the code to run when the container is started. Your Dockerfile must copy the code that needs to be run into the location expected by a container that is compatible with SageMaker.
- Other information that a container needs to manage deployments for training and inference.

### SageMaker Toolkits Containers Structure

When SageMaker trains a model, it creates the following file folder structure in the container's `/opt/ml` directory.

```
/opt/ml
### input
#### config
#   #### hyperparameters.json
#   #### resourceConfig.json
#### data
#       #### <channel_name>
#           #### <input data>
### model
#   #### code
#```

When you run a model *training* job, the SageMaker container uses the `/opt/ml/input/` directory, which contains the JSON files that configure the hyperparameters for the algorithm and the network layout used for distributed training. The `/opt/ml/input/` directory also contains files that specify the channels through which SageMaker accesses the data, which is stored in Amazon Simple Storage Service (Amazon S3). The SageMaker containers library places the scripts that the container will run in the `/opt/ml/code/` directory. Your script should write the model generated by your algorithm to the `/opt/ml/model/` directory. For more information, see [Use Your Own Training Algorithms](#).

When you *host* a trained model on SageMaker to make inferences, you deploy the model to an HTTP endpoint. The model makes real-time predictions in response to inference requests. The container must contain a serving stack to process these requests.

In a hosting or batch transform container, the model files are located in the same folder to which they were written during training.

```
/opt/ml/model
#
### <model files>
```

For more information, see [Use your own inference code](#).

**Single Versus Multiple Containers**

You can either provide separate Docker images for the training algorithm and inference code or you can use a single Docker image for both. When creating Docker images for use with SageMaker, consider the following:

- Providing two Docker images can increase storage requirements and cost because common libraries might be duplicated.
- In general, smaller containers start faster for both training and hosting. Models train faster and the hosting service can react to increases in traffic by automatically scaling more quickly.
- You might be able to write an inference container that is significantly smaller than the training container. This is especially common when you use GPUs for training, but your inference code is optimized for CPUs.
• SageMaker requires that Docker containers run without privileged access.

• Both Docker containers that you build and those provided by SageMaker can send messages to the Stdout and Stderr files. SageMaker sends these messages to Amazon CloudWatch logs in your AWS account.

For more information about how to create SageMaker containers and how scripts are executed inside them, see the SageMaker Training Toolkit and SageMaker Inference Toolkit repositories on GitHub. They also provide lists of important environmental variables and the environmental variables provided by SageMaker containers.

Adapting your own training container

To run your own training model, build a Docker container using the Amazon SageMaker Training Toolkit through an Amazon SageMaker notebook instance.

Step 1: Create a SageMaker notebook instance

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.

2. In the left navigation pane, choose Notebook, choose Notebook instances, and then choose Create notebook instance.

3. On the Create notebook instance page, provide the following information:

   a. For Notebook instance name, enter RunScriptNotebookInstance.

   b. For Notebook Instance type, choose ml.t2.medium.

   c. In the Permissions and encryption section, do the following:

      i. For IAM role, choose Create a new role. This opens a new window.

      ii. On the Create an IAM role page, choose Specific S3 buckets, specify an Amazon S3 bucket named sagemaker-run-script, and then choose Create role.

         SageMaker creates an IAM role named AmazonSageMaker-ExecutionRole-YYYYMMDDTHHmmSS. For example, AmazonSageMaker-ExecutionRole-20190429T110788. Note that the execution role naming convention uses the date and time at which the role was created, separated by a T.

      d. For Root Access, choose Enable.

      e. Choose Create notebook instance.
4. On the **Notebook instances** page, the **Status** is **Pending**. It can take a few minutes for Amazon SageMaker to launch a machine learning compute instance—in this case, it launches a notebook instance—and attach an ML storage volume to it. The notebook instance has a preconfigured Jupyter notebook server and a set of Anaconda libraries. For more information, see [CreateNotebookInstance](#).

5. Click on the **Name** of the notebook you just created. This opens a new page.

6. In the **Permissions and encryption** section, copy the **IAM role ARN number**, and paste it into a notepad file to save it temporarily. You use this IAM role ARN number later to configure a local training estimator in the notebook instance. The IAM role ARN number looks like the following: 'arn:aws:iam::111122223333:role/service-role/AmazonSageMaker-ExecutionRole-20190429T110788'

7. After the status of the notebook instance changes to **InService**, choose **Open JupyterLab**.

**Step 2: Create and upload the Dockerfile and Python training scripts**

1. After JupyterLab opens, create a new folder in the home directory of your JupyterLab. In the upper-left corner, choose the **New Folder** icon, and then enter the folder name `docker_test_folder`.

2. Create a Dockerfile text file in the `docker_test_folder` directory.
   a. Choose the **New Launcher** icon (+) in the upper-left corner.
   b. In the right pane under the **Other** section, choose **Text File**.
   c. Paste the following Dockerfile sample code into your text file.

   ```
   # Download an open source TensorFlow Docker image
   FROM tensorflow/tensorflow:latest-gpu-jupyter

   # Install sagemaker-training toolkit that contains the common functionality necessary to create a container compatible with SageMaker and the Python SDK.
   RUN pip3 install sagemaker-training

   # Copies the training code inside the container
   COPY train.py /opt/ml/code/train.py

   # Defines train.py as script entrypoint
   ```

   **Adapting your own training container**
ENV SAGEMAKER_PROGRAM train.py

The Dockerfile script performs the following tasks:

- **FROM tensorflow/tensorflow:latest-gpu-jupyter** – Downloads the latest TensorFlow Docker base image. You can replace this with any Docker base image you want to bring to build containers, as well as with AWS pre-built container base images.

- **RUN pip install sagemaker-training** – Installs SageMaker Training Toolkit that contains the common functionality necessary to create a container compatible with SageMaker.

- **COPY train.py /opt/ml/code/train.py** – Copies the script to the location inside the container that is expected by SageMaker. The script must be located in this folder.

- **ENV SAGEMAKER_PROGRAM train.py** – Takes your training script train.py as the entrypoint script copied in the /opt/ml/code folder of the container. This is the only environmental variable that you must specify when you build your own container.

d. On the left directory navigation pane, the text file name might automatically be named untitled.txt. To rename the file, right-click the file, choose **Rename**, rename the file as Dockerfile without the .txt extension, and then press Ctrl+s or Command+s to save the file.

3. Upload a training script train.py to the docker_test_folder. You can use the following example script to create a model that reads handwritten digits trained on the MNIST dataset for this exercise.

```python
import tensorflow as tf
import os

mnist = tf.keras.datasets.mnist

(x_train, y_train), (x_test, y_test) = mnist.load_data()

x_train, x_test = x_train / 255.0, x_test / 255.0

model = tf.keras.models.Sequential([tf.keras.layers.Flatten(input_shape=(28, 28)),
                                    tf.keras.layers.Dense(128, activation='relu'),
                                    tf.keras.layers.Dropout(0.2),
                                    tf.keras.layers.Dense(10, activation='softmax')])
```
model.compile(optimizer='adam',
loss='sparse_categorical_crossentropy',
metrics=['accuracy'])

model.fit(x_train, y_train, epochs=1)
model_save_dir = f'{os.environ.get('SM_MODEL_DIR')}/1'

model.evaluate(x_test, y_test)
tf.saved_model.save(model, model_save_dir)

Step 3: Build the container

1. In the JupyterLab home directory, open a Jupyter notebook. To open a new notebook, choose the New Launch icon and then choose the latest version of conda_tensorflow2 in the Notebook section.

2. Run the following command in the first notebook cell to change to the docker_test_folder directory:

   ```
   cd ~/SageMaker/docker_test_folder
   ```

   This returns your current directory as follows:

   ```
   ! pwd
   ```

   output: /home/ec2-user/SageMaker/docker_test_folder

3. To build the Docker container, run the following Docker build command, including the space followed by a period at the end:

   ```
   ! docker build -t tf-custom-container-test .
   ```

   The Docker build command must be run from the Docker directory you created, in this case docker_test_folder.

   **Note**

   If you get the following error message that Docker cannot find the Dockerfile, make sure the Dockerfile has the correct name and has been saved to the directory.
Remember that docker looks for a file specifically called Dockerfile without any extension within the current directory. If you named it something else, you can pass in the file name manually with the `-f` flag. For example, if you named your Dockerfile as Dockerfile-text.txt, run the following command:

```
! docker build -t tf-custom-container-test -f Dockerfile-text.txt .
```

**Step 4: Test the container**

1. To test the container locally in the notebook instance, open a Jupyter notebook. Choose New Launcher and choose the latest version of conda_tensorflow2 in the Notebook section.
2. Paste the following example script into the notebook code cell to configure a SageMaker Estimator.

   ```python
   import sagemaker
   from sagemaker.estimator import Estimator

   estimator = Estimator(image_uri='tf-custom-container-test',
                          role=sagemaker.get_execution_role(),
                          instance_count=1,
                          instance_type='local')

   estimator.fit()
   ```

   In the preceding code example, sagemaker.get_execution_role() is specified to the role argument to automatically retrieve the role set up for the SageMaker session. You can also replace it with the string value of the IAM role ARN number you used when you configured the notebook instance. The ARN should look like the following: 'arn:aws:iam::111122223333:role/service-role/AmazonSageMaker-ExecutionRole-20190429T110788'.
3. Run the code cell. This test outputs the training environment configuration, the values used for the environmental variables, the source of the data, and the loss and accuracy obtained during training.

**Step 5: Push the container to Amazon Elastic Container Registry (Amazon ECR)**

1. After you successfully run the local mode test, you can push the Docker container to Amazon ECR and use it to run training jobs. If you want to use a private Docker registry instead of Amazon ECR, see [Push your training container to a private registry](#).

Run the following command lines in a notebook cell.

```sh
# Specify an algorithm name
algorithm_name=tf-custom-container-test
account=$(aws sts get-caller-identity --query Account --output text)

# Get the region defined in the current configuration (default to us-west-2 if none defined)
region=$(aws configure get region)
region=${region:-us-west-2}
fullname="${account}.dkr.ecr.${region}.amazonaws.com/${algorithm_name}:latest"

# If the repository doesn't exist in ECR, create it.
aws ecr describe-repositories --repository-names "${algorithm_name}" > /dev/null 2>&1
if [ $? -ne 0 ]
then
    aws ecr create-repository --repository-name "${algorithm_name}" > /dev/null
fi

# Get the login command from ECR and execute it directly
aws ecr get-login-password --region ${region}|docker login --username AWS --password-stdin ${fullname}

# Build the docker image locally with the image name and then push it to ECR
# with the full name.
```
**Note**

This bash shell script may raise a permission issue similar to the following error message:

```
"denied: User: [ARN] is not authorized to perform: ecr:InitiateLayerUpload
  on resource:
  arn:aws:ecr:us-east-1:[id]:repository/tf-custom-container-test"
```

If this error occurs, you need to attach the `AmazonEC2ContainerRegistryFullAccess` policy to your IAM role. Go to the IAM console, choose Roles from the left navigation pane, look up the IAM role you used for the Notebook instance. Under the Permission tab, choose the Attach policies button, and search the `AmazonEC2ContainerRegistryFullAccess` policy. Mark the check box of the policy, and choose Add permissions to finish.

2. Run the following code in a Studio notebook cell to call the Amazon ECR image of your training container.

```python
import boto3

account_id = boto3.client('sts').get_caller_identity().get('Account')
ecr_repository = 'tf-custom-container-test'
tag = ':latest'

region = boto3.session.Session().region_name

uri_suffix = 'amazonaws.com'
if region in ['cn-north-1', 'cn-northwest-1']:
    uri_suffix = 'amazonaws.com.cn'

byoc_image_uri = '{}.dkr.ecr.{}.{}{}'.format(account_id, region, uri_suffix, ecr_repository + tag)
```
3. Use the `ecr_image` retrieved from the previous step to configure a SageMaker estimator object. The following code sample configures a SageMaker estimator with the `byoc_image_uri` and initiates a training job on an Amazon EC2 instance.

**SageMaker Python SDK v1**

```python
import sagemaker
from sagemaker import get_execution_role
from sagemaker.estimator import Estimator

estimator = Estimator(image_uri=byoc_image_uri,
                      role=get_execution_role(),
                      base_job_name='tf-custom-container-test-job',
                      instance_count=1,
                      instance_type='ml.g4dn.xlarge')

#train your model
estimator.fit()
```

**SageMaker Python SDK v2**

```python
import sagemaker
from sagemaker import get_execution_role
from sagemaker.estimator import Estimator

estimator = Estimator(image_uri=byoc_image_uri,
                      role=get_execution_role(),
                      base_job_name='tf-custom-container-test-job',
                      instance_count=1,
                      instance_type='ml.g4dn.xlarge')

#train your model
estimator.fit()
```

4. If you want to deploy your model using your own container, refer to [Adapting Your Own Inference Container](#). You can also use an AWSframework container that can deploy a TensorFlow model. To deploy the example model to read handwritten digits, enter the
following example script into the same notebook that you used to train your model in
the previous sub-step to obtain the image URIs (universal resource identifiers) needed for
deployment, and deploy the model.

```python
import boto3
import sagemaker

#obtain image uris
from sagemaker import image_uris
container = image_uris.retrieve(framework='tensorflow', region='us-west-2', version='2.11.0',
                                 image_scope='inference', instance_type='ml.g4dn.xlarge')

#create the model entity, endpoint configuration and endpoint
predictor = estimator.deploy(1, instance_type='ml.g4dn.xlarge', image_uri=container)
```

Test your model using a sample handwritten digit from the MNIST dataset using the following
code example.

```python
#Retrieve an example test dataset to test
import numpy as np
import matplotlib.pyplot as plt
from keras.datasets import mnist

# Load the MNIST dataset and split it into training and testing sets
(x_train, y_train), (x_test, y_test) = mnist.load_data()

# Select a random example from the training set
example_index = np.random.randint(0, x_train.shape[0])
example_image = x_train[example_index]
example_label = y_train[example_index]

# Print the label and show the image
print(f"Label: {example_label}")
plt.imshow(example_image, cmap='gray')
plt.show()
```

Convert the test handwritten digit into a form that TensorFlow can ingest and make a test
prediction.

```python
from sagemaker.serializers import JSONSerializer
data = {'instances': example_image.tolast()}
```
predictor.serializer=JSONSerializer()  # update the predictor to use the JSONSerializer
predictor.predict(data)  # make the prediction

For a full example that shows how to test a custom container locally and push it to an Amazon ECR image, see the Building Your Own TensorFlow Container example notebook.

Tip
To profile and debug training jobs to monitor system utilization issues (such as CPU bottlenecks and GPU underutilization) and identify training issues (such as overfitting, overtraining, exploding tensors, and vanishing gradients), use Amazon SageMaker Debugger. For more information, see Use Debugger with Custom Training Containers.

Step 6: Clean up resources

To clean up resources when done with the get started example

1. Open the SageMaker console, choose the notebook instance RunScriptNotebookInstance, choose Actions, and choose Stop. It can take a few minutes for the instance to stop.

2. After the instance Status changes to Stopped, choose Actions, choose Delete, and then choose Delete in the dialog box. It can take a few minutes for the instance to be deleted. The notebook instance disappears from the table when it has been deleted.

3. Open the Amazon S3 console and delete the bucket that you created for storing model artifacts and the training dataset.

4. Open the IAM console and delete the IAM role. If you created permission policies, you can delete them, too.

Note
The Docker container shuts down automatically after it has run. You don't need to delete it.
Blogs and Case Studies

The following blogs discuss case studies about using custom training containers in Amazon SageMaker.

- Why bring your own container to Amazon SageMaker and how to do it right, Medium (January 20th, 2023)

Adapt your training job to access images in a private Docker registry

You can use a private Docker registry instead of an Amazon Elastic Container Registry (Amazon ECR) to host your images for SageMaker Training. The following instructions show you how to create a Docker registry, configure your virtual private cloud (VPC) and training job, store images, and give SageMaker access to the training image in the private docker registry. These instructions also show you how to use a Docker registry that requires authentication for a SageMaker training job.

Create and store your images in a private Docker registry

Create a private Docker registry to store your images. Your registry must:

- use the Docker Registry HTTP API protocol
- be accessible from the same VPC specified in the VpcConfig parameter in the CreateTrainingJob API. Input VpcConfig when you create your training job.
- secured with a TLS certificate from a known public certificate authority.

For more information about creating a Docker registry, see Deploy a registry server.

Configure your VPC and SageMaker training job

SageMaker uses a network connection within your VPC to access images in your Docker registry. To use the images in your Docker registry for training, the registry must be accessible from an Amazon VPC in your account. For more information, see Use a Docker registry that requires authentication for training.

You must also configure your training job to connect to the same VPC to which your Docker registry has access. For more information, see Configure a Training Job for Amazon VPC Access.
Create a training job using an image from your private Docker registry

To use an image from your private Docker registry for training, use the following guide to configure your image, configure and create a training job. The code examples that follow use the AWS SDK for Python (Boto3) client.

1. Create a training image configuration object and input Vpc the TrainingRepositoryAccessMode field as follows.

```python
training_image_config = {
    'TrainingRepositoryAccessMode': 'Vpc'
}
```

**Note**

If your private Docker registry requires authentication, you must add a TrainingRepositoryAuthConfig object to the training image configuration object. You must also specify the Amazon Resource Name (ARN) of an AWS Lambda function that provides access credentials to SageMaker using the TrainingRepositoryCredentialsProviderArn field of the TrainingRepositoryAuthConfig object. For more information, see the example code structure below.

```python
training_image_config = {
    'TrainingRepositoryAccessMode': 'Vpc',
    'TrainingRepositoryAuthConfig': {
        'TrainingRepositoryCredentialsProviderArn':
            'arn:aws:lambda:Region:Acct:function:FunctionName'
    }
}
```

For information about how to create the Lambda function to provide authentication, see Use a Docker registry that requires authentication for training.

2. Use a Boto3 client to create a training job and pass the correct configuration to the create_training_job API. The following instructions show you how to configure the components and create a training job.
a. Create the AlgorithmSpecification object that you want to pass to
create_training_job. Use the training image configuration object that you created in the
previous step, as shown in the following code example.

```python
algorithm_specification = {
    'TrainingImage': 'myteam.myorg.com/docker-local/my-training-image:<IMAGE-TAG>',
    'TrainingImageConfig': training_image_config,
    'TrainingInputMode': 'File'
}
```

**Note**
To use a fixed, rather than an updated version of an image, refer to the image’s digest
instead of by name or tag.

b. Specify the name of the training job and role that you want to pass to
create_training_job, as shown in the following code example.

```python
training_job_name = 'private-registry-job'
execution_role_arn = 'arn:aws:iam::123456789012:role/SageMakerExecutionRole'
```

c. Specify a security group and subnet for the VPC configuration for your training job. Your
private Docker registry must allow inbound traffic from the security groups that you specify,
as shown in the following code example.

```python
vpc_config = {
    'SecurityGroupIds': ['sg-0123456789abcdef0'],
    'Subnets': ['subnet-0123456789abcdef0', 'subnet-0123456789abcdef1']
}
```

**Note**
If your subnet is not in the same VPC as your private Docker registry, you must set up
a networking connection between the two VPCs. SeeConnect VPCs using VPC peering
for more information.

d. Specify the resource configuration, including machine learning compute instances and
storage volumes to use for training, as shown in the following code example.
resource_config = {
    'InstanceType': 'ml.m4.xlarge',
    'InstanceCount': 1,
    'VolumeSizeInGB': 10,
}

e. Specify the input and output data configuration, where the training dataset is stored, and where you want to store model artifacts, as shown in the following code example.

input_data_config = [
    {
        "ChannelName": "training",
        "DataSource": {
            "S3DataSource": {
                "S3DataDistributionType": "FullyReplicated",
                "S3DataType": "S3Prefix",
                "S3Uri": "s3://your-training-data-bucket/training-data-folder"
            }
        }
    }
]

output_data_config = {
    'S3OutputPath': 's3://your-output-data-bucket/model-folder'
}

f. Specify the maximum number of seconds that a model training job can run as shown in the following code example.

stopping_condition = {
    'MaxRuntimeInSeconds': 1800
}

g. Finally, create the training job using the parameters you specified in the previous steps as shown in the following code example.

import boto3
sm = boto3.client('sagemaker')
try:
Use a SageMaker estimator to run a training job

You can also use an estimator from the SageMaker Python SDK to handle the configuration and running of your SageMaker training job. The following code examples show how to configure and run an estimator using images from a private Docker registry.

1. Import the required libraries and dependencies, as shown in the following code example.

```python
import boto3
import sagemaker
from sagemaker.estimator import Estimator

session = sagemaker.Session()
role = sagemaker.get_execution_role()
```

2. Provide a Uniform Resource Identifier (URI) to your training image, security groups and subnets for the VPC configuration for your training job, as shown in the following code example.

```python
image_uri = "myteam.myorg.com/docker-local/my-training-image:<IMAGE-TAG>"
security_groups = ["sg-0123456789abcdef0"]
subnets = ["subnet-0123456789abcdef0", "subnet-0123456789abcdef0"]
```

For more information about security_group_ids and subnets, see the appropriate parameter description in the Estimators section of the SageMaker Python SDK.
Note

SageMaker uses a network connection within your VPC to access images in your Docker registry. To use the images in your Docker registry for training, the registry must be accessible from an Amazon VPC in your account.

3. Optionally, if your Docker registry requires authentication, you must also specify the Amazon Resource Name (ARN) of an AWS Lambda function that provides access credentials to SageMaker. The following code example shows how to specify the ARN.

```python
```

For more information about using images in a Docker registry requiring authentication, see Use a Docker registry that requires authentication for training below.

4. Use the code examples from the previous steps to configure an estimator, as shown in the following code example.

```python
# The training repository access mode must be 'Vpc' for private docker registry jobs
training_repository_access_mode = "Vpc"

# Specify the instance type, instance count you want to use
instance_type="ml.m5.xlarge"
instance_count=1

# Specify the maximum number of seconds that a model training job can run
max_run_time = 1800

# Specify the output path for the model artifacts
output_path = "s3://your-output-bucket/your-output-path"

estimator = Estimator(
    image_uri=image_uri,
    role=role,
    subnets=subnets,
    security_group_ids=security_groups,
    training_repository_access_mode=training_repository_access_mode,
    training_repository_credentials_provider_arn=training_repository_credentials_provider_arn,
    # remove this line if auth is not needed
```
5. Start your training job by calling `estimator.fit` with your job name and input path as parameters, as shown in the following code example.

```python
input_path = "s3://your-input-bucket/your-input-path"
job_name = "your-job-name"

estimator.fit(
    inputs=input_path,
    job_name=job_name
)
```

### Use a Docker registry that requires authentication for training

If your Docker registry requires authentication, you must create an AWS Lambda function that provides access credentials to SageMaker. Then, create a training job and provide the ARN of this Lambda function inside the `create_training_job` API. Lastly, you can optionally create an interface VPC endpoint so that your VPC can communicate with your Lambda function without sending traffic over the internet. The following guide shows how to create a Lambda function, assign it the correct role and create an interface VPC endpoint.

#### Create the Lambda function

Create an AWS Lambda function that passes access credentials to SageMaker and returns a response. The following code example creates the Lambda function handler, as follows.

```python
def handler(event, context):
    response = {
        "Credentials": {"Username": "username", "Password": "password"}
    }
    return response
```

The type of authentication used to set up your private Docker registry determines the contents of the response returned by your Lambda function as follows.
• If your private Docker registry uses basic authentication, the Lambda function will return the username and password needed in order to authenticate to the registry.

• If your private Docker registry uses bearer token authentication, the username and password are sent to your authorization server, which then returns a bearer token. This token is then used to authenticate to your private Docker registry.

**Note**
If you have more than one Lambda functions for your registries in the same account, and the execution role is the same for your training jobs, then training jobs for registry one would have access to the Lambda functions for other registries.

**Grant the correct role permission to your Lambda function**

The IAMrole that you use in the create_training_job API must have permission to call an AWS Lambda function. The following code example shows how to extend permissions policy of an IAM role to call myLambdaFunction.

```json
{
    "Effect": "Allow",
    "Action": [
        "lambda:InvokeFunction"
    ],
    "Resource": [
        "arn:aws:lambda:*::*:function:*myLambdaFunction*"
    ]
}
```

For information about editing a role permissions policy, see [Modifying a role permissions policy (console)](console) in the AWS Identity and Access Management User Guide.

**Note**
An IAM role with an attached AmazonSageMakerFullAccess managed policy has permission to call any Lambda function with "SageMaker" in its name.
Create an interface VPC endpoint for Lambda

If you create an interface endpoint, your Amazon VPC can communicate with your Lambda function without sending traffic over the internet. For more information, see Configuring interface VPC endpoints for Lambda in the AWS Lambda Developer Guide.

After your interface endpoint is created, SageMaker training will call your Lambda function by sending a request through your VPC to lambda.region.amazonaws.com. If you select Enable DNS Name when you create your interface endpoint, Amazon Route 53 routes the call to the Lambda interface endpoint. If you use a different DNS provider, you must map lambda.region.amazonaws.com to your Lambda interface endpoint.

Adapting Your Own Inference Container

If you can't use any of the images listed in Use Pre-built SageMaker Docker images Amazon SageMaker for your use case, you can build your own Docker container and use it inside SageMaker for training and inference. To be compatible with SageMaker, your container must have the following characteristics:

- Your container must have a web server listing on port 8080.
- Your container must accept POST requests to the /invocations and /ping real-time endpoints. The requests that you send to these endpoints must be returned with 60 seconds and have a maximum size of 6 MB.

For more information and an example of how to build your own Docker container for training and inference with SageMaker, see Building your own algorithm container.

The following guide shows you how to use a JupyterLab space with Amazon SageMaker Studio Classic to adapt an inference container to work with SageMaker hosting. The example uses an NGINX web server, Gunicorn as a Python web server gateway interface, and Flask as a web application framework. You can use different applications to adapt your container as long as it meets the previous listed requirements. For more information about using your own inference code, see Use Your Own Inference Code with Hosting Services.

Adapt your inference container

Use the following steps to adapt your own inference container to work with SageMaker hosting. The example shown in the following steps uses a pre-trained Named Entity Recognition (NER) model that uses the spaCy natural language processing (NLP) library for Python and the following:
• A Dockerfile to build the container that contains the NER model.
• Inference scripts to serve the NER model.

If you adapt this example for your use case, you must use a Dockerfile and inference scripts that are needed to deploy and serve your model.

1. Create JupyterLab space with Amazon SageMaker Studio Classic (optional).

   You can use any notebook to run scripts to adapt your inference container with SageMaker hosting. This example shows you how to use a JupyterLab space within Amazon SageMaker Studio Classic to launch a JupyterLab application that comes with a SageMaker Distribution image. For more information, see SageMaker JupyterLab.

2. Upload a Docker file and inference scripts.

   1. Create a new folder in your home directory. If you’re using JupyterLab, in the upper-left corner, choose the **New Folder** icon, and enter a folder name to contain your Dockerfile. In this example, the folder is called **docker_test_folder**.

   2. Upload a Dockerfile text file into your new folder. The following is an example Dockerfile that creates a Docker container with a pre-trained Named Entity Recognition (NER) model from *spaCy*, the applications and environment variables needed to run the example:

```python
FROM python:3.8

RUN apt-get -y update && apt-get install -y --no-install-recommends \
    wget \
    python3 \
    nginx \
    ca-certificates \
    && rm -rf /var/lib/apt/lists/*

RUN wget https://bootstrap.pypa.io/get-pip.py && python3 get-pip.py && \
    pip install flask gevent gunicorn && \
    rm -rf /root/.cache

#pre-trained model package installation
RUN pip install spacy
RUN python -m spacy download en

# Set environment variables
```

Adapting Your Own Inference Container
In the previous code example, the environment variable `PYTHONUNBUFFERED` keeps Python from buffering the standard output stream, which allows for faster delivery of logs to the user. The environment variable `PYTHONDONTWRITEBYTECODE` keeps Python from writing compiled bytecode `.pyc` files, which are unnecessary for this use case. The environment variable `PATH` is used to identify the location of the `train` and `serve` programs when the container is invoked.

3. Create a new directory inside your new folder to contain scripts to serve your model. This example uses a directory called `NER`, which contains the following scripts necessary to run this example:
   - `predictor.py` – A Python script that contains the logic to load and perform inference with your model.
   - `nginx.conf` – A script to configure a web server.
   - `serve` – A script that starts an inference server.
   - `wsgi.py` – A helper script to serve a model.

⚠️ **Important**

If you copy your inference scripts into a notebook ending in `.ipynb` and rename them, your script may contain formatting characters that will prevent your endpoint from deploying. Instead, create a text file and rename them.

4. Upload a script to make your model available for inference. The following is an example script called `predictor.py` that uses Flask to provide the `/ping` and `/invocations` endpoints:

```python
from flask import Flask
import flask
import spacy
import os
import json
import logging
```
#Load in model
nlp = spacy.load('en_core_web_sm')
#If you plan to use a your own model artifacts,
#your model artifacts should be stored in /opt/ml/model/

# The flask app for serving predictions
app = Flask(__name__)
@app.route('/ping', methods=['GET'])
def ping():
    # Check if the classifier was loaded correctly
    health = nlp is not None
    status = 200 if health else 404
    return flask.Response(response='\n', status=status, mimetype='application/json')

@app.route('/invocations', methods=['POST'])
def transformation():
    #Process input
    input_json = flask.request.get_json()
    resp = input_json['input']

    #NER
    doc = nlp(resp)
    entities = [(X.text, X.label_) for X in doc.ents]

    # Transform predictions to JSON
    result = {
        'output': entities
    }

    resultjson = json.dumps(result)
    return flask.Response(response=resultjson, status=200, mimetype='application/json')

The /ping endpoint in the previous script example returns a status code of 200 if the model is loaded correctly, and 404 if the model is loaded incorrectly. The /invocations endpoint processes a request formatted in JSON, extracts the input field, and uses the NER model to identify and store entities in the variable entities. The Flask application returns
the response that contains these entities. For more information about these required health requests, see [How Your Container Should Respond to Health Check (Ping) Requests](#).

5. Upload a script to start an inference server. The following script example calls `serve` using Gunicorn as an application server, and Nginx as a web server:

```python
#!/usr/bin/env python

# This file implements the scoring service shell. You don't necessarily need to modify it for various
# algorithms. It starts nginx and gunicorn with the correct configurations and then simply waits until
# gunicorn exits.
#
# The flask server is specified to be the app object in wsgi.py
#
# We set the following parameters:
#
# Parameter                Environment Variable              Default Value
# ---------                --------------------              -------------
# number of workers        MODEL_SERVER_WORKERS              the number of CPU cores
# timeout                  MODEL_SERVER_TIMEOUT              60 seconds

import multiprocessing
import os
import signal
import subprocess
import sys

cpu_count = multiprocessing.cpu_count()

model_server_timeout = os.environ.get('MODEL_SERVER_TIMEOUT', 60)
model_server_workers = int(os.environ.get('MODEL_SERVER_WORKERS', cpu_count))

def sigterm_handler(nginx_pid, gunicorn_pid):
    try:
        os.kill(nginx_pid, signal.SIGQUIT)
    except OSError:
        pass
    try:
        os.kill(gunicorn_pid, signal.SIGTERM)
    except OSError:
        pass
```
sys.exit(0)

def start_server():
    print('Starting the inference server with {} workers.'.format(model_server_workers))

    # link the log streams to stdout/err so they will be logged to the container logs
    subprocess.check_call(['ln', '-sf', '/dev/stdout', '/var/log/nginx/access.log'])
    subprocess.check_call(['ln', '-sf', '/dev/stderr', '/var/log/nginx/error.log'])

    nginx = subprocess.Popen(['nginx', '-c', '/opt/program/nginx.conf'])
    gunicorn = subprocess.Popen(['gunicorn',
                                 '--timeout', str(model_server_timeout),
                                 '-k', 'sync',
                                 '-b', 'unix:/tmp/gunicorn.sock',
                                 '-w', str(model_server_workers),
                                 'wsgi:app'])

    signal.signal(signal.SIGTERM, lambda a, b: sigterm_handler(nginx.pid, gunicorn.pid))

    # Exit the inference server upon exit of either subprocess
    pids = set([nginx.pid, gunicorn.pid])
    while True:
        pid, _ = os.wait()
        if pid in pids:
            break

    sigterm_handler(nginx.pid, gunicorn.pid)
    print('Inference server exiting')

# The main routine to invoke the start function.

if __name__ == '__main__':
    start_server()

The previous script example defines a signal handler function `sigterm_handler`, which shuts down the Nginx and Gunicorn sub-processes when it receives a SIGTERM signal.
A `start_server` function starts the signal handler, starts and monitors the Nginx and Gunicorn sub-processes, and captures log streams.

6. Upload a script to configure your web server. The following script example called `nginx.conf`, configures a Nginx web server using Gunicorn as an application server to serve your model for inference:

```plaintext
worker_processes 1;
daemon off; # Prevent forking

pid /tmp/nginx.pid;
error_log /var/log/nginx/error.log;

events {
    # defaults
}

http {
    include /etc/nginx/mime.types;
default_type application/octet-stream;
access_log /var/log/nginx/access.log combined;

    upstream gunicorn {
        server unix:/tmp/gunicorn.sock;
    }

    server {
        listen 8080 deferred;
        client_max_body_size 5m;

        keepalive_timeout 5;
        proxy_read_timeout 1200s;

        location ~ ^/(ping|invocations) {
            proxy_set_header X-Forwarded-For $proxy_add_x_forwarded_for;
            proxy_set_header Host $http_host;
            proxy_redirect off;
            proxy_pass http://gunicorn;
        }

        location / {
            return 404 "{}";
        }
```
The previous script example configures Nginx to run in the foreground, sets the location to capture the `error_log`, and defines `upstream` as the Gunicorn server's socket sock. The server configures the server block to listen on port 8080, sets limits on client request body size and timeout values. The server block, forwards requests containing either `/ping` or `/invocations` paths to the Gunicorn server `http://gunicorn`, and returns a 404 error for other paths.

7. Upload any other scripts needed to serve your model. This example needs the following example script called `wsgi.py` to help Gunicorn find your application:

```python
import predictor as myapp

# This is just a simple wrapper for gunicorn to find your app.
# If you want to change the algorithm file, simply change "predictor" above to the
# new file.
app = myapp.app
```

From the folder `docker_test_folder`, your directory structure should contain a Dockerfile and the folder NER. The NER folder should contain the files `nginx.conf`, `predictor.py`, `serve`, and `wsgi.py` as follows:

```
/docker_test_folder
|---Dockerfile
|---NER
 | |---nginx.conf
 | |---predictor.py
 | |---serve
 | |---wsgi.py
```

3. Build your own container.
From the folder `docker_test_folder`, build your Docker container. The following example command will build the Docker container that is configured in your Dockerfile:

```bash
docker build -t byo-container-test .
```

The previous command will build a container called `byo-container-test` in the current working directory. For more information about the Docker build parameters, see Build arguments.

**Note**

If you get the following error message that Docker cannot find the Dockerfile, make sure the Dockerfile has the correct name and has been saved to the directory.

```bash
unable to prepare context: unable to evaluate symlinks in Dockerfile path: lstat /home/ec2-user/SageMaker/docker_test_folder/Dockerfile: no such file or directory
```

Docker looks for a file specifically called Dockerfile without any extension within the current directory. If you named it something else, you can pass in the file name manually with the `-f` flag. For example, if you named your Dockerfile as `Dockerfile-text.txt`, build your Docker container using the `-f` flag followed by your file as follows:

```bash
docker build -t byo-container-test -f Dockerfile-text.txt .
```

4. **Push your Docker Image to an Amazon Elastic Container Registry (Amazon ECR)**

In a notebook cell, push your Docker image to an ECR. The following code example shows you how to build your container locally, login and push it to an ECR:

```bash
# Name of algo -> ECR
algorithm_name=sm-pretrained-spacy

#make serve executable
chmod +x NER/serve
account=$(aws sts get-caller-identity --query Account --output text)
# Region, defaults to us-west-2
```
region=$(aws configure get region)
region=${region:-us-east-1}
fullname="${account}.dkr.ecr.${region}.amazonaws.com/${algorithm_name}:latest"

# If the repository doesn't exist in ECR, create it.
aws ecr describe-repositories --repository-names "${algorithm_name}" > /dev/null 2>&1
if [ $? -ne 0 ]
then
  aws ecr create-repository --repository-name "${algorithm_name}" > /dev/null
fi

# Get the login command from ECR and execute it directly
aws ecr get-login-password --region ${region} | docker login --username AWS --password-stdin ${fullname}

# Build the docker image locally with the image name and then push it to ECR
# with the full name.

docker build -t ${algorithm_name} .
docker tag ${algorithm_name} ${fullname}
docker push ${fullname}

In the previous example shows how to do the following steps necessary to push the example Docker container to an ECR:

a. Define the algorithm name as sm-pretrained-spacy.
b. Make the serve file inside the NER folder executable.
c. Set the AWS Region.
d. Create an ECR if it doesn’t already exist.
e. Login to the ECR.
f. Build the Docker container locally.
g. Push the Docker image to the ECR.

5. Set up the SageMaker client

If you want to use SageMaker hosting services for inference, you must create a model, create an endpoint config and create an endpoint. In order to get inferences from your endpoint, you can use the SageMaker boto3 Runtime client to invoke your endpoint. The following code shows you how to set up both the SageMaker client and the SageMaker Runtime client using the SageMaker boto3 client:

```python
import boto3
```
from sagemaker import get_execution_role

sm_client = boto3.client(service_name='sagemaker')
runtime_sm_client = boto3.client(service_name='sagemaker-runtime')

account_id = boto3.client('sts').get_caller_identity()['Account']
region = boto3.Session().region_name

#used to store model artifacts which SageMaker will extract to /opt/ml/model in the container,
#in this example case we will not be making use of S3 to store the model artifacts
#s3_bucket = '<S3Bucket>'

role = get_execution_role()

In the previous code example, the Amazon S3 bucket is not used, but inserted as a comment to show how to store model artifacts.

If you receive a permission error after you run the previous code example, you may need to add permissions to your IAM role. For more information about IAM roles, see Amazon SageMaker Role Manager. For more information about adding permissions to your current role, see AWS Managed Policies for Amazon SageMaker.

6. Create your model.

If you want to use SageMaker hosting services for inference, you must create a model in SageMaker. The following code example shows you how to create the spaCy NER model inside of SageMaker:

from time import gmtime, strftime

model_name = 'spacy-nermodel-' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
# MODEL S3 URL containing model artifacts as either model.tar.gz or extracted artifacts.
# Here we are not
#model_url = 's3://{}/spacy/'.format(s3_bucket)

container = '{}.dkr.ecr.{}.amazonaws.com/sm-pretrained-spacy:latest'.format(account_id, region)
instance_type = 'ml.c5d.18xlarge'

print('Model name: ' + model_name)
```python
# print('Model data Url: ' + model_url)
print('Container image: ' + container)

container = {
    'Image': container
}

create_model_response = sm_client.create_model(
    ModelName = model_name,
    ExecutionRoleArn = role,
    Containers = [container])

print("Model Arn: " + create_model_response['ModelArn'])
```

The previous code example shows how to define a `model_url` using the `s3_bucket` if you were to use the Amazon S3 bucket from the comments in Step 5, and defines the ECR URI for the container image. The previous code examples defines `ml.c5d.18xlarge` as the instance type. You can also choose a different instance type. For more information about available instance types, see Amazon EC2 instance types.

In the previous code example, The `Image` key points to the container image URI. The `create_model_response` definition uses the `create_model` method to create a model, and return the model name, role and a list containing the container information.

Example output from the previous script follows:

```
Model name: spacy-nermodel-YYYY-MM-DD-HH-MM-SS
Model data Url: s3://spacy-sagemaker-us-east-1-bucket/spacy/
Container image: 123456789012.dkr.ecr.us-east-2.amazonaws.com/sm-pretrained-spacy:latest
```

7. a. **Configure and create an endpoint**

To use SageMaker hosting for inference, you must also configure and create an endpoint. SageMaker will use this endpoint for inference. The following configuration example shows how to generate and configure an endpoint with the instance type and model name that you defined previously:
In the previous configuration example, `create_endpoint_config_response` associates the `model_name` with a unique endpoint configuration name `endpoint_config_name` that is created with a timestamp.

Example output from the previous script follows:

```
Endpoint config name: spacy-ner-configYYYY-MM-DD-HH-MM-SS
```

For more information about endpoint errors, see **Why does my Amazon SageMaker endpoint go into the failed state when I create or update an endpoint?**

b. **Create an endpoint and wait for the endpoint to be in service.**

The following code example creates the endpoint using the configuration from the previous configuration example and deploys the model:

```
%%time
import time

endpoint_name = 'spacy-ner-endpoint' + strftime("%Y-%m-%d-%H-%M-%S", gmtime())
print('Endpoint name: ' + endpoint_name)
```
create_endpoint_response = sm_client.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name)
print('Endpoint Arn: ' + create_endpoint_response['EndpointArn'])

resp = sm_client.describe_endpoint(EndpointName=endpoint_name)
status = resp['EndpointStatus']
print("Endpoint Status: " + status)

print('Waiting for {} endpoint to be in service...'.format(endpoint_name))
waiter = sm_client.get_waiter('endpoint_in_service')
waiter.wait(EndpointName=endpoint_name)

In the previous code example, the create_endpoint method creates the endpoint with the generated endpoint name created in the previous code example, and prints the Amazon Resource Name of the endpoint. The describe_endpoint method returns information about the endpoint and its status. A SageMaker waiter waits for the endpoint to be in service.

8. Test your endpoint.

Once your endpoint is in service, send an invocation request to your endpoint. The following code example shows how to send a test request to your endpoint:

```python
import json
content_type = "application/json"
request_body = {
    "input": "This is a test with NER in America with Amazon and Microsoft in Seattle, writing random stuff."
}

#Serialize data for endpoint
#data = json.loads(json.dumps(request_body))
payload = json.dumps(request_body)

#Endpoint invocation
response = runtime_sm_client.invoke_endpoint(
    EndpointName=endpoint_name,
    ContentType=content_type,
    Body=payload)

#Parse results
result = json.loads(response['Body'].read().decode())['output']
result
```

Adapting Your Own Inference Container
In the previous code example, the method `json.dumps` serializes the `request_body` into a string formatted in JSON and saves it in the variable `payload`. Then SageMaker Runtime client uses the `invoke endpoint` method to send payload to your endpoint. The result contains the response from your endpoint after extracting the output field.

The previous code example should return the following output:

```
[['NER', 'ORG'],
 ['America', 'GPE'],
 ['Amazon', 'ORG'],
 ['Microsoft', 'ORG'],
 ['Seattle', 'GPE']]```

9. **Delete your endpoint**

After you have completed your invocations, delete your endpoint to conserve resources. The following code example shows you how to delete your endpoint:

```
sm_client.delete_endpoint(EndpointName=endpoint_name)
sm_client.delete_endpoint_config(EndpointConfigName=endpoint_config_name)
sm_client.delete_model(ModelName=model_name)
```

For a complete notebook containing the code in this example, see [BYOC-Single-Model](#).

**Troubleshooting your container deployment**

If your endpoint did not deploy, check the Amazon CloudWatch Events logs as follows:

1. From the [https://console.aws.amazon.com/sagemaker/](https://console.aws.amazon.com/sagemaker/) SageMaker console navigation pane, choose **Inference**.
2. Under **Inference**, choose **Endpoints**.
3. Find your endpoint under **Name**, and click on the name of the endpoint. In this example, the name would follow the naming convention `spacy-ner-configYYYY-MM-DD-HH-MM-SS`.
4. Under **Endpoint summary**, choose the link under **Model container logs**.
5. Choose the most recent **Log stream** in the **Log streams** box.
Use the following list to troubleshoot deploying your endpoint. If you need further assistance, contact AWS Support or AWS Developer Forums for Amazon SageMaker.

Topics

• Name error
• Insufficient quota
• Upstream timed out error

Name error

If the logs state NameError: name 'null' is not defined, make sure your scripts were not created in a notebook ending in .ipnyb and then renamed to another file name such as Dockerfile. When you create a notebook, formatting characters may prevent your endpoint from deploying. If you get this error and you change your scripts to fix it, you may need to restart your kernel for the changes to take effect.

Insufficient quota

If you receive a ResourceLimitExceeded error, you must request additional quota as follows:

Request an AWS Service Quotas increase

1. Retrieve the instance name, current quota and necessary quota from the on screen error message. For example, in the following sample error:

   • The instance name is ml.c5d.18xlarge.
   • The current quota from the number following current utilization is 1 instances.
   • The additional required quota from the number following request delta is 1 instances.

   The sample error follows:

   ```
   ResourceLimitExceeded: An error occurred (ResourceLimitExceeded) when calling the CreateEndpoint operation: The account-level service limit 'ml.c5d.18xlarge for endpoint usage' is 1 Instances, with current utilization of 1 Instances and a request delta of 1 Instances. Please use AWS Service Quotas to request an increase for this quota. If AWS Service Quotas is not available, contact AWS support to request an increase for this quota.
   ```
2. Sign into the AWS Management Console and open the Service Quotas console.
3. In the navigation pane, under Manage quotas, input Amazon SageMaker.
4. Choose View quotas.
5. In the search bar under Service quotas, input the name of the instance from Step 1. For example, using the information contained in the error message from Step 1, input ml.c5d.18xlarge.
6. Choose the Quota name that appears next to your instance name and ends with for endpoint usage. For example, using the information contained in the error message from Step 1, choose ml.g5.12xlarge for endpoint usage.
7. Choose Request increase at account-level.
8. Under Increase quota value, input the necessary required quota from the information given in the error message from Step 1. Input the total of current utilization and request delta. In the previous example error, the current utilization is 1 Instances, and the request delta is 1 Instances. In this example, request a quota of 2 to supply the required quota.
9. Choose Request.
10. Choose Quota request history from the navigation pane.
11. When the Status changes from Pending to Approved, rerun your job. You may need to refresh your browser to see the change.

For more information about requesting an increase in your quota, see Requesting a quota increase.

Upstream timed out error

If you receive a upstream timed out (110: Connection timed out) error, you can try the following:

- Reduce the latency of the container or increase the container's timeout limit. SageMaker requires that your container respond to a request within 60 seconds.
- Increase the amount of time before your web server waits for a response from the model.

For more information about time out errors, see How can I resolve the Amazon SageMaker inference error "upstream timed out (110: Connection timed out) while reading response header from upstream"?
Create a container with your own algorithms and models

If none of the existing SageMaker containers meet your needs and you don't have an existing container of your own, you may need to create a new Docker container. The following sections show how to create Docker containers with your training and inference algorithms for use with SageMaker.

Topics

- Use Your Own Training Algorithms
- Use your own inference code

Use Your Own Training Algorithms

This section explains how Amazon SageMaker interacts with a Docker container that runs your custom training algorithm. Use this information to write training code and create a Docker image for your training algorithms.

Topics

- How Amazon SageMaker Runs Your Training Image
- How Amazon SageMaker Provides Training Information
- Run Training with EFA
- How Amazon SageMaker Signals Algorithm Success and Failure
- How Amazon SageMaker Processes Training Output

How Amazon SageMaker Runs Your Training Image

You can use a custom entrypoint script to automate infrastructure to train in a production environment. If you pass your entrypoint script into your Docker container, you can also run it as a standalone script without rebuilding your images. SageMaker processes your training image using a Docker container entrypoint script.

This section shows you how to use a custom entrypoint without using the training toolkit. If you want to use a custom entrypoint but are unfamiliar with how to manually configure a Docker container, we recommend that you use the SageMaker training toolkit library instead. For more information about how to use the training toolkit, see Adapting your own training container.
By default, SageMaker looks for a script called `train` inside your container. You can also manually provide your own custom entrypoint by using the `ContainerArguments` and `ContainerEntrypoint` parameters of the `AlgorithmSpecification` API.

You have the following two options to manually configure your Docker container to run your image:

- Use the `CreateTrainingJob` API and a Docker container with an entrypoint instruction contained inside of it.
- Use the `CreateTrainingJob` API, and pass your training script from outside of your Docker container.

If you pass your training script from outside your Docker container, you don’t need to rebuild the Docker container when you update your script. You can also use several different scripts to run in the same container.

Your entrypoint script should contain training code for your image. If you use the optional `source_dir` parameter inside an `estimator`, it should reference the relative Amazon S3 path to the folder containing your entrypoint script. You can reference multiple files using the `source_dir` parameter. If you do not use `source_dir`, you can specify the entrypoint using the `entry_point` parameter. For an example of a custom entrypoint script that contains an estimator, see Bring Your Own Model with SageMaker Script Mode.

SageMaker model training supports high-performance S3 Express One Zone directory buckets as a data input location for file mode, fast file mode, and pipe mode. You can also use S3 Express One Zone directory buckets to store your training output. To use S3 Express One Zone, provide the URI of an S3 Express One Zone directory bucket instead of an Amazon S3 general purpose bucket. For more information, see S3 Express One Zone.

**Run a training job with an entrypoint script bundled inside the Docker container**

SageMaker can run an entrypoint script bundled inside your Docker container.

- By default, Amazon SageMaker runs the following container.

  ```bash
docker run image train
  ```
Amazon SageMaker

- SageMaker overrides any default `CMD` statements in a container by specifying the `train` argument after the image name. In your Docker container, use the following `exec` form of the `ENTRYPOINT` instruction.

```
ENTRYPOINT ["executable", "param1", "param2", ...]
```

The following example shows how to specify a python entrypoint instruction called `k-means-algorithm.py`.

```
ENTRYPOINT ["python", "k-means-algorithm.py"]
```

The `exec` form of the `ENTRYPOINT` instruction starts the executable directly, not as a child of `/bin/sh`. This enables it to receive signals like SIGTERM and SIGKILL from SageMaker APIs. The following conditions apply when using the SageMaker APIs.

- The `CreateTrainingJob` API has a stopping condition that directs SageMaker to stop model training after a specific time.
- The following shows the `StopTrainingJob` API. This API issues the equivalent of the `docker stop`, with a 2-minute timeout command to gracefully stop the specified container.

```
docker stop -t 120
```

The command attempts to stop the running container by sending a SIGTERM signal. After the 2-minute timeout, the API sends SIGKILL and forcibly stops the containers. If the container handles the SIGTERM gracefully and exits within 120 seconds from receiving it, no SIGKILL is sent.

If you want access to the intermediate model artifacts after SageMaker stops the training, add code to handle saving artifacts in your SIGTERM handler.

- If you plan to use GPU devices for model training, make sure that your containers are `nvidia-docker` compatible. Include only the CUDA toolkit on containers; don't bundle NVIDIA drivers with the image. For more information about `nvidia-docker`, see `NVIDIA/nvidia-docker`.
- You can't use the `tini` initializer as your entrypoint script in SageMaker containers because it gets confused by the `train` and `serve` arguments.
- `/opt/ml` and all subdirectories are reserved by SageMaker training. When building your algorithm's Docker image, make sure that you don't place any data that's required by your algorithm in this directory. Because if you do, the data may no longer be visible during training.
To bundle your shell or Python scripts inside your Docker image, or to provide the script in an Amazon S3 bucket or by using the AWS Command Line Interface (CLI), continue to the following section.

**Bundle your shell script in a Docker container**

If you want to bundle a custom shell script inside your Docker image, use the following steps.

1. Copy your shell script from your working directory to inside your Docker container. The following code snippet copies a custom entrypoint script `custom_entrypoint.sh` from the current working directory to a Docker container located in `mydir`. The following example assumes that the base Docker image has Python installed.

   ```
   FROM <base-docker-image>:<tag>
   
   # Copy custom entrypoint from current dir to /mydir on container
   COPY ./custom_entrypoint.sh /mydir/
   ```

2. Build and push a Docker container to the Amazon Elastic Container Registry (Amazon ECR) by following the instructions at [Pushing a Docker image](https://docs.aws.amazon.com/AmazonECR/latest/userguide/pushing-images.html) in the Amazon ECR User Guide.

3. Launch the training job by running the following AWS CLI command.

   ```
   aws --region <your-region> sagemaker create-training-job 
   --training-job-name <your-training-job-name> 
   --role-arn <your-execution-role-arn> 
   --algorithm-specification '{ 
     "TrainingInputMode": "File", 
     "TrainingImage": "<your-ecr-image>", 
     "ContainerEntrypoint": ["/bin/sh"], 
     "ContainerArguments": ["/mydir/custom_entrypoint.sh"]}' 
   --output-data-config '{"S3OutputPath": "s3://custom-entrypoint-output-bucket/"}' 
   --resource-config '{"VolumeSizeInGB":10,"InstanceCount":1,"InstanceType":"ml.m5.2xlarge"}' 
   --stopping-condition '{"MaxRuntimeInSeconds": 180}'
   ```

**Bundle your Python script in a Docker container**

To bundle a custom Python script inside your Docker image, use the following steps.
1. Copy your Python script from your working directory to inside your Docker container. The following code snippet copies a custom entrypoint script `custom_entrypoint.py` from the current working directory to a Docker container located in `mydir`.

```
FROM <base-docker-image>:<tag>
# Copy custom entrypoint from current dir to /mydir on container
COPY ./custom_entrypoint.py /mydir/
```

2. Launch the training job by running the following AWS CLI command.

```
--algorithm-specification '{
  "TrainingInputMode": "File",
  "TrainingImage": "<your-ecr-image>",
  "ContainerEntrypoint": ["python"],
  "ContainerArguments": ["/mydir/custom_entrypoint.py"]
}'
```

### Run a training job with an entrypoint script outside the Docker container

You can use your own Docker container for training and pass in an entrypoint script from outside the Docker container. There are some benefits to structuring your entrypoint script outside the container. If you update your entrypoint script, you don’t need to rebuild the Docker container. You can also use several different scripts to run in the same container.

Specify the location of your training script using the `ContainerEntrypoint` and `ContainerArguments` parameters of the `AlgorithmSpecification` API. These entrypoints and arguments behave in the same manner as Docker entrypoints and arguments. The values in these parameters override the corresponding `ENTRYPOINT` or `CMD` provided as part of the Docker container.

When you pass your custom entrypoint script to your Docker training container, the inputs that you provide determine the behavior of the container.

- For example, if you provide only `ContainerEntrypoint`, the request syntax using the `CreateTrainingJob` API is as follows.

```
{
  "AlgorithmSpecification": {
    "ContainerEntrypoint": ["string"],
    ...
  }
```

Then, the SageMaker training backend runs your custom entrypoint as follows.

```
docker run --entrypoint <ContainerEntrypoint> image
```

**Note**

If `ContainerEntrypoint` is provided, the SageMaker training backend runs the image with the given entrypoint and overrides the default `ENTRYPOINT` in the image.

- If you provide only `ContainerArguments`, SageMaker assumes that the Docker container contains an entrypoint script. The request syntax using the `CreateTrainingJob` API is as follows.

```
{
   "AlgorithmSpecification": {
      "ContainerArguments": ["arg1", "arg2"],
      ...
   }
}
```

The SageMaker training backend runs your custom entrypoint as follows.

```
docker run image <ContainerArguments>
```

- If you provide both the `ContainerEntrypoint` and `ContainerArguments`, then the request syntax using the `CreateTrainingJob` API is as follows.

```
{
   "AlgorithmSpecification": {
      "ContainerEntrypoint": ["string"],
      "ContainerArguments": ["arg1", "arg2"],
      ...
   }
}
```

The SageMaker training backend runs your custom entrypoint as follows.
You can use any supported InputDataConfig source in the CreateTrainingJob API to provide an entrypoint script to run your training image.

**Provide your entrypoint script in an Amazon S3 bucket**

To provide a custom entrypoint script using an S3 bucket, use the S3DataSource parameter of the [DataSource](https://docs.aws.amazon.com/sagemaker/latest/dg/API_DataSource.html) API to specify the location of the script. If you use the S3DataSource parameter, the following are required.

- The [InputMode](https://docs.aws.amazon.com/sagemaker/latest/dg/API_InputMode.html) must be of the type File.
- The [S3DataDistributionType](https://docs.aws.amazon.com/sagemaker/latest/dg/API_S3DataDistributionType.html) must be FullyReplicated.

The following example has a script called `custom_entrypoint.sh` placed in a path to an S3 bucket `s3://<bucket-name>/<bucket prefix>/custom_entrypoint.sh`.

```bash
#!/bin/bash
echo "Running custom_entrypoint.sh"
echo "Hello you have provided the following arguments: " "$@
```

Next, you must set the configuration of the input data channel to run a training job. Do this either by using the AWS CLI directly or with a JSON file.

**Configure the input data channel using AWS CLI with a JSON file**

To configure your input data channel with a JSON file, use AWS CLI as shown in the following code structure. Ensure that all of the following fields use the request syntax defined in the [CreateTrainingJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateTrainingJob.html) API.

```json
// run-my-training-job.json
{
   "AlgorithmSpecification": {
      "ContainerEntrypoint": ["/bin/sh"],
      "ContainerArguments": ["/opt/ml/input/data/<your_channel_name>/custom_entrypoint.sh"],
      ...
```

**Use Your Own Training Algorithms**
Next, run the AWS CLI command to launch the training job from the JSON file as follows.

```bash
aws sagemaker create-training-job --cli-input-json file://run-my-training-job.json
```

Configure the input data channel using AWS CLI directly

To configure your input data channel without a JSON file, use the following AWS CLI code structure.

```bash
aws --region <your-region> sagemaker create-training-job \
--training-job-name <your-training-job-name> \
--role-arn <your-execution-role-arn> \
--algorithm-specification '{ \
  "TrainingInputMode": "File", \
  "TrainingImage": "<your-ecr-image>", \
  "ContainerEntrypoint": ["/bin/sh"], \
  "ContainerArguments": ["/opt/ml/input/data/<your_channel_name>/custom_entrypoint.sh"]}' \
--input-data-config '{[ \
  "ChannelName": "<your_channel_name>", \
  "DataSource":{ \
    "S3DataSource":{ \
      "S3DataDistributionType": "FullyReplicated", \
      "S3DataType": "S3Prefix", \
      "S3Uri": "s3://<bucket-name>/<bucket_prefix>" \
    }, \
    "InputMode": "File", \
  }, \
...]

Use Your Own Training Algorithms
How Amazon SageMaker Provides Training Information

This section explains how SageMaker makes training information, such as training data, hyperparameters, and other configuration information, available to your Docker container.

When you send a `CreateTrainingJob` request to SageMaker to start model training, you specify the Amazon Elastic Container Registry (Amazon ECR) path of the Docker image that contains the training algorithm. You also specify the Amazon Simple Storage Service (Amazon S3) location where training data is stored and algorithm-specific parameters. SageMaker makes this information available to the Docker container so that your training algorithm can use it. This section explains how we make this information available to your Docker container. For information about creating a training job, see `CreateTrainingJob`. For more information on the way that SageMaker containers organize information, see Using the SageMaker Training and Inference Toolkits.

Topics

- Hyperparameters
- Environment Variables
- Input Data Configuration
- Training Data
- Distributed Training Configuration

Hyperparameters

SageMaker makes the hyperparameters in a `CreateTrainingJob` request available in the Docker container in the `/opt/ml/input/config/hyperparameters.json` file.

The following is an example of a hyperparameter configuration in `hyperparameters.json` to specify the `num_round` and `eta` hyperparameters in the `CreateTrainingJob` operation for XGBoost.

```json
{
    "num_round": "128",
    "eta": "0.001"
}
```

Use Your Own Training Algorithms
For a complete list of hyperparameters that can be used for the SageMaker built-in XGBoost algorithm, see [XGBoost Hyperparameters](#).

The hyperparameters that you can tune depend on the algorithm that you are training. For a list of hyperparameters available for a SageMaker built-in algorithm, find them listed in **Hyperparameters** under the algorithm link in [Use Amazon SageMaker Built-in Algorithms or Pre-trained Models](#).

### Environment Variables

SageMaker sets the following environment variables in your container:

- **TRAINING_JOB_NAME** – Specified in the `TrainingJobName` parameter of the `CreateTrainingJob` request.
- **TRAINING_JOB_ARN** – The Amazon Resource Name (ARN) of the training job returned as the `TrainingJobArn` in the `CreateTrainingJob` response.
- All environment variables specified in the `Environment` parameter in the `CreateTrainingJob` request.

### Input Data Configuration

SageMaker makes the data channel information in the `InputDataConfig` parameter from your `CreateTrainingJob` request available in the `/opt/ml/input/config/inputdataconfig.json` file in your Docker container.

For example, suppose that you specify three data channels (train, evaluation, and validation) in your request. SageMaker provides the following JSON:

```json
{
    "train": {
        "ContentType": "trainingContentType",
        "TrainingInputMode": "File",
        "S3DistributionType": "FullyReplicated",
        "RecordWrapperType": "None"},
    "evaluation": {
        "ContentType": "evalContentType",
        "TrainingInputMode": "File",
        "S3DistributionType": "FullyReplicated",
        "RecordWrapperType": "None"}
}
```
"validation": {
  "TrainingInputMode": "File",
  "S3DistributionType": "FullyReplicated",
  "RecordWrapperType": "None"}
}

**Note**

SageMaker provides only relevant information about each data channel (for example, the channel name and the content type) to the container, as shown in the previous example. S3DistributionType will be set as FullyReplicated if you specify EFS or FSxLustre as input data sources.

**Training Data**

The TrainingInputMode parameter in the AlgorithmSpecification of the CreateTrainingJob request specifies how the training dataset is made available to your container. The following input modes are available.

- **File mode**

  If you use File mode as your TrainingInputMode value, SageMaker sets the following parameters in your container.

  - Your TrainingInputMode parameter is written to inputdataconfig.json as "File".
  - Your data channel directory is written to /opt/ml/input/data/channel_name.

  If you use File mode, SageMaker creates a directory for each channel. For example, if you have three channels named training, validation, and testing, SageMaker makes the following three directories in your Docker container:

  - /opt/ml/input/data/training
  - /opt/ml/input/data/validation
  - /opt/ml/input/data/testing

  File mode also supports the following data sources.

  - Amazon Simple Storage Service (Amazon S3)
  - Amazon Elastic File System (Amazon EFS)
Channels that use file system data sources such as Amazon EFS and Amazon FSx must use File mode. In this case, the directory path provided in the channel is mounted at /opt/ml/input/data/channel_name.

**FastFile mode**

If you use FastFile mode as your TrainingInputNodeParameter, SageMaker sets the following parameters in your container.

- Similar to File mode, in FastFile mode, your TrainingInputMode parameter is written to inputdataconfig.json as "File".
- Your data channel directory is written to /opt/ml/input/data/channel_name.

FastFile mode supports the following data sources.

- Amazon S3

If you use FastFile mode, the channel directory is mounted with read-only permission.

Historically, File mode preceded FastFile mode. To ensure backwards compatibility, algorithms that support File mode can also seamlessly work with FastFile mode as long as the TrainingInputMode parameter is set to File in inputdataconfig.json.

**Note**

Channels that use FastFile mode must use a S3DataType of "S3Prefix".

FastFile mode presents a folder view that uses the forward slash (/) as the delimiter for grouping Amazon S3 objects into folders. S3Uri prefixes must not correspond to a partial folder name. For example, if an Amazon S3 dataset contains s3://my-bucket/train-01/data.csv, then neither s3://my-bucket/train nor s3://my-bucket/train-01 are allowed as S3Uri prefixes.

A trailing forward slash is recommended to define a channel corresponding to a folder. For example, the s3://my-bucket/train-01/ channel for the train-01 folder. Without the trailing forward slash, the channel would be ambiguous if there existed another folder s3://my-bucket/train-011/ or file s3://my-bucket/train-01.txt/.
• **Pipe mode**

  - TrainingInputMode parameter written to inputdataconfig.json: "Pipe"
  
  - Data channel directory in the Docker container: `/opt/ml/input/data/<channel_name_epoch_number>`

  - Supported data sources: Amazon S3

You need to read from a separate pipe for each channel. For example, if you have three channels named training, validation, and testing, you need to read from the following pipes:

  - `/opt/ml/input/data/training_0`, `/opt/ml/input/data/training_1`, ...
  - `/opt/ml/input/data/validation_0`, `/opt/ml/input/data/validation_1`, ...
  - `/opt/ml/input/data/testing_0`, `/opt/ml/input/data/testing_1`, ...

Read the pipes sequentially. For example, if you have a channel called training, read the pipes in this sequence:

1. Open `/opt/ml/input/data/training_0` in read mode and read it to end-of-file (EOF) or, if you are done with the first epoch, close the pipe file early.

2. After closing the first pipe file, look for `/opt/ml/input/data/training_1` and read it until you have completed the second epoch, and so on.

If the file for a given epoch doesn't exist yet, your code may need to retry until the pipe is created. There is no sequencing restriction across channel types. For example, you can read multiple epochs for the training channel and only start reading the validation channel when you are ready. Or, you can read them simultaneously if your algorithm requires that.

For an example of a Jupyter notebook that shows how to use Pipe mode when bringing your own container, see [Bring your own pipe-mode algorithm to Amazon SageMaker](#).

SageMaker model training supports high-performance S3 Express One Zone directory buckets as a data input location for file mode, fast file mode, and pipe mode. To use S3 Express One Zone, input the location of the S3 Express One Zone directory bucket instead of an Amazon S3 general purpose bucket. Provide the ARN for the IAM role with the required access control and permissions policy. Refer to [AmazonSageMakerFullAccesspolicy](#) for details. For more information, see [S3 Express One Zone](#).
Distributed Training Configuration

If you're performing distributed training with multiple containers, SageMaker makes information about all containers available in the /opt/ml/input/config/resourceconfig.json file.

To enable inter-container communication, this JSON file contains information for all containers. SageMaker makes this file available for both File and Pipe mode algorithms. The file provides the following information:

- **current_host**—The name of the current container on the container network. For example, algo-1. Host values can change at any time. Don't write code with specific values for this variable.
- **hosts**—The list of names of all containers on the container network, sorted lexicographically. For example, ["algo-1", "algo-2", "algo-3"] for a three-node cluster. Containers can use these names to address other containers on the container network. Host values can change at any time. Don't write code with specific values for these variables.
- **network_interface_name**—The name of the network interface that is exposed to your container. For example, containers running the Message Passing Interface (MPI) can use this information to set the network interface name.
- Do not use the information in /etc/hostname or /etc/hosts because it might be inaccurate.
- Hostname information may not be immediately available to the algorithm container. We recommend adding a retry policy on hostname resolution operations as nodes become available in the cluster.

The following is an example file on node 1 in a three-node cluster:

```json
{
    "current_host": "algo-1",
    "hosts": ["algo-1","algo-2","algo-3"],
    "network_interface_name":"eth1"
}
```

Run Training with EFA

SageMaker provides integration with EFA devices to accelerate High Performance Computing (HPC) and machine learning applications. This integration allows you to leverage an EFA device when running your distributed training jobs. You can add EFA integration to an existing Docker
container that you bring to SageMaker. The following information outlines how to configure your own container to use an EFA device for your distributed training jobs.

Prerequisites

Your container must satisfy the SageMaker Training container specification.

Install EFA and required packages

Your container must download and install the EFA software. This allows your container to recognize the EFA device, and provides compatible versions of Libfabric and Open MPI.

Any tools like MPI and NCCL must be installed and managed inside the container to be used as part of your EFA-enabled training job. For a list of all available EFA versions, see Verify the EFA installer using a checksum. The following example shows how to modify the Dockerfile of your EFA-enabled container to install EFA, MPI, OFI, NCCL, and NCCL-TEST.

Note

When using PyTorch with EFA on your container, the NCCL version of your container should match the NCCL version of your PyTorch installation. To verify the PyTorch NCCL version, use the following command:

torch.cuda.nccl.version()
Considerations when creating your container

The EFA device is mounted to the container as /dev/infiniband/uverbs0 under the list of devices accessible to the container. On P4d instances, the container has access to 4 EFA devices. The EFA devices can be found in the list of devices accessible to the container as:

- /dev/infiniband/uverbs0
- /dev/infiniband/uverbs1
- /dev/infiniband/uverbs2
- /dev/infiniband/uverbs3

To get information about hostname, peer hostnames, and network interface (for MPI) from the resourceconfig.json file provided to each container instances, see [Distributed Training](#).
Configuration. Your container handles regular TCP traffic among peers through the default Elastic Network Interfaces (ENI), while handling OFI (kernel bypass) traffic through the EFA device.

Verify that your EFA device is recognized

To verify that the EFA device is recognized, run the following command from within your container.

```
/opt/amazon/efa/bin/fi_info -p efa
```

Your output should look similar to the following.

```
provider: efa
    fabric: EFA-fe80::e5:56ff:fe34:56a8
    domain: efa_0-rdm
    version: 2.0
    type: FI_EP_RDM
    protocol: FIPROTO_EFA
provider: efa
    fabric: EFA-fe80::e5:56ff:fe34:56a8
    domain: efa_0-dgrm
    version: 2.0
    type: FI_EP_DGRAM
    protocol: FIPROTO_EFA
provider: efa;ofi_rx
    fabric: EFA-fe80::e5:56ff:fe34:56a8
    domain: efa_0-dgrm
    version: 1.0
    type: FI_EP_RDM
    protocol: FIPROTO_RXD
```

Running a training job with EFA

Once you’ve created an EFA-enabled container, you can run a training job with EFA using a SageMaker Estimator the same way as you would with any other Docker image. For more information on registering your container and using it for training, see Adapting Your Own Training Container.

How Amazon SageMaker Signals Algorithm Success and Failure

A training algorithm indicates whether it succeeded or failed using the exit code of its process.
A successful training execution should exit with an exit code of 0 and an unsuccessful training execution should exit with a non-zero exit code. These will be converted to Completed and Failed in the TrainingJobStatus returned by DescribeTrainingJob. This exit code convention is standard and is easily implemented in all languages. For example, in Python, you can use `sys.exit(1)` to signal a failure exit, and simply running to the end of the main routine will cause Python to exit with code 0.

In the case of failure, the algorithm can write a description of the failure to the failure file. See next section for details.

**How Amazon SageMaker Processes Training Output**

As your algorithm runs in a container, it generates output including the status of the training job and model and output artifacts. Your algorithm should write this information to the following files, which are located in the container's `/output` directory. Amazon SageMaker processes the information contained in this directory as follows:

- `/opt/ml/model` – Your algorithm should write all final model artifacts to this directory. SageMaker copies this data as a single object in compressed tar format to the S3 location that you specified in the CreateTrainingJob request. If multiple containers in a single training job write to this directory they should ensure no file/directory names clash. SageMaker aggregates the result in a TAR file and uploads to S3 at the end of the training job.

- `/opt/ml/output/data` – Your algorithm should write artifacts you want to store other than the final model to this directory. SageMaker copies this data as a single object in compressed tar format to the S3 location that you specified in the CreateTrainingJob request. If multiple containers in a single training job write to this directory they should ensure no file/directory names clash. SageMaker aggregates the result in a TAR file and uploads to S3 at the end of the training job.

- `/opt/ml/output/failure` – If training fails, after all algorithm output (for example, logging) completes, your algorithm should write the failure description to this file. In a DescribeTrainingJob response, SageMaker returns the first 1024 characters from this file as `FailureReason`.

You can specify either an S3 general purpose or S3 directory bucket to store your training output. Directory buckets use only the Amazon S3 Express One Zone storage class, which is designed for workloads or performance-critical applications that require consistent single-digit millisecond latency. Choose the bucket type that best fits your application and performance requirements. For
more information on S3 directory buckets, see Directory buckets in the Amazon Simple Storage Service User Guide.

Use your own inference code

You can use Amazon SageMaker to interact with Docker containers and run your own inference code in one of two ways:

- To use your own inference code with a persistent endpoint to get one prediction at a time, use SageMaker hosting services.
- To use your own inference code to get predictions for an entire dataset, use SageMaker batch transform.

Topics

- Use Your Own Inference Code with Hosting Services
- Use Your Own Inference Code with Batch Transform

Use Your Own Inference Code with Hosting Services

This section explains how Amazon SageMaker interacts with a Docker container that runs your own inference code for hosting services. Use this information to write inference code and create a Docker image.

Topics

- How SageMaker Runs Your Inference Image
- How SageMaker Loads Your Model Artifacts
- How Your Container Should Respond to Inference Requests
- How Your Container Should Respond to Health Check (Ping) Requests
- Use a Private Docker Registry for Real-Time Inference Containers

How SageMaker Runs Your Inference Image

To configure a container to run as an executable, use an ENTRYPOINT instruction in a Dockerfile. Note the following:

- For model inference, SageMaker runs the container as:
Amazon SageMaker

SageMaker overrides default CMD statements in a container by specifying the serve argument after the image name. The serve argument overrides arguments that you provide with the CMD command in the Dockerfile.

- SageMaker expects all containers to run with root users. Create your container so that it uses only root users. When SageMaker runs your container, users that do not have root-level access can cause permissions issues.

- We recommend that you use the exec form of the ENTRYPOINT instruction:

  ```
  ENTRYPOINT ["executable", "param1", "param2"]
  ```

  For example:

  ```
  ENTRYPOINT ["python", "k_means_inference.py"]
  ```

  The exec form of the ENTRYPOINT instruction starts the executable directly, not as a child of /bin/sh. This enables it to receive signals like SIGTERM and SIGKILL from the SageMaker API operations, which is a requirement.

  For example, when you use the CreateEndpoint API to create an endpoint, SageMaker provisions the number of ML compute instances required by the endpoint configuration, which you specify in the request. SageMaker runs the Docker container on those instances.

  If you reduce the number of instances backing the endpoint (by calling the UpdateEndpointWeightsAndCapacities API), SageMaker runs a command to stop the Docker container on the instances that are being terminated. The command sends the SIGTERM signal, then it sends the SIGKILL signal thirty seconds later.

```
If you update the endpoint (by calling the `UpdateEndpoint` API), SageMaker launches another set of ML compute instances and runs the Docker containers that contain your inference code on them. Then it runs a command to stop the previous Docker containers. To stop a Docker container, command sends the SIGTERM signal, then it sends the SIGKILL signal 30 seconds later.

- SageMaker uses the container definition that you provided in your `CreateModel` request to set environment variables and the DNS hostname for the container as follows:
  - It sets environment variables using the `ContainerDefinition.Environment` string-to-string map.
  - It sets the DNS hostname using the `ContainerDefinition(ContainerHostname).

- If you plan to use GPU devices for model inferences (by specifying GPU-based ML compute instances in your `CreateEndpointConfig` request), make sure that your containers are `nvidia-docker` compatible. Don't bundle NVIDIA drivers with the image. For more information about `nvidia-docker`, see `NVIDIA/nvidia-docker`.

- You can't use the `tini` initializer as your entry point in SageMaker containers because it gets confused by the `train` and `serve` arguments.

**How SageMaker Loads Your Model Artifacts**

In your `CreateModel` API request, you can use either the `ModelDataUrl` or `S3DataSource` parameter to identify the S3 location where model artifacts are stored. SageMaker copies your model artifacts from the S3 location to the `/opt/ml/model` directory for use by your inference code. Your container has read-only access to `/opt/ml/model`. Do not write to this directory.

The `ModelDataUrl` must point to a tar.gz file. Otherwise, SageMaker won't download the file.
If you trained your model in SageMaker, the model artifacts are saved as a single compressed tar file in Amazon S3. If you trained your model outside SageMaker, you need to create this single compressed tar file and save it in a S3 location. SageMaker decompresses this tar file into /opt/ml/model directory before your container starts.

For deploying large models, we recommend that you follow [Deploying uncompressed models](#).

**How Your Container Should Respond to Inference Requests**

To obtain inferences, the client application sends a POST request to the SageMaker endpoint. SageMaker passes the request to the container, and returns the inference result from the container to the client.

For more information about the inference requests that your container will receive, see the following actions in the [Amazon SageMaker API Reference](#):

- `InvokeEndpoint`
- `InvokeEndpointAsync`
- `InvokeEndpointWithResponseStream`

**Requirements for inference containers**

To respond to inference requests, your container must meet the following requirements:

- SageMaker strips all POST headers except those supported by `InvokeEndpoint`. SageMaker might add additional headers. Inference containers must be able to safely ignore these additional headers.
- To receive inference requests, the container must have a web server listening on port 8080 and must accept POST requests to the /invocations and /ping endpoints.
- A customer's model containers must accept socket connection requests within 250 ms.
- A customer's model containers must respond to requests within 60 seconds. The model itself can have a maximum processing time of 60 seconds before responding to the /invocations. If your model is going to take 50-60 seconds of processing time, the SDK socket timeout should be set to be 70 seconds.
Example invocation functions

The following examples demonstrate how the code in your container can process inference requests. These examples handle requests that client applications send by using the InvokeEndpoint action.

FastAPI

FastAPI is a web framework for building APIs with Python.

```python
from fastapi import FastAPI, status, Request, Response
...
app = FastAPI()
...
@app.post('/invocations')
async def invocations(request: Request):
    # model() is a hypothetical function that gets the inference output:
    model_resp = await model(Request)

    response = Response(
        content=model_resp,
        status_code=status.HTTP_200_OK,
        media_type="text/plain",
    )
    return response
...
```

In this example, the `invocations` function handles the inference request that SageMaker sends to the `/invocations` endpoint.

Flask

Flask is a framework for developing web applications with Python.

```python
import flask
...
app = flask.Flask(__name__)
...
@app.route('/invocations', methods=['POST'])
def invoke(request):
    # model() is a hypothetical function that gets the inference output:
    resp_body = model(request)
    return flask.Response(resp_body, mimetype='text/plain')
```
In this example, the `invoke` function handles the inference request that SageMaker sends to the `/invocations` endpoint.

**Example invocation functions for streaming requests**

The following examples demonstrate how the code in your inference container can process streaming inference requests. These examples handle requests that client applications send by using the `InvokeEndpointWithResponseStream` action.

When a container handles a streaming inference request, it returns the model's inference as a series of parts incrementally as the model generates them. Client applications start receiving responses immediately when they're available. They don't need to wait for the model to generate the entire response. You can implement streaming to support fast interactive experiences, such as chatbots, virtual assistants, and music generators.

**FastAPI**

FastAPI is a web framework for building APIs with Python.

```python
from starlette.responses import StreamingResponse
from fastapi import FastAPI, status, Request

app = FastAPI()

@app.post('/invocations')
async def invocations(request: Request):
    async def generate():
        # model() is a hypothetical function that gets the inference output:
        yield await model(Request)
        yield "\n"

    response = StreamingResponse(
        content=generate(),
        status_code=status.HTTP_200_OK,
        media_type="text/plain",
    )
    return response
```

**Use Your Own Inference Code**
In this example, the invocations function handles the inference request that SageMaker sends to the /invocations endpoint. To stream the response, the example uses the StreamingResponse class from the Starlette framework.

**Flask**

Flask is a framework for developing web applications with Python.

```python
import flask

app = flask.Flask(__name__)

@app.route('/invocations', methods=['POST'])
def invocations(request):
    # Streams inference response using HTTP chunked encoding

    def generate():
        # model() is a hypothetical function that gets the inference output:
        yield model(request)
        yield '\n'
    return flask.Response(flask.stream_with_context(generate()), mimetype='text/plain')
```

In this example, the invocations function handles the inference request that SageMaker sends to the /invocations endpoint. To stream the response, the example uses the `flask.stream_with_context` function from the Flask framework.

**How Your Container Should Respond to Health Check (Ping) Requests**

SageMaker launches new inference containers in the following situations:

- Responding to `CreateEndpoint`, `UpdateEndpoint`, and `UpdateEndpointWeightsAndCapacities` API calls
- Security patching
- Replacing unhealthy instances

Soon after container startup, SageMaker starts sending periodic GET requests to the /ping endpoint.
The simplest requirement on the container is to respond with an HTTP 200 status code and an empty body. This indicates to SageMaker that the container is ready to accept inference requests at the /invocations endpoint.

If the container does not begin to pass health checks by consistently responding with 200s during the 8 minutes after startup, the new instance launch fails. This causes CreateEndpoint to fail, leaving the endpoint in a failed state. The update requested by UpdateEndpoint isn't completed, security patches aren't applied, and unhealthy instances aren't replaced.

While the minimum bar is for the container to return a static 200, a container developer can use this functionality to perform deeper checks. The request timeout on /ping attempts is 2 seconds.

### Use a Private Docker Registry for Real-Time Inference Containers

Amazon SageMaker hosting enables you to use images stored in Amazon ECR to build your containers for real-time inference by default. Optionally, you can build containers for real-time inference from images in a private Docker registry. The private registry must be accessible from an Amazon VPC in your account. Models that you create based on the images stored in your private Docker registry must be configured to connect to the same VPC where the private Docker registry is accessible. For information about connecting your model to a VPC, see [Give SageMaker Hosted Endpoints Access to Resources in Your Amazon VPC](#).

Your Docker registry must be secured with a TLS certificate from a known public certificate authority (CA).

**Note**

Your private Docker registry must allow inbound traffic from the security groups you specify in the VPC configuration for your model, so that SageMaker hosting is able to pull model images from your registry.

SageMaker can pull model images from DockerHub if there's a path to the open internet inside your VPC.

### Topics

- [Store Images in a Private Docker Registry other than Amazon Elastic Container Registry](#)
- [Use an Image from a Private Docker Registry for Real-time Inference](#)
- [Allow SageMaker to authenticate to a private Docker registry](#)
- [Create the Lambda function](#)
• **Give your execution role permission to Lambda**

• **Create an interface VPC endpoint for Lambda**

**Store Images in a Private Docker Registry other than Amazon Elastic Container Registry**

To use a private Docker registry to store your images for SageMaker real-time inference, create a private registry that is accessible from your Amazon VPC. For information about creating a Docker registry, see [Deploy a registry server](#) in the Docker documentation. The Docker registry must comply with the following:

- The registry must be a [Docker Registry HTTP API V2](#) registry.
- The Docker registry must be accessible from the same VPC that you specify in the `VpcConfig` parameter that you specify when you create your model.

**Use an Image from a Private Docker Registry for Real-time Inference**

When you create a model and deploy it to SageMaker hosting, you can specify that it use an image from your private Docker registry to build the inference container. Specify this in the `ImageConfig` object in the `PrimaryContainer` parameter that you pass to a call to the `create_model` function.

**To use an image stored in your private Docker registry for your inference container**

1. Create the image configuration object and specify a value of `Vpc` for the `RepositoryAccessMode` field.

   ```python
   image_config = {
       'RepositoryAccessMode': 'Vpc'
   }
   ```

2. If your private Docker registry requires authentication, add a `RepositoryAuthConfig` object to the image configuration object. For the `RepositoryCredentialsProviderArn` field of the `RepositoryAuthConfig` object, specify the Amazon Resource Name (ARN) of an AWS Lambda function that provides credentials that allows SageMaker to authenticate to your private Docker Registry. For information about how to create the Lambda function to provide authentication, see [Allow SageMaker to authenticate to a private Docker registry](#).

   ```python
   image_config = {
   ```
3. Create the primary container object that you want to pass to create_model, using the image configuration object that you created in the previous step.

Provide your image in digest form. If you provide your image using the :latest tag, there is a risk that SageMaker pulls a newer version of the image than intended. Using the digest form ensures that SageMaker pulls the intended image version.

```python
primary_container = {
    'ContainerHostname': 'ModelContainer',
    'Image': 'myteam.myorg.com/docker-local/my-inference-image:<IMAGE-TAG>',
    'ImageConfig': image_config
}
```

4. Specify the model name and the execution role that you want to pass to create_model.

```python
model_name = 'vpc-model'
exeution_role_arn = 'arn:aws:iam::123456789012:role/SageMakerExecutionRole'
```

5. Specify one or more security groups and subnets for the VPC configuration for your model. Your private Docker registry must allow inbound traffic from the security groups that you specify. The subnets that you specify must be in the same VPC as your private Docker registry.

```python
vpc_config = {
    'SecurityGroupIds': ['sg-0123456789abcdef0'],
    'Subnets': ['subnet-0123456789abcdef0','subnet-0123456789abcdef1']
}
```


```python
import boto3
sm = boto3.client('sagemaker')
```

7. Create the model by calling create_model, using the values you specified in the previous steps for the PrimaryContainer and VpcConfig parameters.
try:
    resp = sm.create_model(
        ModelName=model_name,
        PrimaryContainer=primary_container,
        ExecutionRoleArn=execution_role_arn,
        VpcConfig=vpc_config,
    )
except Exception as e:
    print(f'error calling CreateModel operation: {e}')
else:
    print(resp)

8. Finally, call `create_endpoint_config` and `create_endpoint` to create the hosting endpoint, using the model that you created in the previous step.

```python
endpoint_config_name = 'my-endpoint-config'
sm.create_endpoint_config(
    EndpointConfigName=endpoint_config_name,
    ProductionVariants=[
        {
            'VariantName': 'MyVariant',
            'ModelName': model_name,
            'InitialInstanceCount': 1,
            'InstanceType': 'ml.t2.medium'
        },
    ],
)

endpoint_name = 'my-endpoint'
sm.create_endpoint(
    EndpointName=endpoint_name,
    EndpointConfigName=endpoint_config_name,
)

sm.describe_endpoint(EndpointName=endpoint_name)
```

**Allow SageMaker to authenticate to a private Docker registry**

To pull an inference image from a private Docker registry that requires authentication, create an AWS Lambda function that provides credentials, and provide the Amazon Resource Name (ARN) of
the Lambda function when you call `create_model`. When SageMaker runs `create_model`, it calls the Lambda function that you specified to get credentials to authenticate to your Docker registry.

**Create the Lambda function**

Create an AWS Lambda function that returns a response with the following form:

```python
def handler(event, context):
    response = {
        "Credentials": {
            "Username": "username",
            "Password": "password"
        }
    }
    return response
```

Depending on how you set up authentication for your private Docker registry, the credentials that your Lambda function returns can mean either of the following:

- If you set up your private Docker registry to use basic authentication, provide the sign-in credentials to authenticate to the registry.
- If you set up your private Docker registry to use bearer token authentication, the sign-in credentials are sent to your authorization server, which returns a Bearer token that can then be used to authenticate to the private Docker registry.

**Give your execution role permission to Lambda**

The execution role that you use to call `create_model` must have permissions to call AWS Lambda functions. Add the following to the permissions policy of your execution role.

```json
{
    "Effect": "Allow",
    "Action": [
        "lambda:InvokeFunction"
    ],
    "Resource": [
        "arn:aws:lambda:*::*:function:*myLambdaFunction*"
    ]
}
```

Where `myLambdaFunction` is the name of your Lambda function. For information about editing a role permissions policy, see [Modifying a role permissions policy (console)](https://docs.aws.amazon.com/IAM/latest/UserGuide/access-service-role-permissions-console.html) in the *AWS Identity and Access Management User Guide*. 

Use Your Own Inference Code
An execution role with the AmazonSageMakerFullAccess managed policy attached to it has permission to call any Lambda function with SageMaker in its name.

Create an interface VPC endpoint for Lambda

Create an interface endpoint so that your Amazon VPC can communicate with your AWS Lambda function without sending traffic over the internet. For information about how to do this, see Configuring interface VPC endpoints for Lambda in the AWS Lambda Developer Guide.

SageMaker hosting sends a request through your VPC to lambda.region.amazonaws.com, to call your Lambda function. If you choose Private DNS Name when you create your interface endpoint, Amazon Route 53 routes the call to the Lambda interface endpoint. If you use a different DNS provider, make sure to map lambda.region.amazonaws.com to your Lambda interface endpoint.

Use Your Own Inference Code with Batch Transform

This section explains how Amazon SageMaker interacts with a Docker container that runs your own inference code for batch transform. Use this information to write inference code and create a Docker image.

Topics

- How SageMaker Runs Your Inference Image
- How SageMaker Loads Your Model Artifacts
- How Containers Serve Requests
- How Your Container Should Respond to Inference Requests
- How Your Container Should Respond to Health Check (Ping) Requests

How SageMaker Runs Your Inference Image

To configure a container to run as an executable, use an ENTRYPOINT instruction in a Dockerfile. Note the following:

- For batch transforms, SageMaker invokes the model on your behalf. SageMaker runs the container as:
The input to batch transforms must be of a format that can be split into smaller files to process in parallel. These formats include CSV, JSON, JSON Lines, TFRecord and RecordIO.

SageMaker overrides default CMD statements in a container by specifying the serve argument after the image name. The serve argument overrides arguments that you provide with the CMD command in the Dockerfile.

- We recommend that you use the exec form of the ENTRYPOINT instruction:

  ```
  ENTRYPOINT ["executable", "param1", "param2"]
  ```

  For example:

  ```
  ENTRYPOINT ["python", "k_means_inference.py"]
  ```

- SageMaker sets environment variables specified in CreateModel and CreateTransformJob on your container. Additionally, the following environment variables are populated:
  - SAGEMAKER_BATCH is set to true when the container runs batch transforms.
  - SAGEMAKER_MAX_PAYLOAD_IN_MB is set to the largest size payload that is sent to the container via HTTP.
  - SAGEMAKER_BATCH_STRATEGY is set to SINGLE_RECORD when the container is sent a single record per call to invocations and MULTI_RECORD when the container gets as many records as will fit in the payload.
  - SAGEMAKER_MAX_CONCURRENT_TRANSFORMS is set to the maximum number of / invocations requests that can be opened simultaneously.

**Note**

The last three environment variables come from the API call made by the user. If the user doesn't set values for them, they aren't passed. In that case, either the default values or
the values requested by the algorithm (in response to the /execution-parameters) are used.

- If you plan to use GPU devices for model inferences (by specifying GPU-based ML compute instances in your CreateTransformJob request), make sure that your containers are nvidia-docker compatible. Don't bundle NVIDIA drivers with the image. For more information about nvidia-docker, see NVIDIA/nvidia-docker.

- You can't use the init initializer as your entry point in SageMaker containers because it gets confused by the train and serve arguments.

**How SageMaker Loads Your Model Artifacts**

In a CreateModel request, container definitions include the ModelDataUrl parameter, which identifies the location in Amazon S3 where model artifacts are stored. When you use SageMaker to run inferences, it uses this information to determine from where to copy the model artifacts. It copies the artifacts to the /opt/ml/model directory in the Docker container for use by your inference code.

The ModelDataUrl parameter must point to a tar.gz file. Otherwise, SageMaker can't download the file. If you train a model in SageMaker, it saves the artifacts as a single compressed tar file in Amazon S3. If you train a model in another framework, you need to store the model artifacts in Amazon S3 as a compressed tar file. SageMaker decompresses this tar file and saves it in the /opt/ml/model directory in the container before the batch transform job starts.

**How Containers Serve Requests**

Containers must implement a web server that responds to invocations and ping requests on port 8080. For batch transforms, you have the option to set algorithms to implement execution-parameters requests to provide a dynamic runtime configuration to SageMaker. SageMaker uses the following endpoints:

- ping—Used to periodically check the health of the container. SageMaker waits for an HTTP 200 status code and an empty body for a successful ping request before sending an invocations request. You might use a ping request to load a model into memory to generate inference when invocations requests are sent.
• (Optional) execution-parameters—Allows the algorithm to provide the optimal tuning parameters for a job during runtime. Based on the memory and CPUs available for a container, the algorithm chooses the appropriate MaxConcurrentTransforms, BatchStrategy, and MaxPayloadInMB values for the job.

Before calling the invocations request, SageMaker attempts to invoke the execution-parameters request. When you create a batch transform job, you can provide values for the MaxConcurrentTransforms, BatchStrategy, and MaxPayloadInMB parameters. SageMaker determines the values for these parameters using this order of precedence:

1. The parameter values that you provide when you create the CreateTransformJob request.
2. The values that the model container returns when SageMaker invokes the execution-parameters endpoint>
3. The default parameter values, listed in the following table.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Default Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>MaxConcurrentTransforms</td>
<td>1</td>
</tr>
<tr>
<td>BatchStrategy</td>
<td>MULTI_RECORD</td>
</tr>
<tr>
<td>MaxPayloadInMB</td>
<td>6</td>
</tr>
</tbody>
</table>

The response for a GET execution-parameters request is a JSON object with keys for MaxConcurrentTransforms, BatchStrategy, and MaxPayloadInMB parameters. This is an example of a valid response:

```json
{
  "MaxConcurrentTransforms": 8,
  "BatchStrategy": "MULTI_RECORD",
  "MaxPayloadInMB": 6
}
```

**How Your Container Should Respond to Inference Requests**

To obtain inferences, Amazon SageMaker sends a POST request to the inference container. The POST request body contains data from Amazon S3. Amazon SageMaker passes the request to the
container, and returns the inference result from the container, saving the data from the response to Amazon S3.

To receive inference requests, the container must have a web server listening on port 8080 and must accept POST requests to the /invocations endpoint. The inference request timeout and max retries can be configured through ModelClientConfig.

How Your Container Should Respond to Health Check (Ping) Requests

The simplest requirement on the container is to respond with an HTTP 200 status code and an empty body. This indicates to SageMaker that the container is ready to accept inference requests at the /invocations endpoint.

While the minimum bar is for the container to return a static 200, a container developer can use this functionality to perform deeper checks. The request timeout on /ping attempts is 2 seconds.

Examples and More Information: Use Your Own Algorithm or Model

The following Jupyter notebooks and added information show how to use your own algorithms or pretrained models from an Amazon SageMaker notebook instance. For links to the GitHub repositories with the prebuilt Dockerfiles for the TensorFlow, MXNet, Chainer, and PyTorch frameworks and instructions on using the AWS SDK for Python (Boto3) estimators to run your own training algorithms on SageMaker Learner and your own models on SageMaker hosting, see Prebuilt SageMaker Docker Images for Deep Learning

Setup

1. Create a SageMaker notebook instance. For instructions on how to create and access Jupyter notebook instances, see Amazon SageMaker Notebook Instances.

2. Open the notebook instance you created.

3. Choose the SageMaker Examples tab for a list of all SageMaker example notebooks.

4. Open the sample notebooks from the Advanced Functionality section in your notebook instance or from GitHub using the provided links. To open a notebook, choose its Use tab, then choose Create copy.
Host models trained in Scikit-learn

To learn how to host models trained in Scikit-learn for making predictions in SageMaker by injecting them into first-party k-means and XGBoost containers, see the following sample notebooks.

- kmeansBringYourOwnModel
- xgboostBringYourOwnModel

Package TensorFlow and Scikit-learn models for use in SageMaker

To learn how to package algorithms that you have developed in TensorFlow and scikit-learn frameworks for training and deployment in the SageMaker environment, see the following notebooks. They show you how to build, register, and deploy your own Docker containers using Dockerfiles.

- tensorflowBringYourOwn
- scikitBringYourOwn

Train and deploy a neural network on SageMaker

To learn how to train a neural network locally using MXNet or TensorFlow, and then create an endpoint from the trained model and deploy it on SageMaker, see the following notebooks. The MXNet model is trained to recognize handwritten numbers from the MNIST dataset. The TensorFlow model is trained to classify irises.

- mxnet_mnist_byom
- tensorflow_BYOM_iris

Training using pipe mode

To learn how to use a Dockerfile to build a container that calls the train.py script and uses pipe mode to custom train an algorithm, see the following notebook. In pipe mode, the input data is transferred to the algorithm while it is training. This can decrease training time compared to using file mode.

- pipeBringYourOwn
Bring your own R model

To learn how to use add a custom R image to build and train a model in a AWS SMS notebook, see the following blog post. This blog post uses a sample R Dockerfile from a library of SageMaker Studio Classic Custom Image Samples.

- Bringing your own R environment to Amazon SageMaker Studio Classic

Extend a pre-built PyTorch container Image

To learn how to extend a prebuilt SageMaker PyTorch container image when you have additional functional requirements for your algorithm or model that the prebuilt Docker image doesn't support, see the following notebook.

- BERTtopic_extending_container

For more information about extending a container, see Extend a Pre-built Container.

Train and debug training jobs on a custom container

To learn how to train and debug training jobs using SageMaker Debugger, see the following notebook. A training script provided through this example uses the TensorFlow Keras ResNet 50 model and the CIFAR10 dataset. A Docker custom container is built with the training script and pushed to Amazon ECR. While the training job is running, Debugger collects tensor outputs and identifies debugging problems. With smdebug client library tools, you can set a smdebug trial object that calls the training job and debugging information, check the training and Debugger rule status, and retrieve tensors saved in an Amazon S3 bucket to analyze training issues.

- build_your_own_container_with_debugger

Troubleshooting your Docker containers

The following are common errors that you might run into when using Docker containers with SageMaker. Each error is followed by a solution to the error.

- Error: SageMaker has lost the Docker daemon.
  
  To fix this error, restart Docker using the following command.
sudo service docker restart

- **Error: The /tmp directory of your Docker container has run out of space.**

  Docker containers use the / and /tmp partitions to store code. These partitions can fill up easily when using large code modules in local mode. The SageMaker Python SDK supports specifying a custom temp directory for your local mode root directory to avoid this issue.

  To specify the custom temp directory in the Amazon Elastic Block Store volume storage, create a file at the following path ~/.sagemaker/config.yaml and add the following configuration. The directory that you specify as container_root must already exist. The SageMaker Python SDK will not try to create it.

  ```yaml
  local:
      container_root: /home/ec2-user/SageMaker/temp
  ```

  With this configuration, local mode uses the /temp directory and not the default /tmp directory.

- **Low space errors on SageMaker notebook instances**

  A Docker container that runs on SageMaker notebook instances uses the root Amazon EBS volume of the notebook instance by default. To resolve low space errors, provide the path of the Amazon EBS volume attached to the notebook instance as part of the volume parameter of Docker commands.

  ```bash
docker run -v EBS-volume-path:container-path
  ```
Configure security in Amazon SageMaker

Cloud security at AWS is the highest priority. As an AWS customer, you benefit from a data center and network architecture that is built to meet the requirements of the most security-sensitive organizations.

Security is a shared responsibility between AWS and you. The shared responsibility model describes this as security of the cloud and security in the cloud:

- **Security of the cloud** – AWS is responsible for protecting the infrastructure that runs AWS services in the AWS Cloud. AWS also provides you with services that you can use securely. Third-party auditors regularly test and verify the effectiveness of our security as part of the AWS compliance programs. To learn about the compliance programs that apply to Amazon SageMaker, see AWS Services in Scope by Compliance Program.

- **Security in the cloud** – Your responsibility is determined by the AWS service that you use. You are also responsible for other factors including the sensitivity of your data, your company’s requirements, and applicable laws and regulations.

This documentation helps you understand how to apply the shared responsibility model when using SageMaker. The following topics show you how to configure SageMaker to meet your security and compliance objectives. You also learn how to use other AWS services that help you to monitor and secure your SageMaker resources.

**Topics**

- Data Privacy in Amazon SageMaker
- Data Protection in Amazon SageMaker
- Identity and Access Management for Amazon SageMaker
- Logging and Monitoring
- Compliance validation for Amazon SageMaker
- Resilience in Amazon SageMaker
- Infrastructure Security in Amazon SageMaker
Data Privacy in Amazon SageMaker

Amazon SageMaker collects aggregate information about the use of AWS-owned and open source libraries used during training. SageMaker uses this aggregate metadata to improve services and customer experience.

The following sections provide explanations for the type of metadata that SageMaker collects and how to opt out of metadata collection.

Types of information collected

Usage Information

Metadata from AWS-owned and open source libraries that are used with SageMaker training, such as those used for distributed training, compilation, and quantization.

Errors

Errors from unexpected behavior including failures, crashes, cascades, and failures that result from interacting with the SageMaker training platform.

How to opt out of metadata collection

You can opt out of sharing aggregated metadata with SageMaker training when creating a training job using the CreateTrainingJob API. If you are using the console to create training jobs, metadata collection is disabled by default.

⚠️ Important

You must choose to opt out of metadata collection for each training job that you submit. You must also choose to opt out in an API call as shown in the following examples. You cannot choose to opt out inside a training script.

The following section shows how you can opt out of metadata collection using the AWS CLI, AWS SDK for Python (Boto3), or the SageMaker Python SDK.
Opt out of metadata collection using the AWS Command Line Interface (AWS CLI)

To opt out of metadata collection using the AWS CLI, set the environment variable `OPT_OUT_TRACKING` to 1 in the `create-training-job` API as shown in the following code example.

```
aws sagemaker create-training-job \
  --training-job-name your_job_name \
  --algorithm-specification AlgorithmName=your_algorithm_name\ 
  --output-data-config S3OutputPath=s3://bucket-name/key-name-prefix \ 
  --resource-config InstanceType=ml.c5.xlarge, InstanceCount=1 \ 
  --stopping-condition MaxRuntimeInSeconds=100 \ 
  --environment OPT_OUT_TRACKING=1
```

Opt out of metadata collection using the AWS SDK for Python (Boto3)

To opt out of metadata collection using the SDK for Python (Boto3), set the environment variable `OPT_OUT_TRACKING` to 1 in the `create_training_job` API as shown in the following code example.

```
boto3.client('sagemaker').create_training_job(
    TrainingJobName='your_training_job',
    AlgorithmSpecification={
        'AlgorithmName': 'your_algorithm_name',
        'TrainingInputMode': 'File',
    },
    RoleArn='your_arn',
    OutputDataConfig={
        'S3OutputPath': 's3://bucket-name/key-name-prefix',
    },
    ResourceConfig={
        'InstanceType': 'ml.m4.xlarge',
        'InstanceCount': 1,
        'VolumeSizeInGB': 123,
    },
    StoppingCondition={
        'MaxRuntimeIn Seconds': 123,
    },
    Environment={
        'OPT_OUT_TRACKING': '1',
    },
)
```
Opt out of metadata collection using the SageMaker Python SDK

To opt out of metadata collection using the SageMaker Python SDK, set the environment variable OPT_OUT_TRACKING to 1 inside a SageMaker estimator as shown in the following code example.

```python
sagemaker.estimator(
    image_uri='path_to_container',
    role='rolearn',
    instance_count=1,
    instance_type='ml.c5.xlarge',
    environment=
    {
        'OPT_OUT_TRACKING': '1'
    },
)
```

Opt out of metadata collection account-wide

If you want to opt-out of metadata collection for several accounts, you can set an environment variable to opt-out of tracking account-wide. You must use the SageMaker Python SDK to opt out of metadata collection at an account level.

The following code example shows how opt out of tracking account-wide.

```json
SchemaVersion: '1.0'
SageMaker:
    TrainingJob:
        Environment:
            'OPT_OUT_TRACKING': '1'
```

For more information about how to opt out of tracking account-wide, see Configuring and using defaults with the SageMaker Python SDK.

Additional information

If your downstream service depends on SageMaker training

If you operate a service that relies on SageMaker training, it is highly recommended that you inform your customer about aggregate metadata collection in the SageMaker Training platform and present them with the choice to opt out. Alternatively, you can opt out of metadata collection on behalf of your customer.

If you are a client or a customer of a service that uses SageMaker training
If you are a client or customer of a service that uses SageMaker training, use your preferred method in the previous section to opt out of metadata collection.

**Data Protection in Amazon SageMaker**

The AWS [shared responsibility model](#) applies to data protection in Amazon SageMaker. As described in this model, AWS is responsible for protecting the global infrastructure that runs all of the AWS Cloud. You are responsible for maintaining control over your content that is hosted on this infrastructure. You are also responsible for the security configuration and management tasks for the AWS services that you use. For more information about data privacy, see the [Data Privacy FAQ](#). For information about data protection in Europe, see the [AWS Shared Responsibility Model and GDPR](#) blog post on the [AWS Security Blog](#).

For data protection purposes, we recommend that you protect AWS account credentials and set up individual users with AWS IAM Identity Center or AWS Identity and Access Management (IAM). That way, each user is given only the permissions necessary to fulfill their job duties. We also recommend that you secure your data in the following ways:

- Use multi-factor authentication (MFA) with each account.
- Use SSL/TLS to communicate with AWS resources. We require TLS 1.2 and recommend TLS 1.3.
- Set up API and user activity logging with AWS CloudTrail.
- Use AWS encryption solutions, along with all default security controls within AWS services.
- Use advanced managed security services such as Amazon Macie, which assists in discovering and securing sensitive data that is stored in Amazon S3.
- If you require FIPS 140-2 validated cryptographic modules when accessing AWS through a command line interface or an API, use a FIPS endpoint. For more information about the available FIPS endpoints, see [Federal Information Processing Standard (FIPS) 140-2](#).

We strongly recommend that you never put confidential or sensitive information, such as your customers' email addresses, into tags or free-form text fields such as a **Name** field. This includes when you work with Amazon SageMaker or other AWS services using the console, API, AWS CLI, or AWS SDKs. Any data that you enter into tags or free-form text fields used for names may be used for billing or diagnostic logs. If you provide a URL to an external server, we strongly recommend that you do not include credentials information in the URL to validate your request to that server.

**Topics**
Protect Data at Rest Using Encryption

To protect your Amazon SageMaker Studio notebooks and SageMaker notebook instances, along with your model-building data and model artifacts, SageMaker encrypts the notebooks, as well as output from Training and Batch Transform jobs. SageMaker encrypts these by default using the AWS Managed Key for Amazon S3. This AWS Managed Key for Amazon S3 cannot be shared for cross-account access. For cross-account access, specify your customer managed key while creating SageMaker resources so that it can be shared for cross-account access. For data output to Amazon S3 Express One Zone, the data is encrypted with server-side encryption with Amazon S3 managed keys (SSE-S3). For more information on AWS KMS, see What is AWS Key Management Service?

Topics

- Studio notebooks
- Notebook instances, SageMaker jobs, and Endpoints
- SageMaker geospatial capabilities

Studio notebooks

In Amazon SageMaker Studio, your SageMaker Studio notebooks and data can be stored in the following locations:

- An S3 bucket – When you onboard to Studio and enable shareable notebook resources, SageMaker shares notebook snapshots and metadata in an Amazon Simple Storage Service (Amazon S3) bucket.
- An EFS volume – When you onboard to Studio, SageMaker attaches an Amazon Elastic File System (Amazon EFS) volume to your domain for storing your Studio notebooks and data files. The EFS volume persists after the domain is deleted.
- An EBS volume – When you open a notebook in Studio, an Amazon Elastic Block Store (Amazon EBS) is attached to the instance that the notebook runs on. The EBS volume persists for the duration of the instance.
SageMaker uses the AWS Key Management Service (AWS KMS) to encrypt the S3 bucket and both volumes. By default, it uses a KMS key managed in an AWS service account. For more control, you can specify your own customer managed key when you onboard to Studio or through the SageMaker API. For more information, see Amazon SageMaker domain overview and CreateDomain.

In the CreateDomain API, you use the S3KmsKeyId parameter to specify the customer managed key for shareable notebooks. You use the KmsKeyId parameter to specify the customer managed key for the EFS and EBS volumes. The same customer managed key is used for both volumes. The customer managed key for shareable notebooks can be the same customer managed key as used for the volumes or a different customer managed key.

**Notebook instances, SageMaker jobs, and Endpoints**

To encrypt the machine learning (ML) storage volume that is attached to notebooks, processing jobs, training jobs, hyperparameter tuning jobs, batch transform jobs, and endpoints, you can pass a AWS KMS key to SageMaker. If you don't specify a KMS key, SageMaker encrypts storage volumes with a transient key and discards it immediately after encrypting the storage volume. For notebook instances, if you don't specify a KMS key, SageMaker encrypts both OS volumes and ML data volumes with a system-managed KMS key.

You can use an AWS managed AWS KMS key to encrypt all instance OS volumes. You can encrypt all ML data volumes for all SageMaker instances with a AWS KMS key that you specify. ML storage volumes are mounted as follows:

- **Notebooks** - `/home/ec2-user/SageMaker`
- **Processing** - `/opt/ml/processing` and `/tmp/`
- **Training** - `/opt/ml/` and `/tmp/`
- **Batch** - `/opt/ml/` and `/tmp/`
- **Endpoints** - `/opt/ml/` and `/tmp/`

Processing, batch transform, and training job containers and their storage are ephemeral in nature. When the job completes, output is uploaded to Amazon S3 using AWS KMS encryption with an optional AWS KMS key that you specify and the instance is torn down. If an AWS KMS Key is not provided in the job request, SageMaker uses the default AWS KMS key for Amazon S3 for your role's account. If the output data is stored in Amazon S3 Express One Zone, it is encrypted with server-side encryption with Amazon S3 managed keys (SSE-S3).
Note

The key policy for an AWS Managed Key for Amazon S3 cannot be edited, so cross-account permissions cannot be granted for these key policies. If the output Amazon S3 bucket for the request is from another account, specify your own AWS KMS Customer Key in the job request and ensure that the job’s execution role has permissions to encrypt data with it.

Important

Sensitive data that needs to be encrypted with a KMS key for compliance reasons should be stored in the ML storage volume or in Amazon S3, both of which can be encrypted using a KMS key you specify.

When you open a notebook instance, SageMaker saves it and any files associated with it in the SageMaker folder in the ML storage volume by default. When you stop a notebook instance, SageMaker creates a snapshot of the ML storage volume. Any customizations to the operating system of the stopped instance, such as installed custom libraries or operating system level settings, are lost. Consider using a lifecycle configuration to automate customizations of the default notebook instance. When you terminate an instance, the snapshot and the ML storage volume are deleted. Any data that you need to persist beyond the lifespan of the notebook instance should be transferred to an Amazon S3 bucket.

Note

Certain Nitro-based SageMaker instances include local storage, depending on the instance type. Local storage volumes are encrypted using a hardware module on the instance. You can’t use a KMS key on an instance type with local storage. For a list of instance types that support local instance storage, see Instance Store Volumes. For more information about storage volumes on Nitro-based instances, see Amazon EBS and NVMe on Linux Instances. For more information about local instance storage encryption, see SSD Instance Store Volumes.

SageMaker geospatial capabilities

You can protect your data at rest using encryption for SageMaker geospatial.
Server-Side Encryption with Amazon SageMaker geospatial owned key (Default)

Amazon SageMaker geospatial capabilities encrypts all your data, including computational results from your EarthObservationJobs and VectorEnrichmentJobs along with all your service metadata. There is no data that is stored within Amazon SageMaker unencrypted. It uses a default AWS owned key to encrypt all your data.

Server-Side Encryption with KMS Keys Stored in AWS Key Management Service (SSE-KMS)

Amazon SageMaker geospatial capabilities supports encryption using a customer-owned KMS key. For more information, see Use AWS KMS Permissions for Amazon SageMaker geospatial capabilities.

Protecting Data in Transit with Encryption

All internetwork data in transit supports TLS 1.2 encryption. We recommend that you use TLS 1.3.

With Amazon SageMaker, machine learning (ML) model artifacts and other system artifacts are encrypted in transit and at rest. Requests to the SageMaker API and console are made over a secure (SSL) connection. You pass AWS Identity and Access Management roles to SageMaker to provide permissions to access resources on your behalf for training and deployment.

Some intranetwork data in transit (inside the service platform) is unencrypted. This includes:

- Command and control communications between the service control plane and training job instances (not customer data).
- Communications between nodes in distributed processing jobs (intranetwork).
- Communications between nodes in distributed training jobs (intranetwork).

There are no inter-node communications for batch processing.

You can choose to encrypt communication between nodes in a training cluster.

Note

For use cases in the healthcare sector, the best practice for security is to encrypt communication between the nodes.
For information about how to encrypt communication, see the next topic at Protect Communications Between ML Compute Instances in a Distributed Training Job.

**Note**

Encrypting inter-container traffic can increase training time, especially if you use distributed deep learning algorithms. For affected algorithms, this additional level of security also increases cost. The training time for most SageMaker built-in algorithms, such as XGBoost, DeepAR, and linear learner, typically aren't affected.

FIPS validated endpoints are available for the SageMaker API and request router for hosted models (runtime). For information about FIPS compliant endpoints, see Federal Information Processing Standard (FIPS) 140-2.

**Protect Communications with RStudio on Amazon SageMaker**

RStudio on Amazon SageMaker provides encryption for all communications that involve SageMaker components. However, the previous version did not support encryption between the RStudioServerPro and RSession apps.

RStudio released version 2022.02.2-485.pro2 in April 2022. This version supports encryption between RStudioServerPro and RSession apps to enable end-to-end encryption. The version upgrade, however, is not completely backward compatible. As a result, you must update all of your RStudioServerPro and RSession apps. For information about how to update your apps, see Upgrade the RStudio Version.

**Protect Communications Between ML Compute Instances in a Distributed Training Job**

By default, Amazon SageMaker runs training jobs in an Amazon Virtual Private Cloud (Amazon VPC) to help keep your data secure. You can add another level of security to protect your training containers and data by configuring a *private* VPC. Distributed ML frameworks and algorithms usually transmit information that is directly related to the model, such as weights, not the training dataset. When performing distributed training, you can further protect data that is transmitted between instances. This can help you to comply with regulatory requirements. To do this, use inter-container traffic encryption.
For use cases in the healthcare sector, the best practice for security is to encrypt communication between the nodes.

Enabling inter-container traffic encryption can increase training time, especially if you are using distributed deep learning algorithms. Enabling inter-container traffic encryption doesn't affect training jobs with a single compute instance. However, for training jobs with several compute instances, the effect on training time depends on the amount of communication between compute instances. For affected algorithms, adding this additional level of security also increases cost. The training time for most SageMaker built-in algorithms, such as XGBoost, DeepAR, and linear learner, typically aren't affected.

You can enable inter-container traffic encryption for training jobs or hyperparameter tuning jobs. You can use SageMaker APIs or console to enable inter-container traffic encryption.

For information about running training jobs in a private VPC, see Give SageMaker Training Jobs Access to Resources in Your Amazon VPC.

Enable Inter-container Traffic Encryption (API)

Before enabling inter-container traffic encryption on training or hyperparameter tuning jobs with APIs, add inbound and outbound rules to your private VPC's security group.

To enable inter-container traffic encryption (API)

1. Add the following inbound and outbound rules in the security group for your private VPC:

<table>
<thead>
<tr>
<th>Protocol</th>
<th>Port Range</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>UDP</td>
<td>500</td>
<td>Self Security Group ID</td>
</tr>
<tr>
<td>ESP</td>
<td>50</td>
<td>Self Security Group ID</td>
</tr>
</tbody>
</table>

2. When you send a request to the CreateTrainingJob or CreateHyperParameterTuningJob API, specify True for the EnableInterContainerTrafficEncryption parameter.
For the ESP 50 protocol, the AWS Security Group Console might display the port range as "All". However, Amazon EC2 ignores the specified port range because it is not applicable for the ESP 50 IP protocol.

Enable Inter-container Traffic Encryption (Console)

Enable Inter-container Traffic Encryption in a Training Job

To enable inter-container traffic encryption in a training job

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the navigation pane, choose Training, then choose Training jobs.
3. Choose Create training job.
4. Under Network, choose a VPC. You can use the default VPC or one that you have created.
5. Choose Enable inter-container traffic encryption.

After you enable inter-container traffic encryption, finish creating the training job. For more information, see Step 4: Train a Model.

Enable Inter-container Traffic Encryption in a Hyperparameter Tuning Job

To enable inter-container traffic encryption in a hyperparameter tuning job

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. In the navigation pane, choose Training, then choose Hyperparameter tuning jobs.
3. Choose Create hyperparameter tuning job.
4. Under Network, choose a VPC. You can use the default VPC or one that you have created.
5. Choose Enable inter-container traffic encryption.

After enabling inter-container traffic encryption, finish creating the hyperparameter tuning job. For more information, see Configure and Launch a Hyperparameter Tuning Job.
Key Management

Customers can specify AWS KMS keys, including bring your own keys (BYOK), to use for envelope encryption with Amazon S3 input/output buckets and machine learning (ML) Amazon EBS volumes. ML volumes for notebook instances and for processing, training, and hosted model Docker containers can be optionally encrypted by using AWS KMS customer-owned keys. All instance OS volumes are encrypted with an AWS-managed AWS KMS key.

Note

Certain Nitro-based instances include local storage, dependent on the instance type. Local storage volumes are encrypted using a hardware module on the instance. You can't request a VolumeKmsKeyId when using an instance type with local storage.

For a list of instance types that support local instance storage, see Instance Store Volumes.

For more information about local instance storage encryption, see SSD Instance Store Volumes.

For more information about storage volumes on nitro-based instances, see Amazon EBS and NVMe on Linux Instances.

For information about AWS KMS keys see What is AWS Key Management Service? in the AWS Key Management Service Developer Guide.

Internetwork Traffic Privacy

This topic describes how Amazon SageMaker secures connections from the service to other locations.

Internetwork communications support TLS 1.2 encryption between all components and clients. We recommend TLS 1.3.

Instances can be connected to Customer VPC, providing access to S3 VPC endpoints or customer repositories. Internet egress can be managed through this interface by the customer if service platform internet egress is disabled for notebooks. For training and hosting, egress through the service platform is not available when connected to the customer's VPC.

By default, API calls made to published endpoints traverse the public network to the request router. SageMaker supports Amazon Virtual Private Cloud interface endpoints powered by AWS
PrivateLink for private connectivity between the customer's VPC and the request router to access hosted model endpoints. For information about Amazon VPC, see Connect to SageMaker Within your VPC

Identity and Access Management for Amazon SageMaker

AWS Identity and Access Management (IAM) is an AWS service that helps an administrator securely control access to AWS resources. IAM administrators control who can be authenticated (signed in) and authorized (have permissions) to use SageMaker resources. IAM is an AWS service that you can use with no additional charge.

Topics

- **Audience**
- **Authenticating with Identities**
- **Managing Access Using Policies**
- **How Amazon SageMaker Works with IAM**
- **Amazon SageMaker Identity-Based Policy Examples**
- **Cross-Service Confused Deputy Prevention**
- **SageMaker Roles**
- **Amazon SageMaker Role Manager**
- **Access control for notebooks**
- **Amazon SageMaker API Permissions: Actions, Permissions, and Resources Reference**
- **AWS Managed Policies for Amazon SageMaker**
- **Troubleshooting Amazon SageMaker Identity and Access**

**Audience**

How you use AWS Identity and Access Management (IAM) differs, depending on the work that you do in SageMaker.

**Service user** – If you use the SageMaker service to do your job, then your administrator provides you with the credentials and permissions that you need. As you use more SageMaker features to do your work, you might need additional permissions. Understanding how access is managed can
help you request the right permissions from your administrator. If you cannot access a feature in SageMaker, see Troubleshooting Amazon SageMaker Identity and Access.

**Service administrator** – If you're in charge of SageMaker resources at your company, you probably have full access to SageMaker. It's your job to determine which SageMaker features and resources your service users should access. You must then submit requests to your IAM administrator to change the permissions of your service users. Review the information on this page to understand the basic concepts of IAM. To learn more about how your company can use IAM with SageMaker, see How Amazon SageMaker Works with IAM.

**IAM administrator** – If you're an IAM administrator, you might want to learn details about how you can write policies to manage access to SageMaker. To view example SageMaker identity-based policies that you can use in IAM, see Amazon SageMaker Identity-Based Policy Examples.

**Authenticating with Identities**

Authentication is how you sign in to AWS using your identity credentials. You must be *authenticated* (signed in to AWS) as the AWS account root user, as an IAM user, or by assuming an IAM role.

You can sign in to AWS as a federated identity by using credentials provided through an identity source. AWS IAM Identity Center (IAM Identity Center) users, your company's single sign-on authentication, and your Google or Facebook credentials are examples of federated identities. When you sign in as a federated identity, your administrator previously set up identity federation using IAM roles. When you access AWS by using federation, you are indirectly assuming a role.

Depending on the type of user you are, you can sign in to the AWS Management Console or the AWS access portal. For more information about signing in to AWS, see How to sign in to your AWS account in the AWS Sign-In User Guide.

If you access AWS programmatically, AWS provides a software development kit (SDK) and a command line interface (CLI) to cryptographically sign your requests by using your credentials. If you don't use AWS tools, you must sign requests yourself. For more information about using the recommended method to sign requests yourself, see Signing AWS API requests in the IAM User Guide.

Regardless of the authentication method that you use, you might be required to provide additional security information. For example, AWS recommends that you use multi-factor authentication (MFA) to increase the security of your account. To learn more, see Multi-factor authentication in the

AWS account root user

When you create an AWS account, you begin with one sign-in identity that has complete access to all AWS services and resources in the account. This identity is called the AWS account root user and is accessed by signing in with the email address and password that you used to create the account. We strongly recommend that you don't use the root user for your everyday tasks. Safeguard your root user credentials and use them to perform the tasks that only the root user can perform. For the complete list of tasks that require you to sign in as the root user, see Tasks that require root user credentials in the IAM User Guide.

Federated identity

As a best practice, require human users, including users that require administrator access, to use federation with an identity provider to access AWS services by using temporary credentials.

A federated identity is a user from your enterprise user directory, a web identity provider, the AWS Directory Service, the Identity Center directory, or any user that accesses AWS services by using credentials provided through an identity source. When federated identities access AWS accounts, they assume roles, and the roles provide temporary credentials.

For centralized access management, we recommend that you use AWS IAM Identity Center. You can create users and groups in IAM Identity Center, or you can connect and synchronize to a set of users and groups in your own identity source for use across all your AWS accounts and applications. For information about IAM Identity Center, see What is IAM Identity Center? in the AWS IAM Identity Center User Guide.

IAM Users and Groups

An IAM user is an identity within your AWS account that has specific permissions for a single person or application. Where possible, we recommend relying on temporary credentials instead of creating IAM users who have long-term credentials such as passwords and access keys. However, if you have specific use cases that require long-term credentials with IAM users, we recommend that you rotate access keys. For more information, see Rotate access keys regularly for use cases that require long-term credentials in the IAM User Guide.

An IAM group is an identity that specifies a collection of IAM users. You can't sign in as a group. You can use groups to specify permissions for multiple users at a time. Groups make permissions easier
to manage for large sets of users. For example, you could have a group named `IAMAdmins` and give that group permissions to administer IAM resources.

Users are different from roles. A user is uniquely associated with one person or application, but a role is intended to be assumable by anyone who needs it. Users have permanent long-term credentials, but roles provide temporary credentials. To learn more, see [When to create an IAM user (instead of a role)] in the **IAM User Guide**.

**IAM Roles**

An *IAM role* is an identity within your AWS account that has specific permissions. It is similar to an IAM user, but is not associated with a specific person. You can temporarily assume an IAM role in the AWS Management Console by *switching roles*. You can assume a role by calling an AWS CLI or AWS API operation or by using a custom URL. For more information about methods for using roles, see [Using IAM roles](#) in the **IAM User Guide**.

IAM roles with temporary credentials are useful in the following situations:

- **Federated user access** – To assign permissions to a federated identity, you create a role and define permissions for the role. When a federated identity authenticates, the identity is associated with the role and is granted the permissions that are defined by the role. For information about roles for federation, see [Creating a role for a third-party Identity Provider](#) in the **IAM User Guide**. If you use IAM Identity Center, you configure a permission set. To control what your identities can access after they authenticate, IAM Identity Center correlates the permission set to a role in IAM. For information about permissions sets, see [Permission sets](#) in the **AWS IAM Identity Center User Guide**.

- **Temporary IAM user permissions** – An IAM user or role can assume an IAM role to temporarily take on different permissions for a specific task.

- **Cross-account access** – You can use an IAM role to allow someone (a trusted principal) in a different account to access resources in your account. Roles are the primary way to grant cross-account access. However, with some AWS services, you can attach a policy directly to a resource (instead of using a role as a proxy). To learn the difference between roles and resource-based policies for cross-account access, see [How IAM roles differ from resource-based policies](#) in the **IAM User Guide**.

- **Cross-service access** – Some AWS services use features in other AWS services. For example, when you make a call in a service, it's common for that service to run applications in Amazon EC2 or store objects in Amazon S3. A service might do this using the calling principal's permissions, using a service role, or using a service-linked role.
• **Forward access sessions (FAS)** – When you use an IAM user or role to perform actions in AWS, you are considered a principal. When you use some services, you might perform an action that then initiates another action in a different service. FAS uses the permissions of the principal calling an AWS service, combined with the requesting AWS service to make requests to downstream services. FAS requests are only made when a service receives a request that requires interactions with other AWS services or resources to complete. In this case, you must have permissions to perform both actions. For policy details when making FAS requests, see [Forward access sessions](https://docs.aws.amazon.com/IAM/latest/userguide/forward-access.html).

• **Service role** – A service role is an IAM role that a service assumes to perform actions on your behalf. An IAM administrator can create, modify, and delete a service role from within IAM. For more information, see [Creating a role to delegate permissions to an AWS service](https://docs.aws.amazon.com/IAM/latest/userguide/iam-create-service-linked-role.html) in the IAM User Guide.

• **Service-linked role** – A service-linked role is a type of service role that is linked to an AWS service. The service can assume the role to perform an action on your behalf. Service-linked roles appear in your AWS account and are owned by the service. An IAM administrator can view, but not edit the permissions for service-linked roles.

• **Applications running on Amazon EC2** – You can use an IAM role to manage temporary credentials for applications that are running on an EC2 instance and making AWS CLI or AWS API requests. This is preferable to storing access keys within the EC2 instance. To assign an AWS role to an EC2 instance and make it available to all of its applications, you create an instance profile that is attached to the instance. An instance profile contains the role and enables programs that are running on the EC2 instance to get temporary credentials. For more information, see [Using an IAM role to grant permissions to applications running on Amazon EC2 instances](https://docs.aws.amazon.com/IAM/latest/userguide/iam-create-instance-profile.html) in the IAM User Guide.

To learn whether to use IAM roles or IAM users, see [When to create an IAM role (instead of a user)](https://docs.aws.amazon.com/IAM/latest/userguide/changing-to-iam-roles.html) in the IAM User Guide.

### Managing Access Using Policies

You control access in AWS by creating policies and attaching them to AWS identities or resources. A policy is an object in AWS that, when associated with an identity or resource, defines their permissions. AWS evaluates these policies when a principal (user, root user, or role session) makes a request. Permissions in the policies determine whether the request is allowed or denied. Most policies are stored in AWS as JSON documents. For more information about the structure and contents of JSON policy documents, see [Overview of JSON policies](https://docs.aws.amazon.com/IAM/latest/userguide/policies-overview.html) in the IAM User Guide.
Administrators can use AWS JSON policies to specify who has access to what. That is, which **principal** can perform **actions** on what **resources**, and under what **conditions**.

By default, users and roles have no permissions. To grant users permission to perform actions on the resources that they need, an IAM administrator can create IAM policies. The administrator can then add the IAM policies to roles, and users can assume the roles.

IAM policies define permissions for an action regardless of the method that you use to perform the operation. For example, suppose that you have a policy that allows the `iam:GetRole` action. A user with that policy can get role information from the AWS Management Console, the AWS CLI, or the AWS API.

**Identity-Based Policies**

Identity-based policies are JSON permissions policy documents that you can attach to an identity, such as an IAM user, group of users, or role. These policies control what actions users and roles can perform, on which resources, and under what conditions. To learn how to create an identity-based policy, see [Creating IAM policies](https://docs.aws.amazon.com/iam/latest/userguide/create-policy.html) in the IAM User Guide.

Identity-based policies can be further categorized as **inline policies** or **managed policies**. Inline policies are embedded directly into a single user, group, or role. Managed policies are standalone policies that you can attach to multiple users, groups, and roles in your AWS account. Managed policies include AWS managed policies and customer managed policies. To learn how to choose between a managed policy or an inline policy, see [Choosing between managed policies and inline policies](https://docs.aws.amazon.com/iam/latest/userguide/choosing-between-managed-inline-policies.html) in the IAM User Guide.

**Resource-Based Policies**

Resource-based policies are JSON policy documents that you attach to a resource. Examples of resource-based policies are IAM **role trust policies** and Amazon S3 **bucket policies**. In services that support resource-based policies, service administrators can use them to control access to a specific resource. For the resource where the policy is attached, the policy defines what actions a specified principal can perform on that resource and under what conditions. You must **specify a principal** in a resource-based policy. Principals can include accounts, users, roles, federated users, or AWS services.

Resource-based policies are inline policies that are located in that service. You can't use AWS managed policies from IAM in a resource-based policy.
Access Control Lists (ACLs)

Access control lists (ACLs) control which principals (account members, users, or roles) have permissions to access a resource. ACLs are similar to resource-based policies, although they do not use the JSON policy document format.

Amazon S3, AWS WAF, and Amazon VPC are examples of services that support ACLs. To learn more about ACLs, see Access control list (ACL) overview in the Amazon Simple Storage Service Developer Guide.

Other Policy Types

AWS supports additional, less-common policy types. These policy types can set the maximum permissions granted to you by the more common policy types.

- **Permissions boundaries** – A permissions boundary is an advanced feature in which you set the maximum permissions that an identity-based policy can grant to an IAM entity (IAM user or role). You can set a permissions boundary for an entity. The resulting permissions are the intersection of an entity's identity-based policies and its permissions boundaries. Resource-based policies that specify the user or role in the Principal field are not limited by the permissions boundary. An explicit deny in any of these policies overrides the allow. For more information about permissions boundaries, see Permissions boundaries for IAM entities in the IAM User Guide.

- **Service control policies (SCPs)** – SCPs are JSON policies that specify the maximum permissions for an organization or organizational unit (OU) in AWS Organizations. AWS Organizations is a service for grouping and centrally managing multiple AWS accounts that your business owns. If you enable all features in an organization, then you can apply service control policies (SCPs) to any or all of your accounts. The SCP limits permissions for entities in member accounts, including each AWS account root user. For more information about Organizations and SCPs, see How SCPs work in the AWS Organizations User Guide.

- **Session policies** – Session policies are advanced policies that you pass as a parameter when you programmatically create a temporary session for a role or federated user. The resulting session's permissions are the intersection of the user or role's identity-based policies and the session policies. Permissions can also come from a resource-based policy. An explicit deny in any of these policies overrides the allow. For more information, see Session policies in the IAM User Guide.
Multiple Policy Types

When multiple types of policies apply to a request, the resulting permissions are more complicated to understand. To learn how AWS determines whether to allow a request when multiple policy types are involved, see Policy evaluation logic in the IAM User Guide.

How Amazon SageMaker Works with IAM

Before you use IAM to manage access to SageMaker, you should understand what IAM features are available to use with SageMaker. To get a high-level view of how SageMaker and other AWS services work with IAM, see AWS Services That Work with IAM in the IAM User Guide.

Topics

- SageMaker Identity-Based Policies

SageMaker Identity-Based Policies

With IAM identity-based policies, you can specify allowed or denied actions and resources as well as the conditions under which actions are allowed or denied. SageMaker supports specific actions, resources, and condition keys. To learn about all of the elements that you use in a JSON policy, see IAM JSON Policy Elements Reference in the IAM User Guide.

Actions

Administrators can use AWS JSON policies to specify who has access to what. That is, which principal can perform actions on what resources, and under what conditions.

The Action element of a JSON policy describes the actions that you can use to allow or deny access in a policy. Policy actions usually have the same name as the associated AWS API operation. There are some exceptions, such as permission-only actions that don't have a matching API operation. There are also some operations that require multiple actions in a policy. These additional actions are called dependent actions.

Include actions in a policy to grant permissions to perform the associated operation.

Policy actions in SageMaker use the following prefix before the action: sagemaker:. For example, to grant someone permission to run a SageMaker training job with the SageMaker CreateTrainingJob API operation, you include the sagemaker:CreateTrainingJob action in
their policy. Policy statements must include either an Action or NotAction element. SageMaker defines its own set of actions that describe tasks that you can perform with this service.

To specify multiple actions in a single statement, separate them with commas as follows:

```
"Action": [
    "sagemaker:action1",
    "sagemaker:action2"
]
```

You can specify multiple actions using wildcards (*). For example, to specify all actions that begin with the word Describe, include the following action:

```
"Action": "sagemaker:Describe*"
```

To see a list of SageMaker actions, see Actions, resources, and condition keys for Amazon SageMaker in the Service Authorization Reference.

Resources

SageMaker does not support specifying resource ARNs in a policy.

Condition Keys

Administrators can use AWS JSON policies to specify who has access to what. That is, which principal can perform actions on what resources, and under what conditions.

The Condition element (or Condition block) lets you specify conditions in which a statement is in effect. The Condition element is optional. You can create conditional expressions that use condition operators, such as equals or less than, to match the condition in the policy with values in the request.

If you specify multiple Condition elements in a statement, or multiple keys in a single Condition element, AWS evaluates them using a logical AND operation. If you specify multiple values for a single condition key, AWS evaluates the condition using a logical OR operation. All of the conditions must be met before the statement's permissions are granted.

You can also use placeholder variables when you specify conditions. For example, you can grant an IAM user permission to access a resource only if it is tagged with their IAM user name. For more information, see IAM policy elements: variables and tags in the IAM User Guide.


SageMaker supports a number of service-specific condition keys that you can use for fine-grained access control for the following operations:

- [CreateProcessingJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateProcessingJob.html)
- [CreateTrainingJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateTrainingJob.html)
- [CreateModel](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateModel.html)
- [CreateEndpointConfig](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateEndpointConfig.html)
- [CreateTransformJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateTransformJob.html)
- [CreateHyperParameterTuningJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateHyperParameterTuningJob.html)
- [CreateLabelingJob](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateLabelingJob.html)
- [CreateNotebookInstance](https://docs.aws.amazon.com/sagemaker/latest/dg/API_CreateNotebookInstance.html)
- [UpdateNotebookInstance](https://docs.aws.amazon.com/sagemaker/latest/dg/API_UpdateNotebookInstance.html)

To see a list of SageMaker condition keys, see [Condition keys for Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-iam-policy-reference.html) in the [IAM User Guide](https://docs.aws.amazon.com/IAM/latest/UserGuide/acl_condition-keys.html). To learn with which actions and resources you can use a condition key, see [Actions defined by Amazon SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-iam-policy-reference.html).

For examples of using SageMaker condition keys, see the following: [Control Creation of SageMaker Resources with Condition Keys](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-iam-policy-reference.html).

**Examples**

To view examples of SageMaker identity-based policies, see [Amazon SageMaker Identity-Based Policy Examples](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-iam-policy-reference.html).

**SageMaker Resource-Based Policies**

SageMaker does not support resource-based policies.
Authorization Based on SageMaker Tags

You can attach tags to SageMaker resources or pass tags in a request to SageMaker. To control access based on tags, you provide tag information in the condition element of a policy using the `sagemaker:ResourceTag/key-name`, `aws:RequestTag/key-name`, or `aws:TagKeys` condition keys. For more information about tagging SageMaker resources, see Control Access to SageMaker Resources by Using Tags.

To view an example identity-based policy for limiting access to a resource based on the tags on that resource, see Control Access to SageMaker Resources by Using Tags.

SageMaker IAM Roles

An IAM role is an entity within your AWS account that has specific permissions.

Using Temporary Credentials with SageMaker

You can use temporary credentials to sign in with federation, assume an IAM role, or to assume a cross-account role. You obtain temporary security credentials by calling AWS STS API operations such as `AssumeRole` or `GetFederationToken`.

SageMaker supports using temporary credentials.

Service-Linked Roles

SageMaker partially supports service-linked roles. Service-linked roles are currently available for SageMaker Studio Classic.

Service Roles

This feature allows a service to assume a service role on your behalf. This role allows the service to access resources in other services to complete an action on your behalf. Service roles appear in your IAM account and are owned by the account. This means that an IAM administrator can change the permissions for this role. However, doing so might break the functionality of the service.

SageMaker supports service roles.

Choosing an IAM Role in SageMaker

When you create a notebook instance, processing job, training job, hosted endpoint, or batch transform job resource in SageMaker, you must choose a role to allow SageMaker to access SageMaker on your behalf. If you have previously created a service role or service-linked role, then SageMaker provides you with a list of roles to choose from. It's important to choose a role that
allows access to the AWS operations and resources you need. For more information, see SageMaker Roles.

Amazon SageMaker Identity-Based Policy Examples

By default, IAM users and roles don't have permission to create or modify SageMaker resources. They also can't perform tasks using the AWS Management Console, AWS CLI, or AWS API. An IAM administrator must create IAM policies that grant users and roles permission to perform specific API operations on the specified resources they need. The administrator must then attach those policies to the IAM users or groups that require those permissions. To learn how to attach policies to an IAM user or group, see Adding and Removing IAM Identity Permissions in the IAM User Guide.

To learn how to create an IAM identity-based policy using these example JSON policy documents, see Creating Policies on the JSON Tab in the IAM User Guide.

Topics

- Policy Best Practices
- Using the SageMaker Console
- Allow Users to View Their Own Permissions
- Control Creation of SageMaker Resources with Condition Keys
- Control Access to the SageMaker API by Using Identity-based Policies
- Limit Access to SageMaker API and Runtime Calls by IP Address
- Limit Access to a Notebook Instance by IP Address
- Control Access to SageMaker Resources by Using Tags
- Provide Permissions for Tagging SageMaker Resources Upon Creation
- Limit Access to Searchable Resources with Visibility Conditions

Policy Best Practices

Identity-based policies determine whether someone can create, access, or delete SageMaker resources in your account. These actions can incur costs for your AWS account. When you create or edit identity-based policies, follow these guidelines and recommendations:

- Get started with AWS managed policies and move toward least-privilege permissions – To get started granting permissions to your users and workloads, use the AWS managed policies that grant permissions for many common use cases. They are available in your AWS account. We
recommend that you reduce permissions further by defining AWS customer managed policies that are specific to your use cases. For more information, see AWS managed policies or AWS managed policies for job functions in the IAM User Guide.

- **Apply least-privilege permissions** – When you set permissions with IAM policies, grant only the permissions required to perform a task. You do this by defining the actions that can be taken on specific resources under specific conditions, also known as least-privilege permissions. For more information about using IAM to apply permissions, see Policies and permissions in IAM in the IAM User Guide.

- **Use conditions in IAM policies to further restrict access** – You can add a condition to your policies to limit access to actions and resources. For example, you can write a policy condition to specify that all requests must be sent using SSL. You can also use conditions to grant access to service actions if they are used through a specific AWS service, such as AWS CloudFormation. For more information, see IAM JSON policy elements: Condition in the IAM User Guide.

- **Use IAM Access Analyzer to validate your IAM policies to ensure secure and functional permissions** – IAM Access Analyzer validates new and existing policies so that the policies adhere to the IAM policy language (JSON) and IAM best practices. IAM Access Analyzer provides more than 100 policy checks and actionable recommendations to help you author secure and functional policies. For more information, see IAM Access Analyzer policy validation in the IAM User Guide.

- **Require multi-factor authentication (MFA)** – If you have a scenario that requires IAM users or a root user in your AWS account, turn on MFA for additional security. To require MFA when API operations are called, add MFA conditions to your policies. For more information, see Configuring MFA-protected API access in the IAM User Guide.

For more information about best practices in IAM, see Security best practices in IAM in the IAM User Guide.

### Using the SageMaker Console

To access the Amazon SageMaker console, you must have a minimum set of permissions. These permissions must allow you to list and view details about the SageMaker resources in your AWS account. If you create an identity-based policy that is more restrictive than the minimum required permissions, the console won't function as intended for entities (users or roles) with that policy.

To ensure that those entities can still use the SageMaker console, also attach the following AWS managed policy to the entities. For more information, see Adding Permissions to a User in the IAM User Guide:
You don't need to allow minimum console permissions for users that are making calls only to the AWS CLI or the AWS API. Instead, allow access to only the actions that match the API operation that you're trying to perform.

**Topics**

- Permissions Required to Use the Amazon SageMaker Console
- Permissions Required to Use the Amazon SageMaker Ground Truth Console
- Permissions Required to Use the Amazon Augmented AI (Preview) Console

**Permissions Required to Use the Amazon SageMaker Console**

The permissions reference table lists the Amazon SageMaker API operations and shows the required permissions for each operation. For more information about Amazon SageMaker API operations, see [Amazon SageMaker API Permissions: Actions, Permissions, and Resources Reference](#).

To use the Amazon SageMaker console, you need to grant permissions for additional actions. Specifically, the console needs permissions that allow the `ec2` actions to display subnets, VPCs, and security groups. Optionally, the console needs permission to create execution roles for tasks such as `CreateNotebook`, `CreateTrainingJob`, and `CreateModel`. Grant these permissions with the following permissions policy:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "SageMakerApis",
      "Effect": "Allow",
      "Action": ["sagemaker:*"],
      "Resource": "*"
    },
    {
      "Sid": "VpcConfigurationForCreateForms",
      "Effect": "Allow",
      "Action": ["ec2:DescribeVpcs",
                  "ec2:DescribeSubnets",
                  "ec2:DescribeSecurityGroups"
                ],
      "Resource": "*"
    }
  ]
}
```
Identity-Based Policy Examples

```
],
"Resource": "*
],
{
  "Sid":"KmsKeysForCreateForms",
  "Effect": "Allow",
  "Action": [
    "kms:DescribeKey",
    "kms:ListAliases"
  ],
  "Resource": "*
],
{
  "Sid": "AccessAwsMarketplaceSubscriptions",
  "Effect": "Allow",
  "Action": [
    "aws-marketplace:ViewSubscriptions"
  ],
  "Resource": "*
],
{
  "Effect": "Allow",
  "Action": [
    "codecommit:BatchGetRepositories",
    "codecommit:CreateRepository",
    "codecommit:GetRepository",
    "codecommit:ListRepositories",
    "codecommit:ListBranches",
    "secretsmanager:CreateSecret",
    "secretsmanager:DescribeSecret",
    "secretsmanager:ListSecrets"
  ],
  "Resource": "*
],
{
  "Sid": "ListAndCreateExecutionRoles",
  "Effect": "Allow",
  "Action": [
    "iam:ListRoles",
    "iam:CreateRole",
    "iam:CreatePolicy",
    "iam:AttachRolePolicy"
  ],
  "Resource": "*
}
Permissions Required to Use the Amazon SageMaker Ground Truth Console

To use the Amazon SageMaker Ground Truth console, you need to grant permissions for additional resources. Specifically, the console needs permissions for the AWS Marketplace to view subscriptions, Amazon Cognito operations to manage your private workforce, Amazon S3 actions for access to your input and output files, and AWS Lambda actions to list and invoke functions. Grant these permissions with the following permissions policy:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "GroundTruthConsole",
            "Effect": "Allow",
            "Action": [
                "aws-marketplace:DescribeListings",
                "aws-marketplace:ViewSubscriptions",
                "iam:PassRole"
            ],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "iam:PassedToService": "sagemaker.amazonaws.com"
                }
            }
        }
    ]
}
```
Permissions Required to Use the Amazon Augmented AI (Preview) Console

To use the Augmented AI console, you need to grant permissions for additional resources. Grant these permissions with the following permissions policy:
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:*Algorithm",
                "sagemaker:*Algorithms",
                "sagemaker:*App",
                "sagemaker:*Apps",
                "sagemaker:*AutoMLJob",
                "sagemaker:*AutoMLJobs",
                "sagemaker:*CodeRepositories",
                "sagemaker:*CodeRepository",
                "sagemaker:*CompilationJob",
                "sagemaker:*CompilationJobs",
                "sagemaker:*Endpoint",
                "sagemaker:*EndpointConfig",
                "sagemaker:*EndpointConfigs",
                "sagemaker:*EndpointWeightsAndCapacities",
                "sagemaker:*Endpoints",
                "sagemaker:*Environment",
                "sagemaker:*EnvironmentVersion",
                "sagemaker:*EnvironmentVersions",
                "sagemaker:*Environments",
                "sagemaker:*Experiment",
                "sagemaker:*Experiments",
                "sagemaker:*FlowDefinitions",
                "sagemaker:*HumanLoop",
                "sagemaker:*HumanLoops",
                "sagemaker:*HumanTaskUi",
                "sagemaker:*HumanTaskUis",
                "sagemaker:*HyperParameterTuningJob",
                "sagemaker:*HyperParameterTuningJobs",
                "sagemaker:*LabelingJob",
                "sagemaker:*LabelingJobs",
                "sagemaker:*Metrics",
                "sagemaker:*Model",
                "sagemaker:*ModelPackage",
                "sagemaker:*ModelPackages",
                "sagemaker:*Models",
                "sagemaker:*MonitoringExecutions",
                "sagemaker:*MonitoringSchedule",
            ]
        }
    ]
}
"sagemaker:*MonitoringSchedules",
"sagemaker:*NotebookInstance",
"sagemaker:*NotebookInstanceLifecycleConfig",
"sagemaker:*NotebookInstanceLifecycleConfigs",
"sagemaker:*NotebookInstanceUrl",
"sagemaker:*NotebookInstances",
"sagemaker:*ProcessingJob",
"sagemaker:*ProcessingJobs",
"sagemaker:*RenderUiTemplate",
"sagemaker:*Search",
"sagemaker:*SearchSuggestions",
"sagemaker:*Tags",
"sagemaker:*TrainingJob",
"sagemaker:*TrainingJobs",
"sagemaker:*TransformJob",
"sagemaker:*TransformJobs",
"sagemaker:*Trial",
"sagemaker:*TrialComponent",
"sagemaker:*TrialComponents",
"sagemaker:*Trials",
"sagemaker:*Workteam",
"sagemaker:*Workteams"
],
"Resource": "*
",
{
  "Effect": "Allow",
  "Action": [
    "sagemaker:*FlowDefinition"
  ],
  "Resource": "*
",
  "Condition": {
    "StringEqualsIfExists": {
      "sagemaker:WorkteamType": [
        "private-crowd",
        "vendor-crowd"
      ]
    }
  }
},
{
  "Effect": "Allow",
  "Action": [
    "application-autoscaling:DeleteScalingPolicy",
  ]
}
"application-autoscaling:DeleteScheduledAction",
"application-autoscaling:DeregisterScalableTarget",
"application-autoscaling:DescribeScalableTargets",
"application-autoscaling:DescribeScalingActivities",
"application-autoscaling:DescribeScalingPolicies",
"application-autoscaling:DescribeScheduledActions",
"application-autoscaling:PutScalingPolicy",
"application-autoscaling:PutScheduledAction",
"application-autoscaling:RegisterScalableTarget",
"aws-marketplace:ViewSubscriptions",
"cloudwatch:DeleteAlarms",
"cloudwatch:DescribeAlarms",
"cloudwatch:GetMetricData",
"cloudwatch:GetMetricStatistics",
"cloudwatch:ListMetrics",
"cloudwatch:PutMetricAlarm",
"cloudwatch:PutMetricData",
"codecommit:BatchGetRepositories",
"codecommit:CreateRepository",
"codecommit:GetRepository",
"codecommit:ListBranches",
"codecommit:ListRepositories",
"cognito-idp:AdminAddUserToGroup",
"cognito-idp:AdminCreateUser",
"cognito-idp:AdminDeleteUser",
"cognito-idp:AdminDisableUser",
"cognito-idp:AdminEnableUser",
"cognito-idp:AdminRemoveUserFromGroup",
"cognito-idp:CreateGroup",
"cognito-idp:CreateUserPool",
"cognito-idp:CreateUserPoolClient",
"cognito-idp:CreateUserPoolDomain",
"cognito-idp:DescribeUserPool",
"cognito-idp:DescribeUserPoolClient",
"cognito-idp:ListGroups",
"cognito-idp:ListIdentityProviders",
"cognito-idp:ListUserPoolClients",
"cognito-idp:ListUserPools",
"cognito-idp:ListUsers",
"cognito-idp:ListUsersInGroup",
"cognito-idp:UpdateUserPool",
"cognito-idp:UpdateUserPoolClient",
"ec2:CreateNetworkInterface",
"ec2:CreateNetworkInterfacePermission",
"ec2:CreateVpcEndpoint",
"ec2:DeleteNetworkInterface",
"ec2:DeleteNetworkInterfacePermission",
"ec2:DescribeDhcpOptions",
"ec2:DescribeNetworkInterfaces",
"ec2:DescribeRouteTables",
"ec2:DescribeSecurityGroups",
"ec2:DescribeSubnets",
"ec2:DescribeVpcEndpoints",
"ec2:DescribeVpcs",
"ecr:BatchCheckLayerAvailability",
"ecr:BatchGetImage",
"ecr:CreateRepository",
"ecr:Describe*",
"ecr:GetAuthorizationToken",
"ecr:GetDownloadUrlForLayer",
"elastic-inference:Connect",
"elasticfilesystem:DescribeFileSystems",
"elasticfilesystem:DescribeMountTargets",
"fsx:DescribeFileSystems",
"glue:CreateJob",
"glue:DeleteJob",
"glue:GetJob",
"glue:GetJobRun",
"glue:GetJobRuns",
"glue:GetJobs",
"glue:ResetJobBookmark",
"glue:StartJobRun",
"glue:UpdateJob",
"groundtruthlabeling:*",
"iam:ListRoles",
"kms:DescribeKey",
"kms:ListAliases",
"lambda:ListFunctions",
"logs:CreateLogGroup",
"logs:CreateLogStream",
"logs:DescribeLogGroups",
"logs:DescribeLogStreams",
"logs:GetLogEvents",
"logs:PutLogEvents",
"sns:ListTopics"
],
"Resource": "*
}
["Effect": "Allow",
"Action": [
"logs:CreateLogDelivery",
"logs:DeleteLogDelivery",
"logs:DescribeResourcePolicies",
"logs:GetLogDelivery",
"logs:ListLogDeliveries",
"logs:PutResourcePolicy",
"logs:UpdateLogDelivery"
],
"Resource": "*"]
},

["Effect": "Allow",
"Action": [
"ecr:SetRepositoryPolicy",
"ecr:CompleteLayerUpload",
"ecr:BatchDeleteImage",
"ecr:UploadLayerPart",
"ecr:DeleteRepositoryPolicy",
"ecr:InitiateLayerUpload",
"ecr:DeleteRepository",
"ecr:PutImage"
],
"Resource": "arn:aws:ecr:*:*:repository/*sagemaker*"
],

["Effect": "Allow",
"Action": [
"codecommit:GitPull",
"codecommit:GitPush"
],
"Resource": [
"arn:aws:codecommit:*:*:*sagemaker*",
"arn:aws:codecommit:*:*:*SageMaker*",
"arn:aws:codecommit:*:*:*Sagemaker*"
]
},

["Effect": "Allow",
"Action": [
"secretsmanager:ListSecrets"
],}
Identity-Based Policy Examples

```
"Resource": "*",
{
  "Effect": "Allow",
  "Action": [
    "secretsmanager:DescribeSecret",
    "secretsmanager:GetSecretValue",
    "secretsmanager:CreateSecret"
  ],
  "Resource": [
    "arn:aws:secretsmanager:*:*:secret:AmazonSageMaker-*"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "secretsmanager:DescribeSecret",
    "secretsmanager:GetSecretValue"
  ],
  "Resource": "*",
  "Condition": {
    "StringEquals": {
      "secretsmanager:ResourceTag/SageMaker": "true"
    }
  }
},
{
  "Effect": "Allow",
  "Action": [
    "robomaker:CreateSimulationApplication",
    "robomaker:DescribeSimulationApplication",
    "robomaker:DeleteSimulationApplication"
  ],
  "Resource": [
    "*"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "robomaker:CreateSimulationJob",
    "robomaker:DescribeSimulationJob",
    "robomaker:CancelSimulationJob"
  ],
```
Identity-Based Policy Examples

```json

"Resource": [
  "*
]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:GetObject",
    "s3:PutObject",
    "s3:DeleteObject",
    "s3:AbortMultipartUpload",
    "s3:GetBucketCors",
    "s3:PutBucketCors"
  ],
  "Resource": [
    "arn:aws:s3:::*SageMaker*",
    "arn:aws:s3:::*Sagemaker*",
    "arn:aws:s3:::*sagemaker*",
    "arn:aws:s3:::*aws-glue*"
  ],
},
{
  "Effect": "Allow",
  "Action": [
    "s3:CreateBucket",
    "s3:GetBucketLocation",
    "s3:ListBucket",
    "s3:ListAllMyBuckets"
  ],
  "Resource": "*
},
{
  "Effect": "Allow",
  "Action": [
    "s3:GetObject"
  ],
  "Resource": "*
  "Condition": {
    "StringEqualsIgnoreCase": {
      "s3:ExistingObjectTag/SageMaker": "true"
    }
  }
},
{

```
"Effect": "Allow",
"Action": [
   "lambda:InvokeFunction"
],
"Resource": [
   "arn:aws:lambda:*:*:function:*SageMaker*",
   "arn:aws:lambda:*:*:function:*sagemaker*",
   "arn:aws:lambda:*:*:function:*Sagemaker*",
   "arn:aws:lambda:*:*:function:*LabelingFunction*"
]
},
{
   "Action": "iam:CreateServiceLinkedRole",
   "Effect": "Allow",
   "Resource": "arn:aws:iam::*:role/aws-service-role/sagemaker.application-autoscaling.amazonaws.com/AWSServiceRoleForApplicationAutoScaling_SageMakerEndpoint",
   "Condition": {
      "StringLike": {
         "iam:AWSServiceName": "sagemaker.application-autoscaling.amazonaws.com"
      }
   }
}
},
{
   "Effect": "Allow",
   "Action": "iam:CreateServiceLinkedRole",
   "Resource": "*",
   "Condition": {
      "StringEquals": {
         "iam:AWSServiceName": "robomaker.amazonaws.com"
      }
   }
}
},
{
   "Effect": "Allow",
   "Action": [
      "sns:Subscribe",
      "sns:CreateTopic"
   ],
   "Resource": [
      "arn:aws:sns:*:*:*SageMaker*",
      "arn:aws:sns:*:*:*sagemaker*"
   ]
}
Allow Users to View Their Own Permissions

This example shows how you might create a policy that allows IAM users to view the inline and managed policies that are attached to their user identity. This policy includes permissions to complete this action on the console or programmatically using the AWS CLI or AWS API.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "ViewOwnUserInfo",
            "Effect": "Allow",
            "Action": [
                "iam:GetUserPolicy",
                "iam:ListGroupsForUser",
                "iam:ListAttachedUserPolicies",
                "iam:ListUserPolicies",
                "iam:GetUser"
            ],
            "Resource": ["arn:aws:iam::*:user/${aws:username}"],
        }
    ]
}
```
Control Creation of SageMaker Resources with Condition Keys

Control fine-grained access to allow the creation of SageMaker resources by using SageMaker-specific condition keys. For information about using condition keys in IAM policies, see IAM JSON Policy Elements: Condition in the IAM User Guide.

The condition keys, along with related API actions, and links to relevant documentation are listed in Condition Keys for SageMaker in the IAM User Guide.

The following examples show how to use the SageMaker condition keys to control access.

Topics

- Control Access to SageMaker Resources by Using File System Condition Keys
- Restrict Training to a Specific VPC
- Restrict Access to Workforce Types for Ground Truth Labeling Jobs and Amazon A2I Human Review Workflows
- Enforce Encryption of Input Data
- Enforce Encryption of Notebook Instance Storage Volume
- Enforce Network Isolation for Training Jobs
- Enforce a Specific Instance Type for Training Jobs
- Enforce a Specific EL Accelerator for Training Jobs
- Enforce Disabling Internet Access and Root Access for Creating Notebook Instances
Control Access to SageMaker Resources by Using File System Condition Keys

SageMaker training provides a secure infrastructure for the training algorithm to run in, but for some cases you may want increased defense in depth. For example, you minimize the risk of running untrusted code in your algorithm, or you have specific security mandates in your organization. For these scenarios, you can use the service-specific condition keys in the Condition element of an IAM policy to scope down the user to specific file systems, directories, access modes (read-write, read-only) and security groups.

Topics

-_restrict an IAM User to Specific Directories and Access Modes
-_restrict a User to a Specific File System

Restrict an IAM User to Specific Directories and Access Modes

The policy below restricts a user to the /sagemaker/xgboost-dm/train and /sagemaker/xgboost-dm/validation directories of an EFS file system to ro (read-only) AccessMode:

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "AccessToElasticFileSystem",
      "Effect": "Allow",
      "Action": ["sagemaker:CreateTrainingJob", "sagemaker:CreateHyperParameterTuningJob"],
      "Resource": "*",
      "Condition": {
        "StringEquals": {
          "sagemaker:FileSystemId": "fs-12345678",
          "sagemaker:FileSystemAccessMode": "ro",
        }
      }
    }
  ]
}
```

### Note

When a directory is allowed, all of its subdirectories are also accessible by the training algorithm. POSIX permissions are ignored.
"sagemaker:FileSystemType": "EFS",
"sagemaker:FileSystemDirectoryPath": "/sagemaker/xgboost-dm/train"
]
},
{
"Sid": "AccessToElasticFileSystemValidation",
"Effect": "Allow",
"Action": [

"sagemaker:CreateTrainingJob",
"sagemaker:CreateHyperParameterTuningJob"
],
"Resource": "/",
"Condition": {
"StringEquals": {
"sagemaker:FileSystemId": "fs-12345678",
"sagemaker:FileSystemAccessMode": "ro",
"sagemaker:FileSystemType": "EFS",
"sagemaker:FileSystemDirectoryPath": "/sagemaker/xgboost-dm/validation"
}
}
}
}

Restrict a User to a Specific File System

To prevent a malicious algorithm using a user space client from accessing any file system directly in your account, you can restrict networking traffic by allowing ingress from a specific security group. In the following example, the user can only use the specified security group to access the file system:

{
"Version": "2012-10-17",
"Statement": [

{ "Sid": "AccessToLustreFileSystem",
"Effect": "Allow",
"Action": [

"sagemaker:CreateTrainingJob",
"sagemaker:CreateHyperParameterTuningJob"
],
Although the above example can restrict an algorithm to a specific file system, it does not prevent an algorithm from accessing any directory within that file system using the user space client. To mitigate this, you can:

- Ensure that the file system only contains data that you trust your users to access
- Create an IAM role that restricts your users to launching training jobs with algorithms from approved ECR repositories

For more information on how to use roles with SageMaker, see SageMaker Roles.

**Restrict Training to a Specific VPC**

Restrict an AWS user to creating training jobs from within a Amazon VPC. When a training job is created within a VPC, you can use VPC flow logs to monitor all traffic to and from the training cluster. For information about using VPC flow logs, see VPC Flow Logs in the Amazon Virtual Private Cloud User Guide.

The following policy enforces that a training job is created by a user calling CreateTrainingJob from within a VPC:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "sagemaker:FileSystemId": "fs-12345678",
                    "sagemaker:FileSystemAccessMode": "ro",
                    "sagemaker:FileSystemType": "FSxLustre",
                    "sagemaker:FileSystemDirectoryPath": "/fsx/sagemaker/xgboost/train"
                },
                "ForAllValues:StringEquals": {
                    "sagemaker:VpcSecurityGroupIds": [
                        "sg-12345678"
                    ]
                }
            }
        }
    ]
}
```
restrict access to workforce types for ground truth labeling jobs and amazon a2i human review workflows

amazon sagemaker ground truth and amazon augmented ai work teams fall into one of three workforce types: public (with amazon mechanical turk), private, and vendor. to restrict user access to a specific work team using one of these types or the work team arn, use the sagemaker:workteamtype and/or the sagemaker:workteamarn condition keys. for the sagemaker:workteamtype condition key, use string condition operators. for the sagemaker:workteamarn condition key, use amazon resource name (arn) condition operators. if the user attempts to create a labeling job with a restricted work team, sagemaker returns an access denied error.

the policies below demonstrate different ways to use the sagemaker:workteamtype and sagemaker:workteamarn condition keys with appropriate condition operators and valid condition values.

the following example uses the sagemaker:workteamtype condition key with the stringequals condition operator to restrict access to a public work team. it accepts condition
values in the following format: workforcetype-crowd, where workforcetype can equal public, private, or vendor.

```
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "RestrictWorkteamType",
         "Effect": "Deny",
         "Action": "sagemaker:CreateLabelingJob",
         "Resource": "*",
         "Condition": {
            "StringEquals": {
               "sagemaker:WorkteamType": "public-crowd"
            }
         }
      }
   ]
}
```

The following policies show how to restrict access to a public work team using the `sagemaker:WorkteamArn` condition key. The first shows how to use it with a valid IAM regex-variant of the work team ARN and the `ArnLike` condition operator. The second shows how to use it with the `ArnEquals` condition operator and the work team ARN.

```
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "RestrictWorkteamType",
         "Effect": "Deny",
         "Action": "sagemaker:CreateLabelingJob",
         "Resource": "*",
         "Condition": {
            "ArnLike": {
               "sagemaker:WorkteamArn": "arn:aws:sagemaker:*:*:workteam/public-crowd/**"
            }
         }
      }
   ]
}
```
Enforce Encryption of Input Data

The following policy restricts a user to specify an AWS KMS key to encrypt input data when creating training, hyperparameter tuning, and labeling jobs by using the `sagemaker:VolumeKmsKey` condition key:

```
{
    "Version": "2012-10-17",
    "Statement": [ 
        {
            "Sid": "EnforceEncryption",
            "Effect": "Allow",
            "Action": [ 
                "sagemaker:CreateTrainingJob",
                "sagemaker:CreateHyperParameterTuningJob",
                "sagemaker:CreateLabelingJob",
                "sagemaker:CreateFlowDefiniton"
            ],
            "Resource": "*",
            "Condition": { 
                "ArnEquals": { 
                    "sagemaker:VolumeKmsKey": "arn:aws:kms:us-west-2:111122223333:key/1234abcd-12ab-34cd-56ef-1234567890ab"
                }
            }
        }
    ]
}
```
Enforce Encryption of Notebook Instance Storage Volume

The following policy restricts a user to specify an AWS KMS key to encrypt the attached storage volume when creating or updating a notebook instance by using the `sagemaker:VolumeKmsKey` condition key:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "EnforceEncryption",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateNotebookInstance"
            ],
            "Resource": "*",
            "Condition": {
                "ArnLike": {
                    "sagemaker:VolumeKmsKey": "*key/volume-kms-key-12345"
                }
            }
        }
    ]
}
```

Enforce Network Isolation for Training Jobs

The following policy restricts a user to enable network isolation when creating training jobs by using the `sagemaker:NetworkIsolation` condition key:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "EnforceIsolation",
        }
    ]
}
```
"Effect": "Allow",
"Action": [
    "sagemaker:CreateTrainingJob",
    "sagemaker:CreateHyperParameterTuningJob"
],
"Resource": "*",
"Condition": {
    "Bool": {
        "sagemaker:NetworkIsolation": "true"
    }
}
}

Enforce a Specific Instance Type for Training Jobs

The following policy restricts a user to use a specific instance type when creating training jobs by using the `sagemaker:InstanceTypes` condition key:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "EnforceInstanceType",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateTrainingJob",
                "sagemaker:CreateHyperParameterTuningJob"
            ],
            "Resource": "*",
            "Condition": {
                "ForAllValues:StringLike": {
                    "sagemaker:InstanceTypes": ["ml.c5.*"]
                }
            }
        }
    ]
}
```
Enforce a Specific EI Accelerator for Training Jobs

The following policy restricts a user to use a specific elastic inference (EI) accelerator, if an accelerator is provided, when creating or updating notebook instances and when creating endpoint configurations by using the `sagemaker:AcceleratorTypes` condition key:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "EnforceAcceleratorType",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateNotebookInstance",
                "sagemaker:UpdateNotebookInstance",
                "sagemaker:CreateEndpointConfig"
            ],
            "Resource": "*",
            "Condition": {
                "ForAllValues:StringEquals": {
                    "sagemaker:AcceleratorTypes": ["ml.eia1.medium"]
                }
            }
        }
    ]
}
```

Enforce Disabling Internet Access and Root Access for Creating Notebook Instances

You can disable both internet access and root access to notebook instances to help make them more secure. For information about controlling root access to a notebook instance, see Control root access to a SageMaker notebook instance. For information about disabling internet access for a notebook instance, see Connect a Notebook Instance in a VPC to External Resources.

The following policy requires a user to disable network access when creating instance, and disable root access when creating or updating a notebook instance.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
```

Identity-Based Policy Examples
Control Access to the SageMaker API by Using Identity-based Policies

To control access to SageMaker API calls and calls to SageMaker hosted endpoints, use identity-based IAM policies.

Topics

- Restrict Access to SageMaker API and Runtime to Calls from Within Your VPC
Restrict Access to SageMaker API and Runtime to Calls from Within Your VPC

If you set up an interface endpoint in your VPC, individuals outside the VPC can still connect to the SageMaker API and runtime over the internet unless you attach an IAM policy that restricts access to calls coming from within the VPC to all users and groups that have access to your SageMaker resources. For information about creating a VPC interface endpoint for the SageMaker API and runtime, see [Connect to SageMaker Within your VPC](#).

⚠️ Important

If you apply an IAM policy similar to one of the following, users can't access the specified SageMaker APIs through the console.

To restrict access to only connections made from within your VPC, create an AWS Identity and Access Management policy that restricts access to only calls that come from within your VPC. Then add that policy to every AWS Identity and Access Management user, group, or role used to access the SageMaker API or runtime.

🔍 Note

This policy allows connections only to callers within a subnet where you created an interface endpoint.

```json
{
    "Id": "api-example-1",
    "Version": "2012-10-17",
    "Statement": [
    {
        "Sid": "EnableAPIAccess",
        "Effect": "Allow",
        "Action": ["sagemaker:*"],
        "Resource": "*",
        "Condition": {
            "StringEquals": {
                "aws:SourceVpc": "vpc-111bbaaa"
            }
        }
    }

    ]
}
```
If you want to restrict access to the API to only calls made using the interface endpoint, use the `aws:SourceVpce` condition key instead of `aws:SourceVpc`:

```
{
    "Id": "api-example-1",
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "EnableAPIAccess",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreatePresignedNotebookInstanceUrl"
            ],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "aws:sourceVpce": [
                        "vpce-111bbccc",
                        "vpce-111bbddd"
                    ]
                }
            }
        }
    ]
}
```

**Limit Access to SageMaker API and Runtime Calls by IP Address**

To allow access to SageMaker API calls and runtime invocations only from IP addresses in a list that you specify, attach an IAM policy that denies access to the API unless the call comes from an IP address in the list to every AWS Identity and Access Management user, group, or role used to access the API or runtime. For information about creating IAM policies, see [Creating IAM Policies](#) in the *AWS Identity and Access Management User Guide*. To specify the list of IP addresses that you want to have access to the API call, use the `IpAddress` condition operator and the `aws:SourceIP` condition context key. For information about IAM condition operators, see [IAM Identity-Based Policy Examples](#).
**JSON Policy Elements: Condition Operators** in the *AWS Identity and Access Management User Guide*. For information about IAM condition context keys, see *AWS Global Condition Context Keys*.

For example, the following policy allows access to the `CreateTrainingJob` only from IP addresses in the ranges 192.0.2.0-192.0.2.255 and 203.0.113.0-203.0.113.255:

```json
{
   "Version": "2012-10-17",
   "Statement": [
       {
           "Effect": "Allow",
           "Action": "sagemaker:CreateTrainingJob",
           "Resource": "*",
           "Condition": {
               "IpAddress": {
                   "aws:SourceIp": ["192.0.2.0/24", "203.0.113.0/24"]
               }
           }
       }
   ]
}
```

**Limit Access to a Notebook Instance by IP Address**

To allow access to a notebook instance only from IP addresses in a list that you specify, attach an IAM policy that denies access to `CreatePresignedNotebookInstanceUrl` unless the call comes from an IP address in the list to every AWS Identity and Access Management user, group, or role used to access the notebook instance. For information about creating IAM policies, see *Creating IAM Policies* in the *AWS Identity and Access Management User Guide*. To specify the list of IP addresses that you want to have access to the notebook instance, use the `IpAddress` condition operator and the `aws:SourceIp` condition context key. For information about IAM condition operators, see *IAM JSON Policy Elements: Condition Operators* in the *AWS Identity and Access Management User Guide*. For information about IAM condition context keys, see *AWS Global Condition Context Keys*.

For example, the following policy allows access to a notebook instance only from IP addresses in the ranges 192.0.2.0-192.0.2.255 and 203.0.113.0-203.0.113.255:
The policy restricts access to both the call to `CreatePresignedNotebookInstanceUrl` and to the URL that the call returns. The policy also restricts access to opening a notebook instance in the console and is enforced for every HTTP request and WebSocket frame that attempts to connect to the notebook instance.

### Note
Using this method to filter by IP address is incompatible when connecting to SageMaker through a VPC interface endpoint. For information about restricting access to a notebook instance when connecting through a VPC interface endpoint, see Connect to a Notebook Instance Through a VPC Interface Endpoint.

## Control Access to SageMaker Resources by Using Tags

Specify tags within an IAM policy to control access to groups of SageMaker resources. Use tags to implement attribute based access control (ABAC). Using tags helps you partition access to resources to specific groups of users. You can have one team with access to one group of resources and a different team with access to another set of resources. You can provide `ResourceTag` conditions in IAM policies to provide access for each group.
Note

Tag-based policies don't work to restrict the following API calls:

- DeleteImageVersion
- DescribeImageVersion
- ListAlgorithms
- ListCodeRepositories
- ListCompilationJobs
- ListEndpointConfigs
- ListEndpoints
- ListFlowDefinitions
- ListHumanTaskUis
- ListHyperparameterTuningJobs
- ListLabelingJobs
- ListLabelingJobsForWorkteam
- ListModelPackages
- ListModels
- ListNotebookInstanceLifecycleConfigs
- ListNotebookInstances
- ListSubscribedWorkteams
- ListTags
- ListProcessingJobs
- ListTrainingJobs
- ListTrainingJobsForHyperParameterTuningJob
- ListTransformJobs
- ListWorkteams
- Search

A simple example can help you understand how you can use tags to partition resources. Suppose that you've defined two different IAM groups, named DevTeam1 and DevTeam2, in your AWS
account. You’ve created 10 notebook instances as well. You’re using 5 of the notebook instances for one project. You’re using the other 5 for a second project. You can provide DevTeam1 with permissions to make API calls on the notebook instances that you’re using for the first project. You can provide DevTeam2 to make API calls on notebook instances used for the second project.

The following procedure provides a simple example that helps you understand the concept of adding tags. You can use it to implement the solution described in the preceding paragraph.

To control access to API calls (example)

1. Add a tag with the key Project and value A to the notebook instances used for the first project. For information about adding tags to SageMaker resources, see AddTags.
2. Add a tag with the key Project and value B to the notebook instances used for the second project.
3. Create an IAM policy with a ResourceTag condition that denies access to the notebook instances used for the second project, and attach that policy to DevTeam1. The following is an example of a policy that denying all API calls on any notebook instance that has a tag with a key of Project and a value of B:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": "sagemaker:*",
         "Resource": "*"
      },
      {
         "Effect": "Deny",
         "Action": "sagemaker:*",
         "Resource": "*",
         "Condition": {
            "StringEquals": {
               "sagemaker:ResourceTag/Project": "B"
            }
         }
      },
      {
         "Effect": "Deny",
         "Action": [
            "sagemaker:AddTags",
            // ...
         ],
         "Resource": "*"
      }
   ]
}
```

4. Create an IAM policy with a `ResourceTag` condition that denies access to the notebook instances used for the first project, and attach that policy to DevTeam2. The following is an example of a policy that denies all API calls on any notebook instance that has a tag with a key of `Project` and a value of `A`:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": "sagemaker:*",
      "Resource": "*"
    },
    {
      "Effect": "Deny",
      "Action": "sagemaker:*",
      "Resource": "*",
      "Condition": {
        "StringEquals": {
          "sagemaker:ResourceTag/Project": "A"
        }
      }
    },
    {
      "Effect": "Deny",
      "Action": ["sagemaker:AddTags", "sagemaker:DeleteTags"],
      "Resource": "*"
    }
  ]
}
```
Provide Permissions for Tagging SageMaker Resources Upon Creation

There are many Amazon SageMaker operations that let your users specify tags when you create a resource. Use resource tags to implement attributed based access control (ABAC). Using tags helps you partition access to resources to specific groups of users. You can have one team with access to one group of resources and a different team with access to another set of resources. You can provide ResourceTag conditions in IAM policies to provide access for each group.

You must give your users access to the permissions that they're using to create the resource. For users that need to create a processing job, you must give them access to sagemaker:CreateProcessingJob within the policy. If your users are tagging the resources that they create, you must also give tagging permissions.

⚠️ Important

You can only give your users permissions to add tags to resources that they create. They can't add tags to resources that have already been created.

The following are operations where you can't give them resources to add tags:

- DeleteImageVersion
- DescribeImageVersion
- ListAlgorithms
- ListCodeRepositories
- ListCompilationJobs
- ListEndpointConfigs
- ListEndpoints
- ListFlowDefinitions
- ListHumanTaskUis
- ListHyperparameterTuningJobs
- ListLabelingJobs
- ListLabelingJobsForWorkteam
- ListModelPackages
- ListModels
You can control access to the operations that the user can tag or the tags that they can use. You can specify permissions for the following use cases:

- Adding permissions to tag the resources created using any operation and permissions to use any tag
- Adding permissions to tag the resources created using specific operations and permissions to use any tag
- Adding permissions to tag the resources created using any operation and permissions to specific tags
- Adding permissions to tag the resources created using specific operations and permissions to specific tags

Permissions for any operation and any tag

Add the following statement to an IAM policy to grant your users permissions to add any tag for any operation:

```json
{
    "Statement": [
        {
            "Effect": "Allow",
            "Action": ["sagemaker:Create*"]
        }
    ]
}
```
Permissions for a specific operation with specific tags

The following statement gives the user permission to add the cc123 tag to resources created using the CreateModel operation. You can modify the statement to suit your own needs and add it to the IAM policy attached to the role of your users.

```json
{
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateModel"
            ],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "aws:RequestTag/cost-center": [ "cc123" ]
                },
                "ForAnyValue:StringEquals": {
                    "aws:TagKeys": [ "purpose" ]
                }
            }
        }
    ]
}
```
Permissions for a specific operation and any tag

The following statement gives your users permissions to add any tag to resources created using the CreateModel operation.

```json
{
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateModel"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:AddTags"
            ],
            "Condition": {
                "String": {
                    "sagemaker:TaggingAction": [ "CreateModel" ]
                }
            }
        }
    ]
}
```
Limit Access to Searchable Resources with Visibility Conditions

Use visibility conditions to limit the access of your users to specific tagged resources within an AWS account. Your users can access only those resources for which they have permissions. When your users are searching through their resources, they can limit the search results to specific resources.

You might want your users to only see and interact with the resources associated with specific Amazon SageMaker Studio or Amazon SageMaker Studio Classic domains. You can use visibility conditions to limit their access to a single domain or multiple domains.

The general format of a visibility condition is "sagemaker:SearchVisibilityCondition/Tags.key": "value". You can provide the key-value pair for any tagged resource.

```json
[
  {
    "Sid": "SageMakerApis",
    "Effect": "Allow",
    "Action": "sagemaker:Search",
    "Resource": "*",
    "Condition": {
      "StringEquals": {
      }
    }
  }
]
```

```json
{
  "MaxResults": number,
  "NextToken": "string",
  "Resource": "string", # Required Parameter
  "SearchExpression": {
    "Filters": [?
```
The visibility condition within uses the same "sagemaker:SearchVisibilityCondition/Tags.key": "value" formatting specified in the policy. Your users can specify the key-value pairs used for any tagged resource.

If a user includes the VisibilityConditions parameter in their Search request, but the access policy that applies to that user doesn't contain any matching conditions keys that were specified in VisibilityConditions, the Search request is still allowed and will run.
If a VisibilityConditions parameter is not specified in the user's Search API request, but the access policy that applies to that user contains condition keys related to VisibilityConditions, that user's Search request is denied.

**Cross-Service Confused Deputy Prevention**

The confused deputy problem is a security issue where an entity that doesn't have permission to perform an action can coerce a more-privileged entity to perform the action. In AWS, the confused deputy problem can arise due to cross-service impersonation. Cross-service impersonation can occur when one service (the calling service) invokes another service (the called service) and leverages the called service's elevated permissions to act on resources the calling service has no authorization to access. To prevent unauthorized access through the confused deputy problem, AWS provides tools to help secure your data across services. These tools help you control the permissions granted to service principals, limiting their access to only the resources in your account that are required. By carefully managing the access privileges of service principals, you can help mitigate the risk of services improperly accessing data or resources to which they should not have permissions.

Read on for general guidance or navigate to an example for a specific SageMaker feature:

**Topics**

- [Limit Permissions With Global Condition Keys](#)
- [SageMaker Edge Manager](#)
- [SageMaker Images](#)
- [SageMaker Inference](#)
- [SageMaker Batch Transform Jobs](#)
- [SageMaker Marketplace](#)
- [SageMaker Neo](#)
- [SageMaker Pipelines](#)
- [SageMaker Processing Jobs](#)
- [SageMaker Studio](#)
- [SageMaker Training Jobs](#)
Limit Permissions With Global Condition Keys

We recommend using the **aws:SourceArn** and **aws:SourceAccount** global condition keys in resource policies to limit the permissions to the resource that Amazon SageMaker gives another service. If you use both global condition keys and the **aws:SourceArn** value contains the account ID, the **aws:SourceAccount** value and the account in the **aws:SourceArn** value must use the same account ID when used in the same policy statement. Use **aws:SourceArn** if you want only one resource to be associated with the cross-service access. Use **aws:SourceAccount** if you want to allow any resource in that account to be associated with the cross-service use.

The most effective way to protect against the confused deputy problem is to use the **aws:SourceArn** global condition key with the full ARN of the resource. If you don't know the full ARN of the resource or if you are specifying multiple resources, use the **aws:SourceArn** global condition key with wildcards (*) for the unknown portions of the ARN. For example, `arn:aws:sagemaker::*:123456789012:*`.

The following example shows how you can use the **aws:SourceArn** and **aws:SourceAccount** global condition keys in SageMaker to prevent the confused deputy problem.

```json
{
    "Version": "2012-10-17",
    "Statement": {
        "Sid": "ConfusedDeputyPreventionExamplePolicy",
        "Effect": "Allow",
        "Principal": {
            "Service": "sagemaker.amazonaws.com"
        },
        # Specify an action and resource policy for another service
        "Action": "service:ActionName",
        "Resource": [
            "arn:aws:service:::ResourceName/*"
        ],
        "Condition": {
            "ArnLike": {
                "aws:SourceArn": "arn:partition:sagemaker:region:123456789012:"*
            },
            "StringEquals": {
                "aws:SourceAccount": "123456789012"
            }
        }
    }
}
```
SageMaker Edge Manager

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker Edge Manager created by account number `123456789012` in the `us-west-2` Region.

```json
{
    "Version": "2012-10-17",
    "Statement": {
        "Effect": "Allow",
        "Principal": { "Service": "sagemaker.amazonaws.com" },
        "Action": "sts:AssumeRole",
        "Condition": {
            "ArnLike": {
                "aws:SourceArn": "arn:aws:sagemaker:us-west-2:123456789012:*"
            }
        }
    }
}
```

You can replace the `aws:SourceArn` in this template with the full ARN of one specific packaging job to further limit permissions.

SageMaker Images

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker Images. Use this template with either `Image` or `ImageVersion`. This example uses an ImageVersion record ARN with the account number `123456789012`. Note that because the account number is part of the `aws:SourceArn` value, you do not need to specify an `aws:SourceAccount` value.

```json
{
    "Version": "2012-10-17",
    "Statement": {
        "Effect": "Allow",
        "Principal": { "Service": "sagemaker.amazonaws.com" },
        "Action": "sts:AssumeRole",
        "Condition": {
            "ArnLike": {
                "aws:SourceArn": "arn:aws:sagemaker:us-west-2:123456789012:*"
            }
        }
    }
}
```
Do not replace the `aws:SourceArn` in this template with the full ARN of a specific image or image version. The ARN must be in the format provided above and specify either image or image version. The partition placeholder should designate either an AWS commercial partition (`aws`) or an AWS in China partition (`aws-cn`), depending on where the image or image version is running. Similarly, the region placeholder in the ARN can be any valid Region where SageMaker images are available.

**SageMaker Inference**

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker **real-time**, **serverless**, and **asynchronous** inference. Note that because the account number is part of the `aws:SourceArn` value, you do not need to specify an `aws:SourceAccount` value.

```json
{
    "Version": "2012-10-17",
    "Statement": {
        "Effect": "Allow",
        "Principal": { "Service": "sagemaker.amazonaws.com" },
        "Action": "sts:AssumeRole",
        "Condition": {
            "ArnLike": {
                "aws:SourceArn": "arn:aws:sagemaker:*"  
            }
        }
    }
}
```

Do not replace the `aws:SourceArn` in this template with the full ARN of a specific model or endpoint. The ARN must be in the format provided above. The asterisk in the ARN template does not stand for wildcard and should not be changed.
SageMaker Batch Transform Jobs

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker batch transform jobs created by account number 123456789012 in the `us-west-2` Region. Note that because the account number is in the ARN, you do not need to specify an `aws:SourceAccount` value.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Service": "sagemaker.amazonaws.com"
            },
            "Action": "sts:AssumeRole",
            "Condition": {
                "ArnLike": {
                }
            }
        }
    ]
}
```

You can replace the `aws:SourceArn` in this template with the full ARN of one specific batch transform job to further limit permissions.

SageMaker Marketplace

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker Marketplace resources created by account number 123456789012 in the `us-west-2` Region. Note that because the account number is in the ARN, you do not need to specify an `aws:SourceAccount` value.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Service": "sagemaker.amazonaws.com"
            },
            "Action": "sts:AssumeRole",
            "Condition": {
                "ArnLike": {
                    "aws:SourceArn": "arn:aws:sagemaker:us-west-2:123456789012:*"
                }
            }
        }
    ]
}
```
Do not replace the `aws:SourceArn` in this template with the full ARN of a specific algorithm or model package. The ARN must be in the format provided above. The asterisk in the ARN template does stand for wildcard and covers all training jobs, models, and batch transform jobs from validation steps, as well as algorithm and model packages published to SageMaker Marketplace.

**SageMaker Neo**

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker Neo compilation jobs created by account number 123456789012 in the `us-west-2` Region. Note that because the account number is in the ARN, you do not need to specify an `aws:SourceAccount` value.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Service": "sagemaker.amazonaws.com"
            },
            "Action": "sts:AssumeRole",
            "Condition": {
                "ArnLike": {
                }
            }
        }
    ]
}
```
You can replace the `aws:SourceArn` in this template with the full ARN of one specific compilation job to further limit permissions.

**SageMaker Pipelines**

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker Pipelines using pipeline execution records from one or more pipelines. Note that because the account number is in the ARN, you do not need to specify an `aws:SourceAccount` value.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Principal": {
            "Service": "sagemaker.amazonaws.com"
         },
         "Action": "sts:AssumeRole",
         "Condition": {
            "ArnLike": {
               "aws:SourceArn": "arn:partition:sagemaker:region:123456789012:pipeline/mypipeline/*"
            }
         }
      }
   ]
}
```

Do not replace the `aws:SourceArn` in this template with the full ARN of a specific pipeline execution. The ARN must be in the format provided above. The `partition` placeholder should designate either an AWS commercial partition (`aws`) or an AWS in China partition (`aws-cn`), depending on where the pipeline is running. Similarly, the `region` placeholder in the ARN can be any valid Region where SageMaker Pipelines is available.

The asterisk in the ARN template does stand for wildcard and covers all pipeline executions of a pipeline named `mypipeline`. If you want to allow the AssumeRole permissions for all pipelines in account `123456789012` rather than one specific pipeline, then the `aws:SourceArn` would be `arn:aws:sagemaker::*:123456789012:pipeline/*`. 
SageMaker Processing Jobs

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker processing jobs created by account number 123456789012 in the `us-west-2` Region. Note that because the account number is in the ARN, you do not need to specify an `aws:SourceAccount` value.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Principal": {
            "Service": "sagemaker.amazonaws.com"
         },
         "Action": "sts:AssumeRole",
         "Condition": {
            "ArnLike": {
            }
         }
      }
   ]
}
```

You can replace the `aws:SourceArn` in this template with the full ARN of one specific processing job to further limit permissions.

SageMaker Studio

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker Studio created by account number 123456789012 in the `us-west-2` Region. Note that because the account number is part of the `aws:SourceArn` value, you do not need to specify an `aws:SourceAccount` value.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Principal": {
            "Service": "sagemaker.amazonaws.com"
         },
         "Action": "sts:AssumeRole",
         "Condition": {
            "ArnLike": {
            }
         }
      }
   ]
}
```
Do not replace the `aws:SourceArn` in this template with the full ARN of a specific Studio application, user profile, or domain. The ARN must be in the format provided in the previous example. The asterisk in the ARN template does not stand for wildcard and should not be changed.

**SageMaker Training Jobs**

The following example shows how you can use the `aws:SourceArn` global condition key to prevent the cross-service confused deputy problem for SageMaker training jobs created by account number `123456789012` in the `us-west-2` Region. Note that because the account number is in the ARN, you do not need to specify an `aws:SourceAccount` value.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Principal": {
                "Service": "sagemaker.amazonaws.com"
            },
            "Action": "sts:AssumeRole",
            "Condition": {
                "ArnLike": {
                }
            }
        }
    ]
}
```

You can replace the `aws:SourceArn` in this template with the full ARN of one specific training job to further limit permissions.
Next Up

For more information on managing execution roles, see SageMaker Roles.

SageMaker Roles

Amazon SageMaker performs operations on your behalf using other AWS services. You must grant SageMaker permissions to use these services and the resources they act upon. You grant SageMaker these permissions using an AWS Identity and Access Management (IAM) execution role. For more information on IAM roles, see IAM roles.

To create and use an execution role, you can use the following procedures.

Create execution role

Use the following procedure to create an execution role with the IAM managed policy, AmazonSageMakerFullAccess, attached. If your use case requires more granular permissions, use other sections on this page to create an execution role that meets your business needs. You can create an execution role using the SageMaker console or the AWS CLI.

⚠️ Important

The IAM managed policy, AmazonSageMakerFullAccess, used in the following procedure only grants the execution role permission to perform certain Amazon S3 actions on buckets or objects with SageMaker, Sagemaker, sagemaker, or aws-glue in the name. To learn how to add an additional policy to an execution role to grant it access to other Amazon S3 buckets and objects, see Add Additional Amazon S3 Permissions to a SageMaker Execution Role.

🔍 Note

You can create an execution role directly when you create a SageMaker domain or a notebook instance.

- For information on how to create a SageMaker domain, see Custom onboarding to Amazon SageMaker domain using IAM Identity Center.
- For information on how to create a notebook instance, see Step 1: Create an Amazon SageMaker Notebook Instance.
To create a new execution role from the SageMaker console

1. Open the IAM console at https://console.aws.amazon.com/iam/.
2. Choose Roles and then choose Create role.
3. Keep AWS service as the Trusted entity type and then use the down arrow to find SageMaker in Use cases for other AWS services.
4. Choose SageMaker – Execution and then choose Next.
5. The IAM managed policy, AmazonSageMakerFullAccess, is automatically attached to the role. To see the permissions included in this policy, choose the plus (+) sign next to the policy name. Choose Next.
6. Enter a Role name and a Description.
7. (Optional) Add additional permissions and tags to the role.
8. Choose Create role.
9. On the Roles section of the IAM console, find the role you just created. If needed, use the text box to search for the role using the role name.
10. On the role summary page, make note of the ARN.

To create a new execution role from the AWS CLI

Before you create an execution role using the AWS CLI, make sure to update and configure it by following the instructions in AWS CLI Prerequisites, then continue with the instructions in Onboard from the AWS CLI.

Once you have created an execution role, you can associate it with a SageMaker domain, a user profile, or with a Jupyter notebook instance.

- To learn about how to associate an execution role with an existing SageMaker domain, see Edit domain settings.
- To learn about how to associate an execution role with an existing user profile, see Add and Remove User Profiles.
- To learn about how to associate an execution role with an existing notebook instance, see Update a Notebook Instance.

You can also pass the ARN of an execution role to your API call. For example, using Amazon SageMaker Python SDK, you can pass the ARN of your execution role to an estimator. In the code...
sample that follows, we create an estimator using the XGBoost algorithm container and pass the ARN of the execution role as a parameter. For the full example on GitHub, see Customer Churn Prediction with XGBoost.

```python
import sagemaker, boto3
from sagemaker import image_uris

text = sagemaker.Session()
region = text.boto_region_name
bucket = text.default_bucket()
prefix = "sagemaker/DEMO-xgboost-churn"
container = sagemaker.image_uris.retrieve("xgboost", region, "1.7-1")

xgb = sagemaker.estimator.Estimator(
    container,
    execution-role-ARN,
    instance_count=1,
    instance_type="ml.m4.xlarge",
    output_path="s3://{0}/{1}/output".format(bucket, prefix),
    sagemaker_session=text,
)
...
```

**Add Additional Amazon S3 Permissions to a SageMaker Execution Role**

When you use a SageMaker feature with resources in Amazon S3, such as input data, the execution role you specify in your request (for example CreateTrainingJob) is used to access these resources.

If you attach the IAM managed policy, AmazonSageMakerFullAccess, to an execution role, that role has permission to perform certain Amazon S3 actions on buckets or objects with SageMaker, Sagemaker, sagemaker, or aws-glue in the name. It also has permission to perform the following actions on any Amazon S3 resource:

```
"s3:CreateBucket",
"s3:GetBucketLocation",
"s3:ListBucket",
"s3:ListAllMyBuckets",
"s3:GetBucketCors",
"s3:PutBucketCors"
```
To give an execution role permissions to access one or more specific buckets in Amazon S3, you can attach a policy similar to the following to the role. This policy grants an IAM role permission to perform all actions that AmazonSageMakerFullAccess allows but restricts this access to the buckets DOC-EXAMPLE-BUCKET1 and DOC-EXAMPLE-BUCKET2. Refer to the security documentation for the specific SageMaker feature you are using to learn more about the Amazon S3 permissions required for that feature.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "s3:GetObject",
                "s3:PutObject",
                "s3:DeleteObject",
                "s3:AbortMultipartUpload"
            ],
            "Resource": [
                "arn:aws:s3:::DOC-EXAMPLE-BUCKET1/*",
                "arn:aws:s3:::DOC-EXAMPLE-BUCKET2/*"
            ]
        },
        {
            "Effect": "Allow",
            "Action": [
                "s3:CreateBucket",
                "s3:GetBucketLocation",
                "s3:ListBucket",
                "s3:ListAllMyBuckets",
                "s3:GetBucketCors",
                "s3:PutBucketCors"
            ],
            "Resource": "*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "s3:GetBucketAcl",
                "s3:PutObjectAcl"
            ],
            "Resource": ["arn:aws:s3:::DOC-EXAMPLE-BUCKET1",
                          "arn:aws:s3:::DOC-EXAMPLE-BUCKET2"
                      ]
        }
    ]
}
```
Get execution role

You can use the SageMaker console or the AWS CLI to retrieve the ARN of the execution role attached to a SageMaker domain, a user profile, or a notebook instance.

- To find the ARN of the IAM execution role attached to a SageMaker domain, see View and edit domains.
- To find the ARN of the IAM execution role attached to a user profile, see View User Profiles and User Profile Details.
- To find the ARN of the IAM execution role attached to a notebook instance:
  1. Open the IAM console at https://console.aws.amazon.com/iam/.
  2. On the left navigation pane, choose Notebook then Notebook instances.
  3. From the list of notebooks, select the notebook that you want to view.
  4. The ARN is in the Permissions and encryption section.

Alternatively, Amazon SageMaker Python SDK users can retrieve the ARN of the execution role attached to their user profile or a notebook instance by running the following code:

```python
import sagemaker
sagemaker_session = sagemaker.Session()
role = sagemaker.get_execution_role()
```

**Note**

The execution role is available only when running a notebook within SageMaker. If you run get_execution_role in a notebook not on SageMaker, expect a "region" error.
Passing Roles

Actions like passing a role between services are a common function within SageMaker. You can find more details on [Actions, Resources, and Condition Keys for SageMaker](#) in the [IAM User Guide](#).

You pass the role (iam:PassRole) when making these API calls: `CreateAutoMLJob`, `CreateCompilationJob`, `CreateDomain`, `CreateFeatureGroup`, `CreateFlowDefinition`, `CreateHyperParameterTuningJob`, `CreateImage`, `CreateLabelingJob`, `CreateModel`, `CreateMonitoringSchedule`, `CreateNotebookInstance`, `CreateProcessingJob`, `CreateTrainingJob`, `CreateUserProfile`, `RenderUiTemplate`, `UpdateImage`, and `UpdateNotebookInstance`.

You attach the following trust policy to the IAM role which grants SageMaker principal permissions to assume the role, and is the same for all of the execution roles:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Principal": {
        "Service": "sagemaker.amazonaws.com"
      },
      "Action": "sts:AssumeRole"
    }
  ]
}
```

The permissions that you need to grant to the role vary depending on the API that you call. The following sections explain these permissions.

**Note**

Instead of managing permissions by crafting a permission policy, you can use the AWS-managed `AmazonSageMakerFullAccess` permission policy. The permissions in this policy are fairly broad, to allow for any actions you might want to perform in SageMaker. For a listing of the policy including information about the reasons for adding many of the permissions, see [AWS managed policy: AmazonSageMakerFullAccess](#). If you prefer to create custom policies and manage permissions to scope the permissions only to the actions you need to perform with the execution role, see the following topics.
Important

If you're running into issues, see Troubleshooting Amazon SageMaker Identity and Access.

For more information about IAM roles, see IAM Roles in the IAM User Guide.

Topics

• CreateAutoMLJob API: Execution Role Permissions
• CreateDomain API: Execution Role Permissions
• CreateImage and UpdateImage APIs: Execution Role Permissions
• CreateNotebookInstance API: Execution Role Permissions
• CreateHyperParameterTuningJob API: Execution Role Permissions
• CreateProcessingJob API: Execution Role Permissions
• CreateTrainingJob API: Execution Role Permissions
• CreateModel API: Execution Role Permissions
• SageMaker geospatial capabilities roles

CreateAutoMLJob API: Execution Role Permissions

For an execution role that you can pass in a CreateAutoMLJob API request, you can attach the following minimum permission policy to the role:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": ["iam:PassRole"],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "iam:PassedToService": "sagemaker.amazonaws.com"
                }
            }
        }
    ]
}
```
If you specify a private VPC for your AutoML job, add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "ec2:CreateNetworkInterface",
        "ec2:CreateNetworkInterfacePermission",
        "ec2:DeleteNetworkInterface",
        "ec2:DeleteNetworkInterfacePermission",
        "ec2:DescribeNetworkInterfaces",
        "ec2:DeleteNetworkInterface",
        "ec2:DeleteNetworkInterfacePermission",
        "ec2:DescribeNetworkInterfaces",
    ],
    "Resource": "*"
}
```
If your input is encrypted using server-side encryption with an AWS KMS–managed key (SSE-KMS), add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "kms:Decrypt"
    ]
}
```

If you specify a KMS key in the output configuration of your AutoML job, add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "kms:Encrypt"
    ]
}
```

If you specify a volume KMS key in the resource configuration of your AutoML job, add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "kms:CreateGrant"
    ]
}
```

**CreateDomain API: Execution Role Permissions**

The execution role for domains with IAM Identity Center and the user/execution role for IAM domains need the following permissions when you pass an AWS KMS customer managed key as the
KmsKeyId in the CreateDomain API request. The permissions are enforced during the CreateApp API call.

For an execution role that you can pass in the CreateDomain API request, you can attach the following permission policy to the role:

```
{
   "Version": "2012-10-17",
   "Statement": [
       {
           "Effect": "Allow",
           "Action": [
               "kms:CreateGrant",
               "kms:DescribeKey"
           ],
           "Resource": "arn:aws:kms:region:account-id:key/kms-key-id"
       }
   ]
}
```

Alternatively, if the permissions are specified in a KMS policy, you can attach the following policy to the role:

```
{
   "Version": "2012-10-17",
   "Statement": [
       {
           "Sid": "Allow use of the key",
           "Effect": "Allow",
           "Principal": {
               "AWS": [
                   "arn:aws:iam::account-id:role/ExecutionRole"
               ]
           },
           "Action": [
               "kms:CreateGrant",
               "kms:DescribeKey"
           ],
           "Resource": "*"
       }
   ]
}
```
CreateImage and UpdateImage APIs: Execution Role Permissions

For an execution role that you can pass in a CreateImage or UpdateImage API request, you can attach the following permission policy to the role:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "ecr:BatchGetImage",
        "ecr:GetDownloadUrlForLayer"
      ],
      "Resource": "*"
    },
    {
      "Effect": "Allow",
      "Action": [
        "iam:PassRole"
      ],
      "Resource": "*",
      "Condition": {
        "StringEquals": {
          "iam:PassedToService": "sagemaker.amazonaws.com"
        }
      }
    }
  ]
}
```

CreateNotebookInstance API: Execution Role Permissions

The permissions that you grant to the execution role for calling the CreateNotebookInstance API depend on what you plan to do with the notebook instance. If you plan to use it to invoke SageMaker APIs and pass the same role when calling the CreateTrainingJob and CreateModel APIs, attach the following permissions policy to the role:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "sagemaker:CreateTrainingJob",
        "sagemaker:CreateModel"
      ],
      "Resource": "*",
      "Condition": {
        "StringEquals": {
          "iam:PassedToService": "sagemaker.amazonaws.com"
        }
      }
    }
  ]
}
```
"Effect": "Allow",
"Action": [
  "sagemaker:*",
  "ecr:GetAuthorizationToken",
  "ecr:GetDownloadUrlForLayer",
  "ecr:BatchGetImage",
  "ecr:BatchCheckLayerAvailability",
  "ecr:SetRepositoryPolicy",
  "ecr:CompleteLayerUpload",
  "ecr:BatchDeleteImage",
  "ecr:UploadLayerPart",
  "ecr:DeleteRepositoryPolicy",
  "ecr:InitiateLayerUpload",
  "ecr:DeleteRepository",
  "ecr:PutImage",
  "ecr:CreateRepository",
  "cloudwatch:PutMetricData",
  "cloudwatch:GetMetricData",
  "cloudwatch:GetMetricStatistics",
  "cloudwatch:ListMetrics",
  "logs:CreateLogGroup",
  "logs:CreateLogStream",
  "logs:DescribeLogStreams",
  "logs:PutLogEvents",
  "logs:GetLogEvents",
  "s3:CreateBucket",
  "s3:ListBucket",
  "s3:GetBucketLocation",
  "s3:GetObject",
  "s3:PutObject",
  "s3:DeleteObject",
  "robomaker:CreateSimulationApplication",
  "robomaker:DescribeSimulationApplication",
  "robomaker:DeleteSimulationApplication",
  "robomaker:CreateSimulationJob",
  "robomaker:DescribeSimulationJob",
  "robomaker:CancelSimulationJob",
  "ec2:CreateVpcEndpoint",
  "ec2:DescribeRouteTables",
  "elasticfilesystem:DescribeMountTargets"
],
"Resource": "*
}
To tighten the permissions, limit them to specific Amazon S3 and Amazon ECR resources, by restricting "Resource": "*", as follows:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:*",
                "ecr:GetAuthorizationToken",
                "cloudwatch:PutMetricData",
                "logs:CreateLogGroup",
                "logs:CreateLogStream",
                "logs:DescribeLogStreams",
                "logs:PutLogEvents",
                "logs:GetLogEvents"
            ]
        }
    ]
}
```
SageMaker Roles

Amazon SageMaker

5770
If you plan to access other resources, such as Amazon DynamoDB or Amazon Relational Database Service, add the relevant permissions to this policy.

In the preceding policy, you scope the policy as follows:

- **Scope the s3:ListBucket permission to the specific bucket that you specify as**
  
  InputDataConfig.DataSource.S3DataSource.S3Uri in a CreateTrainingJob request.

- **Scope s3:GetObject, s3:PutObject, and s3:DeleteObject permissions as follows:**
  
  - **Scope to the following values that you specify in a CreateTrainingJob request:**
    
    InputDataConfig.DataSource.S3DataSource.S3Uri
    
    OutputDataConfig.S3OutputPath
  
  - **Scope to the following values that you specify in a CreateModel request:**
    
    PrimaryContainer.ModelDataUrl
    
    SuplementalContainers.ModelDataUrl

- **Scope ecr permissions as follows:**
  
  - Scope to the AlgorithmSpecification.TrainingImage value that you specify in a CreateTrainingJob request.
  
  - Scope to the PrimaryContainer.Image value that you specify in a CreateModel request.

The cloudwatch and logs actions are applicable for "*" resources. For more information, see CloudWatch Resources and Operations in the Amazon CloudWatch User Guide.
CreateHyperParameterTuningJob API: Execution Role Permissions

For an execution role that you can pass in a CreateHyperParameterTuningJob API request, you can attach the following permission policy to the role:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "cloudwatch:PutMetricData",
            "logs:CreateLogStream",
            "logs:PutLogEvents",
            "logs:CreateLogGroup",
            "logs:DescribeLogStreams",
            "s3:GetObject",
            "s3:PutObject",
            "s3:ListBucket",
            "ecr:GetAuthorizationToken",
            "ecr:BatchCheckLayerAvailability",
            "ecr:GetDownloadUrlForLayer",
            "ecr:BatchGetImage"
         ],
         "Resource": "*"
      }
   ]
}
```

Instead of specifying "Resource": "*", you could scope these permissions to specific Amazon S3, Amazon ECR, and Amazon CloudWatch Logs resources:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "cloudwatch:PutMetricData",
            "ecr:GetAuthorizationToken"
         ],
         "Resource": "*"
      }
   ]
}
```
{
    "Effect": "Allow",
    "Action": [
      "s3:ListBucket"
    ],
    "Resource": ["arn:aws:s3:::inputbucket"
  ]
},
{
    "Effect": "Allow",
    "Action": ["s3:GetObject", "s3:PutObject"
    ],
    "Resource": ["arn:aws:s3:::inputbucket/object",
                "arn:aws:s3:::outputbucket/path"
    ]
},
{
    "Effect": "Allow",
    "Action": ["ecr:BatchCheckLayerAvailability", "ecr:GetDownloadUrlForLayer", "ecr:BatchGetImage"
    ],
    "Resource": "arn:aws:ecr:region::repository/my-repo"
},
{
    "Effect": "Allow",
    ],
    "Resource": "arn:aws:logs:*:*:log-group:/aws/sagemaker/TrainingJobs"
}
If the training container associated with the hyperparameter tuning job needs to access other data sources, such as DynamoDB or Amazon RDS resources, add relevant permissions to this policy.

In the preceding policy, you scope the policy as follows:

- Scope the `s3:ListBucket` permission to a specific bucket that you specify as the `InputDataConfig.DataSource.S3DataSource.S3Uri` in a `CreateTrainingJob` request.
- Scope the `s3:GetObject` and `s3:PutObject` permissions to the following objects that you specify in the input and output data configuration in a `CreateHyperParameterTuningJob` request:
  
  - `InputDataConfig.DataSource.S3DataSource.S3Uri`
  - `OutputDataConfig.S3OutputPath`

- Scope Amazon ECR permissions to the registry path (`AlgorithmSpecification.TrainingImage`) that you specify in a `CreateHyperParameterTuningJob` request.
- Scope Amazon CloudWatch Logs permissions to log group of SageMaker training jobs.

The `cloudwatch` actions are applicable for "*" resources. For more information, see CloudWatch Resources and Operations in the Amazon CloudWatch User Guide.

If you specify a private VPC for your hyperparameter tuning job, add the following permissions:

```json
{
   "Effect": "Allow",
   "Action": [
      "ec2:CreateNetworkInterface",
      "ec2:CreateNetworkInterfacePermission",
      "ec2:DeleteNetworkInterface",
      "ec2:DeleteNetworkInterfacePermission",
      "ec2:DescribeNetworkInterfaces",
      "ec2:DescribeVpcs",
      "ec2:DescribeDhcpOptions",
      "ec2:DescribeSubnets",
      "ec2:DescribeSecurityGroups"
   ]
}
```
If your input is encrypted using server-side encryption with an AWS KMS–managed key (SSE-KMS), add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "kms:Decrypt"
    ]
}
```

If you specify a KMS key in the output configuration of your hyperparameter tuning job, add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "kms:Encrypt"
    ]
}
```

If you specify a volume KMS key in the resource configuration of your hyperparameter tuning job, add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "kms:CreateGrant"
    ]
}
```

**CreateProcessingJob API: Execution Role Permissions**

For an execution role that you can pass in a `CreateProcessingJob` API request, you can attach the following permission policy to the role:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
            
            ]
        }
    ]
}
```
Instead of specifying "Resource": "*", you could scope these permissions to specific Amazon S3 and Amazon ECR resources:

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "cloudwatch:PutMetricData",
            "logs:CreateLogStream",
            "logs:PutLogEvents",
            "logs:CreateLogGroup",
            "logs:DescribeLogStreams",
            "s3:GetObject",
            "s3:PutObject",
            "s3:ListBucket",
            "ecr:GetAuthorizationToken",
            "ecr:BatchCheckLayerAvailability",
            "ecr:GetDownloadUrlForLayer",
            "ecr:BatchGetImage"
         ],
         "Resource": "*"
      },
      {
         "Effect": "Allow",
         "Action": [
            "s3:ListBucket"
         ],
         "Resource": [
            "arn:aws:s3:::inputbucket"
         ]
      }
   ]
}
```
If `CreateProcessingJob.AppSpecification.ImageUri` needs to access other data sources, such as DynamoDB or Amazon RDS resources, add relevant permissions to this policy.

In the preceding policy, you scope the policy as follows:

- Scope the `s3:ListBucket` permission to a specific bucket that you specify as the `ProcessingInputs` in a `CreateProcessingJob` request.
- Scope the `s3:GetObject` and `s3:PutObject` permissions to the objects that will be downloaded or uploaded in the `ProcessingInputs` and `ProcessingOutputConfig` in a `CreateProcessingJob` request.
- Scope Amazon ECR permissions to the registry path (`AppSpecification.ImageUri`) that you specify in a `CreateProcessingJob` request.

The `cloudwatch` and `logs` actions are applicable for "*" resources. For more information, see [CloudWatch Resources and Operations](https://docs.aws.amazon.com/AmazonCloudWatch/latest/monitoring/cloudwatch-resources-and-operations.html) in the Amazon CloudWatch User Guide.
If you specify a private VPC for your processing job, add the following permissions. Don't scope in the policy with any conditions or resource filters. Otherwise, the validation checks that occur during the creation of the processing job fail.

```
{
  "Effect": "Allow",
  "Action": [
    "ec2:CreateNetworkInterface",
    "ec2:CreateNetworkInterfacePermission",
    "ec2:DeleteNetworkInterface",
    "ec2:DeleteNetworkInterfacePermission",
    "ec2:DescribeNetworkInterfaces",
    "ec2:DescribeVpcs",
    "ec2:DescribeDhcpOptions",
    "ec2:DescribeSubnets",
    "ec2:DescribeSecurityGroups"
  ]
}
```

If your input is encrypted using server-side encryption with an AWS KMS–managed key (SSE-KMS), add the following permissions:

```
{
  "Effect": "Allow",
  "Action": [
    "kms:Decrypt"
  ]
}
```

If you specify a KMS key in the output configuration of your processing job, add the following permissions:

```
{
  "Effect": "Allow",
  "Action": [
    "kms:Encrypt"
  ]
}
```

If you specify a volume KMS key in the resource configuration of your processing job, add the following permissions:
CreateTrainingJob API: Execution Role Permissions

For an execution role that you can pass in a CreateTrainingJob API request, you can attach the following permission policy to the role:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "cloudwatch:PutMetricData",
                "logs:CreateLogStream",
                "logs:PutLogEvents",
                "logs:CreateLogGroup",
                "logs:DescribeLogStreams",
                "s3:GetObject",
                "s3:PutObject",
                "s3:ListBucket",
                "ecr:GetAuthorizationToken",
                "ecr:BatchCheckLayerAvailability",
                "ecr:GetDownloadUrlForLayer",
                "ecr:BatchGetImage"
            ],
            "Resource": "*"
        }
    ]
}
```

Instead of specifying "Resource": "*", you could scope these permissions to specific Amazon S3 and Amazon ECR resources:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "cloudwatch:PutMetricData",
                "logs:CreateLogStream",
                "logs:PutLogEvents",
                "logs:CreateLogGroup",
                "logs:DescribeLogStreams",
                "s3:GetObject",
                "s3:PutObject",
                "s3:ListBucket",
                "ecr:GetAuthorizationToken",
                "ecr:BatchCheckLayerAvailability",
                "ecr:GetDownloadUrlForLayer",
                "ecr:BatchGetImage"
            ],
            "Resource": "*"
        }
    ]
}
```
SageMaker Roles

```json
[  
    {
      "Effect": "Allow",
      "Action": [
        "cloudwatch:PutMetricData",
        "logs:CreateLogStream",
        "logs:PutLogEvents",
        "logs:CreateLogGroup",
        "logs:DescribeLogStreams",
        "ecr:GetAuthorizationToken"
      ],
      "Resource": "*
    },
    {
      "Effect": "Allow",
      "Action": ["s3:ListBucket"
      ],
      "Resource": ["
        arn:aws:s3:::inputbucket"
      ]
    },
    {
      "Effect": "Allow",
      "Action": ["s3:GetObject", "s3:PutObject"
      ],
      "Resource": ["
        arn:aws:s3:::inputbucket/object",
        arn:aws:s3:::outputbucket/path"
      ]
    },
    {
      "Effect": "Allow",
      "Action": ["ecr:BatchCheckLayerAvailability", "ecr:GetDownloadUrlForLayer", "ecr:BatchGetImage"
      ],
      "Resource": "arn:aws:ecr:region::repository/my-repo"
    }
  ]
]
If `CreateTrainingJob.AlgorithmSpecifications.TrainingImage` needs to access other data sources, such as DynamoDB or Amazon RDS resources, add relevant permissions to this policy.

In the preceding policy, you scope the policy as follows:

- **Scope the s3:ListBucket permission to a specific bucket that you specify as the `InputDataConfig.DataSource.S3DataSource.S3Uri` in a `CreateTrainingJob` request.**

- **Scope the s3:GetObject and s3:PutObject permissions to the following objects that you specify in the input and output data configuration in a `CreateTrainingJob` request:**
  
  `InputDataConfig.DataSource.S3DataSource.S3Uri`

  `OutputDataConfig.S3OutputPath`

- **Scope Amazon ECR permissions to the registry path**
  
  `(AlgorithmSpecification.TrainingImage)` that you specify in a `CreateTrainingJob` request.

The `cloudwatch` and `logs` actions are applicable for "*" resources. For more information, see [CloudWatch Resources and Operations](https://docs.aws.amazon.com/AmazonCloudWatch/latest/monitoring/CloudWatch-Resources-Operations.html) in the Amazon CloudWatch User Guide.

If you specify a private VPC for your training job, add the following permissions:

```json
{
   "Effect": "Allow",
   "Action": [
      "ec2:CreateNetworkInterface",
      "ec2:CreateNetworkInterfacePermission",
      "ec2:DeleteNetworkInterface",
      "ec2:DeleteNetworkInterfacePermission",
      "ec2:DescribeNetworkInterfaces",
      "ec2:DescribeVpcs",
      "ec2:DescribeDhcpOptions",
      "ec2:DescribeSubnets",
      "ec2:DescribeSecurityGroups"
   ]
}
```

If your input is encrypted using server-side encryption with an AWS KMS–managed key (SSE-KMS), add the following permissions:
If you specify a KMS key in the output configuration of your training job, add the following permissions:

```
{
    "Effect": "Allow",
    "Action": [
        "kms:Decrypt"
    ]
}
```

If you specify a volume KMS key in the resource configuration of your training job, add the following permissions:

```
{
    "Effect": "Allow",
    "Action": [
        "kms:Encrypt"
    ]
}
```

**CreateModel API: Execution Role Permissions**

For an execution role that you can pass in a `CreateModel` API request, you can attach the following permission policy to the role:

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "cloudwatch:PutMetricData",
                "logs:CreateLogStream",
                "logs:PutLogEvents",
                "kms:Decrypt",
                "kms:Encrypt",
                "kms:CreateGrant",
                "logs:GetLogEvents",
                "logs:FilterLogEvents",
                "logs:PutLogRetentionPeriod",
                "logs:PutLogRetentionPeriod"
            ]
        }
    ]
}
```
Instead of specifying "Resource": "*", you can scope these permissions to specific Amazon S3 and Amazon ECR resources:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "cloudwatch:PutMetricData",
        "logs:CreateLogStream",
        "logs:PutLogEvents",
        "logs:CreateLogGroup",
        "logs:DescribeLogStreams",
        "ecr:GetAuthorizationToken"
      ],
      "Resource": "*"
    },
    {
      "Effect": "Allow",
      "Action": [
        "s3:GetObject"
      ],
      "Resource": [
        "arn:aws:s3:::inputbucket/object"
      ]
    },
    {
      "Effect": "Allow",
      "Action": [
        "s3:GetObject"
      ],
      "Resource": [
        "arn:aws:s3:::inputbucket/object"
      ]
    }
  ]
}
```
If `CreateModel.PrimaryContainer.Image` need to access other data sources, such as Amazon DynamoDB or Amazon RDS resources, add relevant permissions to this policy.

In the preceding policy, you scope the policy as follows:

- Scope S3 permissions to objects that you specify in the `PrimaryContainer.ModelDataUrl` in a `CreateModel` request.
- Scope Amazon ECR permissions to a specific registry path that you specify as the `PrimaryContainer.Image` and `SecondaryContainer.Image` in a `CreateModel` request.

The `cloudwatch` and `logs` actions are applicable for "*" resources. For more information, see CloudWatch Resources and Operations in the Amazon CloudWatch User Guide.

**Note**

If you plan to use the SageMaker deployment guardrails feature for model deployment in production, ensure that your execution role has permission to perform the `cloudwatch:DescribeAlarms` action on your auto-rollback alarms.

If you specify a private VPC for your model, add the following permissions:

```json
{
    "Effect": "Allow",
    "Action": [
        "ec2:CreateNetworkInterface",
        "ec2:CreateNetworkInterfacePermission",
    ]
}
```
SageMaker geospatial capabilities roles

As a managed service, Amazon SageMaker geospatial capabilities performs operations on your behalf on the AWS hardware that is managed by SageMaker. Use AWS Identity and Access Management to grant users, groups, and roles access to SageMaker geospatial.

An IAM Administrator can grant these permissions to user, group, or role using the AWS Management Console, AWS CLI, or one of the AWS SDKs.

To use SageMaker geospatial you need the following IAM permissions.

1. **An SageMaker execution role.**

   To use the SageMaker geospatial specific API operations your SageMaker execution role must include the SageMaker geospatial service principal, sagemaker-geospatial.amazonaws.com in the execution role's trust policy. This allows the SageMaker execution role to perform actions in your AWS account on your behalf.

2. **A user, group, or role that has access Amazon SageMaker Studio Classic and SageMaker geospatial**

   To get started with SageMaker geospatial you can use the AWS managed policy: AmazonSageMakerGeospatialFullAccess. This grants will grant a user, group, or role full access to SageMaker geospatial. To see the policy and learn more about which actions, resources, and conditions are available, see [AWS managed policy: AmazonSageMakerFullAccess](https://docs.aws.amazon.com/IAM/latest/userguide/managed-policies-list.html#managed-policies-amazonsagemakeraccess).

   To get started with Studio Classic and creating a Amazon SageMaker domain, see [Amazon SageMaker domain overview](https://docs.aws.amazon.com/sagemaker/latest/dg/creating-domain.html).
Use the following topics to create a new SageMaker execution role, update an existing SageMaker execution role, and learn how to manage permissions using SageMaker geospatial specific IAM actions, resources, and conditions.

Topics

- **Creating an new SageMaker execution role**
- **Adding the SageMaker geospatial service principal to an existing SageMaker execution role**
- **StartEarthObservationJob API: Execution role permissions**
- **StartVectorEnrichmentJob API: Execution role permissions**
- **ExportEarthObservationJob API: Execution role permissions**
- **ExportVectorEnrichmentJob API: Execution Role Permissions**

**Creating an new SageMaker execution role**

To work with SageMaker geospatial capabilities, you must set up a user, group, or role, and an execution role. A user role is an AWS identity with permissions policies that determine what the user can and cannot do within AWS. An execution role is an IAM role that grants the service permission to access your AWS resources. An execution role consists of permissions and trust policy. The trust policy specifies which principals have the permission to assume the role.

SageMaker geospatial also requires a different service principal, sagemaker-geospatial.amazonaws.com. If you are an existing SageMaker customer, you must add this additional service principal to your trust policy.

Use the following procedure to create an new execution role with the IAM managed policy, AmazonSageMakerGeospatialFullAccess, attached. If your use case requires more granular permissions, use other sections of this guide to create an execution role that meets your business needs.

⚠️ **Important**

The IAM managed policy, AmazonSageMakerGeospatialFullAccess, used in the following procedure, only grants the execution role permission to perform certain Amazon S3 actions on buckets or objects with SageMaker, Sagemaker, sagemaker, or aws-glue in the name. To learn how to update the execution role’s policy to grant it access to
To create a new role

1. Open the IAM console at https://console.aws.amazon.com/iam/.
2. Select Roles and then select Create role.
4. Select Next: Permissions.
5. The IAM managed policy, AmazonSageMakerGeospatialFullAccess is automatically attached to this role. To see the permissions included in this policy, select the sideways arrow next to the policy name. Select Next: Tags.
6. (Optional) Add tags and select Next: Review.
7. Give the role a name in the text field under Role name and select Create role.
8. In the Roles section of the IAM console, select the role you just created in step 7. If needed, use the text box to search for the role using the role name you entered in step 7.
9. On the role summary page, make note of the ARN.

Adding the SageMaker geospatial service principal to an existing SageMaker execution role

To use the SageMaker geospatial specific API operations your SageMaker execution role must include the SageMaker geospatial service principal, sagemaker-geospatial.amazonaws.com in the execution role's trust policy. This allows the SageMaker execution role to perform actions in your AWS account on your behalf.

Actions like passing a role between services are common within SageMaker. For more details,

To add the SageMaker geospatial service principal to an existing SageMaker execution role update the existing policy to include the SageMaker geospatial service principal as shown in the following trust policy. By attaching the service principal to the trust policy a SageMaker execution role can now run the SageMaker geospatial specific APIs on your behalf.

To learn more about SageMaker geospatial specific IAM actions, resources, and conditions, see Actions, Resources, and Condition Keys for SageMaker in the IAM User Guide.
For an execution role that you can pass in a StartEarthObservationJob API request, you can attach the following minimum permissions policy to the role:

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "s3:AbortMultipartUpload",
        "s3:PutObject",
        "s3:GetObject",
        "s3:ListBucketMultipartUploads"
      ],
      "Resource": [
        "arn:aws:s3:::*SageMaker*",
        "arn:aws:s3:::*Sagemaker*",
        "arn:aws:s3:::*sagemaker*"
      ]
    },
    {
      "Effect": "Allow",
      "Action": "sagemaker-geospatial:GetEarthObservationJob",
      "Resource": "arn:aws:sagemaker-geospatial:*:*:earth-observation-job/*"
    },
    {
      "Effect": "Allow",
      "Action": "sts:AssumeRole"
    }
  ]
}
```
"Effect": "Allow",
"Action": "sagemaker-geospatial:GetRasterDataCollection",
}
]
}

If your input Amazon S3 bucket is encrypted using server-side encryption with an AWS KMS managed key (SSE-KMS), see Using Amazon S3 Bucket Keys for more information.

**StartVectorEnrichmentJob API: Execution role permissions**

For an execution role that you can pass in a StartVectorEnrichmentJob API request, you can attach the following minimum permissions policy to the role:

```
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "s3:AbortMultipartUpload",
        "s3:PutObject",
        "s3:GetObject",
        "s3:ListBucketMultipartUploads"
      ],
      "Resource": [
        "arn:aws:s3:::*SageMaker*",
        "arn:aws:s3:::*Sagemaker*",
        "arn:aws:s3:::*sagemaker*"
      ]
    },
    {
      "Effect": "Allow",
      "Action": "sagemaker-geospatial:GetVectorEnrichmentJob",
      "Resource": "arn:aws:sagemaker-geospatial:*:*:vector-enrichment-job/*"
    }
  ]
}
```

If your input Amazon S3 bucket is encrypted using server-side encryption with an AWS KMS managed key (SSE-KMS), see Using Amazon S3 Bucket Keys for more information.
ExportEarthObservationJob API: Execution role permissions

For an execution role that you can pass in a ExportEarthObservationJob API request, you can attach the following minimum permissions policy to the role:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "s3:AbortMultipartUpload",
                "s3:PutObject",
                "s3:GetObject",
                "s3:ListBucketMultipartUploads"
            ],
            "Resource": [
                "arn:aws:s3:::*SageMaker*",
                "arn:aws:s3:::*Sagemaker*",
                "arn:aws:s3:::*sagemaker*"
            ]
        },
        {
            "Effect": "Allow",
            "Action": "sagemaker-geospatial:GetEarthObservationJob",
            "Resource": "arn:aws:sagemaker-geospatial:*:*:earth-observation-job/*"
        }
    ]
}
```

If your input Amazon S3 bucket is encrypted using server-side encryption with an AWS KMS managed key (SSE-KMS), see [Using Amazon S3 Bucket Keys](https://docs.aws.amazon.com/AmazonS3/latest/userguide/using-bucket-keys.html) for more information.

ExportVectorEnrichmentJob API: Execution Role Permissions

For an execution role that you can pass in a ExportVectorEnrichmentJob API request, you can attach the following minimum permissions policy to the role:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": "sagemaker-geospatial:GetEarthObservationJob",
            "Resource": "arn:aws:sagemaker-geospatial:*:*:earth-observation-job/*"
        }
    ]
}
```
"Action": [ 
  "s3:AbortMultipartUpload",
  "s3:PutObject",
  "s3:GetObject",
  "s3:ListBucketMultipartUploads"
],
"Resource": [ 
  "arn:aws:s3:::*SageMaker*",
  "arn:aws:s3:::*Sagemaker*",
  "arn:aws:s3:::*sagemaker*"
]
},
{
  "Effect": "Allow",
  "Action": "sagemaker-geospatial:GetVectorEnrichmentJob",
  "Resource":  "arn:aws:sagemaker-geospatial:*:*:vector-enrichment-job/*"
}
]

If your input Amazon S3 bucket is encrypted using server-side encryption with an AWS KMS managed key (SSE-KMS), see Using Amazon S3 Bucket Keys.

**Amazon SageMaker Role Manager**

Machine learning (ML) administrators striving for least-privilege permissions with Amazon SageMaker must account for a diversity of industry perspectives, including the unique least-privilege access needs required for personas such as data scientists, machine learning operation (MLOps) engineers, and more. Use Amazon SageMaker Role Manager to build and manage persona-based IAM roles for common machine learning needs directly through the Amazon SageMaker console.

Amazon SageMaker Role Manager provides 3 preconfigured role personas and predefined permissions for 12 common ML activities. Explore the provided personas and their suggested policies, or create and maintain roles for personas unique to your business needs. If you require additional customization, specify networking and encryption permissions for Amazon Virtual Private Cloud resources and AWS Key Management Service encryption keys in Step 1. Enter role information of the Amazon SageMaker Role Manager.

**Topics**

- Using the role manager (console)
Using the role manager (console)

You can use the Amazon SageMaker Role Manager from the following locations on the left-hand navigation of the Amazon SageMaker console:

- **Getting started** – Quickly add permissions policies for your users.
- **domains** – Add permissions policies for users within a Amazon SageMaker domain.
- **Notebooks** – Add least permissions for users who create and run notebooks.
- **Training** – Add least permissions for users who create and manage training jobs.
- **Inference** – Add least permissions for users who deploy and manage models for inference.

You can use the following are procedures to start the process of creating a role from different locations in the SageMaker console.

**Getting started**

If you're using SageMaker for the first time, we recommend creating a role from the *Getting started* section.

To create a role using Amazon SageMaker Role Manager, do the following.

1. Open the Amazon SageMaker console.
2. On the left navigation pane, choose **Admin configurations**.
3. Under **Admin configurations**, choose **Role manager**.
4. Choose **Create a role**.

**domains**

You can create a role using Amazon SageMaker Role Manager when you start the process of creating a Amazon SageMaker domain.
To create a role using Amazon SageMaker Role Manager, do the following.

1. Open the Amazon SageMaker console.
2. On the left navigation pane, choose Admin configurations.
3. Under Admin configurations, choose domains.
4. Choose Create domain.
5. Choose Create role using the role creation wizard.

**Notebook**

You can create a role using Amazon SageMaker Role Manager when you start the process of creating a notebook.

To create a role using Amazon SageMaker Role Manager, do the following.

1. Open the Amazon SageMaker console.
2. On the left-hand navigation, select Notebook.
3. Choose Notebook instances.
4. Choose Create notebook instance.
5. Choose Create role using the role creation wizard.

**Training**

You can create a role using Amazon SageMaker Role Manager when you start the process of creating a training job.

To create a role using Amazon SageMaker Role Manager, do the following.

1. Open the Amazon SageMaker console.
2. On the left-hand navigation, choose Training.
3. Select Training jobs.
4. Choose Create training job.
5. Choose Create role using the role creation wizard.
Inference

You can create a role using Amazon SageMaker Role Manager when you start the process of deploying a model for inference.

To create a role using Amazon SageMaker Role Manager, do the following.

1. Open the Amazon SageMaker console.
2. On the left-hand navigation, choose **Inference**.
3. Select **Models**.
4. Choose **Create model**.
5. Choose **Create role using the role creation wizard**.

After you've completed one of the preceding procedures, use the information in the following sections to help you create the role.

Prerequisites

To use Amazon SageMaker Role Manager, you must have permission to create an IAM role. This permission is usually available to ML administrators and roles with least-privilege permissions for ML practitioners.

You can temporarily assume an IAM role in the AWS Management Console by switching roles. For more information about methods for using roles, see Using IAM roles in the IAM User Guide.

Step 1. Enter role information

Provide a name to use as the unique suffix of your new SageMaker role. By default, the prefix "sagemaker-" is added to every role name for easier search in the IAM console. For example, if you name your role test-123 during role creation, your role shows up as sagemaker-test-123 in the IAM console. You can optionally add a description of your role to provide additional details.

Then, choose from one of the available personas to get suggested permissions for personas such as data scientists, data engineers, or machine learning operations (MLOps) engineers. For information on available personas and their suggested permissions, see Persona reference. To create a role without any suggested permissions to guide you, choose Custom Role Settings.
Note

We recommend that you first use the role manager to create a SageMaker Compute Role so that SageMaker compute resources have the ability to perform tasks such as training and inference. Use the SageMaker Compute Role persona to create this role with the role manager. After creating a SageMaker Compute Role, take note of its ARN for future use.

Network and encryption conditions

We recommend that you activate VPC customization to use VPC configurations, subnets, and security groups with IAM policies associated with your new role. When VPC customization is activated, IAM policies for ML activities that interact with VPC resources are scoped down for least-privilege access. VPC customization is not activated by default. For more details on recommended networking architecture, see Networking architecture in the AWS Technical Guide.

You can also use a KMS key to encrypt, decrypt, and re-encrypt data for regulated workloads with highly sensitive data. When AWS KMS customization is activated, IAM policies for ML activities that support custom encryption keys are scoped down for least-privilege access. For more information, see Encryption with AWS KMS in the AWS Technical Guide.

Step 2. Configure ML activities

Each Amazon SageMaker Role Manager ML activity includes suggested IAM permissions to provide access to relevant AWS resources. Some ML activities require that you add service role ARNs to complete setup. For information on predefined ML activities and their permissions, see ML activity reference. For information on adding service roles, see Service roles.

Based on the chosen persona, certain ML activities are already selected. You can deselect any suggested ML activities or select additional activities to create your own role. If you selected the Custom Role Settings persona, then no ML activities are preselected in this step.

You can add any additional AWS or customer-managed IAM policies to your role in Step 3: Add additional policies and tags.

Service roles

Some AWS services require a service role to perform actions on your behalf. If the ML activity that you selected requires you to pass a service role, then you must provide the ARN for that service role.
You can either create a new service role or use an existing one, such as a service role created with the SageMaker Compute Role persona. You can find the ARN of an existing role by selecting the role name in the Roles section of the IAM console. To learn more about service roles, see Creating a role for an AWS service.

**Step 3: Add additional policies and tags**

You can add any existing AWS or customer-managed IAM policies to your new role. For information on existing SageMaker policies, see AWS Managed Policies for Amazon SageMaker. You can also check your existing policies in the Roles section of the IAM console.

Optionally, use tag-based policy conditions to assign metadata information to categorize and manage AWS resources. Each tag is represented by a key-value pair. For more information, see Controlling access to AWS resources using tags.

**Review role**

Take the time to review all of the information associated with your new role. Choose Previous to go back and edit any of the information. When you are ready to create your role, choose Create role. This generates a role with permissions for your selected ML activities. You can view your new role in the Roles section of the IAM console.

**Using the role manager (AWS CDK)**

Use the AWS Cloud Development Kit (AWS CDK) with Amazon SageMaker Role Manager to programmatically create roles and set permissions. You can use the AWS CDK to accomplish any task that you could perform using the AWS Management Console. The programmatic access of the CDK makes it easier to provide permissions that give your users access to specific resources. For more information about the AWS CDK, see What is AWS CDK?

---

**Important**

You must use the SageMaker Compute Role persona to create a SageMaker Compute Role. For more information about the compute persona, see SageMaker compute persona. For code that you can use to create the compute role within the AWS CDK, see Grant permissions to a Compute persona.

The following are examples of tasks that you can perform in the AWS CDK:
• Create IAM roles with granular permissions for machine learning (ML) personas, such as Data Scientists and MLOps Engineers.
• Grant permissions to CDK constructs from ML personas or ML activities.
• Set ML activity condition parameters.
• Enable global Amazon VPC and AWS Key Management Service conditions and set values for them.
• Choose from all versions of the ML activities for your users without causing disruptions in their access.

There are common AWS tasks related to machine learning (ML) with SageMaker that require specific IAM permissions. The permissions to perform the tasks are defined as ML activities in Amazon SageMaker Role Manager. ML activities specify a set of permissions that are linked to the IAM role. For example, the ML activity for Amazon SageMaker Studio Classic has all of the permissions that a user needs to access Studio Classic. For more information about ML activities, see ML activity reference.

When you're creating roles, you first define the constructs for the ML persona or the ML activity. A construct is a resource within the AWS CDK stack. For example, a construct could be an Amazon S3 bucket, an Amazon VPC subnet, or an IAM role.

As you're creating the persona or activity, you can limit the permissions associated with that persona or activity to specific resources. For example, you can customize the activity to only provide permissions for a specific subnet within an Amazon VPC.

After you've defined permissions, you can create roles and then pass those roles to create other resources, such as SageMaker notebook instances.

The following are code examples in Typescript for tasks that you can accomplish using the CDK. When you create an activity, you specify an ID and the options for the activity's construct. The options are dictionaries that specify the required parameters for the activities, such as an Amazon S3. You pass an empty dictionary for activities that don't have required parameters.

**Grant permissions to a Compute persona**

The following code creates a Data Scientist ML persona with a set of ML activities specific to the persona. The permissions from ML activities only apply to the Amazon VPC and AWS KMS configurations specified in the persona construct. The following code creates a class for a Data
Grant permissions to a Data Scientist persona

The following code creates a Data Scientist ML persona with a set of ML activities specific to the persona. The permissions from ML activities only apply to the VPC and KMS configurations specified in the persona construct. The following code creates a class for a Data Scientist persona. The ML activities are defined in the activities list. The Amazon VPC permissions and the AWS KMS permissions are defined as optional parameters outside of the activities list.

After you’ve defined the class, you can create a role as a construct within the AWS CDK stack. You can also create a notebook instance. The person who is using the IAM role that you’ve created in the following code can access the notebook instance when they log in to their AWS account.

```typescript
export class myCDKStack extends cdk.Stack {
    constructor(scope: cdk.App, id: string, props?: cdk.StackProps) {
        super(scope, id, props);

        const persona = new Persona(this, 'example-persona-id', {
            activities: [
                Activity.accessAwsServices(this, 'example-id1', {})
            ]
        });

        const role = persona.createRole(this, 'example-IAM-role-id', 'example-IAM-role-name');
    }
}
```
const persona = new Persona(this, 'example-persona-id', {
  activities: [
    Activity.runStudioAppsV2(this, 'example-id1', {}),
    Activity.manageJobs(this, 'example-id2', {rolesToPass:
      [iam.Role.fromRoleName('example-IAM-role-name')]})],
    Activity.manageModels(this, 'example-id3', {rolesToPass:
      [iam.Role.fromRoleName('example-IAM-role-name')]})],
    Activity.manageExperiments(this, 'example-id4', {}),
    Activity.visualizeExperiments(this, 'example-id5', {}),
    Activity.accessS3Buckets(this, 'example-id6', {s3buckets: [s3.S3Bucket.fromBucketName('DOC-EXAMPLE-BUCKET')]})
  ],
  // optional: to configure VPC permissions
  subnets: [ec2.Subnet.fromSubnetId('example-VPC-subnet-id')],
  securityGroups: [ec2.SecurityGroup.fromSecurityGroupId('example-VPC-security-group-id')],
  // optional: to configure KMS permissions
  dataKeys: [kms.Key.fromKeyArn('example-KMS-key-ARN')],
  volumeKeys: [kms.Key.fromKeyArn('example-KMS-key-ARN')],
});

const role = persona.createRole(this, 'example-IAM-role-id', 'example-IAM-role-name');

const notebookInstance = new CfnNotebookInstance(this, 'example-notebook-instance-name', { RoleArn: role.RoleArn, ...});

Grant permissions to an ML Ops persona

The following code creates an ML Ops persona with a set of ML activities specific to the persona. The permissions from ML activities only apply to the Amazon VPC and AWS KMS configurations specified in the persona construct. The following code creates a class for an ML Ops persona. The ML activities are defined in the activities list. The VPC permissions and the KMS permissions are defined as optional parameters outside of the activities list.

After you’ve defined the class, you can create a role as a construct within the AWS CDK stack. You can also create an Amazon SageMaker Studio Classic user profile. The person who is using the IAM role that you’ve created in the following code can open SageMaker Studio Classic when they log in to their AWS account.
Grant permissions to a construct

The following code creates an ML Ops persona with a set of ML activities specific to the persona. The following code creates a class for a ML Ops persona. The ML activities are defined in the activities list.

After you’ve defined the class, you can create a role as a construct within the AWS CDK stack. You can also create a notebook instance. The code grants permissions from the ML activities to the IAM role of the Lambda function.
Grant permissions for a single ML activity

The following code creates an ML activity and creates a role from the activity. The permissions from the activity only apply to the VPC and KMS configuration that you specify for the user.

```typescript
export class myCDKStack extends cdk.Stack {
  constructor(scope: cdk.App, id: string, props?: cdk.StackProps) {
    super(scope, id, props);

    const activity = Activity.manageJobs(this, 'example-activity-id', {
      rolesToPass: [iam.Role.fromRoleName('example-IAM-role-name')],
      subnets: [ec2.Subnet.fromSubnetId('example-VPC-subnet-id')],
      securityGroups: [ec2.SecurityGroup.fromSecurityGroupId('example-VPC-security-group-id')],
      dataKeys: [kms.Key.fromKeyArn('example-KMS-key-ARN')],
      volumeKeys: [kms.Key.fromKeyArn('example-KMS-key-ARN')],
    });
  }
}
```
Create a role and give it permissions for a single activity

The following code creates an IAM role for a single ML activity.

```javascript
export class myCDKStack extends cdk.Stack {
  constructor(scope: cdk.App, id: string, props?: cdk.StackProps) {
    super(scope, id, props);

    const activity = Activity.manageJobs(this, 'example-activity-id', {
      rolesToPass: [iam.Role.fromRoleName('example-IAM-role-name')],
    });

    activity.create_role(this, 'example-IAM-role-id', 'example-IAM-role-name')
  }
}
```

Persona reference

Amazon SageMaker Role Manager provides suggested permissions for a number of ML personas. These include user execution roles for common ML practitioner responsibilities as well as service execution roles for common AWS service interactions needed to work with SageMaker.

Each persona has suggested permissions in the form of selected ML activities. For information on predefined ML activities and their permissions, see ML activity reference.

Data scientist persona

Use this persona to configure permissions to perform general machine learning development and experimentation in a SageMaker environment. This persona includes the following preselected ML activities:

- Run Studio Classic Applications
• Manage ML Jobs
• Manage Models
• Manage Experiments
• Search and Visualize Experiments
• Amazon S3 Bucket Access

**MLOps persona**

Choose this persona to configure permissions for operational activities. This persona includes the following preselected ML activities:

• Run Studio Classic Applications
• Manage Models
• Manage Endpoints
• Manage Pipelines
• Search and Visualize Experiments

**SageMaker compute persona**

**Note**

We recommend that you first use the role manager to create a SageMaker Compute Role so that SageMaker compute resources can perform tasks such as training and inference. Use the SageMaker Compute Role persona to create this role with the role manager. After creating a SageMaker Compute Role, take note of its ARN for future use.

This persona includes the following preselected ML activity:

• Access Required AWS Services

**ML activity reference**

ML activities are common AWS tasks related to machine learning with SageMaker that require specific IAM permissions. Each **persona** suggests related ML activities when creating a role with
Amazon SageMaker Role Manager. You can select any additional ML activities or deselect any suggested ML activities to create a role that meets your unique business needs.

Amazon SageMaker Role Manager provides predefined permissions for the following ML activities:

<table>
<thead>
<tr>
<th>ML activity</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Access Required AWS Services</td>
<td>Permissions to access Amazon S3, Amazon ECR, Amazon CloudWatch, and Amazon EC2. Required for execution roles for jobs and endpoints.</td>
</tr>
<tr>
<td>Run Studio Classic Applications</td>
<td>Permissions to operate within a Studio Classic environment. Required for domain and user profile execution roles.</td>
</tr>
<tr>
<td>Manage ML Jobs</td>
<td>Permissions to audit, query lineage, and visualize experiments.</td>
</tr>
<tr>
<td>Manage Models</td>
<td>Permissions to manage SageMaker jobs across their lifecycles.</td>
</tr>
<tr>
<td>Manage Endpoints</td>
<td>Permissions to manage SageMaker endpoint deployments and updates.</td>
</tr>
<tr>
<td>Manage Pipelines</td>
<td>Permissions to manage SageMaker pipelines and pipeline executions.</td>
</tr>
<tr>
<td>Manage Experiments</td>
<td>Permissions to manage SageMaker experiments and trials.</td>
</tr>
<tr>
<td>Search and Visualize Experiments</td>
<td>Permissions to audit, query lineage, and visualize experiments.</td>
</tr>
<tr>
<td>Manage Model Monitoring</td>
<td>Permissions to manage monitoring schedules for SageMaker Model Monitor.</td>
</tr>
<tr>
<td>S3 Full Access</td>
<td>Permissions to perform all Amazon S3 operations.</td>
</tr>
<tr>
<td>ML activity</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------</td>
<td>--------------------------------------------------------------</td>
</tr>
<tr>
<td>S3 Bucket Access</td>
<td>Permissions to perform operations on specified S3 buckets.</td>
</tr>
<tr>
<td>Query Athena Workgroups</td>
<td>Permissions to run and manage Amazon Athena queries.</td>
</tr>
</tbody>
</table>

**Launch Studio Classic**

Use your persona-focused roles to launch Studio Classic. If you are an administrator, you can give your users access to Studio Classic and have them assume their persona role either directly through the AWS Management Console or through the AWS IAM Identity Center.

**Launch Studio Classic with AWS Management Console**

For data scientists or other users to assume their given persona through the AWS Management Console, they require a console role to get to the Studio Classic environment.

You cannot use Amazon SageMaker Role Manager to create a role that grants permissions to the AWS Management Console. However, after creating a service role in the role manager, you can go to the IAM console to edit the role and add a user access role. The following is an example of a role that provides user access to the AWS Management Console:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "DescribeCurrentDomain",
      "Effect": "Allow",
      "Action": "sagemaker:DescribeDomain",
    },
    {
      "Sid": "RemoveErrorMessagesFromConsole",
      "Effect": "Allow",
      "Action": [
```
In the Studio Classic control panel, choose **Add User** to create a new user. In the **General Settings** section, give your user a name and set the **Default execution role** for the user to be the role that you created using Amazon SageMaker Role Manager.

On the next screen, choose the appropriate Jupyter Lab version, and whether to turn on SageMaker Jumpstart and SageMaker Project templates. Then choose **Next**. On the SageMaker Canvas settings page, choose whether to turn on SageMaker Canvas support, and additionally whether to allow for timeseries forecasting in SageMaker Canvas. Then choose **Submit**.

Your new user should now be visible in the Studio Classic control panel. To test this user, choose **Studio** from the **Launch app** dropdown list in the same row as the user’s name.
Launch Studio Classic with IAM Identity Center

To assign IAM Identity Center users to execution roles, the user must first exist in the IAM Identity Center directory. For more information, see Manage identities in IAM Identity Center in the AWS IAM Identity Center.

Note

Your IAM Identity Center Authentication directory and Studio Classic domain must be in the same AWS Region.

1. To assign IAM Identity Center users to your Studio Classic domain, choose Assign users and Groups in the Studio Classic control panel. On the Assign users and groups screen select your data scientist user, and then choose Assign Users and Groups.
2. After the user is added to the Studio Classic control panel, choose the user to open the user details screen.
3. On the User details screen, choose Edit.
4. On the Edit user profile screen, under General settings, modify the Default execution role to match the user execution role you've created for your data scientists.
5. Choose Next through the rest of the settings pages, and choose Submit to save your changes.

When your data scientist or other user logs into the IAM Identity Center portal, they see a tile for this Studio Classic domain. Choosing that tile logs them into Studio Classic with their assigned user execution role.

Role Manager FAQs

Refer to the following FAQ items for answers to commonly asked questions about Amazon SageMaker Role Manager.

Q. How can I access Amazon SageMaker Role Manager?

A: You can access Amazon SageMaker Role Manager through multiple location in the Amazon SageMaker console. For information about accessing role manager and using it to create a role, see Using the role manager (console).
Q. What are personas?

A: Personas are preconfigured groups of permissions based on common machine learning (ML) responsibilities. For example, the data science persona suggests permissions for general machine learning development and experimentation in a SageMaker environment, while the MLOps persona suggests permissions for ML activities related to operations.

Q. What are ML activities?

A: ML activities are common AWS tasks related to machine learning with SageMaker that require specific IAM permissions. Each persona suggests related ML activities when creating a role with Amazon SageMaker Role Manager. ML activities include tasks such as Amazon S3 full access or searching and visualizing experiments. For more information, see ML activity reference.

Q. Are the roles that I create with the role manager AWS Identity and Access Management (IAM) roles?

A: Yes. Roles created using the Amazon SageMaker Role Manager are IAM roles with customized access policies. You can view created roles in the Roles section of the IAM console.

Q. How can I view the roles that I created using Amazon SageMaker Role Manager?

A: You can view created roles in the Roles section of the IAM console. By default, the prefix "sagemaker-" is added to every role name for easier search in the IAM console. For example, if you named your role test-123 during role creation, your role shows up as sagemaker-test-123 in the IAM console.

Q. Can I modify a role made with Amazon SageMaker Role Manager once it is created?

A: Yes. You can modify the roles and policies created by Amazon SageMaker Role Manager through the IAM console. For more information, see Modifying a role in the AWS Identity and Access Management User Guide.

Q. Can I attach my own policies to roles created using Amazon SageMaker Role Manager?

A: Yes. You can attach any AWS or customer-managed IAM policies from your account to the role that you create using Amazon SageMaker Role Manager.

Q. How many policies can I add to a role that I create with Amazon SageMaker Role Manager?

A: The maximum limit for attaching managed policies to an IAM role or user is 20. The maximum character size limit for managed policies is 6,144. For more information, see IAM object quotas and IAM and AWS Security Token Service quotas name requirements, and character limits.
Q. Can I add conditions to ML activities?

A: Any conditions that you provide in Step 1. Enter role information of the Amazon SageMaker Role Manager, such as subnets, security groups, or KMS keys, are automatically passed to any ML activities selected in Step 2. Configure ML activities. You can also add additional conditions to ML activities if necessary. For example, you might also add InstanceTypes or IntercontainerTrafficEncryption conditions to the Manage Training Jobs activity.

Q. Can I use tagging to manage access to any AWS resource?

A: You can add tags to your role in Step 3: Add additional policies and tags of the Amazon SageMaker Role Manager. To successfully manage AWS resources using tags, you must add the same tag to both the role and any associated policies. For example, you can add a tag to a role and to an Amazon S3 bucket. Then, because the role passes the tag to the SageMaker session, only a user with that role can access that S3 bucket. You can add tags to a policy through the IAM console.

For more information, see Tagging IAM roles in the AWS Identity and Access Management User Guide.

Q. Can I use Amazon SageMaker Role Manager to create a role to access the AWS Management Console?

A: No. However, after creating a service role in the role manager, you can go to the IAM console to edit the role and add a human access role in IAM console.

Q. What is difference between a user federation role and a SageMaker execution role?

A: A user federation role is directly assumed by a user to access AWS resources such as access to the AWS Management Console. A SageMaker execution role is assumed by the SageMaker service to perform a function on behalf of a user or an automation tool. For example, when a user opens a Studio Classic instance, Studio Classic assumes the execution role associated with the user profile in order to access AWS resources on the behalf of the user. If the user profile does not specify an execution role, then the execution role is specified at the Amazon SageMaker domain level.

Q. If I am using a custom web application that accesses Studio Classic through a presigned url, what role is used?

A: If you use a custom web application to access Studio Classic, then you have a hybrid user federation role and SageMaker execution role. Be sure that this role has least privilege permissions for both what the user can do and what Studio Classic can do on the associated user's behalf.
Q: Can I use Amazon SageMaker Role Manager with AWS IAM Identity Center authentication for my Studio Classic domain?

A: AWS IAM Identity Center Studio Classic Cloud Applications use a Studio Classic execution role to grant permissions to federated users. This execution role can be specified at the Studio Classic IAM Identity Center user profile level or the default domain level. User identities and groups must be synchronized into IAM Identity Center and the Studio Classic user profile must be created with IAM Identity Center user assignment using CreateUserProfile. For more information, see Launch Studio Classic with IAM Identity Center.

Access control for notebooks

You must use different procedures to control access to Amazon SageMaker Studio Classic notebooks and SageMaker notebook instances because they have different runtime environments. Studio Classic uses file system permissions and containers to control access to Studio Classic notebooks and isolation of users. A SageMaker notebook instance gives users that login to the notebook instance default root access. The following topics describe how to change permissions for both kinds of notebooks.

Topics

- Access control and setting permissions for SageMaker Studio notebooks
- Control root access to a SageMaker notebook instance

Access control and setting permissions for SageMaker Studio notebooks

Amazon SageMaker Studio uses filesystem and container permissions for access control and isolation of Studio users and notebooks. This is one of the major differences between Studio notebooks and SageMaker notebook instances. This topic describes how permissions are set up to avoid security threats, what SageMaker does by default, and how the customer can customize the permissions. For more information about Studio notebooks and their runtime environment, see Use Amazon SageMaker Studio Classic Notebooks.

SageMaker app permissions

A run-as user is a POSIX user/group which is used to run the JupyterServer app and KernelGateway apps inside the container.

The run-as user for the JupyterServer app is sagemaker-user (1000) by default. This user has sudo permissions to enable the installation of dependencies such as yum packages.
The run-as user for the KernelGateway apps is root (0) by default. This user is able to install dependencies using pip/apt-get/conda.

Due to user remapping, neither user is able to access resources or make changes to the host instance.

**User remapping**

SageMaker performs user-remapping to map a user inside the container to a user on the host instance outside the container. The range of user IDs (0 - 65535) in the container are mapped to non-privileged user IDs above 65535 on the instance. For example, sagemaker-user (1000) inside the container might map to user (200001) on the instance, where the number in parentheses is the user ID. If the customer creates a new user/group inside the container, it won't be privileged on the host instance regardless of the user/group ID. The root user of the container is also mapped to a non-privileged user on the instance. For more information, see *Isolate containers with a user namespace*.

ℹ️ **Note**

Files created by the user sagemaker-user may look like they are owned by sagemaker-studio (uid 65534). This is a side effect of a fast app creation mode where SageMaker container images are pre-pulled, allowing applications to start in under a minute. If your application requires the file owner uid and the process owner uid to match, ask the customer service to remove your account number from the image pre-pull feature.

**Custom image permissions**

Customers can bring their own custom SageMaker images. These images can specify a different run-as user/group to launch the KernelGateway app. The customer can implement fine grained permission control inside the image, for example, to disable root access or perform other actions. The same user remapping applies here. For more information, see *Bring your own SageMaker image*.

**Container isolation**

Docker keeps a list of default capabilities that the container can use. SageMaker doesn’t add additional capabilities. SageMaker adds specific route rules to block requests to Amazon EFS and the *instance metadata service* (IMDS) from the container. Customers can’t change these route rules from the container. For more information, see *Runtime privilege and Linux capabilities*.
App metadata access

Metadata used by running apps are mounted to the container with read-only permission. Customers aren't able to modify this metadata from the container. For the available metadata, see Get Studio Classic Notebook and App Metadata.

User isolation on EFS

When you onboard to Studio, SageMaker creates an Amazon Elastic File System (EFS) volume for your domain that is shared by all Studio users in the domain. Each user gets their own private home directory on the EFS volume. This home directory is used to store the user's notebooks, Git repositories, and other data. To prevent other users in the domain from accessing the user's data, SageMaker creates a globally unique user ID for the user's profile and applies it as a POSIX user/group ID for the user's home directory.

EBS access

An Amazon Elastic Block Store (Amazon EBS) volume is attached to the host instance and shared across all images. It's used for the root volume of the notebooks and stores temporary data that's generated inside the container. The storage isn't persisted when the instance running the notebooks is deleted. The root user inside the container can't access the EBS volume.

IMDS access

Due to security concerns, access to the Amazon Elastic Compute Cloud (Amazon EC2) Instance Metadata Service (IMDS) is unavailable in SageMaker Studio. For more information on IMDS, see Instance metadata and user data.

Control root access to a SageMaker notebook instance

By default, when you create a notebook instance, users that log into that notebook instance have root access. Data science is an iterative process that might require the data scientist to test and use different software tools and packages, so many notebook instance users need to have root access to be able to install these tools and packages. Because users with root access have administrator privileges, users can access and edit all files on a notebook instance with root access enabled.

If you don't want users to have root access to a notebook instance, when you call CreateNotebookInstance or UpdateNotebookInstance operations, set the RootAccess field to Disabled. You can also disable root access for users when you create or update a notebook instance in the Amazon SageMaker console. For information, see Step 1: Create an Amazon SageMaker Notebook Instance.
Note
Lifecycle configurations need root access to be able to set up a notebook instance. Because of this, lifecycle configurations associated with a notebook instance always run with root access even if you disable root access for users.

Note
For security reasons, Rootless Docker is installed on root-disabled notebook instances instead of regular Docker. For more information, see Run the Docker daemon as a non-root user (Rootless mode)

Amazon SageMaker API Permissions: Actions, Permissions, and Resources Reference

When you are setting up access control and writing a permissions policy that you can attach to an IAM identity (an identity-based policy), use the following as a reference. The each Amazon SageMaker API operation, the corresponding actions for which you can grant permissions to perform the action, and the AWS resource for which you can grant the permissions. You specify the actions in the policy's Action field, and you specify the resource value in the policy's Resource field.

Note
Except for the ListTags API, resource-level restrictions are not available on List- calls. Any user calling a List- API will see all resources of that type in the account.

To express conditions in your Amazon SageMaker policies, you can use AWS-wide condition keys. For a complete list of AWS-wide keys, see Available Keys in the IAM User Guide.

Warning
Some SageMaker API actions may still be accessible through the Search API. For example, if a user has an IAM policy that denies permissions to a Describe call for a particular SageMaker resource, that user can still access the description information through the
Amazon SageMaker API Operations and Required Permissions for Actions

<table>
<thead>
<tr>
<th>Amazon SageMaker API Operations</th>
<th>Required Permissions (API Actions)</th>
<th>Resources</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Amazon SageMaker API Operations</strong></td>
<td><strong>Required Permissions (API Actions)</strong></td>
<td><strong>Resources</strong></td>
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<tr>
<td><strong>ListEarthObservationJobs</strong></td>
<td>sagemaker-geospatial:ListEarthObservationJobs</td>
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<td><strong>ListRasterDataCollections</strong></td>
<td>sagemaker-geospatial:ListRasterDataCollections</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<td>sagemaker-geospatial:StartVectorEnrichmentJob</td>
<td>**arn:**aws:sagemaker-geospatial l: <code>region:account-id</code> d: <code>vector-enrichment-job/id</code></td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<td>StopEarth ObservationJob</td>
<td>sagemaker-geospatial:StopEarthObservationJob</td>
<td>arn:aws:sagemaker-geospatial\n\l: region:account-id:earth-observation-job/id</td>
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<td>StopVectorEnrichmentJob</td>
<td>sagemaker-geospatial:StopVectorEnrichmentJob</td>
<td>arn:aws:sagemaker-geospatial\n\l: region:account-id:vector-enrichment-job/id</td>
</tr>
<tr>
<td></td>
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<td>arn:aws:sagemaker-geospatial\n\l: region:account-id:vector-enrichment-job/id</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
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</tr>
<tr>
<td>AddTags</td>
<td>sagemaker:AddTags</td>
<td>arn:aws:sagemaker: region:account-id : *</td>
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<tr>
<td>CreateApp</td>
<td>sagemaker:CreateApp</td>
<td>arn:aws:sagemaker: region:account-id : app/domain-id /user-profile-name /app-type/appName</td>
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<td>CreateApp ImageConfig</td>
<td>sagemaker:CreateAppImageConfig</td>
<td>arn:aws:sagemaker: region:account-id : app-image-config/appImageConfig/appImageConfigName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<tr>
<td><code>CreateAutoMLJob</code></td>
<td><code>sagemaker:CreateAutoMLJob</code></td>
<td><code>arn:aws:sagemaker:</code> region:account-id :autoMLJob Name <code>iam:PassRole</code></td>
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<tr>
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<td>The following permission is required only if any of the associated ResourceConfig have a specified VolumeKms KeyId and the associated role does not have a policy that permits this action:</td>
<td></td>
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<tr>
<td></td>
<td><code>kms:CreateGrant</code></td>
<td></td>
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<tr>
<td><code>CreateAutoMLJobV2</code></td>
<td><code>sagemaker:CreateAutoMLJobV2</code></td>
<td><code>arn:aws:sagemaker:</code> region:account-id :autoMLJob Name <code>iam:PassRole</code></td>
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<td></td>
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<td><code>kms:CreateGrant</code></td>
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<td><strong>CreateDomain</strong></td>
<td>sagemaker:CreateDomain</td>
<td>arn:aws:sagemaker: region:account-id :domain/domain-id</td>
</tr>
<tr>
<td></td>
<td>iam:GetServiceLinkedRole</td>
<td></td>
</tr>
<tr>
<td></td>
<td>iam:PassRole</td>
<td></td>
</tr>
<tr>
<td></td>
<td>elasticfilesystem:</td>
<td></td>
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<tr>
<td></td>
<td>CreateFileSystem</td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:CreateGrant</td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:Decrypt</td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:DescribeKey</td>
<td></td>
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<tr>
<td></td>
<td>kms:GenerateDataKeyWithoutPlainText</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Required if a KMS customer managed key is specified for KmsKeyId:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>sagemaker:CreateApp</td>
<td></td>
</tr>
<tr>
<td><strong>CreateEndpoint</strong></td>
<td>sagemaker:CreateEndpoint</td>
<td>arn:aws:sagemaker: region:account-id :endpoint/endpointName</td>
</tr>
<tr>
<td></td>
<td>kms:CreateGrant</td>
<td>arn:aws:sagemaker: region:account-id :endpointConfig/endpointConfigName</td>
</tr>
<tr>
<td></td>
<td>(required only if the associated EndPointConfig has a KmsKeyId specified)</td>
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</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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</tr>
<tr>
<td>CreateEndpointConfig</td>
<td>sagemaker:CreateEndpointConfig</td>
<td>arn:aws:sagemaker: region:account-id :endpoint-config/endpointConfigName</td>
</tr>
<tr>
<td>CreateFlowDefinition</td>
<td>sagemaker:CreateFlowDefinition</td>
<td>arn:aws:sagemaker: region:account-id :flow-definition/ flowDefinitionName</td>
</tr>
<tr>
<td>CreateHumanTaskUi</td>
<td>sagemaker:CreateHumanTaskUi</td>
<td>arn:aws:sagemaker: region:account-id :human-task-ui/ humanTaskUiName</td>
</tr>
</tbody>
</table>

The following permissions are required only if you specify an encryption key:

- kms:CreateGrant
- kms:Decrypt
- kms:DescribeKey
- kms:GenerateDataKey
<table>
<thead>
<tr>
<th>Amazon SageMaker API Operations</th>
<th>Required Permissions (API Actions)</th>
<th>Resources</th>
</tr>
</thead>
<tbody>
<tr>
<td>CreateHyperParameterTuningJob</td>
<td>sagemaker:CreateHyperParameterTuningJob, iam:PassRole</td>
<td>arn:aws:sagemaker: region:account-id :hyper-parameter-tuning-job/hyperParameterTuningJobName</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CreateModel</td>
<td>sagemaker:CreateModel, iam:PassRole</td>
<td>arn:aws:sagemaker: region:account-id :model/modelName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<tr>
<td>--------------------------------</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<tr>
<td>-------------------------------</td>
<td>-----------------------------------</td>
<td>-----------</td>
</tr>
<tr>
<td><strong>CreateNotebookInstance</strong></td>
<td>sagemaker:CreateNotebookInstance</td>
<td>arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName</td>
</tr>
<tr>
<td></td>
<td>iam:PassRole</td>
<td></td>
</tr>
</tbody>
</table>

The following permissions are required only if you specify a VPC for your notebook instance:

- ec2:CreateNetworkInterface
- ec2:DescribeSecurityGroups
- ec2:DescribeSubnets
- ec2:DescribeVpcs

The following permission is required only if you specify a VPC and an elastic inference accelerator for your notebook instance:

- ec2:DescribeVpcEndpoints

The following permissions are required only if you specify an encryption key:

- kms:DescribeKey
- kms:CreateGrant
The following permission is required only if you specify an AWS Secrets Manager secret to access a private Git repository:

```
secretsmanager:GetSecretValue
```

<table>
<thead>
<tr>
<th>Amazon SageMaker API Operations</th>
<th>Required Permissions (API Actions)</th>
<th>Resources</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CreatePipeline</strong></td>
<td>sagemaker:CreatePipeline</td>
<td>arn:aws-partition:sagemaker:region:account-id:pipeline/pipeline-name</td>
</tr>
<tr>
<td></td>
<td>iam:PassRole</td>
<td>arn:aws-partition:iam::account-id:role/role-name</td>
</tr>
<tr>
<td><strong>CreatePresignedDomainUrl</strong></td>
<td>sagemaker:CreatePresignedDomainUrl</td>
<td>arn:aws:sagemaker:region:account-id:app/domain-id/userProfileName /*</td>
</tr>
<tr>
<td><strong>CreatePresignedNotebookInstanceUrl</strong></td>
<td>sagemaker:CreatePresignedNotebookInstanceUrl</td>
<td>arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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</tr>
<tr>
<td><strong>CreateProcessingJob</strong></td>
<td>sagemaker:CreateProcessingJob</td>
<td>arn:aws:sagemaker:region:account-id:processing-job/processingJobName</td>
</tr>
<tr>
<td></td>
<td>iam:PassRole</td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:CreateGrant</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(required only if the associated ProcessingResources has a specified VolumeKmsKeyId and the associated role does not have a policy that permits this action)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ec2:CreateNetworkInterface</td>
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<tr>
<td></td>
<td>(required only if you specify a VPC)</td>
<td></td>
</tr>
<tr>
<td><strong>CreateSpace</strong></td>
<td>sagemaker:CreateSpace</td>
<td>arn:aws:sagemaker:region:account-id:space/domain-id/spaceName</td>
</tr>
<tr>
<td><strong>CreateStudioLifecycleConfig</strong></td>
<td>sagemaker:CreateStudioLifecycleConfig</td>
<td>arn:aws:sagemaker:region:account-id:studio-lifecycle-config/*</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<td>---------------------------------</td>
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</tr>
<tr>
<td>&quot;CreateTrainingJob&quot;</td>
<td>sagemaker:CreateTrainingJob</td>
<td>arn:aws:sagemaker: region:account-id :training-job/ trainingJobName</td>
</tr>
<tr>
<td></td>
<td>iam:PassRole</td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:CreateGrant</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(required only if the associated ResourceConfig has a specified VolumeKmsKeyId and the associated role does not have a policy that permits this action)</td>
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<tr>
<td></td>
<td>kms:CreateGrant</td>
<td></td>
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<tr>
<td></td>
<td>(required only if the associated TransformResources has a specified VolumeKmsKeyId and the associated role does not have a policy that permits this action)</td>
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</tr>
<tr>
<td>&quot;CreateUserProfile&quot;</td>
<td>sagemaker:CreateUserProfile</td>
<td>arn:aws:sagemaker: region:account-id :user-profile/domain-id/userProfileName</td>
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<tr>
<td></td>
<td>iam:PassRole</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
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<tr>
<td>CreateWorkforce</td>
<td>sagemaker:CreateWorkforce</td>
<td>arn:aws:sagemaker: region:account-id :workforce/*</td>
</tr>
<tr>
<td></td>
<td>cogito-idp:DescribeUserPoolClient</td>
<td></td>
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<tr>
<td></td>
<td>cogito-idp:UpdateUserPool</td>
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<tr>
<td></td>
<td>cogito-idp:DescribeUserPool</td>
<td></td>
</tr>
<tr>
<td></td>
<td>cogito-idp:UpdateUserPoolClient</td>
<td></td>
</tr>
<tr>
<td>CreateWorkteam</td>
<td>sagemaker:CreateWorkteam</td>
<td>arn:aws:sagemaker: region:account-id :workteam/private-crowd/ work team name</td>
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<tr>
<td></td>
<td>cogito-idp:DescribeUserPoolClient</td>
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<td></td>
<td>cogito-idp:UpdateUserPool</td>
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<tr>
<td></td>
<td>cogito-idp:DescribeUserPool</td>
<td></td>
</tr>
<tr>
<td></td>
<td>cogito-idp:UpdateUserPoolClient</td>
<td></td>
</tr>
<tr>
<td>DeleteApp</td>
<td>sagemaker:DeleteApp</td>
<td>arn:aws:sagemaker: region:account-id :app domain-id /user-profile-name /app-type/appName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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</tr>
<tr>
<td>DeleteDomain</td>
<td>sagemaker:DeleteDomain</td>
<td>arn:aws:sagemaker:region:account-id :domain/domainId</td>
</tr>
<tr>
<td>DeleteEndpoint</td>
<td>sagemaker:DeleteEndpoint</td>
<td>arn:aws:sagemaker:region:account-id :endpoint/endpointName</td>
</tr>
<tr>
<td>DeleteEndpointConfig</td>
<td>sagemaker:DeleteEndpointConfig</td>
<td>arn:aws:sagemaker:region:account-id :endpoint-config/endpointConfigName</td>
</tr>
<tr>
<td>DeleteHumanLoop</td>
<td>sagemaker:DeleteHumanLoop</td>
<td>arn:aws:sagemaker:region:account-id :human-loop/humanLoopName</td>
</tr>
<tr>
<td>DeleteImage</td>
<td>sagemaker:DeleteImage</td>
<td>arn:aws:sagemaker:region:account-id :image/imageName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<tr>
<td><strong>DeleteModel</strong></td>
<td>sagemaker:DeleteModel</td>
<td>arn:aws:sagemaker:region:account-id:model/modelName</td>
</tr>
<tr>
<td><strong>DeleteModelPackage</strong></td>
<td>sagemaker:DeleteModelPackage</td>
<td>arn:aws:sagemaker:region:account-id:model-package/modelPackageName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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</tr>
<tr>
<td>DeleteNotebookInstance</td>
<td>sagemaker:DeleteNotebookInstance</td>
<td>arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName</td>
</tr>
<tr>
<td></td>
<td>ec2:DeleteNetworkInterface</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The following permission is required only if you specified a VPC for your notebook instance:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:DescribeKey</td>
<td></td>
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<tr>
<td></td>
<td>The following permissions are required only if you specified an encryption key when you created the notebook instance:</td>
<td></td>
</tr>
<tr>
<td>DeletePipeline</td>
<td>sagemaker:DeletePipeline</td>
<td>arn:aws-partition:sagemaker:region:account-id:pipeline/pipeline-name</td>
</tr>
<tr>
<td>DeleteSpace</td>
<td>sagemaker:DeleteSpace</td>
<td>arn:aws:sagemaker:region:account-id:space/domain-id/spaceName</td>
</tr>
<tr>
<td>DeleteTags</td>
<td>sagemaker:DeleteTags</td>
<td>arn:aws:sagemaker:region:account-id:*</td>
</tr>
<tr>
<td>DeleteUserProfile</td>
<td>sagemaker:DeleteUserProfile</td>
<td>arn:aws:sagemaker:region:account-id:user-profile/domain-id/userProfileName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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</tr>
<tr>
<td>DeleteWorkforce</td>
<td>sagemaker:DeleteWorkforce</td>
<td>arn:aws:sagemaker:region:account-id:workforce/*</td>
</tr>
<tr>
<td>DescribeApp</td>
<td>sagemaker:DescribeApp</td>
<td>arn:aws:sagemaker:region:account-id:app/domain-id/user-profile-name/app-type/appName</td>
</tr>
<tr>
<td>DescribeAutoMLJob</td>
<td>sagemaker:DescribeAutoMLJob</td>
<td>arn:aws:sagemaker:region:account-id:automl-job/automlJobName</td>
</tr>
<tr>
<td>DescribeAutoMLJobV2</td>
<td>sagemaker:DescribeAutoMLJobV2</td>
<td>arn:aws:sagemaker:region:account-id:automl-job/automlJobName</td>
</tr>
<tr>
<td>DescribeDomain</td>
<td>sagemaker:DescribeDomain</td>
<td>arn:aws:sagemaker:region:account-id:domain/domainId</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
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<tr>
<td>DescribeEndpoint</td>
<td>sagemaker:Describe Endpoint</td>
<td>arn:aws:sagemaker:region:account-id:endpoint/endpointName</td>
</tr>
<tr>
<td>DescribeEndpointConfig</td>
<td>sagemaker:Describe EndpointConfig</td>
<td>arn:aws:sagemaker:region:account-id:endpoint-config/endpointConfigName</td>
</tr>
<tr>
<td>DescribeHumanLoop</td>
<td>sagemaker:Describe HumanLoop</td>
<td>arn:aws:sagemaker:region:account-id:human-loop/humanLoopName</td>
</tr>
<tr>
<td>DescribeHyperParameterTuningJob</td>
<td>sagemaker:Describe HyperParameterTuningJob</td>
<td>arn:aws:sagemaker:region:account-id:hyper-parameter-tuning-job/hyperParameterTuningJob</td>
</tr>
<tr>
<td>DescribeImage</td>
<td>sagemaker:DescribeImage</td>
<td>arn:aws:sagemaker:region:account-id:image/imageName</td>
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<tr>
<td><strong>Amazon SageMaker API Operations</strong></td>
<td><strong>Required Permissions (API Actions)</strong></td>
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<tr>
<td>DescribeModel</td>
<td>sagemaker:DescribeModel</td>
<td>arn:aws:sagemaker: region:account-id :model/modelName</td>
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<tr>
<td>DescribeNotebookInstance</td>
<td>sagemaker:DescribeNotebookInstance</td>
<td>arn:aws:sagemaker: region:account-id :notebook-instance / notebookInstanceName</td>
</tr>
<tr>
<td>DescribePipeline</td>
<td>sagemaker:DescribePipeline</td>
<td>arn:aws-partition :sagemaker: region:account-id :pipeline/ pipeline-name</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<tr>
<td><strong>DescribePipelineDefinitionForExecution</strong></td>
<td>sagemaker:DescribePipelineDefinitionForExecution</td>
<td>arn:aws-partition:sagemaker:region:account-id: pipeline/ pipeline-name /execution/ execution-id</td>
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<tr>
<td><strong>DescribePipelineExecution</strong></td>
<td>sagemaker:DescribePipelineExecution</td>
<td>arn:aws-partition:sagemaker:region:account-id: pipeline/ pipeline-name /execution/ execution-id</td>
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<tr>
<td><strong>DescribeProcessingJob</strong></td>
<td>sagemaker:DescribeProcessingJob</td>
<td>arn:aws:sagemaker:region:account-id:processing-job/processingjobname</td>
</tr>
<tr>
<td><strong>DescribeSpace</strong></td>
<td>sagemaker:DescribeSpace</td>
<td>arn:aws:sagemaker:region:account-id:space/domain-id/spaceName</td>
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<tr>
<td><strong>DescribeSubscribedWorkteam</strong></td>
<td>sagemaker:DescribeSubscribedWorkteam</td>
<td>arn:aws:sagemaker:region:account-id:workteam/vendor-crowd/*</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
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<tr>
<td>DescribeUserProfile</td>
<td>sagemaker:DescribeUserProfile</td>
<td>arn:aws:sagemaker:region:account-id:user-profile/domain-id/userProfileName</td>
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<tr>
<td>DescribeWorkforce</td>
<td>sagemaker:DescribeWorkforce</td>
<td>arn:aws:sagemaker:region:account-id:workforce/*</td>
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<tr>
<td>InvokeEndpoint</td>
<td>sagemaker:InvokeEndpoint</td>
<td>arn:aws:sagemaker:region:account-id:endpoint/endpointName</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<tr>
<td>ListEndpointConfigs</td>
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<td>ListEndpoints</td>
<td>sagemaker:ListEndpoints</td>
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<tr>
<td>ListFlowDefinitions</td>
<td>sagemaker:ListFlowDefinitions</td>
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<td>ListHumanLoops</td>
<td>sagemaker:ListHumanLoops</td>
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<tr>
<td>ListHumanTaskUis</td>
<td>sagemaker:ListHumanTaskUis</td>
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<tr>
<td>ListImages</td>
<td>sagemaker:ListImages</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
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<td><strong>ListLabelingJobs</strong></td>
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<tr>
<td><strong>ListLabelingJobsForWorkteam</strong></td>
<td>sagemaker:ListLabelingJobForWorkteam</td>
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<tr>
<td><strong>ListModels</strong></td>
<td>sagemaker:ListModels</td>
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<tr>
<td><strong>ListNotebookInstances</strong></td>
<td>sagemaker:ListNotebookInstances</td>
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<tr>
<td><strong>ListPipelineExecutions</strong></td>
<td>sagemaker:ListPipelineExecutions</td>
<td>arn:aws-partition:sagemaker:region:account-id:mode:partition:pipeline/executionId</td>
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<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
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<td>ListPipelineParametersForExecution</td>
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<td>ListPipelines</td>
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</tr>
<tr>
<td>ListProcessingJobs</td>
<td>sagemaker:ListProcessingJobs</td>
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<tr>
<td>ListSpaces</td>
<td>sagemaker:ListSpaces</td>
<td>arn:aws:sagemaker:region:account-id:space/domain-id /*</td>
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<tr>
<td>ListSubscribedWorkteams</td>
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<td>aws-marketplace:ViewSubscriptions</td>
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<td>ListTags</td>
<td>sagemaker:ListTags</td>
<td>arn:aws:sagemaker:region:account-id :*</td>
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<td>ListTrainingJobs</td>
<td>sagemaker:ListTrainingJobs</td>
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<td>ListTransformJobs</td>
<td>sagemaker:ListTransformJobs</td>
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<tr>
<td>ListWorkteams</td>
<td>sagemaker:ListWorkteams</td>
<td>*</td>
</tr>
<tr>
<td>RetryPipelineExecution</td>
<td>sagemaker:RetryPipelineExecution</td>
<td>arn:aws-partition :sagemaker /:region:account-id :pipeline/ pipeline-name /execution/ execution-id</td>
</tr>
<tr>
<td>Search</td>
<td>sagemaker:Search</td>
<td>*</td>
</tr>
<tr>
<td>SendPipelineExecutionStepFailure</td>
<td>sagemaker:SendPipelineExecutionStepFailure</td>
<td>*</td>
</tr>
<tr>
<td>SendPipelineExecutionStepSuccess</td>
<td>sagemaker:SendPipelineExecutionStepSuccess</td>
<td>*</td>
</tr>
<tr>
<td>StartHumanLoop</td>
<td>sagemaker:StartHumanLoop</td>
<td>arn:aws:sagemaker: region:account-id :human-loop/ humanLoopName</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>------------------------------------</td>
<td>-----------</td>
</tr>
<tr>
<td><strong>StartNotebookInstance</strong></td>
<td>sagemaker:StartNotebookInstance</td>
<td>arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName</td>
</tr>
<tr>
<td></td>
<td>ec2:CreateNetworkInterface</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ec2:DescribeNetworkInterfaces</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ec2:DescribeSecurityGroups</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ec2:DescribeSubnets</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ec2:DescribeVpcs</td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>The following permission is</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>required only if you specify a</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>a VPC and an elastic inference</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>accelerator for your notebook</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>instance:</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ec2:DescribeVpcEndpoints</td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>The following permissions are</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>required only if you specified</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>an encryption key when you created</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>the notebook instance:</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td>kms:DescribeKey</td>
<td></td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td><code>kms:CreateGrant</code></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The following permission is required only if you specified an AWS Secrets Manager secret to access a private Git repository when you created the notebook instance:</td>
<td></td>
</tr>
<tr>
<td></td>
<td><code>secretsmanager:GetSecretValue</code></td>
<td></td>
</tr>
<tr>
<td><strong>StartPipelineExecution</strong></td>
<td><code>sagemaker:StartPipelineExecution</code></td>
<td><code>arn:aws-partition:sagemaker:region:account-id:pipeline/pipeline-name</code></td>
</tr>
<tr>
<td><strong>StopHumanLoop</strong></td>
<td><code>sagemaker:StopHumanLoop</code></td>
<td><code>arn:aws:sagemaker:region:account-id:human-loop/humanLoopName</code></td>
</tr>
<tr>
<td><strong>StopHyperParameterTuningJob</strong></td>
<td><code>sagemaker:StopHyperParameterTuningJob</code></td>
<td><code>arn:aws:sagemaker:region:account-id:hyper-parameter-tuning-job/hyperParameterTuningJob</code></td>
</tr>
<tr>
<td><strong>StopLabelingJob</strong></td>
<td><code>sagemaker:StopLabelingJob</code></td>
<td><code>arn:aws:sagemaker:region:account-id:labeling-job/labelingJobName</code></td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
</tr>
<tr>
<td>------------------------------------------------</td>
<td>------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------</td>
</tr>
<tr>
<td>StopNotebookInstance</td>
<td>sagemaker:StopNotebookInstance</td>
<td>arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName</td>
</tr>
<tr>
<td>StopPipelineExecution</td>
<td>sagemaker:StopPipelineExecution</td>
<td>arn:aws:partition:sagemaker:region:account-id:partition/pipeline/pipeline-name/excution/excution-execution-executionExecutionId</td>
</tr>
<tr>
<td>StopProcessingJob</td>
<td>sagemaker:StopProcessingJob</td>
<td>arn:aws:sagemaker:region:account-id:processing-job/processingJobName</td>
</tr>
<tr>
<td>UpdateDomain</td>
<td>sagemaker:UpdateDomain</td>
<td>arn:aws:sagemaker:region:account-id:domain/domainId</td>
</tr>
<tr>
<td>Amazon SageMaker API Operations</td>
<td>Required Permissions (API Actions)</td>
<td>Resources</td>
</tr>
<tr>
<td>--------------------------------</td>
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<td>-----------</td>
</tr>
<tr>
<td><strong>UpdateEndpoint</strong></td>
<td>sagemaker:UpdateEndpoint</td>
<td>arn:aws:sagemaker:region:account-id:endpoint/endpointName</td>
</tr>
<tr>
<td><strong>UpdateEndpointWeightsAndCapacities</strong></td>
<td>sagemaker:UpdateEndpointWeightsAndCapacities</td>
<td>arn:aws:sagemaker:region:account-id:endpoint/endpointName</td>
</tr>
<tr>
<td><strong>Amazon SageMaker API Operations</strong></td>
<td><strong>Required Permissions (API Actions)</strong></td>
<td><strong>Resources</strong></td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>--------------------------------------</td>
<td>---------------</td>
</tr>
</tbody>
</table>
| `UpdatePipelineExecution`         | sagemaker:UpdatePipelineExecution    | arn:aws:partition:sagemaker:region:account-id:pipeline/pipeline-name/exe
   cution/execution-id |
| `UpdateSpace`                     | sagemaker:UpdateSpace                | arn:aws:sagemaker:region:account-id:space/domain-id/spaceName |
| `UpdateUserProfile`               | sagemaker:UpdateUserProfile          | arn:aws:sagemaker:region:account-id:user-profile/domain-id/userProfileName |

**Amazon SageMaker API and Required Permissions for Actions**

**API Operation:** AddTags

Required Permissions (API Action): sagemaker:AddTags

Resources: *
**API Operation:** CreateEndpoint

Required Permissions (API Action): sagemaker:CreateEndpoint

Resources: arn:aws:sagemaker:region:account-id:endpoint/endpointName

**API Operation:** CreateEndpointConfig

Required Permissions (API Action): sagemaker:CreateEndpointConfig

Resources: arn:aws:sagemaker:region:account-id:endpoint-config/endpointConfigName

**API Operation:** CreateModel

Required Permissions (API Action): sagemaker:CreateModel, iam:PassRole

Resources: arn:aws:sagemaker:region:account-id:model/modelName

**API Operation:** CreateLabelingJob

Required Permissions (API Action): sagemaker:CreateLabelingJob, iam:PassRole

Resources: arn:aws:sagemaker:region:account-id:labeling-job/labelingJobName

**API Operation:** CreateNotebookInstance


Resources: arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName

**API Operation:** CreateTrainingJob

Required Permissions (API Action): sagemaker:CreateTrainingJob, iam:PassRole

Resources: arn:aws:sagemaker:region:account-id:training-job/trainingJobName

**API Operation:** CreateWorkforce


API Operation: CreateWorkteam


API Operation: DeleteEndpoint

Required Permissions (API Action): sagemaker:DeleteEndpoint

Resources: arn:aws:sagemaker:region:account-id:endpoint/endpointName

API Operation: DeleteEndpointConfig

Required Permissions (API Action): sagemaker:DeleteEndpointConfig

Resources: arn:aws:sagemaker:region:account-id:endpoint-config/endpointConfigName

API Operation: DeleteModel

Required Permissions (API Action): sagemaker:DeleteModel

Resources: arn:aws:sagemaker:region:account-id:model/modelName

API Operation: DeleteNotebookInstance


Resources: arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName

API Operation: DeleteTags

Required Permissions (API Action): sagemaker:DeleteTags

Resources: *
API Operation: **DeleteWorkteam**

Required Permissions (API Action): sagemaker:DeleteWorkforce

Resources: arn:aws:sagemaker:*:workforce/private-crowd/*

API Operation: **DeleteWorkteam**

Required Permissions (API Action): sagemaker:DeleteWorkteam

Resources: arn:aws:sagemaker:*:workteam/private-crowd/*

API Operation: **DescribeEndpoint**

Required Permissions (API Action): sagemaker:DescribeEndpoint

Resources: arn:aws:sagemaker:*:endpoint/endpointName

API Operation: **DescribeEndpointConfig**

Required Permissions (API Action): sagemaker:DescribeEndpointConfig

Resources: arn:aws:sagemaker:*:endpoint-config/endpointConfigName

API Operation: **DescribeLabelingJob**

Required Permissions (API Action): sagemaker:DescribeLabelingJob

Resources: arn:aws:sagemaker:*:labeling-job/labelingJobName

API Operation: **DescribeModel**

Required Permissions (API Action): sagemaker:DescribeModel

Resources: arn:aws:sagemaker:*:model/modelName

API Operation: **DescribeNotebookInstance**

Required Permissions (API Action): sagemaker:DescribeNotebookInstance

Resources: arn:aws:sagemaker:*:notebook-instance/notebookInstanceName

API Operation: **DescribeSubscribedWorkforce**

Required Permissions (API Action): sagemaker:DescribeSubscribedWorkforce, aws-marketplace:ViewSubscriptions
Resources: arn:aws:sagemaker:*region:account-id*:workforce/*

API Operation: DescribeSubscribedWorkteam

Required Permissions (API Action): sagemaker:DescribeSubscribedWorkteam, aws-marketplace:ViewSubscriptions

Resources: arn:aws:sagemaker:*region:account-id*:workteam/vendor-crowd/*

API Operation: DescribeTrainingJob

Required Permissions (API Action): sagemaker:DescribeTrainingJob

Resources: arn:aws:sagemaker:*region:account-id*:training-job/trainingJobName

API Operation: DescribeWorkteam

Required Permissions (API Action): sagemaker:DescribeWorkteam


API Operation: CreatePresignedNotebookInstanceUrl

Required Permissions (API Action): sagemaker:CreatePresignedNotebookInstanceUrl

Resources: arn:aws:sagemaker:*region:account-id*:notebook-instance/notebookInstanceName

API Operation: runtime_InvokeEndpoint

Required Permissions (API Action): sagemaker:InvokeEndpoint

Resources: arn:aws:sagemaker:*region:account-id*:endpoint/endpointName

API Operation: ListEndpointConfigs

Required Permissions (API Action): sagemaker:ListEndpointConfigs

Resources: *

API Operation: ListEndpoints

Required Permissions (API Action): sagemaker:ListEndpoints

Resources: *

API Operation: ListLabelingJobs

Required Permissions (API Action): sagemaker:ListLabelingJobs
Resources: *

**API Operation:** [ListLabelingJobsForWorkteam](#)

Required Permissions (API Action): sagemaker:ListLabelingJobsForWorkteam

Resources: *

**API Operation:** [ListModels](#)

Required Permissions (API Action): sagemaker:ListModels

Resources: *

**API Operation:** [ListNotebookInstances](#)

Required Permissions (API Action): sagemaker:ListNotebookInstances

Resources: *

**API Operation:** [ListSubscribedWorkteams](#)


Resources: *

**API Operation:** [ListTags](#)

Required Permissions (API Action): sagemaker:ListTags

Resources: *

**API Operation:** [ListTrainingJobs](#)

Required Permissions (API Action): sagemaker:ListTrainingJobs

Resources: *

**API Operation:** [ListWorkteams](#)

Required Permissions (API Action): sagemaker:ListWorkforces

Resources: *

**API Operation:** [ListWorkteams](#)

Required Permissions (API Action): sagemaker:ListWorkteams
API Operation: **StartNotebookInstance**


Resources: arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName

API Operation: **StopLabelingJob**

Required Permissions (API Action): sagemaker:StopLabelingJob

Resources: arn:aws:sagemaker:region:account-id:labeling-job/labelingJobName

API Operation: **StopNotebookInstance**

Required Permissions (API Action): sagemaker:StopNotebookInstance

Resources: arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName

API Operation: **StopTrainingJob**

Required Permissions (API Action): sagemaker:StopTrainingJob

Resources: arn:aws:sagemaker:region:account-id:training-job/trainingJobName

API Operation: **UpdateEndpoint**

Required Permissions (API Action): sagemaker:UpdateEndpoints

Resources: arn:aws:sagemaker:region:account-id:endpoint/endpointName

API Operation: **UpdateNotebookInstance**

Required Permissions (API Action): sagemaker:UpdateNotebookInstance, iam:PassRole

Resources: arn:aws:sagemaker:region:account-id:notebook-instance/notebookInstanceName
API Operation: **UpdateWorkteam**

Required Permissions (API Action): sagemaker:UpdateWorkteam

Resources: arn:aws:sagemaker:*:account-id:workteam/private-crowd/*

### AWS Managed Policies for Amazon SageMaker

To add permissions to users, groups, and roles, it is easier to use AWS managed policies than to write policies yourself. It takes time and expertise to create IAM customer managed policies that provide your team with only the permissions they need. To get started quickly, you can use our AWS managed policies. These policies cover common use cases and are available in your AWS account. For more information about AWS managed policies, see AWS managed policies in the IAM User Guide.

AWS services maintain and update AWS managed policies. You can't change the permissions in AWS managed policies. Services occasionally add additional permissions to an AWS managed policy to support new features. This type of update affects all identities (users, groups, and roles) to which the policy is attached. Services are most likely to update an AWS managed policy when a new feature is launched or when new operations become available. Services do not remove permissions from an AWS managed policy, so policy updates won't break your existing permissions.

Additionally, AWS supports managed policies for job functions that span multiple services. For example, the ReadOnlyAccess AWS managed policy provides read-only access to all AWS services and resources. When a service launches a new feature, AWS adds read-only permissions for new operations and resources. For a list and descriptions of job function policies, see AWS managed policies for job functions in the IAM User Guide.

⚠️ **Important**

We recommend that you use the most restricted policy that allows you to perform your use case.

The following AWS managed policies, which you can attach to users in your account, are specific to Amazon SageMaker:

- **AmazonSageMakerFullAccess** – Grants full access to Amazon SageMaker and SageMaker geospatial resources and the supported operations. This does not provide unrestricted
Amazon S3 access, but supports buckets and objects with specific sagemaker tags. This policy allows all IAM roles to be passed to Amazon SageMaker, but only allows IAM roles with "AmazonSageMaker" in them to be passed to the AWS Glue, AWS Step Functions, and AWS RoboMaker services.

- **AmazonSageMakerReadOnly** – Grants read-only access to Amazon SageMaker resources.

The following AWS managed policies can be attached to users in your account but are not recommended:

- **AdministratorAccess** – Grants all actions for all AWS services and for all resources in the account.
- **DataScientist** – Grants a wide range of permissions to cover most of the use cases (primarily for analytics and business intelligence) encountered by data scientists.

You can review these permissions policies by signing in to the IAM console and searching for them.

You can also create your own custom IAM policies to allow permissions for Amazon SageMaker actions and resources as you need them. You can attach these custom policies to the users or groups that require them.

**Topics**

- [AWS managed policy: AmazonSageMakerFullAccess](#)
- [AWS managed policy: AmazonSageMakerReadOnly](#)
- [AWS managed policies for Amazon SageMaker Canvas](#)
- [AWS managed policies for Amazon SageMaker Cluster](#)
- [AWS managed policies for Amazon SageMaker Feature Store](#)
- [AWS managed policies for Amazon SageMaker geospatial](#)
- [AWS Managed Policies for Amazon SageMaker Ground Truth](#)
- [AWS Managed Policies for SageMaker Model Governance](#)
- [AWS Managed Policies for Model Registry](#)
- [AWS Managed Policies for SageMaker Notebooks](#)
- [AWS Managed Policies for SageMaker Pipelines](#)
- [AWS Managed Policies for SageMaker projects and JumpStart](#)
AWS managed policy: AmazonSageMakerFullAccess

This policy grants administrative permissions that allow a principal full access to all Amazon SageMaker and SageMaker geospatial resources and operations. The policy also provides select access to related services. This policy allows all IAM roles to be passed to Amazon SageMaker, but only allows IAM roles with "AmazonSageMaker" in them to be passed to the AWS Glue, AWS Step Functions, and AWS RoboMaker services. This policy does not include permissions to create an Amazon SageMaker domain. For information on the policy needed to create a domain, see Create an Administrative User and Group.

Permissions details

This policy includes the following permissions.

- application-autoscaling – Allows principals to automatically scale a SageMaker real-time inference endpoint.
- athena – Allows principals to query a list of data catalogs, databases, and table metadata from Amazon Athena.
- aws-marketplace – Allows principals to view AWS AI Marketplace subscriptions. You need this if you want to access SageMaker software subscribed in AWS Marketplace.
- cloudformation – Allows principals to get AWS CloudFormation templates for using SageMaker JumpStart solutions and Pipelines. SageMaker JumpStart creates resources necessary to run end-to-end machine learning solutions that tie SageMaker to other AWS services. SageMaker Pipelines creates new projects that are backed by Service Catalog.
- cloudwatch – Allows principals to post CloudWatch metrics, interact with alarms, and upload logs to CloudWatch Logs in your account.
- codebuild – Allows principals to store AWS CodeBuild artifacts for SageMaker Pipeline and Projects.
- codecommit – Needed for AWS CodeCommit integration with SageMaker notebook instances.
- cognito-idp – Needed for Amazon SageMaker Ground Truth to define private workforce and work teams.
- ec2 – Needed for SageMaker to manage Amazon EC2 resources and network interfaces when you specify an Amazon VPC for your SageMaker jobs, models, endpoints, and notebook instances.
• **ecr** – Needed to pull and store Docker artifacts for Amazon SageMaker Studio Classic (custom images), training, processing, batch inference, and inference endpoints. This is also required to use your own container in SageMaker. Additional permissions for SageMaker JumpStart solutions are required to create and remove custom images on behalf of users.

• **elastic-inference** – Allows principals to connect to Amazon Elastic Inference for using SageMaker notebook instances and endpoints.

• **elasticfilesystem** – Allows principals to access Amazon Elastic File System. This is needed for SageMaker to use data sources in Amazon Elastic File System for training machine learning models.

• **fsx** – Allows principals to access Amazon FSx. This is needed for SageMaker to use data sources in Amazon FSx for training machine learning models.

• **glue** – Needed for inference pipeline pre-processing from within SageMaker notebook instances.

• **groundtruthlabeling** – Needed for Ground Truth labeling jobs. The groundtruthlabeling endpoint is accessed by the Ground Truth console.

• **iam** – Needed to give the SageMaker console access to available IAM roles and create service-linked roles.

• **kms** – Needed to give the SageMaker console access to available AWS KMS keys and retrieve them for any specified AWS KMS aliases in jobs and endpoints.

• **lambda** – Allows principals to invoke and get a list of AWS Lambda functions.

• **logs** – Needed to allow SageMaker jobs and endpoints to publish log streams.

• **redshift** – Allows principals to access Amazon Redshift cluster credentials.

• **redshift-data** – Allows principals to use data from Amazon Redshift to run, describe, and cancel statements; get statement results; and list schemas and tables.

• **robomaker** – Allows principals to have full access to create, get descriptions, and delete AWS RoboMaker simulation applications and jobs. This is also needed to run reinforcement learning examples on notebook instances.

• **s3, s3express** – Allows principals to have full access to Amazon S3 and Amazon S3 Express resources pertaining to SageMaker, but not all of Amazon S3 or Amazon S3 Express.

• **sagemaker** – Allows principals to list tags on SageMaker user profiles and add tags to SageMaker apps. Allows access only to SageMaker flow-definitions of sagemaker:WorkteamType "private-crowd" or "vendor-crowd".

• **sagemaker and sagemaker-geospatial** – Allows principals read-only access to SageMaker domains and user profiles.
• secretsmanager – Allows principals to have full access to AWS Secrets Manager. The principals can securely encrypt, store, and retrieve credentials for databases and other services. This is also needed for SageMaker notebook instances with SageMaker code repositories that use GitHub.

• servicecatalog – Allows principals to use Service Catalog. The principals can create, get a list of, update, or terminate provisioned products, such as servers, databases, websites, or applications deployed using AWS resources. This is needed for SageMaker JumpStart and Projects to find and read service catalog products and launch AWS resources in users.

• sns – Allows principals to get a list of Amazon SNS topics. This is needed for endpoints with Async Inference enabled for notifying users that their inference has completed.

• states – Needed for SageMaker JumpStart and Pipelines to use a service catalog to create step function resources.

• tag – Needed for SageMaker Pipelines to render in Studio Classic. Studio Classic needs resources tagged with particular sagemaker:project-id tag-key. This requires the tag:GetResources permission.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "AllowAllNonAdminSageMakerActions",
            "Effect": "Allow",
            "Action": [
                "sagemaker:*",
                "sagemaker-geospatial:*"
            ],
            "NotResource": [
                "arn:aws:sagemaker:*:*:domain/*",
                "arn:aws:sagemaker:*:*:user-profile/*",
                "arn:aws:sagemaker:*:*:app/*",
                "arn:aws:sagemaker:*:*:space/*",
                "arn:aws:sagemaker:*:*:flow-definition/*"
            ]
        },
        {
            "Sid": "AllowAddTagsForApp",
            "Effect": "Allow",
            "Action": [
                "sagemaker:AddTags"
            ],
```
"Resource": [  
  "arn:aws:sagemaker:*::*:app/*"
],
"Sid": "AllowStudioActions",
"Effect": "Allow",
"Action": [  
  "sagemaker:CreatePresignedDomainUrl",
  "sagemaker:DescribeDomain",
  "sagemaker:ListDomains",
  "sagemaker:DescribeUserProfile",
  "sagemaker:ListUserProfiles",
  "sagemaker:DescribeSpace",
  "sagemaker:ListSpaces",
  "sagemaker:DescribeApp",
  "sagemaker:ListApps"
],
"Resource": "*",
"Condition": {}
},
{"Sid": "AllowAppActionsForUserProfile",
"Effect": "Allow",
"Action": [  
  "sagemaker:CreateApp",
  "sagemaker:DeleteApp"
],
"Resource": "arn:aws:sagemaker:*::*:app/*/*/*/*",
"Condition": {  
  "Null": {  
    "sagemaker:OwnerUserProfileArn": "true"
  }
}
},
{"Sid": "AllowAppActionsForSharedSpaces",
"Effect": "Allow",
"Action": [  
  "sagemaker:CreateApp",
  "sagemaker:DeleteApp"
],
"Resource": "arn:aws:sagemaker:*::*:app/${sagemaker:DomainId}/*/*/*/*",
"Condition": {  
  "StringEquals": {  
    "sagemaker:DomainId": ".*"
  }
}
}
"sagemaker:SpaceSharingType": [ "Shared"
 ]
}
}
{
"Sid": "AllowMutatingActionsOnSharedSpacesWithoutOwner",
"Effect": "Allow",
"Action": [ "sagemaker:CreateSpace",
            "sagemaker:UpdateSpace",
            "sagemaker:DeleteSpace"
          ],
"Resource": "arn:aws:sagemaker:*:*:space/${sagemaker:DomainId}/*",
"Condition": { "Null": {
                      "sagemaker:OwnerUserProfileArn": "true"
                  }
          }
}
{
"Sid": "RestrictMutatingActionsOnSpacesToOwnerUserProfile",
"Effect": "Allow",
"Action": [ "sagemaker:CreateSpace",
            "sagemaker:UpdateSpace",
            "sagemaker:DeleteSpace"
          ],
"Resource": "arn:aws:sagemaker:*:*:space/${sagemaker:DomainId}/*",
"Condition": { "ArnLike": {
                      "sagemaker:OwnerUserProfileArn": "arn:aws:sagemaker:*:*:user-profile/${sagemaker:DomainId}/${sagemaker:UserProfileName}"
                  },
            "StringEquals": { "sagemaker:SpaceSharingType": [ "Private",
                                                     "Shared"
                                               ]
                  }
          }
}
}
"Sid": "RestrictMutatingActionsOnPrivateSpaceAppsToOwnerUserProfile",
"Effect": "Allow",
"Action": [
    "sagemaker:CreateApp",
    "sagemaker:DeleteApp"
],
"Resource": "arn:aws:sagemaker:*:*:app/${sagemaker:DomainId}/**/**",
"Condition": {
    "ArnLike": {
        "sagemaker:OwnerUserProfileArn": "arn:aws:sagemaker:*:*:user-profile/${sagemaker:DomainId}/${sagemaker:UserProfileName}"
    },
    "StringEquals": {
        "sagemaker:SpaceSharingType": ["Private"]
    }
}
},
{
    "Sid": "AllowFlowDefinitionActions",
    "Effect": "Allow",
    "Action": "sagemaker:*",
    "Resource": [
        "arn:aws:sagemaker:*:*:flow-definition/**"
    ],
    "Condition": {
        "StringEqualsIfExists": {
            "sagemaker:WorkteamType": [
                "private-crowd",
                "vendor-crowd"
            ]
        }
    }
},
{
    "Sid": "AllowAWSServiceActions",
    "Effect": "Allow",
    "Action": [
        "application-autoscaling:DeleteScalingPolicy",
        "application-autoscaling:DeleteScheduledAction",
        "application-autoscaling:DeregisterScalableTarget",
        "application-autoscaling:DescribeScalableTargets",
        "application-autoscaling:DescribeScalingActivities",
        ...
"application-autoscaling:DescribeScalingPolicies",
"application-autoscaling:DescribeScheduledActions",
"application-autoscaling:PutScalingPolicy",
"application-autoscaling:PutScheduledAction",
"application-autoscaling:RegisterScalableTarget",
"aws-marketplace:ViewSubscriptions",
"cloudformation:GetTemplateSummary",
"cloudwatch:DeleteAlarms",
"cloudwatch:DescribeAlarms",
"cloudwatch:GetMetricData",
"cloudwatch:GetMetricStatistics",
"cloudwatch:ListMetrics",
"cloudwatch:PutMetricAlarm",
"cloudwatch:PutMetricData",
"codecommit:BatchGetRepositories",
"codecommit:CreateRepository",
"codecommit:GetRepository",
"codecommit:List*",
"cognito-idp:AdminAddUserToGroup",
"cognito-idp:AdminCreateUser",
"cognito-idp:AdminDeleteUser",
"cognito-idp:AdminDisableUser",
"cognito-idp:AdminEnableUser",
"cognito-idp:AdminRemoveUserFromGroup",
"cognito-idp:CreateGroup",
"cognito-idp:CreateUserPool",
"cognito-idp:CreateUserPoolClient",
"cognito-idp:CreateUserPoolDomain",
"cognito-idp:DescribeUserPool",
"cognito-idp:DescribeUserPoolClient",
"cognito-idp:List*",
"cognito-idp:UpdateUserPool",
"cognito-idp:UpdateUserPoolClient",
"ec2:CreateNetworkInterface",
"ec2:CreateNetworkInterfacePermission",
"ec2:CreateVpcEndpoint",
"ec2:DeleteNetworkInterface",
"ec2:DeleteNetworkInterfacePermission",
"ec2:DescribeDhcpOptions",
"ec2:DescribeNetworkInterfaces",
"ec2:DescribeRouteTables",
"ec2:DescribeSecurityGroups",
"ec2:DescribeSubnets",
"ec2:DescribeVpcEndpoints"
"ec2:DescribeVpcs",
"ecr:BatchCheckLayerAvailability",
"ecr:BatchGetImage",
"ecr:CreateRepository",
"ecr:Describe**",
"ecr:GetAuthorizationToken",
"ecr:GetDownloadUrlForLayer",
"ecr:StartImageScan",
"elastic-inference:Connect",
"elasticfilesystem:DescribeFileSystems",
"elasticfilesystem:DescribeMountTargets",
"fsx:DescribeFileSystems",
"glue:CreateJob",
"glue:DeleteJob",
"glue:GetJob**",
"glue:GetTable**",
"glue:GetWorkflowRun",
"glue:ResetJobBookmark",
"glue:StartJobRun",
"glue:StartWorkflowRun",
"glue:UpdateJob",
"groundtruthlabeling:**",
"iam:ListRoles",
"kms:DescribeKey",
"kms:ListAliases",
"lambda:ListFunctions",
"logs:CreateLogDelivery",
"logs:CreateLogGroup",
"logs:CreateLogStream",
"logs:DeleteLogDelivery",
"logs:Describe**",
"logs:GetLogDelivery",
"logs:GetLogEvents",
"logs:ListLogDeliveries",
"logs:PutLogEvents",
"logs:PutResourcePolicy",
"logs:UpdateLogDelivery",
"robomaker:CreateSimulationApplication",
"robomaker:DescribeSimulationApplication",
"robomaker:DeleteSimulationApplication",
"robomaker:CreateSimulationJob",
"robomaker:DescribeSimulationJob",
"robomaker:CancelSimulationJob",
"secretsmanager:ListSecrets",
"servicecatalog:Describe*",
"servicecatalog:List*",
"servicecatalog:ScanProvisionedProducts",
"servicecatalog:SearchProducts",
"servicecatalog:SearchProvisionedProducts",
"sns:ListTopics",
"tag:GetResources",
],
"Resource": "*
"},
{
"Sid": "AllowECRActions",
"Effect": "Allow",
"Action": [
"ecr:SetRepositoryPolicy",
"ecr:CompleteLayerUpload",
"ecr:BatchDeleteImage",
"ecr:UploadLayerPart",
"ecr:DeleteRepositoryPolicy",
"ecr:InitiateLayerUpload",
"ecr:DeleteRepository",
"ecr:PutImage"
],
"Resource": [  
"arn:aws:ecr:*:*:repository/*sagemaker*"
]}
},
{
"Sid": "AllowCodeCommitActions",
"Effect": "Allow",
"Action": [
"codecommit:GitPull",
"codecommit:GitPush"
],
"Resource": [  
"arn:aws:codecommit:*:*:sagemaker*",  
"arn:aws:codecommit:*:*:SageMaker*",  
"arn:aws:codecommit:*:*:Sagemaker*"
]}
},
{
"Sid": "AllowCodeBuildActions",
"Action": [
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
"codebuild:BatchGetBuilds",
---
AWS Managed Policies for SageMaker
5862
"codebuild:StartBuild",
[,
"Resource": [
  "arn:aws:codebuild:*::*:project/sagemaker*",
  "arn:aws:codebuild:*::*:build/*"
],
"Effect": "Allow"
},
{
  "Sid": "AllowStepFunctionsActions",
  "Action": [
    "states:DescribeExecution",
    "states:GetExecutionHistory",
    "states:StartExecution",
    "states:StopExecution",
    "states:UpdateStateMachine"
  ],
  "Resource": [
    "arn:aws:states:*::*:statemachine:*sagemaker*",
    "arn:aws:states:*::*:execution:*sagemaker::*"
  ],
  "Effect": "Allow"
},
{
  "Sid": "AllowSecretManagerActions",
  "Effect": "Allow",
  "Action": [
    "secretsmanager:DescribeSecret",
    "secretsmanager:GetSecretValue",
    "secretsmanager:CreateSecret"
  ],
  "Resource": [
    "arn:aws:secretsmanager:*::*:secret:AmazonSageMaker-*"
  ]
},
{
  "Sid": "AllowReadOnlySecretManagerActions",
  "Effect": "Allow",
  "Action": [
    "secretsmanager:DescribeSecret",
    "secretsmanager:GetSecretValue"
  ],
  "Resource": "*",
  "Condition": {
"StringEquals": {
    "secretsmanager:ResourceTag/SageMaker": "true"
  }
},
{
  "Sid": "AllowServiceCatalogProvisionProduct",
  "Effect": "Allow",
  "Action": [
    "servicecatalog:ProvisionProduct"
  ],
  "Resource": "*
"},
{
  "Sid": "AllowServiceCatalogTerminateUpdateProvisionProduct",
  "Effect": "Allow",
  "Action": [
    "servicecatalog:TerminateProvisionedProduct",
    "servicecatalog:UpdateProvisionedProduct"
  ],
  "Resource": "*
",
  "Condition": {
    "StringEquals": {
      "servicecatalog:userLevel": "self"
    }
  }
},
{
  "Sid": "AllowS3ObjectActions",
  "Effect": "Allow",
  "Action": [
    "s3:GetObject",
    "s3:PutObject",
    "s3:DeleteObject",
    "s3:AbortMultipartUpload"
  ],
  "Resource": [
    "arn:aws:s3:::*SageMaker*",
    "arn:aws:s3:::*Sagemaker*",
    "arn:aws:s3:::*sagemaker*",
    "arn:aws:s3:::*aws-glue*"
  ]
}
"Sid": "AllowS3GetObjectWithSageMakerExistingObjectTag",
"Effect": "Allow",
"Action": [ 
    "s3:GetObject"
],
"Resource": [ 
    "arn:aws:s3:::*"
],
"Condition": { 
    "StringEqualsIgnoreCase": { 
        "s3:ExistingObjectTag/SageMaker": "true"
    }
}
},
{
    "Sid": "AllowS3GetObjectWithServiceCatalogProvisioningExistingObjectTag",
    "Effect": "Allow",
    "Action": [ 
        "s3:GetObject"
    ],
    "Resource": [ 
        "arn:aws:s3:::*"
    ],
    "Condition": { 
        "StringEquals": { 
            "s3:ExistingObjectTag/servicecatalog:provisioning": "true"
        }
    }
}
},
{
    "Sid": "AllowS3BucketActions",
    "Effect": "Allow",
    "Action": [ 
        "s3:CreateBucket",
        "s3:GetBucketLocation",
        "s3:ListBucket",
        "s3:ListAllMyBuckets",
        "s3:GetBucketCors",
        "s3:PutBucketCors"
    ],
    "Resource": "*"
},
{
    "Sid": "AllowS3BucketACL",
}
"Effect": "Allow",
"Action": [  
  "s3:GetBucketAcl",
  "s3:PutObjectAcl"
],
"Resource": [  
  "arn:aws:s3:::*SageMaker*",
  "arn:aws:s3:::*Sagemaker*",
  "arn:aws:s3:::*sagemaker*
]
},
{
"Sid": "AllowLambdaInvokeFunction",
"Effect": "Allow",
"Action": [  
  "lambda:InvokeFunction"
],
"Resource": [  
  "arn:aws:lambda:*:*:function:*SageMaker*",
  "arn:aws:lambda:*:*:function:*sagemaker*",
  "arn:aws:lambda:*:*:function:*Sagemaker*",
  "arn:aws:lambda:*:*:function:*LabelingFunction*"
]
},
{
"Sid": "AllowCreateServiceLinkedRoleForSageMakerApplicationAutoscaling",
"Action": "iam:CreateServiceLinkedRole",
"Effect": "Allow",
"Resource": "arn:aws:iam::*:role/aws-service-role/sagemaker.application-autoscaling.amazonaws.com/AWSServiceRoleForApplicationAutoScaling_SageMakerEndpoint",
"Condition": {  
  "StringLike": {  
    "iam:AWSServiceName": "sagemaker.application-autoscaling.amazonaws.com"
  }
}
},
{
"Sid": "AllowCreateServiceLinkedRoleForRobomaker",
"Effect": "Allow",
"Action": "iam:CreateServiceLinkedRole",
"Resource": "*",
"Condition": {  
  "StringEquals": {  
    "iam:AWSServiceName": "robomaker.amazonaws.com"
  }
}
}
{
  "Sid": "AllowSNSActions",
  "Effect": "Allow",
  "Action": [
    "sns:Subscribe",
    "sns:CreateTopic",
    "sns:Publish"
  ],
  "Resource": [
    "arn:aws:sns:*:*:*SageMaker*",
    "arn:aws:sns:*:*:*Sagemaker*",
    "arn:aws:sns:*:*:*sagemaker*"
  ]
},
{
  "Sid": "AllowPassRoleForSageMakerRoles",
  "Effect": "Allow",
  "Action": [
    "iam:PassRole"
  ],
  "Resource": "arn:aws:iam::*:role/*AmazonSageMaker*",
  "Condition": {
    "StringEquals": {
      "iam:PassedToService": [
        "glue.amazonaws.com",
        "robomaker.amazonaws.com",
        "states.amazonaws.com"
      ]
    }
  }
},
{
  "Sid": "AllowPassRoleToSageMaker",
  "Effect": "Allow",
  "Action": [
    "iam:PassRole"
  ],
  "Resource": "arn:aws:iam::*:role/*",
  "Condition": {
    "StringEquals": {
      "iam:PassedToService": "sagemaker.amazonaws.com"
    }
  }
}
"Sid": "AllowAthenaActions",
"Effect": "Allow",
"Action": [
  "athena:ListDataCatalogs",
  "athena:ListDatabases",
  "athena:ListTableMetadata",
  "athena:GetQueryExecution",
  "athena:GetQueryResults",
  "athena:StartQueryExecution",
  "athena:StopQueryExecution"
],
"Resource": [
  "*
]
},

{"Sid": "AllowGlueCreateTable",
"Effect": "Allow",
"Action": [
  "glue:CreateTable"
],
"Resource": [
  "arn:aws:glue:*:*:table/*/sagemaker_tmp_*",
  "arn:aws:glue:*:*:table/sagemaker_featurestore/**",
  "arn:aws:glue:*:*:catalog",
  "arn:aws:glue:*:*:database/**
]
},

{"Sid": "AllowGlueUpdateTable",
"Effect": "Allow",
"Action": [
  "glue:UpdateTable"
],
"Resource": [
  "arn:aws:glue:*:*:table/sagemaker_featurestore/**",
  "arn:aws:glue:*:*:catalog",
  "arn:aws:glue:*:*:database/sagemaker_featurestore"}
}
{  
  "Sid": "AllowGlueDeleteTable",
  "Effect": "Allow",
  "Action": [  
    "glue:DeleteTable"
  ],
  "Resource": [  
    "arn:aws:glue:*:*:table/*/sagemaker_tmp_*",
    "arn:aws:glue:*:*:catalog",
    "arn:aws:glue:*:*:database/*"
  ]
},

{  
  "Sid": "AllowGlueGetTablesAndDatabases",
  "Effect": "Allow",
  "Action": [  
    "glue:GetDatabases",
    "glue:GetTable",
    "glue:GetTables"
  ],
  "Resource": [  
    "arn:aws:glue:*:*:table/*",
    "arn:aws:glue:*:*:catalog",
    "arn:aws:glue:*:*:database/*"
  ]
},

{  
  "Sid": "AllowGlueGetAndCreateDatabase",
  "Effect": "Allow",
  "Action": [  
    "glue:CreateDatabase",
    "glue:GetDatabase"
  ],
  "Resource": [  
    "arn:aws:glue:*:*:catalog",
    "arn:aws:glue:*:*:database/sagemaker_featurestore",
    "arn:aws:glue:*:*:database/sagemaker_processing",
    "arn:aws:glue:*:*:database/default",
    "arn:aws:glue:*:*:database/sagemaker_data_wrangler"
  ]
},

{  
  "Sid": "AllowRedshiftDataActions",
  "Effect": "Allow",
  "Action": [  
    "redshift:GetTableData",
    "redshift:GetTableStatistics",
    "redshift:Query",
    "redshift:CreateTable",
    "redshift:DeleteTable",
    "redshift:DropTable",
    "redshift:UpdateTableStatistics",
    "redshift:DescribeTable",
    "redshift:TagResource",
    "redshift:UntagResource"
  ],
  "Resource": [  
    "arn:aws:redshift:*:*:table/*",
    "arn:aws:redshift:*:*:catalog",
    "arn:aws:redshift:*:*:database/*"
  ]
}
"Action": [
  "redshift-data:ExecuteStatement",
  "redshift-data:DescribeStatement",
  "redshift-data:CancelStatement",
  "redshift-data:GetStatementResult",
  "redshift-data:ListSchemas",
  "redshift-data:ListTables"
],
"Resource": [
  "*"
]
},
{
  "Sid": "AllowRedshiftGetClusterCredentials",
  "Effect": "Allow",
  "Action": [
    "redshift:GetClusterCredentials"
  ],
  "Resource": [
    "arn:aws:redshift:*:*:dbuser/*/sagemaker_access*",
    "arn:aws:redshift:*:*:dbname:*"
  ]
},
{
  "Sid": "AllowListTagsForUserProfile",
  "Effect": "Allow",
  "Action": [
    "sagemaker:ListTags"
  ],
  "Resource": [
    "arn:aws:sagemaker:*:*:user-profile/*"
  ]
},
{
  "Sid": "AllowCloudformationListStackResources",
  "Effect": "Allow",
  "Action": [
    "cloudformation:ListStackResources"
  ],
  "Resource": "arn:aws:cloudformation:*:*:stack/SC-*"
},
{
  "Sid": "AllowS3ExpressObjectActions",
  "Effect": "Allow",
  "Action": ["s3:ListObjects", "s3:GetObject"]
}
"Action": [
    "s3express:CreateSession"
],
"Resource": [
    "arn:aws:s3express:*:*:bucket/*SageMaker*",
    "arn:aws:s3express:*:*:bucket/*Sagemaker*",
    "arn:aws:s3express:*:*:bucket/*sagemaker*",
    "arn:aws:s3express:*:*:bucket/*aws-glue*"
],
"Condition": {
    "StringEquals": {
        "aws:ResourceAccount": "${aws:PrincipalAccount}" 
    }
},
{
    "Sid": "AllowS3ExpressCreateBucketActions",
    "Effect": "Allow",
    "Action": [
        "s3express:CreateBucket"
    ],
    "Resource": [
        "arn:aws:s3express:*:*:bucket/*SageMaker*",
        "arn:aws:s3express:*:*:bucket/*Sagemaker*",
        "arn:aws:s3express:*:*:bucket/*sagemaker*"
    ],
    "Condition": {
        "StringEquals": {
            "aws:ResourceAccount": "${aws:PrincipalAccount}" 
        }
    }
},
{
    "Sid": "AllowS3ExpressListBucketActions",
    "Effect": "Allow",
    "Action": [
        "s3express:ListAllMyDirectoryBuckets"
    ],
    "Resource": "*"
}
AWS managed policy: AmazonSageMakerReadOnly

This policy grants read-only access to Amazon SageMaker through the AWS Management Console and SDK.

Permissions details

This policy includes the following permissions.

- **application-autoscaling** – Allows users to browse descriptions of scalable SageMaker real-time inference endpoints.
- **aws-marketplace** – Allows users to view AWS AI Marketplace subscriptions.
- **cloudwatch** – Allows users to receive CloudWatch alarms.
- **cognito-idp** – Needed for Amazon SageMaker Ground Truth to browse descriptions and lists of private workforce and work teams.
- **ecr** – Needed to read Docker artifacts for training and inference.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:Describe*",
                "sagemaker:List*",
                "sagemaker:BatchGetMetrics",
                "sagemaker:GetDeviceRegistration",
                "sagemaker:GetDeviceFleetReport",
                "sagemaker:GetSearchSuggestions",
                "sagemaker:BatchGetRecord",
                "sagemaker:GetRecord",
                "sagemaker:Search",
                "sagemaker:QueryLineage",
                "sagemaker:GetLineageGroupPolicy",
                "sagemaker:BatchDescribeModelPackage",
                "sagemaker:GetModelPackageGroupPolicy"
            ],
            "Resource": "*"
        },
        {
```
```
AWS managed policies for Amazon SageMaker Canvas

These AWS managed policies add permissions required to use Amazon SageMaker Canvas. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.

Topics

- [AWS managed policy: AmazonSageMakerCanvasFullAccess](#)
- [AWS managed policy: AmazonSageMakerCanvasDataPrepFullAccess](#)
- [AWS managed policy: AmazonSageMakerCanvasDirectDeployAccess](#)
- [AWS managed policy: AmazonSageMakerCanvasAIServicesAccess](#)
- [AWS managed policy: AmazonSageMakerCanvasBedrockAccess](#)
- [AWS managed policy: AmazonSageMakerCanvasForecastAccess](#)
- Amazon SageMaker updates to Amazon SageMaker Canvas managed policies
AWS managed policy: AmazonSageMakerCanvasFullAccess

This policy grants permissions that allow full access to Amazon SageMaker Canvas through the AWS Management Console and SDK. The policy also provides select access to related services [for example, Amazon Simple Storage Service (Amazon S3), AWS Identity and Access Management (IAM), Amazon Virtual Private Cloud (Amazon VPC), Amazon Elastic Container Registry (Amazon ECR), Amazon CloudWatch Logs, Amazon Redshift, AWS Secrets Manager, Amazon SageMaker Autopilot, SageMaker Model Registry, and Amazon Forecast].

This policy is intended to help customers experiment and get started with all the capabilities of SageMaker Canvas. For more fine-grained control, we suggest customers build their own scoped down versions as they move to production workloads. For more information, see IAM policy types: How and when to use them.

Permissions details

This AWS managed policy includes the following permissions.

- sagemaker – Allows principals to create and host SageMaker models on resources whose ARN contains "Canvas", "canvas", or "model-compilation-". Additionally, users can register their SageMaker Canvas model to SageMaker Model Registry in the same AWS account.
- ec2 – Allows principals to create Amazon VPC endpoints.
- ecr – Allows principals to get information about a container image.
- glue – Allows principals to retrieve the tables in the catalog.
- iam – Allows principals to pass an IAM role to Amazon SageMaker and Amazon Forecast. Also allows principals to create a service-linked role.
- logs – Allows principals to publish logs from training jobs and endpoints.
- s3 – Allows principals to add and retrieve objects from Amazon S3 buckets. These objects are limited to those whose name includes "SageMaker", "Sagemaker", or "sagemaker". Also allows principals to retrieve objects from Amazon S3 buckets whose ARN starts with "jumpstart-cache-prod-" in specific regions.
- secretsmanager – Allows principals to store customer credentials to connect to a Snowflake database using Secrets Manager.
- redshift – Allows principals to get credentials for a "sagemaker_access*" dbuser on any Amazon Redshift cluster if that user exists.
- redshift-data – Allows principals to run queries on Amazon Redshift using the Amazon Redshift Data API. This only provides access to the Redshift Data APIs themselves and does not
directly provide access to your Amazon Redshift clusters. For more information, see [Using the Amazon Redshift Data API](#).

- forecast – Allows principals to use Amazon Forecast.
- application-autoscaling – Allows principals to automatically scale a SageMaker inference endpoint.
- rds – Allows principals to return information about provisioned Amazon RDS instances.
- cloudwatch – Allows principals to create and manage Amazon CloudWatch alarms.
- athena – Allows principals to create, read, and manage Amazon Athena queries, catalogs, and executions.

```json
{
   "Version": "2012-10-17",
   "Statement": [
   {
      "Sid": "SageMakerUserDetailsAndPackageOperations",
      "Effect": "Allow",
      "Action": [
      "sagemaker:DescribeDomain",
      "sagemaker:DescribeUserProfile",
      "sagemaker:ListTags",
      "sagemaker:ListModelPackages",
      "sagemaker:ListModelPackageGroups",
      "sagemaker:ListEndpoints"
      ],
      "Resource": "*"
   },
   {
      "Sid": "SageMakerPackageGroupOperations",
      "Effect": "Allow",
      "Action": [
      "sagemaker:CreateModelPackageGroup",
      "sagemaker:CreateModelPackage",
      "sagemaker:DescribeModelPackageGroup",
      "sagemaker:DescribeModelPackage"
      ],
      "Resource": [
      "arn:aws:sagemaker:*::*:model-package/*",
      "arn:aws:sagemaker:*::*:model-package-group/*"
      ]
   }
   ]
}```
{
    "Sid": "SageMakerTrainingOperations",
    "Effect": "Allow",
    "Action": [
        "sagemaker:CreateCompilationJob",
        "sagemaker:CreateEndpoint",
        "sagemaker:CreateEndpointConfig",
        "sagemaker:CreateModel",
        "sagemaker:CreateProcessingJob",
        "sagemaker:CreateAutoMLJob",
        "sagemaker:CreateAutoMLJobV2",
        "sagemaker:DeleteEndpoint",
        "sagemaker:DescribeCompilationJob",
        "sagemaker:DescribeEndpoint",
        "sagemaker:DescribeEndpointConfig",
        "sagemaker:DescribeModel",
        "sagemaker:DescribeProcessingJob",
        "sagemaker:DescribeAutoMLJob",
        "sagemaker:DescribeAutoMLJobV2",
        "sagemaker:ListCandidatesForAutoMLJob",
        "sagemaker:AddTags",
        "sagemaker:DeleteApp"
    ],
    "Resource": [
        "arn:aws:sagemaker:*::*:Canvas*",
        "arn:aws:sagemaker:*::*:canvas*",
        "arn:aws:sagemaker:*::*:model-compilation-*"
    ]
},
{
    "Sid": "SageMakerHostingOperations",
    "Effect": "Allow",
    "Action": [
        "sagemaker:DeleteEndpointConfig",
        "sagemaker:DeleteModel",
        "sagemaker:InvokeEndpoint",
        "sagemaker:UpdateEndpointWeightsAndCapacities",
        "sagemaker:InvokeEndpointAsync"
    ],
    "Resource": [
        "arn:aws:sagemaker:*::*:Canvas*",
        "arn:aws:sagemaker:*::*:canvas*"
    ]
}
{
  "Sid": "EC2VPCOperation",
  "Effect": "Allow",
  "Action": [
    "ec2:CreateVpcEndpoint",
    "ec2:DescribeSecurityGroups",
    "ec2:DescribeSubnets",
    "ec2:DescribeVpcs",
    "ec2:DescribeVpcEndpoints",
    "ec2:DescribeVpcEndpointServices"
  ],
  "Resource": "*"
},
{
  "Sid": "ECROperations",
  "Effect": "Allow",
  "Action": [
    "ecr:BatchGetImage",
    "ecr:GetDownloadUrlForLayer",
    "ecr:GetAuthorizationToken"
  ],
  "Resource": "*"
},
{
  "Sid": "IAMGetOperations",
  "Effect": "Allow",
  "Action": [
    "iam:GetRole"
  ],
  "Resource": "arn:aws:iam::*:role/*"
},
{
  "Sid": "IAMPassOperation",
  "Effect": "Allow",
  "Action": [
    "iam:PassRole"
  ],
  "Resource": "arn:aws:iam::*:role/*",
  "Condition": {
    "StringEquals": {
      "iam:PassedToService": "sagemaker.amazonaws.com"
    }
  }
}
{
  "Sid": "LoggingOperation",
  "Effect": "Allow",
  "Action": [
    "logs:CreateLogGroup",
    "logs:CreateLogStream",
    "logs:PutLogEvents"
  ],
  "Resource": "arn:aws:logs:*:*:log-group:/aws/sagemaker/**"
},
{
  "Sid": "S3Operations",
  "Effect": "Allow",
  "Action": [
    "s3:GetObject",
    "s3:PutObject",
    "s3:DeleteObject",
    "s3:CreateBucket",
    "s3:GetBucketCors",
    "s3:GetBucketLocation"
  ],
  "Resource": [
    "arn:aws:s3:::*Sagemaker*",
    "arn:aws:s3:::*SageMaker*",
    "arn:aws:s3:::*sagemaker*
  ]
},
{
  "Sid": "ReadSageMakerJumpstartArtifacts",
  "Effect": "Allow",
  "Action": "s3:GetObject",
  "Resource": [
    "arn:aws:s3:::jumpstart-cache-prod-us-west-2/*",
    "arn:aws:s3:::jumpstart-cache-prod-us-east-1/*",
    "arn:aws:s3:::jumpstart-cache-prod-us-east-2/*",
    "arn:aws:s3:::jumpstart-cache-prod-eu-west-1/*",
    "arn:aws:s3:::jumpstart-cache-prod-eu-central-1/*",
    "arn:aws:s3:::jumpstart-cache-prod-ap-south-1/*",
    "arn:aws:s3:::jumpstart-cache-prod-ap-northeast-2/*",
    "arn:aws:s3:::jumpstart-cache-prod-ap-northeast-1/*",
    "arn:aws:s3:::jumpstart-cache-prod-ap-southeast-1/*",
    "arn:aws:s3:::jumpstart-cache-prod-ap-southeast-2/*"
  ]
}
{  
  "Sid": "S3ListOperations",
  "Effect": "Allow",
  "Action": [
    "s3:ListBucket",
    "s3:ListAllMyBuckets"
  ],
  "Resource": "*"
},
{  
  "Sid": "GlueOperations",
  "Effect": "Allow",
  "Action": "glue:SearchTables",
  "Resource": [
    "arn:aws:glue:*:*:table/*/*",
    "arn:aws:glue:*:*:database/*",
    "arn:aws:glue:*:*:catalog"
  ]
},
{  
  "Sid": "SecretsManagerARNBasedOperation",
  "Effect": "Allow",
  "Action": [
    "secretsmanager:DescribeSecret",
    "secretsmanager:GetSecretValue",
    "secretsmanager:CreateSecret",
    "secretsmanager:PutResourcePolicy"
  ],
  "Resource": [
    "arn:aws:secretsmanager:*:*:secret:AmazonSageMaker-*"
  ]
},
{  
  "Sid": "SecretManagerTagBasedOperation",
  "Effect": "Allow",
  "Action": [
    "secretsmanager:DescribeSecret",
    "secretsmanager:GetSecretValue"
  ],
  "Resource": "*",
  "Condition": {
    "StringEquals": {
      "secretsmanager:ResourceTag/SageMaker": "true"
    }
  }
}


```json
{}
{
    "Sid": "RedshiftOperations",
    "Effect": "Allow",
    "Action": [
        "redshift-data:ExecuteStatement",
        "redshift-data:DescribeStatement",
        "redshift-data:CancelStatement",
        "redshift-data:GetStatementResult",
        "redshift-data:ListSchemas",
        "redshift-data:ListTables",
        "redshift-data:DescribeTable"
    ],
    "Resource": "*"
},
{
    "Sid": "RedshiftGetCredentialsOperation",
    "Effect": "Allow",
    "Action": [
        "redshift:GetClusterCredentials"
    ],
    "Resource": [
        "arn:aws:redshift:*:*:dbuser:*/sagemaker_access*",
        "arn:aws:redshift:*:*:dbname:*"
    ]
},
{
    "Sid": "ForecastOperations",
    "Effect": "Allow",
    "Action": [
        "forecast:CreateExplainabilityExport",
        "forecast:CreateExplainability",
        "forecast:CreateForecastEndpoint",
        "forecast:CreateAutoPredictor",
        "forecast:CreateDatasetImportJob",
        "forecast:CreateDatasetGroup",
        "forecast:CreateDataset",
        "forecast:CreateForecast",
        "forecast:CreateForecastExportJob",
        "forecast:CreatePredictorBacktestExportJob",
        "forecast:CreatePredictor",
        "forecast:DescribeExplainabilityExport",
        "forecast:DescribeExplainability"
    ]
}
```
"forecast:DescribeAutoPredictor",
"forecast:DescribeForecastEndpoint",
"forecast:DescribeDatasetImportJob",
"forecast:DescribeDataset",
"forecast:DescribeForecast",
"forecast:DescribeForecastExportJob",
"forecast:DescribePredictorBacktestExportJob",
"forecast:GetAccuracyMetrics",
"forecast:InvokeForecastEndpoint",
"forecast:GetRecentForecastContext",
"forecast:DescribePredictor",
"forecast:TagResource",
"forecast:DeleteResourceTree"
],
"Resource": [
  "arn:aws:forecast:*:*:*Canvas*"
],
{
  "Sid": "RDSOperation",
  "Effect": "Allow",
  "Action": "rds:DescribeDBInstances",
  "Resource": "*"
},
{
  "Sid": "IAMPassOperationForForecast",
  "Effect": "Allow",
  "Action": ["iam:PassRole"]
},
"Resource": "arn:aws:iam::*:role/*",
"Condition": {
  "StringEquals": {
    "iam:PassedToService": "forecast.amazonaws.com"
  }
}
],
{
  "Sid": "AutoscalingOperations",
  "Effect": "Allow",
  "Action": ["application-autoscaling:PutScalingPolicy",
    "application-autoscaling:RegisterScalableTarget"
  ]
}
"Resource": "arn:aws:application-autoscaling::*:scalable-target/**",
"Condition": {
    "StringEquals": {
    "application-autoscaling:service-namespace": "sagemaker",
    "application-autoscaling:scalable-dimension": "sagemaker:variant:DesiredInstanceCount"
    }
    
},
{
  "Sid": "AsyncEndpointOperations",
  "Effect": "Allow",
  "Action": [
    "cloudwatch:DescribeAlarms",
    "sagemaker:DescribeEndpointConfig"
  ],
  "Resource": "*"
},
{
  "Sid": "SageMakerCloudWatchUpdate",
  "Effect": "Allow",
  "Action": [
    "cloudwatch:PutMetricAlarm",
    "cloudwatch:DeleteAlarms"
  ],
  "Resource": [
    "arn:aws:cloudwatch::*:*:alarm:*TargetTracking*"
  ],
  "Condition": {
    "StringEquals": {
    "aws:CalledViaLast": "application-autoscaling.amazonaws.com"
    }
  }
}
},
{
  "Sid": "AutoscalingSageMakerEndpointOperation",
  "Action": "iam:CreateServiceLinkedRole",
  "Effect": "Allow",
  "Resource": "arn:aws:iam::*:role/aws-service-role/sagemaker.application-autoscaling.amazonaws.com/AWSServiceRoleForApplicationAutoScaling_SageMakerEndpoint",
  "Condition": {
    "StringLike": {
      "iam:AWSServiceName": "sagemaker.application-autoscaling.amazonaws.com"}
AWS managed policy: AmazonSageMakerCanvasDataPrepFullAccess

This policy grants permissions that allow full access to the data preparation functionality of Amazon SageMaker Canvas. The policy also provides least privilege permissions for the services that integrate with the data preparation functionality [for example, Amazon Simple Storage Service (Amazon S3), AWS Identity and Access Management (IAM), Amazon EMR, Amazon EventBridge, Amazon Redshift, AWS Key Management Service (AWS KMS) and AWS Secrets Manager].

Permissions details

This AWS managed policy includes the following permissions.

- **sagemaker** – Allows principals to access processing jobs, training jobs, inference pipelines, AutoML jobs, and feature groups.
- **athena** – Allows principals to query a list of data catalogs, databases, and table metadata from Amazon Athena.
- **elasticmapreduce** – Allows principals to read and list Amazon EMR clusters.
- **events** – Allows principals to create, read, update, and add targets to Amazon EventBridge rules for scheduled jobs.
- **glue** – Allows principals to get and search tables from databases in the AWS Glue catalog.
- **iam** – Allows principals to pass an IAM role to Amazon SageMaker and EventBridge.
- **kms** – Allows principals to retrieve AWS KMS aliases stored in jobs and endpoints, and access the associated KMS key.
- **logs** – Allows principals to publish logs from training jobs and endpoints.
- **redshift** – Allows principals to get credentials to access an Amazon Redshift database.
- **redshift-data** – Allows principals to run, cancel, describe, list, and get the results of Amazon Redshift queries. Also allows principals to list Amazon Redshift schemas and tables.
- **s3** – Allows principals to add and retrieve objects from Amazon S3 buckets. These objects are limited to those whose name includes "SageMaker", "Sagemaker", or "sagemaker"; or is tagged with "SageMaker", case-insensitive.
• secretsmanager – Allows principals to store and retrieve customer database credentials using Secrets Manager.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "SageMakerListFeatureGroupOperation",
            "Effect": "Allow",
            "Action": "sagemaker:ListFeatureGroups",
            "Resource": "*"
        },
        {
            "Sid": "SageMakerFeatureGroupOperations",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateFeatureGroup",
                "sagemaker:DescribeFeatureGroup"
            ],
            "Resource": "arn:aws:sagemaker:*:*:feature-group/*"
        },
        {
            "Sid": "SageMakerProcessingJobOperations",
            "Effect": "Allow",
            "Action": [
                "sagemaker:CreateProcessingJob",
                "sagemaker:DescribeProcessingJob",
                "sagemaker:AddTags"
            ],
            "Resource": "arn:aws:sagemaker:*:*:processing-job/*canvas-data-prep*"
        },
        {
            "Sid": "SageMakerProcessingJobListOperation",
            "Effect": "Allow",
            "Action": "sagemaker:ListProcessingJobs",
            "Resource": "*"
        },
        {
            "Sid": "SageMakerPipelineOperations",
            "Effect": "Allow",
            "Action": "sagemaker:DescribePipeline",
            "Resource": "*
```
"sagemaker:CreatePipeline",
"sagemaker:UpdatePipeline",
"sagemaker:DeletePipeline",
"sagemaker:StartPipelineExecution",
"sagemaker:ListPipelineExecutionSteps",
"sagemaker:DescribePipelineExecution"
],
"Resource": "arn:aws:sagemaker:*:*:pipeline/*canvas-data-prep*"
},
{
"Sid": "KMSListOperations",
"Effect": "Allow",
"Action": "kms:ListAliases",
"Resource": "*"
},
{
"Sid": "KMSOperations",
"Effect": "Allow",
"Action": "kms:DescribeKey",
"Resource": "arn:aws:kms:*:*:key/*"
},
{
"Sid": "S3Operations",
"Effect": "Allow",
"Action": [
"s3:GetObject",
"s3:PutObject",
"s3:DeleteObject",
"s3:GetBucketCors",
"s3:GetBucketLocation",
"s3:AbortMultipartUpload"
],
"Resource": [
"arn:aws:s3:::*SageMaker*",
"arn:aws:s3:::*Sagemaker*",
"arn:aws:s3:::*sagemaker*"
],
"Condition": {
"StringEquals": {
"aws:ResourceAccount": "${aws:PrincipalAccount}"
}
}
}
"Sid": "S3GetObjectOperation",
"Effect": "Allow",
"Action": "s3:GetObject",
"Resource": "arn:aws:s3:::*",
"Condition": {
    "StringEqualsIgnoreCase": {
        "s3:ExistingObjectTag/SageMaker": "true"
    },
    "StringEquals": {
        "aws:ResourceAccount": "$\{aws:PrincipalAccount\}"
    }
}
],
{
    "Sid": "S3ListOperations",
    "Effect": "Allow",
    "Action": [
        "s3:ListBucket",
        "s3:ListAllMyBuckets"
    ],
    "Resource": "*"
},
{
    "Sid": "IAMListOperations",
    "Effect": "Allow",
    "Action": "iam:ListRoles",
    "Resource": "*"
},
{
    "Sid": "IAMGetOperations",
    "Effect": "Allow",
    "Action": "iam:GetRole",
    "Resource": "arn:aws:iam::*:role/*"
},
{
    "Sid": "IAMPassOperation",
    "Effect": "Allow",
    "Action": "iam:PassRole",
    "Resource": "arn:aws:iam::*:role/*",
    "Condition": {
        "StringEquals": {
            "iam:PassedToService": [
                "sagemaker.amazonaws.com",
                "events.amazonaws.com"
            ]
        }
    }
}
}


```json
  },
  {
    "Sid": "EventBridgePutOperation",
    "Effect": "Allow",
    "Action": [
      "events:PutRule"
    ],
    "Resource": "arn:aws:events:*:*:rule/*",
    "Condition": {
      "StringEquals": {
        "aws:RequestTag/sagemaker:is-canvas-data-prep-job": "true"
      }
    }
  },
  {
    "Sid": "EventBridgeOperations",
    "Effect": "Allow",
    "Action": [
      "events:DescribeRule",
      "events:PutTargets"
    ],
    "Resource": "arn:aws:events:*:*:rule/*",
    "Condition": {
      "StringEquals": {
        "aws:ResourceTag/sagemaker:is-canvas-data-prep-job": "true"
      }
    }
  },
  {
    "Sid": "EventBridgeTagBasedOperations",
    "Effect": "Allow",
    "Action": [
      "events:TagResource"
    ],
    "Resource": "arn:aws:events:*:*:rule/*",
    "Condition": {
      "StringEquals": {
        "aws:RequestTag/sagemaker:is-canvas-data-prep-job": "true",
        "aws:ResourceTag/sagemaker:is-canvas-data-prep-job": "true"
      }
    }
  }
```


},

{  
    "Sid": "EventBridgeListTagOperation",
    "Effect": "Allow",
    "Action": "events:ListTagsForResource",
    "Resource": "*"
},

{  
    "Sid": "GlueOperations",
    "Effect": "Allow",
    "Action": [
        "glue:GetDatabases",
        "glue:GetTable",
        "glue:GetTables",
        "glue:SearchTables"
    ],
    "Resource": [
        "arn:aws:glue:*:*:table/*",
        "arn:aws:glue:*:*:catalog",
        "arn:aws:glue:*:*:database/*"
    ]
},

{  
    "Sid": "EMROperations",
    "Effect": "Allow",
    "Action": [
        "elasticmapreduce:DescribeCluster",
        "elasticmapreduce:ListInstanceGroups"
    ],
    "Resource": "arn:aws:elasticmapreduce:*:*:cluster/*"
},

{  
    "Sid": "EMRListOperation",
    "Effect": "Allow",
    "Action": "elasticmapreduce:ListClusters",
    "Resource": "*"
},

{  
    "Sid": "AthenaListDataCatalogOperation",
    "Effect": "Allow",
    "Action": "athena:ListDataCatalogs",
    "Resource": "*"
},
"Sid": "AthenaQueryExecutionOperations",
"Effect": "Allow",
"Action": [
    "athena:GetQueryExecution",
    "athena:GetQueryResults",
    "athena:StartQueryExecution",
    "athena:StopQueryExecution"
],
"Resource": "arn:aws:athena:*:*:workgroup/**",
},
{
    "Sid": "AthenaDataCatalogOperations",
    "Effect": "Allow",
    "Action": [
        "athena:ListDatabases",
        "athena:ListTableMetadata"
    ],
    "Resource": "arn:aws:athena:*:*:datacatalog/**"
},
{
    "Sid": "RedshiftOperations",
    "Effect": "Allow",
    "Action": [
        "redshift-data:DescribeStatement",
        "redshift-data:CancelStatement",
        "redshift-data:GetStatementResult"
    ],
    "Resource": "*"
},
{
    "Sid": "RedshiftArnBasedOperations",
    "Effect": "Allow",
    "Action": [
        "redshift-data:ExecuteStatement",
        "redshift-data:ListSchemas",
        "redshift-data:ListTables"
    ],
    "Resource": "arn:aws:redshift:*:*:cluster:*"
},
{
    "Sid": "RedshiftGetCredentialsOperation",
    "Effect": "Allow",
    "Action": "redshift:GetClusterCredentials",
    "Resource": [  

"arn:aws:redshift:*:*:dbuser/*/sagemaker_access*",
"arn:aws:redshift:*:*:dbname:*"
],
},
{
"Sid": "SecretsManagerARNBasedOperation",
"Effect": "Allow",
"Action": "secretsmanager:CreateSecret",
"Resource": "arn:aws:secretsmanager:*:*:secret:AmazonSageMaker-*"
},
{
"Sid": "SecretManagerTagBasedOperation",
"Effect": "Allow",
"Action": [  
"secretsmanager:DescribeSecret",
"secretsmanager:GetSecretValue"
],  
"Resource": "arn:aws:secretsmanager:*:*:secret:AmazonSageMaker-*",
"Condition": {  
"StringEquals": {  
"aws:ResourceTag/SageMaker": "true",
"aws:ResourceAccount": "${aws:PrincipalAccount}"
}
}
},
{
"Sid": "RDSOperation",
"Effect": "Allow",
"Action": "rds:DescribeDBInstances",
"Resource": "*"
},
{
"Sid": "LoggingOperation",
"Effect": "Allow",
"Action": [  
"logs:CreateLogGroup",
"logs:CreateLogStream",
"logs:PutLogEvents"
],  
"Resource": "arn:aws:logs:*:*:log-group:/aws/sagemaker/studio:*"
}
]}
}
AWS managed policy: AmazonSageMakerCanvasDirectDeployAccess

This policy grants permissions needed for Amazon SageMaker Canvas to create and manage Amazon SageMaker endpoints.

Permissions details

This AWS managed policy includes the following permissions.

- **sagemaker** – Allows principals to create and manage SageMaker endpoints with an ARN resource name that starts with "Canvas" or "canvas".
- **cloudwatch** – Allows principals to retrieve Amazon CloudWatch metric data.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "SageMakerEndpointPerms",
         "Effect": "Allow",
         "Action": [
            "sagemaker:CreateEndpoint",
            "sagemaker:CreateEndpointConfig",
            "sagemaker:DeleteEndpoint",
            "sagemaker:DescribeEndpoint",
            "sagemaker:DescribeEndpointConfig",
            "sagemaker:InvokeEndpoint",
            "sagemaker:UpdateEndpoint"
         ],
         "Resource": [
            "arn:aws:sagemaker:*:*:Canvas*",
            "arn:aws:sagemaker:*:*:canvas*"
         ]
      },
      {
         "Sid": "ReadCWInvocationMetrics",
         "Effect": "Allow",
         "Action": "cloudwatch:GetMetricData",
         "Resource": "*"
      }
   ]
}
```
AWS managed policy: AmazonSageMakerCanvasAIServicesAccess

This policy grants permissions for Amazon SageMaker Canvas to use Amazon Textract, Amazon Rekognition, Amazon Comprehend, and Amazon Bedrock.

Permissions details

This AWS managed policy includes the following permissions.

- **textract** – Allows principals to use Amazon Textract to detect documents, expenses, and identities within an image.
- **rekognition** – Allows principals to use Amazon Rekognition to detect labels and text within an image.
- **comprehend** – Allows principals to use Amazon Comprehend to detect sentiment and dominant language, and named and personally identifiable information (PII) entities within a text document.
- **bedrock** – Allows principals to use Amazon Bedrock to list and invoke foundation models.
- **iam** – Allows principals to pass an IAM role to Amazon Bedrock.

```json
{
    "Version": "2012-10-17",
    "Statement": [
      {
        "Sid": "Textract",
        "Effect": "Allow",
        "Action": [
          "textract:AnalyzeDocument",
          "textract:AnalyzeExpense",
          "textract:AnalyzeID",
          "textract:StartDocumentAnalysis",
          "textract:StartExpenseAnalysis",
          "textract:GetDocumentAnalysis",
          "textract:GetExpenseAnalysis"
        ],
        "Resource": "*"
      },
      {
        "Sid": "Rekognition",
        "Effect": "Allow",
        "Action": [
          "rekognition:DetectLabels",
          "rekognition:DetectText",
          "rekognition:StartLabelDetection",
          "rekognition:StartTextDetection"
        ]
      }
    ]
}
```
"rekognition:DetectLabels",
"rekognition:DetectText"
],
"Resource": "*
},
{
"Sid": "Comprehend",
"Effect": "Allow",
"Action": [
"comprehend:BatchDetectDominantLanguage",
"comprehend:BatchDetectEntities",
"comprehend:BatchDetectSentiment",
"comprehend:DetectPiiEntities",
"comprehend:DetectEntities",
"comprehend:DetectSentiment",
"comprehend:DetectDominantLanguage"
],
"Resource": "*
},
{
"Sid": "Bedrock",
"Effect": "Allow",
"Action": [
"bedrock:InvokeModel",
"bedrock:ListFoundationModels",
"bedrock:InvokeModelWithResponseStream"
],
"Resource": "*
},
{
"Sid": "CreateBedrockResourcesPermission",
"Effect": "Allow",
"Action": [
"bedrock:CreateModelCustomizationJob",
"bedrock:CreateProvisionedModelThroughput",
"bedrock:TagResource"
],
"Resource": [
"arn:aws:bedrock:*:*:model-customization-job/**",
"arn:aws:bedrock:*:*:custom-model/**",
"arn:aws:bedrock:*:*:provisioned-model/**"
],
"Condition": {
"ForAnyValue:StringEquals": {
"aws:TagKeys": [
    "SageMaker",
    "Canvas"
  ],

  "StringEquals": {
    "aws:RequestTag/SageMaker": "true",
    "aws:RequestTag/Canvas": "true",
    "aws:ResourceTag/SageMaker": "true",
    "aws:ResourceTag/Canvas": "true"
  }
},

{
  "Sid": "GetStopAndDeleteBedrockResourcesPermission",
  "Effect": "Allow",
  "Action": [
    "bedrock:GetModelCustomizationJob",
    "bedrock:GetCustomModel",
    "bedrock:GetProvisionedModelThroughput",
    "bedrock:StopModelCustomizationJob",
    "bedrock:DeleteProvisionedModelThroughput"
  ],
  "Resource": [
    "arn:aws:bedrock:*::model-customization-job/*",
    "arn:aws:bedrock:*::custom-model/*",
    "arn:aws:bedrock:*::provisioned-model/*"
  ],
  "Condition": {
    "StringEquals": {
      "aws:ResourceTag/SageMaker": "true",
      "aws:ResourceTag/Canvas": "true"
    }
  }
},

{
  "Sid": "FoundationModelPermission",
  "Effect": "Allow",
  "Action": [
    "bedrock:CreateModelCustomizationJob"
  ],
  "Resource": [
    "arn:aws:bedrock:*::foundation-model/*"
  ]}
AWS managed policy: AmazonSageMakerCanvasBedrockAccess

This policy grants permissions commonly needed to use Amazon SageMaker Canvas with Amazon Bedrock.

Permissions details

This AWS managed policy includes the following permissions.

- **s3** – Allows principals to add and retrieve objects from Amazon S3 buckets in the "sagemaker-*/Canvas" directory.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "S3CanvasAccess",
      "Effect": "Allow",
      "Action": [
        "s3:GetObject",
        "s3:PutObject"
      ],
      "Resource": [
        "arn:aws:s3:::sagemaker-*/Canvas"
      ],
      "Condition": {
        "StringEquals": {
          "iam:PassedToService": "bedrock.amazonaws.com"
        }
      }
    }
  ]
}
```
AWS managed policy: AmazonSageMakerCanvasForecastAccess

This policy grants permissions commonly needed to use Amazon SageMaker Canvas with Amazon Forecast.

Permissions details

This AWS managed policy includes the following permissions.

- s3 – Allows principals to add and retrieve objects from Amazon S3 buckets. These objects are limited to those whose name starts with "sagemaker-".
Amazon SageMaker updates to Amazon SageMaker Canvas managed policies

View details about updates to AWS managed policies for SageMaker Canvas since this service began tracking these changes.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerCanvasBedrockAccess</td>
<td>1</td>
<td>Initial policy</td>
<td>February 2, 2024</td>
</tr>
<tr>
<td><strong>AmazonSageMakerCanvasFullAccess</strong></td>
<td>9</td>
<td>Add sagemaker:ListEndpoints permission.</td>
<td>January 24, 2024</td>
</tr>
<tr>
<td><strong>AmazonSageMakerCanvasFullAccess</strong></td>
<td>8</td>
<td>Add sagemaker:UpdateEndpointWeightsAndCapacities, sagemaker:DescribeEndpointConfig, sagemaker:InvokeEndpointAsync, athena:ListDataCatalogs, athena:GetQueryExec</td>
<td>December 8, 2023</td>
</tr>
<tr>
<td>Policy</td>
<td>Version</td>
<td>Change</td>
<td>Date</td>
</tr>
<tr>
<td>--------------------------------------------</td>
<td>---------</td>
<td>------------------------------------------------------------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>AmazonSageMakerCanvasDataPrepFullAccess</td>
<td>2</td>
<td>Small update to enforce the intents of the previous policy, version 1; no permissions added or deleted.</td>
<td>December 7, 2023</td>
</tr>
</tbody>
</table>

Amazon SageMaker Canvas Data Prep Full Access - Update to an existing policy.
<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AmazonSageMakerCanvasData Prep Full Access</strong></td>
<td>1</td>
<td>Initial policy</td>
<td>October 26, 2023</td>
</tr>
<tr>
<td><strong>AmazonSageMakerCanvas Direct Deploy Access</strong></td>
<td>1</td>
<td>Initial policy</td>
<td>October 6, 2023</td>
</tr>
<tr>
<td>Policy</td>
<td>Version</td>
<td>Change</td>
<td>Date</td>
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<tr>
<td>--------</td>
<td>---------</td>
<td>--------</td>
<td>------</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasFullAccess - Update to an existing policy</td>
<td>7</td>
<td>Add <code>sagemaker:DeleteEndpointConfig</code>, <code>sagemaker:DeleteModel</code>, and <code>sagemaker:InvokeEndpoint</code> permissions. Also add <code>s3:GetObject</code> permission for SageMaker JumpStart resources in specific regions.</td>
<td>September 29, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasAIServicesAccess - Update to an existing policy</td>
<td>2</td>
<td>Add <code>bedrock:InvokeModel</code> and <code>bedrock:ListFoundationModels</code> permissions.</td>
<td>September 29, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasFullAccess - Update to an existing policy</td>
<td>6</td>
<td>Add <code>rds:DescribeDBInstances</code> permission.</td>
<td>August 29, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasFullAccess - Update to an existing policy</td>
<td>5</td>
<td>Add <code>application-autoscaling:PutScalingPolicy</code> and <code>application-autoscaling:RegisterScalableTarget</code> permissions.</td>
<td>July 24, 2023</td>
</tr>
<tr>
<td>Policy</td>
<td>Version</td>
<td>Change</td>
<td>Date</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>---------</td>
<td>------------------------------------------------------------------------</td>
<td>------------------</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasFullAccess - Update to an existing policy</td>
<td>3</td>
<td>Add sagemaker:CreateAutoMLJobV2, sagemaker:DescribeAutoMLJobV2, and glue:SearchTables permissions.</td>
<td>March 24, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasAIServicesAccess - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>March 23, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerCan vasFullAccess - Update to an existing policy</td>
<td>2</td>
<td>Add forecast:DeleteResourceTree permission.</td>
<td>December 6, 2022</td>
</tr>
</tbody>
</table>
AWS managed policies for Amazon SageMaker Cluster

These AWS managed policies add permissions required to use SageMaker Cluster. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.

Topics

- AWS managed policy: AmazonSageMakerClusterInstanceRolePolicy
- Amazon SageMaker updates to Amazon SageMaker Cluster managed policies

AWS managed policy: AmazonSageMakerClusterInstanceRolePolicy

This policy grants permissions commonly needed to use Amazon SageMaker Cluster.

Permissions details

This AWS managed policy includes the following permissions.

- cloudwatch – Allows principals to post Amazon CloudWatch metrics.
- logs – Allows principals to publish CloudWatch log streams.
- s3 – Allows principals to list and retrieve lifecycle script files from an Amazon S3 bucket in your account. These buckets are limited to those whose name starts with "sagemaker-".
- ssmmessages – Allows principals to open a connection to AWS Systems Manager.

```json
{
    "Version" : "2012-10-17",
    "Statement" : [
```
{
    "Sid" : "CloudwatchLogStreamPublishPermissions",
    "Effect" : "Allow",
    "Action" : [
        "logs:PutLogEvents",
        "logs:CreateLogStream",
        "logs:DescribeLogStreams"
    ],
    "Resource" : [
        "arn:aws:logs:*:*:log-group:/aws/sagemaker/Clusters/*:log-stream:*"
    ],
},
{
    "Sid" : "CloudwatchLogGroupCreationPermissions",
    "Effect" : "Allow",
    "Action" : [
        "logs:CreateLogGroup"
    ],
    "Resource" : [
        "arn:aws:logs:*:*:log-group:/aws/sagemaker/Clusters/*"
    ],
},
{
    "Sid" : "CloudwatchPutMetricDataAccess",
    "Effect" : "Allow",
    "Action" : [
        "cloudwatch:PutMetricData"
    ],
    "Resource" : [
        "*"
    ],
    "Condition" : {
        "StringEquals" : {
            "cloudwatch:namespace" : "/aws/sagemaker/Clusters"
        }
    }
},
{
    "Sid" : "DataRetrievalFromS3BucketPermissions",
    "Effect" : "Allow",
    "Action" : [
        "s3:ListBucket",
        "s3:GetObject"
    ]
}
Amazon SageMaker updates to Amazon SageMaker Cluster managed policies

View details about updates to AWS managed policies for SageMaker Cluster since this service began tracking these changes. For automatic alerts about changes to this page, subscribe to the RSS feed on the SageMaker Document history page.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerClusterInstanceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>November 29, 2023</td>
</tr>
</tbody>
</table>

AWS managed policies for Amazon SageMaker Feature Store

These AWS managed policies add permissions required to use Feature Store. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.
Topics

- AWS managed policy: AmazonSageMakerFeatureStoreAccess
- Amazon SageMaker updates to Amazon SageMaker Feature Store managed policies

AWS managed policy: AmazonSageMakerFeatureStoreAccess

This policy grants permissions required to enable the offline store for an Amazon SageMaker Feature Store feature group.

Permissions details

This AWS managed policy includes the following permissions.

- s3 – Allows principals to write data into an offline store Amazon S3 bucket. These buckets are limited to those whose name includes "SageMaker", "Sagemaker", or "sagemaker".
- s3 – Allows principals to read existing manifest files maintained in the metadata folder of an offline store S3 bucket.
- glue – Allows principals to read and update AWS Glue tables. These permissions are limited to tables in the sagemaker_featurestore folder.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": [
            "s3:PutObject",
            "s3:GetBucketAcl",
            "s3:PutObjectAcl"
         ],
         "Resource": [
            "arn:aws:s3:::*SageMaker*",
            "arn:aws:s3:::*Sagemaker*",
            "arn:aws:s3:::*sagemaker*"
         ]
      },
      {
         "Effect": "Allow",
         "Action": [
          "s3:PutObject",
          "s3:GetBucketAcl",
          "s3:PutObjectAcl"
         ],
         "Resource": [
            "arn:aws:s3:::*SageMaker*",
            "arn:aws:s3:::*Sagemaker*",
            "arn:aws:s3:::*sagemaker*"
         ]
      }
   ]
}
```
Amazon SageMaker updates to Amazon SageMaker Feature Store managed policies

View details about updates to AWS managed policies for Feature Store since this service began tracking these changes. For automatic alerts about changes to this page, subscribe to the RSS feed on the SageMaker Document history page.

<table>
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<tr>
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<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerFeatureStoreAccess - Update to an existing policy</td>
<td>3</td>
<td>Add s3:GetObject, glue:GetTable, and glue:UpdateTable permissions.</td>
<td>December 5, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerFeatureStoreAccess - Update to an existing policy</td>
<td>2</td>
<td>Add s3:PutObjectAcl permission.</td>
<td>February 23, 2021</td>
</tr>
</tbody>
</table>
AWS managed policies for Amazon SageMaker geospatial

These AWS managed policies add permissions required to use SageMaker geospatial. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.

Topics

- AWS managed policy: AmazonSageMakerGeospatialFullAccess
- AWS managed policy: AmazonSageMakerGeospatialExecutionRole
- Amazon SageMaker updates to Amazon SageMaker geospatial managed policies

AWS managed policy: AmazonSageMakerGeospatialFullAccess

This policy grants permissions that allow full access to Amazon SageMaker geospatial through the AWS Management Console and SDK.

Permissions details

This AWS managed policy includes the following permissions.

- sagemaker-geospatial – Allows principals full access to all SageMaker geospatial resources.
- iam – Allows principals to pass an IAM role to SageMaker geospatial.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Action": "sagemaker-geospatial:*",
         "Resource": "*"
      },
      {
```

AWS Managed Policies for SageMaker
AWS managed policy: AmazonSageMakerGeospatialExecutionRole

This policy grants permissions commonly needed to use SageMaker geospatial.

Permissions details

This AWS managed policy includes the following permissions.

- **s3** – Allows principals to add and retrieve objects from Amazon S3 buckets. These objects are limited to those whose name contains "SageMaker", "Sagemaker", or "sagemaker".

- **sagemaker-geospatial** – Allows principals to access Earth observation jobs through the GetEarthObservationJob API.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "s3:AbortMultipartUpload",
        "s3:PutObject",
        "s3:GetObject",
        "s3:ListBucketMultipartUploads"
      ],
      "Resource": [
        "arn:aws:s3:::*SageMaker*",
        "arn:aws:s3:::*Sagemaker*",
        "arn:aws:s3:::*sagemaker*"
      ]
    }
  ]
}
```
Amazon SageMaker updates to Amazon SageMaker geospatial managed policies

View details about updates to AWS managed policies for SageMaker geospatial since this service began tracking these changes.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerGeospatialExecutionRole - Updated policy</td>
<td>2</td>
<td>Add sagemaker-geospatial:GetRasterDataCollection permission.</td>
<td>May 10, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerGeospatialFullAccess - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>November 30, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerGeospatialExecutionRole - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>November 30, 2022</td>
</tr>
</tbody>
</table>

AWS Managed Policies for Amazon SageMaker Ground Truth

These AWS managed policies add permissions required to use SageMaker Ground Truth. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.
Topics

- AWS managed policy: AmazonSageMakerGroundTruthExecution
- Amazon SageMaker updates to SageMaker Ground Truth managed policies

AWS managed policy: AmazonSageMakerGroundTruthExecution

This AWS managed policy grants permissions commonly needed to use SageMaker Ground Truth.

Permissions details

This policy includes the following permissions.

- lambda – Allows principals to invoke Lambda functions whose name includes "sagemaker" (case-insensitive), "GtRecipe", or "LabelingFunction".
- s3 – Allows principals to add and retrieve objects from Amazon S3 buckets. These objects are limited to those whose case-insensitive name contains "groundtruth" or "sagemaker", or are tagged with "SageMaker".
- cloudwatch – Allows principals to post CloudWatch metrics.
- logs – Allows principals to create and access log streams, and post log events.
- sqs – Allows principals to create Amazon SQS queues, and send and receive Amazon SQS messages. These permissions are limited to queues whose name includes "GroundTruth".
- sns – Allows principals to subscribe to and publish messages to Amazon SNS topics whose case-insensitive name contains "groundtruth" or "sagemaker".
- ec2 – Allows principals to create, describe, and delete Amazon VPC endpoints whose VPC endpoint service name contains "sagemaker-task-resources" or "labeling".

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "CustomLabelingJobs",
            "Effect": "Allow",
            "Action": [
                "lambda:InvokeFunction"
            ],
            "Resource": [
                "arn:aws:lambda:*:*:function:*GtRecipe*",
```
"arn:aws:lambda:*:*:function:*LabelingFunction*",
"arn:aws:lambda:*:*:function:*SageMaker*",
"arn:aws:lambda:*:*:function:*sagemaker*",
"arn:aws:lambda:*:*:function:*Sagemaker*"
]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:AbortMultipartUpload",
    "s3:GetObject",
    "s3:PutObject"
  ],
  "Resource": [  
    "arn:aws:s3:::*GroundTruth*",
    "arn:aws:s3:::*Groundtruth*",
    "arn:aws:s3:::*groundtruth*",
    "arn:aws:s3:::*SageMaker*",
    "arn:aws:s3:::*Sagemaker*",
    "arn:aws:s3:::*sagemaker*"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:GetObject"
  ],
  "Resource": "*",
  "Condition": {
    "StringEqualsIgnoreCase": {
      "s3:ExistingObjectTag/SageMaker": "true"
    }
  }
},
{
  "Sid": "CloudWatch",
}
"Effect": "Allow",
"Action": [
  "cloudwatch:PutMetricData",
  "logs:CreateLogStream",
  "logs:CreateLogGroup",
  "logs:DescribeLogStreams",
  "logs:PutLogEvents"
],
"Resource": "*
",

"Sid": "StreamingQueue",
"Effect": "Allow",
"Action": [
  "sqs:CreateQueue",
  "sqs:DeleteMessage",
  "sqs:GetQueueAttributes",
  "sqs:GetQueueUrl",
  "sqs:ReceiveMessage",
  "sqs:SendMessage",
  "sqs:SetQueueAttributes"
],
"Resource": "arn:aws:sqs:*:*:*GroundTruth*"
",

"Sid": "StreamingTopicSubscribe",
"Effect": "Allow",
"Action": "sns:Subscribe",
"Resource": [
  "arn:aws:sns:*::*:*GroundTruth*",
  "arn:aws:sns:*::*:*Groundtruth*",
  "arn:aws:sns:*::*:*groundTruth*",
  "arn:aws:sns:*::*:*groundtruth*",
  "arn:aws:sns:*::*:*SageMaker*",
  "arn:aws:sns:*::*:*Sagemaker*",
  "arn:aws:sns:*::*:*sageMaker*",
  "arn:aws:sns:*::*:*sagemaker*
],
"Condition": {
  "StringEquals": {
    "sns:Protocol": "sqs"
  },
  "StringLike": {
    "sns:Endpoint": "arn:aws:sqs:*::*:*GroundTruth*"
  }
}
"Sid": "StreamingTopic",
"Effect": "Allow",
"Action": [
  "sns:Publish"
],
"Resource": [
  "arn:aws:sns:*:*:*GroundTruth*",
  "arn:aws:sns:*:*:*Groundtruth*",
  "arn:aws:sns:*:*:*groundTruth*",
  "arn:aws:sns:*:*:*groundtruth*",
  "arn:aws:sns:*:*:*SageMaker*",
  "arn:aws:sns:*:*:*Sagemaker*",
  "arn:aws:sns:*:*:*sageMaker*",
  "arn:aws:sns:*:*:*sagemaker*"
]
},
{
  "Sid": "StreamingTopicUnsubscribe",
  "Effect": "Allow",
  "Action": [
    "sns:Unsubscribe"
  ],
  "Resource": "*
",
  "Resource": "*
",
  "Sid": "WorkforceVPC",
  "Effect": "Allow",
  "Action": [
    "ec2:CreateVpcEndpoint",
    "ec2:DescribeVpcEndpoints",
    "ec2:DeleteVpcEndpoints"
  ],
  "Resource": "*
",
  "Condition": {
    "StringLikeIfExists": {
      "ec2:VpceServiceName": [
        "sagemaker-task-resources*",
        "aws.sagemaker*labeling*
      ]
    }
  }
}
## Amazon SageMaker updates to SageMaker Ground Truth managed policies

View details about updates to AWS managed policies for Amazon SageMaker Ground Truth since this service began tracking these changes.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerGroundTruthExecution - Update to an existing policy</td>
<td>3</td>
<td>Add ec2:CreateVpcEndpoint, ec2:DescribeVpcEndpoints, and ec2:DeleteVpcEndpoints permissions.</td>
<td>April 29, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerGroundTruthExecution - Update to an existing policy</td>
<td>2</td>
<td>Remove sqs:SendMessageBatch permission.</td>
<td>April 11, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerGroundTruthExecution - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>July 20, 2020</td>
</tr>
</tbody>
</table>

## AWS Managed Policies for SageMaker Model Governance

This AWS managed policy adds permissions required to use SageMaker Model Governance. The policy is available in your AWS account and is used by execution roles created from the SageMaker console.

### Topics
- AWS managed policy: AmazonSageMakerModelGovernanceUseAccess
- Amazon SageMaker updates to SageMaker Model Governance managed policies
**AWS managed policy: AmazonSageMakerModelGovernanceUseAccess**

This AWS managed policy grants permissions needed to use all Amazon SageMaker Governance features. The policy is available in your AWS account.

This policy includes the following permissions.

- **s3** – Retrieve objects from Amazon S3 buckets. Retrievable objects are limited to those whose case-insensitive name contains the string "sagemaker".
- **kms** – List the AWS KMS keys to use for content encryption.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "sagemaker:ListMonitoringAlerts",
        "sagemaker:ListMonitoringExecutions",
        "sagemaker:UpdateMonitoringAlert",
        "sagemaker:StartMonitoringSchedule",
        "sagemaker:StopMonitoringSchedule",
        "sagemaker:ListMonitoringAlertHistory",
        "sagemaker:DescribeModelPackage",
        "sagemaker:DescribeModelPackageGroup",
        "sagemaker:CreateModelCard",
        "sagemaker:DescribeModelCard",
        "sagemaker:UpdateModelCard",
        "sagemaker:DeleteModelCard",
        "sagemaker:ListModelCards",
        "sagemaker:ListModelCardVersions",
        "sagemaker:CreateModelCardExportJob",
        "sagemaker:DescribeModelCardExportJob",
        "sagemaker:ListModelCardExportJobs"
      ],
      "Resource": "*"
    },
    {
      "Effect": "Allow",
      "Action": [
        "sagemaker:ListTrainingJobs",
        "sagemaker:DescribeTrainingJob",
        "sagemaker:DeleteTrainingJob",
        "sagemaker:StartTrainingJob",
        "sagemaker:StopTrainingJob",
        "sagemaker:CreateTrainingJob",
        "sagemaker:DescribeTrainingJobWithResultStream",
        "sagemaker:DescribeModel`
```
"sagemaker:ListModels",
"sagemaker:DescribeModel",
"sagemaker:Search",
"sagemaker:AddTags",
"sagemaker:DeleteTags",
"sagemaker:ListTags"
],
"Resource": "*",
},
{
"Effect": "Allow",
"Action": [
"kms:ListAliases"
],
"Resource": "*",
},
{
"Effect": "Allow",
"Action": [
"s3:GetObject",
"s3:PutObject",
"s3:CreateBucket",
"s3:GetBucketLocation",
],
"Resource": ["arn:aws:s3:::*SageMaker*",
"arn:aws:s3:::*Sagemaker*",
"arn:aws:s3:::*sagemaker*"
]
},
{
"Effect": "Allow",
"Action": [
"s3:ListBucket",
"s3:ListAllMyBuckets"
],
"Resource": "*"
}
Amazon SageMaker updates to SageMaker Model Governance managed policies

View details about updates to AWS managed policies for SageMaker Model Governance since this service began tracking these changes. For automatic alerts about changes to this page, subscribe to the RSS feed on the SageMaker Document history page.

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<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerModelGovernanceUseAccess - Update to an existing policy</td>
<td>2</td>
<td>Add sagemaker:DescribeModelPackage and DescribeModelPackageGroup permissions.</td>
<td>July 17, 2023</td>
</tr>
<tr>
<td>AmazonSageMakerModelGovernanceUseAccess - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>November 30, 2022</td>
</tr>
</tbody>
</table>

AWS Managed Policies for Model Registry

These AWS managed policies adds permissions required to use Model Registry. The policies are available in your AWS account and are used by execution roles created from the Amazon SageMaker console.

Topics
- AWS managed policy: AmazonSageMakerModelRegistryFullAccess
- Amazon SageMaker updates to Model Registry managed policies

AWS managed policy: AmazonSageMakerModelRegistryFullAccess

This AWS managed policy grants permissions needed to use all Model Registry features inside an Amazon SageMaker domain. This policy is attached to an execution role when configuring Model Registry settings to enable Model Registry permissions.

This policy includes the following permissions.

- `ecr` – Allows principals to retrieve information, including metadata, about Amazon Elastic Container Registry (Amazon ECR) images.
• **iam** – Allows principals to pass the execution role to the Amazon SageMaker service.

• **resource-groups** – Allows principals to create, list, tag, and delete AWS Resource Groups.

• **s3** – Allows principals to retrieve objects from the Amazon Simple Storage Service (Amazon S3) buckets where model versions are stored. Retrievable objects are limited to those whose case-insensitive name contains the string "sagemaker".

• **sagemaker** – Allows principals to catalog, manage, and deploy models using the SageMaker model registry.

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "sagemaker:DescribeAction",
        "sagemaker:DescribeInferenceRecommendationsJob",
        "sagemaker:DescribeModelPackage",
        "sagemaker:DescribeModelPackageGroup",
        "sagemaker:DescribePipeline",
        "sagemaker:DescribePipelineExecution",
        "sagemaker:ListAssociations",
        "sagemaker:ListArtifacts",
        "sagemaker:ListModelMetadata",
        "sagemaker:ListModelPackages",
        "sagemaker:Search",
        "sagemaker:GetSearchSuggestions"
    ],
    "Resource": "**"
  },
  {
    "Effect": "Allow",
    "Action": [
      "sagemaker:AddTags",
      "sagemaker:CreateModel",
      "sagemaker:CreateModelPackage",
      "sagemaker:CreateModelPackageGroup",
      "sagemaker:CreateEndpoint",
      "sagemaker:CreateEndpointConfig",
      "sagemaker:CreateInferenceRecommendationsJob",
      "sagemaker:DeleteModelPackage",
      "sagemaker:DeleteModelPackageGroup",
      "sagemaker:GetModelArtifacts",
      "sagemaker:ListInferenceRecommendationsJobs",
      "sagemaker:PutInferenceRecommendations",
      "sagemaker:PutModelPackageGroupPolicy",
      "sagemaker:PutModelVersionPolicy",
      "sagemaker:PutModelPolicy",
      "sagemaker:PutPipelinePolicy",
      "sagemaker:PutPredictionJobDefinition",
      "sagemaker:PutProvisionedResourceProperties",
      "sagemaker:PutS3DataAccessPolicy",
      "sagemaker:UpdateEndpointGroup",
      "sagemaker:UpdateInferenceRecommendationsJob",
      "sagemaker:UpdateInferenceRecommendationsJobOutputConfig",
      "sagemaker:UpdatePipeline",
      "sagemaker:UpdatePipelineExecution",
      "sagemaker:UpdateS3DataAccessPolicy"
    ],
    "Resource": "**"
  }
}
```
"sagemaker:DeleteTags",
"sagemaker:UpdateModelPackage"
],
"Resource": "*
",
{
    "Effect": "Allow",
    "Action": [
        "s3:GetObject"
    ],
    "Resource": [
        "arn:aws:s3:::*SageMaker*",
        "arn:aws:s3:::*Sagemaker*",
        "arn:aws:s3:::*sagemaker*"
    ]
},
{
    "Effect": "Allow",
    "Action": [
        "s3:ListBucket",
        "s3:ListAllMyBuckets"
    ],
    "Resource": "*
",
},
{
    "Effect": "Allow",
    "Action": [
        "ecr:BatchGetImage",
        "ecr:DescribeImages"
    ],
    "Resource": "*
",
},
{
    "Effect": "Allow",
    "Action": [
        "iam:PassRole"
    ],
    "Resource": "arn:aws:iam::*:role/*",
    "Condition": {
        "StringEquals": {
            "iam:PassedToService": "sagemaker.amazonaws.com"
        }
    }
},
}
["Effect": "Allow",
"Action": [
  "tag:GetResources"
],
"Resource": "*"],

["Effect": "Allow",
"Action": [
  "resource-groups:GetGroupQuery"
],
"Resource": "arn:aws:resource-groups::*:*:group/*"],

["Effect": "Allow",
"Action": [
  "resource-groups:ListGroupResources"
],
"Resource": "*"],

["Effect": "Allow",
"Action": [
  "resource-groups:CreateGroup",
  "resource-groups:Tag"
],
"Resource": "arn:aws:resource-groups::*:*:group/*",
"Condition": {
  "ForAnyValue:StringEquals": {
    "aws:TagKeys": "sagemaker:collection"
  }
}

["Effect": "Allow",
"Action": "resource-groups:DeleteGroup",
"Resource": "arn:aws:resource-groups::*:*:group/*",
"Condition": {
  "StringEquals": {
    "aws:ResourceTag/sagemaker:collection": "true"
  }
}
]
Amazon SageMaker updates to Model Registry managed policies

View details about updates to AWS managed policies for Model Registry since this service began tracking these changes. For automatic alerts about changes to this page, subscribe to the RSS feed on the SageMaker Document history page.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerModelRegistryFullAccess - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>April 12, 2023</td>
</tr>
</tbody>
</table>

AWS Managed Policies for SageMaker Notebooks

These AWS managed policies add permissions required to use SageMaker Notebooks. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.

Topics
- AWS managed policy: AmazonSageMakerNotebooksServiceRolePolicy
- Amazon SageMaker updates to SageMaker Notebooks managed policies

AWS managed policy: AmazonSageMakerNotebooksServiceRolePolicy

This AWS managed policy grants permissions commonly needed to use Amazon SageMaker Notebooks. The policy is added to the AmazonSageMaker-ExecutionRole that is created when you onboard to Amazon SageMaker Studio Classic. For more information on service-linked roles, see Service-Linked Roles.

Permissions details

This policy includes the following permissions.

- elasticfilesystem – Allows principals to create and delete Amazon Elastic File System (EFS) file systems, access points, and mount targets. These are limited to those tagged with the key
`ManagedByAmazonSageMakerResource`. Allows principals to describe all EFS file systems, access points, and mount targets. Allows principals to create or overwrite tags for EFS access points and mount targets.

- **ec2** – Allows principals to create network interfaces and security groups for Amazon Elastic Compute Cloud (EC2) instances. Also allows principals to create and overwrite tags for these resources.
- **sso** – Allows principals to add and delete managed application instances to AWS IAM Identity Center.
- **sagemaker** – Allows principals to create and read SageMaker user profiles.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": "elasticfilesystem:CreateAccessPoint",
            "Resource": "arn:aws:elasticfilesystem:*:*:file-system/**",
            "Condition": {
                "StringLike": {
                    "aws:ResourceTag/ManagedByAmazonSageMakerResource": "*",
                    "aws:RequestTag/ManagedByAmazonSageMakerResource": "*"
                }
            }
        },
        {
            "Effect": "Allow",
            "Action": [ "elasticfilesystem:DeleteAccessPoint" ],
            "Resource": "arn:aws:elasticfilesystem:*:*:access-point/**",
            "Condition": {
                "StringLike": {
                    "aws:ResourceTag/ManagedByAmazonSageMakerResource": "*"
                }
            }
        },
        {
            "Effect": "Allow",
            "Action": "elasticfilesystem:CreateFileSystem",
            "Resource": "*",
            "Condition": {
                "StringLike": {
                    "aws:ResourceTag/ManagedByAmazonSageMakerResource": "*",
                    "aws:RequestTag/ManagedByAmazonSageMakerResource": "*"
                }
            }
        }
    ]
}
```
"Condition": {
    "StringLike": {
        "aws:RequestTag/ManagedByAmazonSageMakerResource": "*"
    }
},

"Effect": "Allow",
"Action": [
    "elasticfilesystem:CreateMountTarget",
    "elasticfilesystem:DeleteFileSystem",
    "elasticfilesystem:DeleteMountTarget"
],
"Resource": "*",
"Condition": {
    "StringLike": {
        "aws:ResourceTag/ManagedByAmazonSageMakerResource": "*"
    }
}]
},

"Effect": "Allow",
"Action": [
    "elasticfilesystem:DescribeAccessPoints",
    "elasticfilesystem:DescribeFileSystems",
    "elasticfilesystem:DescribeMountTargets"
],
"Resource": "*
},

"Effect": "Allow",
"Action": "elasticfilesystem:TagResource",
"Resource": [
    "arn:aws:elasticfilesystem::*:*:access-point/**",
    "arn:aws:elasticfilesystem::*:*:file-system/**"
],
"Condition": {
    "StringLike": {
        "aws:ResourceTag/ManagedByAmazonSageMakerResource": "*"
    }
}]
},

"Effect": "Allow",
"Action": [
    "elasticfilesystem:CreateMountTarget",
    "elasticfilesystem:DeleteFileSystem",
    "elasticfilesystem:DeleteMountTarget"
],
"Resource": "*",
"Condition": {
    "StringLike": {
        "aws:RequestTag/ManagedByAmazonSageMakerResource": "*"
    }
}]
},

"Effect": "Allow",
"Action": [
    "elasticfilesystem:CreateMountTarget",
    "elasticfilesystem:DeleteFileSystem",
    "elasticfilesystem:DeleteMountTarget"
],
"Resource": "*",
"Condition": {
    "StringLike": {
        "aws:RequestTag/ManagedByAmazonSageMakerResource": "*"
    }
}]
},

"Effect": "Allow",
"Action": "ec2:CreateTags",
"Resource": [
   "arn:aws:ec2:*::*:network-interface/*",
   "arn:aws:ec2:*::*:security-group/*"
],
{
   "Effect": "Allow",
   "Action": [
      "ec2:CreateNetworkInterface",
      "ec2:CreateSecurityGroup",
      "ec2:DeleteNetworkInterface",
      "ec2:DescribeDhcpOptions",
      "ec2:DescribeNetworkInterfaces",
      "ec2:DescribeSecurityGroups",
      "ec2:DescribeSubnets",
      "ec2:DescribeVpcs",
      "ec2:ModifyNetworkInterfaceAttribute"
   ],
   "Resource": "*"
},
{
   "Effect": "Allow",
   "Action": [
      "ec2:AuthorizeSecurityGroupEgress",
      "ec2:AuthorizeSecurityGroupIngress",
      "ec2:CreateNetworkInterfacePermission",
      "ec2:DeleteNetworkInterfacePermission",
      "ec2:DeleteSecurityGroup",
      "ec2:RevokeSecurityGroupEgress",
      "ec2:RevokeSecurityGroupIngress"
   ],
   "Resource": "*",
   "Condition": {
      "StringLike": {
         "ec2:ResourceTag/ManagedByAmazonSageMakerResource": "*"
      }
   }
},
{
   "Effect": "Allow",
   "Action": [
      "sso:CreateManagedApplicationInstance",
      "sso:DeleteManagedApplicationInstance",
      "sso:GetManagedApplicationInstance",
      "sso:GetManagedApplicationInstances",
      "sso:ListManagedApplicationInstances",
      "sso:TagManagedApplicationInstance",
      "sso:UntagManagedApplicationInstance"
   ],
   "Resource": "*"
Amazon SageMaker updates to SageMaker Notebooks managed policies

View details about updates to AWS managed policies for Amazon SageMaker since this service began tracking these changes.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerNotebooksServiceRolePolicy</td>
<td>7</td>
<td>Added elasticfilesystem: TagResource permission.</td>
<td>March 9, 2023</td>
</tr>
</tbody>
</table>
AWS Managed Policies for SageMaker Pipelines

These AWS managed policies add permissions required to use SageMaker Pipelines. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.

Topics

- **AWS managed policy:** AmazonSageMakerPipelinesIntegrations
- Amazon SageMaker updates to SageMaker Pipelines managed policies

AWS managed policy: AmazonSageMakerPipelinesIntegrations

This AWS managed policy grants permissions commonly needed to use Callback steps and Lambda steps in SageMaker Pipelines. The policy is added to the AmazonSageMaker-ExecutionRole that is created when you onboard to Amazon SageMaker Studio Classic. The policy can be attached to any role used for authoring or executing a pipeline.

This policy grants appropriate AWS Lambda, Amazon Simple Queue Service (Amazon SQS), Amazon EventBridge, and IAM permissions needed when building pipelines that invoke Lambda functions or include callback steps, which can be used for manual approval steps or running custom workloads.

The Amazon SQS permissions allow you to create the Amazon SQS queue needed for receiving callback messages, and also to send messages to that queue.

The Lambda permissions allow you to create, read, update, and delete the Lambda functions used in the pipeline steps, and also to invoke those Lambda functions.

This policy grants the Amazon EMR permissions needed to run a pipelines Amazon EMR step.

Permissions details

This policy includes the following permissions.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>SageMaker started tracking changes for its AWS managed policies.</td>
<td></td>
<td>June 1, 2021</td>
<td></td>
</tr>
</tbody>
</table>
• **elasticmapreduce** – Read, add, and cancel steps in a running Amazon EMR cluster. Read, create, and terminate a new Amazon EMR cluster.

• **events** – Read, create, update, and add targets to an EventBridge rule named `SageMakerPipelineExecutionEMRStepStatusUpdateRule` and `SageMakerPipelineExecutionEMRClusterStatusUpdateRule`.

• **iam** – Pass an IAM role to the AWS Lambda service, Amazon EMR and Amazon EC2.

• **lambda** – Create, read, update, delete, and invoke Lambda functions. These permissions are limited to functions whose name includes "sagemaker".

• **sqs** – Create an Amazon SQS queue; send an Amazon SQS message. These permissions are limited to queues whose name includes "sagemaker".

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "lambda:CreateFunction",
        "lambda:DeleteFunction",
        "lambda:GetFunction",
        "lambda:InvokeFunction",
        "lambda:UpdateFunctionCode"
      ],
      "Resource": [
        "arn:aws:lambda:*:*:function:*sagemaker*",
        "arn:aws:lambda:*:*:function:*sageMaker*",
        "arn:aws:lambda:*:*:function:*SageMaker*
      ]
    },
    {
      "Effect": "Allow",
      "Action": [
        "sqs:CreateQueue",
        "sqs:SendMessage"
      ],
      "Resource": [
        "arn:aws:sqs:*::*:sagemaker*",
        "arn:aws:sqs:*::*:sageMaker*",
        "arn:aws:sqs:*::*:SageMaker*"
      ]
    }
  ]
}
```
{,
  "Effect": "Allow",
  "Action": [
    "iam:PassRole"
  ],
  "Resource": "arn:aws:iam::*:role/**",
  "Condition": {
    "StringEquals": {
      "iam:PassedToService": [
        "lambda.amazonaws.com",
        "elasticmapreduce.amazonaws.com",
        "ec2.amazonaws.com"
      ]
    }
  }
},
{
  "Effect": "Allow",
  "Action": [
    "events:DescribeRule",
    "events:PutRule",
    "events:PutTargets"
  ],
  "Resource": [
    "arn:aws:events:***:rule/SageMakerPipelineExecutionEMRStepStatusUpdateRule",
    "arn:aws:events:***:rule/SageMakerPipelineExecutionEMRClusterStatusUpdateRule"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "elasticmapreduce:AddJobFlowSteps",
    "elasticmapreduce:CancelSteps",
    "elasticmapreduce:DescribeStep",
    "elasticmapreduce:RunJobFlow",
    "elasticmapreduce:DescribeCluster",
    "elasticmapreduce:TerminateJobFlows",
    "elasticmapreduce:ListSteps"
  ],
  "Resource": [
    "arn:aws:elasticmapreduce:***:cluster/**
  ]
}
Amazon SageMaker updates to SageMaker Pipelines managed policies

View details about updates to AWS managed policies for Amazon SageMaker since this service began tracking these changes.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerPipelinesIntegrations - New policy</td>
<td>1</td>
<td>Initial policy</td>
<td>July 30, 2021</td>
</tr>
</tbody>
</table>
AWS Managed Policies for SageMaker projects and JumpStart

These AWS managed policies add permissions to use built-in Amazon SageMaker project templates and JumpStart solutions. The policies are available in your AWS account and are used by execution roles created from the SageMaker console.

SageMaker projects and JumpStart use AWS Service Catalog to provision AWS resources in customers' accounts. Some created resources need to assume an execution role. For example, if AWS Service Catalog creates a CodePipeline pipeline on behalf of a customer for a SageMaker machine learning CI/CD project, then that pipeline requires an IAM role.

The AmazonSageMakerServiceCatalogProductsLaunchRole role has the permissions required to launch the SageMaker portfolio of products from AWS Service Catalog. The AmazonSageMakerServiceCatalogProductsUseRole role has the permissions required to use the SageMaker portfolio of products from AWS Service Catalog. The AmazonSageMakerServiceCatalogProductsLaunchRole role passes an AmazonSageMakerServiceCatalogProductsUseRole role to the provisioned AWS Service Catalog product resources.

Topics
- AWS managed policy: AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy
- AWS managed policy: AmazonSageMakerPartnerServiceCatalogProductsApiGatewayServiceRolePolicy
- AWS managed policy: AmazonSageMakerPartnerServiceCatalogProductsCloudFormationServiceRolePolicy
- AWS managed policy: AmazonSageMakerPartnerServiceCatalogProductsLambdaServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsApiGatewayServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsCloudformationServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsCodeBuildServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsCodePipelineServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsEventsServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsFirehoseServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsGlueServiceRolePolicy
- AWS managed policy: AmazonSageMakerServiceCatalogProductsLambdaServiceRolePolicy
Amazon SageMaker updates to AWS Service Catalog AWS managed policies

AWS managed policy: AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy

This service role policy is used by the AWS Service Catalog service to provision products from the Amazon SageMaker portfolio. The policy grants permissions to a set of related AWS services including AWS CodePipeline, AWS CodeBuild, AWS CodeCommit, AWS Glue, AWS CloudFormation, and others.

The AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy policy is intended to be used by the AmazonSageMakerServiceCatalogProductsLaunchRole role created from the SageMaker console. The policy adds permissions to provision AWS resources for SageMaker projects and JumpStart using Service Catalog to a customer's account.

Permissions details

This policy includes the following permissions.

- **apigateway** – Allows the role to call API Gateway endpoints that are tagged with `sagemaker:launch-source`.
- **cloudformation** – Allows AWS Service Catalog to create, update and delete CloudFormation stacks.
- **codebuild** – Allows the role assumed by AWS Service Catalog and passed to CloudFormation to create, update and delete CodeBuild projects.
- **codecommit** – Allows the role assumed by AWS Service Catalog and passed to CloudFormation to create, update and delete CodeCommit repositories.
- **codepipeline** – Allows the role assumed by AWS Service Catalog and passed to CloudFormation to create, update and delete CodePipelines.
- **codestar-connections** – Allows the role to pass AWS CodeStar connections.
- **cognito-idp** – Allows the role to create, update, and delete groups and user pools. Also allows tagging resources.
- **ecr** – Allows the role assumed by AWS Service Catalog and passed to CloudFormation to create and delete Amazon ECR repositories. Also allows tagging resources.
- **events** – Allows the role assumed by AWS Service Catalog and passed to CloudFormation to create and delete EventBridge rules. Used for tying together the various components of the CICD pipeline.
• **firehose** – Allows the role to interact with Firehose streams.

• **glue** – Allows the role to interact with AWS Glue.

• **iam** – Allows the role to pass roles prepended with AmazonSageMakerServiceCatalog. This is needed when Projects provisions a AWS Service Catalog product, as a role needs to be passed to AWS Service Catalog.

• **lambda** – Allows the role to interact with AWS Lambda. Also allows tagging resources.

• **logs** – Allows the role to create, delete and access log streams.

• **s3** – Allows the role assumed by AWS Service Catalog and passed to CloudFormation to access Amazon S3 buckets where the Project template code is stored.

• **sagemaker** – Allows the role to interact with various SageMaker services. This is done both in CloudFormation during template provisioning, as well as in CodeBuild during CICD pipeline execution. Also allows tagging the following resources: endpoints, endpoint configurations, models, pipelines, projects, and model packages.

• **states** – Allows the role to create, delete, and update Step Functions prepended with sagemaker.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "apigateway:GET",
                "apigateway:POST",
                "apigateway:PUT",
                "apigateway:PATCH",
                "apigateway:DELETE"
            ],
            "Resource": "*",
            "Condition": {
                "StringLike": {
                    "aws:ResourceTag/sagemaker:launch-source": "*"
                }
            }
        },
        {
            "Effect": "Allow",
            "Action": [
```
"apigateway:POST",

"Resource": "*",

"Condition": {
  "ForAnyValue:StringLike": {
    "aws:TagKeys": [
      "sagemaker:launch-source"
    ]
  }
},

{
  "Effect": "Allow",
  "Action": [
    "apigateway:PATCH"
  ],
  "Resource": [
    "arn:aws:apigateway:*::/account"
  ]
},

{
  "Effect": "Allow",
  "Action": [
    "cloudformation:CreateStack",
    "cloudformation:UpdateStack",
    "cloudformation:DeleteStack"
  ],
  "Resource": "arn:aws:cloudformation:*::*:stack/SC-**",
  "Condition": {
    "ArnLikeIfExists": {
      "cloudformation:RoleArn": [
        "arn:aws:sts::*:assumed-role/AmazonSageMakerServiceCatalog*
      ]
    }
  }
},

{
  "Effect": "Allow",
  "Action": [
    "cloudformation:DescribeStackEvents",
    "cloudformation:DescribeStacks"
  ],
  "Resource": "arn:aws:cloudformation:*::*:stack/SC-**"
}
{ "Effect": "Allow", "Action": [ "cloudformation:GetTemplateSummary", "cloudformation:ValidateTemplate" ], "Resource": "*" },
{
    "Effect": "Allow",
    "Action": [ "codebuild:CreateProject",
                "codebuild:DeleteProject",
                "codebuild:UpdateProject"
    ],
    "Resource": [ "arn:aws:codebuild:*::*:project/sagemaker-*"
    ]
},
{
    "Effect": "Allow",
    "Action": [ "codecommit:CreateCommit",
                "codecommit:CreateRepository",
                "codecommit:DeleteRepository",
                "codecommit:GetRepository",
                "codecommit:TagResource"
    ],
    "Resource": [ "arn:aws:codecommit:*::*:sagemaker-*"
    ]
},
{
    "Effect": "Allow",
    "Action": [ "codecommit:ListRepositories"
    ],
    "Resource": "*
}
"codepipeline:GetPipeline",
"codepipeline:GetPipelineState",
"codepipeline:StartPipelineExecution",
"codepipeline:TagResource",
"codepipeline:UpdatePipeline"
],
"Resource": [
  "arn:aws:codepipeline:*:*:sagemaker-*"
]
},
{
  "Effect": "Allow",
  "Action": [
    "cognito-idp:CreateUserPool",
    "cognito-idp:TagResource"
  ],
  "Resource": "*",
  "Condition": {
    "ForAnyValue:StringLike": {
      "aws:TagKeys": [
        "sagemaker:launch-source"
      ]
    }
  }
},
{
  "Effect": "Allow",
  "Action": [
    "cognito-idp:CreateGroup",
    "cognito-idp:CreateUserPoolDomain",
    "cognito-idp:CreateUserPoolClient",
    "cognito-idp:DeleteGroup",
    "cognito-idp:DeleteUserPool",
    "cognito-idp:DeleteUserPoolClient",
    "cognito-idp:DeleteUserPoolDomain",
    "cognito-idp:DescribeUserPool",
    "cognito-idp:DescribeUserPoolClient",
    "cognito-idp:UpdateUserPool",
    "cognito-idp:UpdateUserPoolClient"
  ],
  "Resource": "*",
  "Condition": {
    "StringLike": {
      "aws:ResourceTag/sagemaker:launch-source": "*"
    }
  }
}
Amazon SageMaker

AWS Managed Policies for SageMaker

5936

"glue:CreateDatabase",
"glue:DeleteDatabase"
],
"Resource": [
"arn:aws:glue:*::*:catalog",
"arn:aws:glue:*::*:database/sagemaker-*",
"arn:aws:glue:*::*:table/sagemaker-*",
"arn:aws:glue:*::*:userDefinedFunction/sagemaker-*"
]
},
{
"Effect": "Allow",
"Action": [
"glue:CreateClassifier",
"glue:DeleteClassifier",
"glue:DeleteCrawler",
"glue:DeleteJob",
"glue:DeleteTrigger",
"glue:DeleteWorkflow",
"glue:StopCrawler"
],
"Resource": [
"*"
]
},
{
"Effect": "Allow",
"Action": [
"glue:CreateWorkflow"
],
"Resource": [
"arn:aws:glue:*::*:workflow/sagemaker-*"
]
},
{
"Effect": "Allow",
"Action": [
"glue:CreateJob"
],
"Resource": [
"arn:aws:glue:*::*:job/sagemaker-*"
]
},
{

AWS Managed Policies for SageMaker
5937
"Effect": "Allow",
"Action": [
  "glue:CreateCrawler",
  "glue:GetCrawler"
],
"Resource": [
  "arn:aws:glue:*:*:crawler/sagemaker-*"
]
},
{
  "Effect": "Allow",
  "Action": [
    "glue:CreateTrigger",
    "glue:GetTrigger"
  ],
  "Resource": [
    "arn:aws:glue:*:*:trigger/sagemaker-*"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "IAM:PassRole"
  ],
  "Resource": [
    "arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalog"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "lambda:AddPermission",
    "lambda:CreateFunction",
    "lambda:DeleteFunction",
    "lambda:GetFunction",
    "lambda:GetFunctionConfiguration",
    "lambda:InvokeFunction",
    "lambda:RemovePermission"
  ],
  "Resource": [
    "arn:aws:lambda:*:*:function:sagemaker-*"
  ]
}
"Effect": "Allow",
"Action": "lambda:TagResource",
"Resource": [
  "arn:aws:lambda::*::*:function:sagemaker-*"
],
"Condition": {
  "ForAllValues:StringLike": {
    "aws:TagKeys": [
      "sagemaker:*"
    ]
  }
}
},
{
  "Effect": "Allow",
  "Action": [
    "logs:CreateLogGroup",
    "logs:CreateLogStream",
    "logs:DeleteLogGroup",
    "logs:DeleteLogStream",
    "logs:DescribeLogGroups",
    "logs:DescribeLogStreams",
    "logs:PutRetentionPolicy"
  ],
  "Resource": [
    "arn:aws:logs::*::*:log-group:/aws/apigateway/AccessLogs/**",
    "arn:aws:logs::*::*:log-group::log-stream:*"
  ]
},
{
  "Effect": "Allow",
  "Action": "s3:GetObject",
  "Resource": "*",
  "Condition": {
    "StringEquals": {
      "s3:ExistingObjectTag/servicecatalog:provisioning": "true"
    }
  }
},
{
  "Effect": "Allow",
  "Action": "s3:GetObject",
  "Resource": [
    "arn:aws:s3:::sagemaker-*"
  ]
}
[  
  
  "Effect": "Allow",
  "Action": [  
    "s3:CreateBucket",
    "s3:DeleteBucket",
    "s3:DeleteBucketPolicy",
    "s3:GetBucketPolicy",
    "s3:PutBucketAcl",
    "s3:PutBucketNotification",
    "s3:PutBucketPolicy",
    "s3:PutBucketPublicAccessBlock",
    "s3:PutBucketLogging",
    "s3:PutEncryptionConfiguration",
    "s3:PutBucketTagging",
    "s3:PutObjectTagging",
    "s3:PutBucketCORS"
  ],
  "Resource": "arn:aws:s3:::sagemaker-*"
],

"Effect": "Allow",

"Action": [  
  "sagemaker:CreateEndpoint",
  "sagemaker:CreateEndpointConfig",
  "sagemaker:CreateModel",
  "sagemaker:CreateWorkteam",
  "sagemaker:DeleteEndpoint",
  "sagemaker:DeleteEndpointConfig",
  "sagemaker:DeleteModel",
  "sagemaker:DeleteWorkteam",
  "sagemaker:DescribeModel",
  "sagemaker:DescribeEndpointConfig",
  "sagemaker:DescribeEndpoint",
  "sagemaker:DescribeWorkteam",
  "sagemaker:CreateCodeRepository",
  "sagemaker:DescribeCodeRepository",
  "sagemaker:UpdateCodeRepository",
  "sagemaker:DeleteCodeRepository"
],

"Resource": [  
  "arn:aws:sagemaker:*:*:*"
]
AWS managed policy:
AmazonSageMakerPartnerServiceCatalogProductsApiGatewayServiceRolePolicy

This policy is used by Amazon API Gateway within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by API Gateway that require a role.

Permissions details

This policy includes the following permissions.

- **lambda** – Invoke a function created by a partner template.
- **sagemaker** – Invoke an endpoint created by a partner template.
"StringEquals": {  
  "aws:ResourceAccount": "${aws:PrincipalAccount}"  
}
},
{
  "Effect": "Allow",
  "Action": "sagemaker:InvokeEndpoint",
  "Resource": "arn:aws:sagemaker:*:*:endpoint/*",
  "Condition": {
    "Null": {
      "aws:ResourceTag/sagemaker:project-name": "false",
      "aws:ResourceTag/sagemaker:partner": "false"
    },
    "StringEquals": {
      "aws:ResourceAccount": "${aws:PrincipalAccount}"
    }
  }
}
]

AWS managed policy:  
AmazonSageMakerPartnerServiceCatalogProductsCloudFormationServiceRolePolicy  

This policy is used by AWS CloudFormation within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by AWS CloudFormation that require a role.

Permissions details

This policy includes the following permissions.

- **iam** – Pass the AmazonSageMakerServiceCatalogProductsLambdaRole and AmazonSageMakerServiceCatalogProductsApiGatewayRole roles.
- **lambda** – Create, update, delete, and invoke AWS Lambda functions; retrieve, publish, and delete versions of a Lambda layer.
- **apigateway** – Create, update, and delete Amazon API Gateway resources.
- **s3** – Retrieve the lambda-auth-code/layer.zip file from an Amazon Simple Storage Service (Amazon S3) bucket.
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": ["iam:PassRole"],
      "Resource": [
        "arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsLambdaRole"
      ],
      "Condition": {
        "StringEquals": {
          "iam:PassedToService": "lambda.amazonaws.com"
        }
      }
    },
    {
      "Effect": "Allow",
      "Action": ["iam:PassRole"],
      "Resource": [
        "arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsApiGatewayRole"
      ],
      "Condition": {
        "StringEquals": {
          "iam:PassedToService": "apigateway.amazonaws.com"
        }
      }
    },
    {
      "Effect": "Allow",
      "Resource": [
        "arn:aws:lambda::*:*:function:sagemaker-*"
      ]
    }
  ]
}
],
      "Condition": {
        "Null": {
          "aws:ResourceTag/sagemaker:project-name": "false",
          "aws:ResourceTag/sagemaker:partner": "false"
        }
      },
    },
    {
      "Effect": "Allow",
      "Action": [
        "lambda:CreateFunction",
        "lambda:TagResource"
      ],
      "Resource": [
        "arn:aws:lambda:*:function:sagemaker-*"
      ],
      "Condition": {
        "Null": {
          "aws:ResourceTag/sagemaker:project-name": "false",
          "aws:ResourceTag/sagemaker:partner": "false"
        },
        "ForAnyValue:StringEquals": {
          "aws:TagKeys": [
            "sagemaker:project-name",
            "sagemaker:partner"
          ]
        }
      }
    },
    {
      "Effect": "Allow",
      "Action": [
        "lambda:PublishLayerVersion",
        "lambda:GetLayerVersion",
        "lambda:DeleteLayerVersion",
        "lambda:GetFunction"
      ],
      "Resource": [
        "arn:aws:lambda:*:layer:sagemaker-*",
        "arn:aws:lambda:*:function:sagemaker-*"
      ]
    }
  ]
}
"Effect": "Allow",
"Action": [
   "apigateway:GET",
   "apigateway:DELETE",
   "apigateway:PATCH",
   "apigateway:POST",
   "apigateway:PUT"
],
"Resource": [
   "arn:aws:apigateway:*::/restapis/*",
   "arn:aws:apigateway:*::/restapis"
],
"Condition": {
   "Null": {
      "aws:ResourceTag/sagemaker:project-name": "false",
      "aws:ResourceTag/sagemaker:partner": "false"
   }
}
],
{
   "Effect": "Allow",
   "Action": [
      "apigateway:POST",
      "apigateway:PUT"
   ],
   "Resource": [
      "arn:aws:apigateway:*::/restapis",
      "arn:aws:apigateway:*::/tags/*"
   ],
   "Condition": {
      "Null": {
         "aws:ResourceTag/sagemaker:project-name": "false",
         "aws:ResourceTag/sagemaker:partner": "false"
      }
   },
   "ForAnyValue: StringEquals": {
      "aws:TagKeys": [
         "sagemaker:project-name",
         "sagemaker:partner"
      ]
   }
}
],
{
   "Effect": "Allow",
   "Action": [
      "apigateway:POST",
      "apigateway:PUT"
   ],
   "Resource": [
      "arn:aws:apigateway:*::/restapis",
      "arn:aws:apigateway:*::/restapis"
   ],
   "Condition": {
      "Null": {
         "aws:ResourceTag/sagemaker:project-name": "false",
         "aws:ResourceTag/sagemaker:partner": "false"
      }
   },
   "ForAnyValue: StringEquals": {
      "aws:TagKeys": [
         "sagemaker:project-name",
         "sagemaker:partner"
      ]
   }
}
AWS managed policy:
AmazonSageMakerPartnerServiceCatalogProductosLambdaServiceRolePolicy

This policy is used by AWS Lambda within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductosLaunchRole passes to the AWS resources created by Lambda that require a role.

Permissions details

This policy includes the following permissions.

- secretsmanager – Retrieve data from partner provided secrets for a partner template.
AWS managed policy: AmazonSageMakerServiceCatalogProductsApiGatewayServiceRolePolicy

This policy is used by Amazon API Gateway within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by API Gateway that require a role.

Permissions details

This policy includes the following permissions.

- logs – Create and read CloudWatch Logs groups, streams, and events; update events; describe various resources.

These permissions are limited to resources whose log group prefix starts with "aws/apigateway/".

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "logs:CreateLogDelivery",
        "logs:CreateLogGroup",
        "logs:CreateLogStream",
        "logs:DeleteLogDelivery",
        "logs:DescribeLogGroups",
        "logs:DescribeLogStreams",
        "logs:DescribeResourcePolicies",
        "logs:DescribeDestinations",
        "logs:DescribeExportTasks",
        "logs:DescribeMetricFilters",
        "logs:DescribeQueries",
        "logs:DescribeQueryDefinitions",
        "logs:DescribeSubscriptionFilters",
        "logs:GetLogDelivery",
```
AWS managed policy:
AmazonSageMakerServiceCatalogProductsCloudformationServiceRolePolicy

This policy is used by AWS CloudFormation within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by AWS CloudFormation that require a role.

Permissions details

This policy includes the following permissions.

- **sagemaker** – Allow access to various SageMaker resources excluding domains, user-profiles, apps, and flow definitions.
- **iam** – Pass the AmazonSageMakerServiceCatalogProductsCodeBuildRole and AmazonSageMakerServiceCatalogProductsExecutionRole roles.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "sagemaker:AddAssociation",
                "sagemaker:AddTags",
                "sagemaker:AssociateTrialComponent",
                "sagemaker:BatchDescribeModelPackage",
                "sagemaker:BatchGetMetrics",
                "sagemaker:BatchGetRecord",
                "sagemaker:BatchPutMetrics",
                "sagemaker:CreateAction",
                "sagemaker:GetInputDataConfig",
                "sagemaker:GetOutputDataConfig",
                "sagemaker:GetPredictions",
                "sagemaker:GetProcessingJobMetrics",
                "sagemaker:GetProcessingOutputConfig",
                "sagemaker:GetProcessingOutputForPrediction",
                "sagemaker:GetProcessingOutputMetrics",
                "sagemaker:GetProcessingOutputRecord",
                "sagemaker:GetProcessingOutputSummary",
                "sagemaker:GetProcessingJobs",
                "sagemaker:GetPredictionsByBatchPredictionJob",
                "sagemaker:GetPredictionsByBatchTransform",
                "sagemaker:GetPredictionsByInvoke",
                "sagemaker:GetPredictionsByRun",
                "sagemaker:GetRecordSetMetrics",
                "sagemaker:GetRecordSets",
                "sagemaker:GetRecordSetsForInvocation",
                "sagemaker:GetReports",
                "sagemaker:GetTransformPredictions",
                "sagemaker:GetTransforms",
                "sagemaker:GetTransformsForMultiModel",
                "sagemaker:GetTransformsForMultiModelOutput",
                "sagemaker:GetTransformsWithDetails",
                "sagemaker:InvokeFunction",
                "sagemaker:ListAssociationNames",
                "sagemaker:ListAutoMLJobs",
                "sagemaker:ListBlueprints",
                "sagemaker:ListClient一身s"
            ],
            "Resource": "arn:aws:logs:*:*:log-group:/aws/apigateway/*"
        }
    ]
}
```
"sagemaker:CreateWorkteam",
"sagemaker:DeleteAction",
"sagemaker:DeleteAlgorithm",
"sagemaker:DeleteApp",
"sagemaker:DeleteAppImageConfig",
"sagemaker:DeleteArtifact",
"sagemaker:DeleteAssociation",
"sagemaker:DeleteCodeRepository",
"sagemaker:DeleteContext",
"sagemaker:DeleteDataQualityJobDefinition",
"sagemaker:DeleteDeviceFleet",
"sagemaker:DeleteDomain",
"sagemaker:DeleteEndpoint",
"sagemaker:DeleteEndpointConfig",
"sagemaker:DeleteExperiment",
"sagemaker:DeleteFeatureGroup",
"sagemaker:DeleteFlowDefinition",
"sagemaker:DeleteHumanLoop",
"sagemaker:DeleteHumanTaskUi",
"sagemaker:DeleteImage",
"sagemaker:DeleteImageVersion",
"sagemaker:DeleteLineageGroupPolicy",
"sagemaker:DeleteModel",
"sagemaker:DeleteModelBiasJobDefinition",
"sagemaker:DeleteModelExplainabilityJobDefinition",
"sagemaker:DeleteModelPackage",
"sagemaker:DeleteModelPackageGroup",
"sagemaker:DeleteModelPackageGroupPolicy",
"sagemaker:DeleteModelQualityJobDefinition",
"sagemaker:DeleteMonitoringSchedule",
"sagemaker:DeleteNotebookInstance",
"sagemaker:DeleteNotebookInstanceLifecycleConfig",
"sagemaker:DeletePipeline",
"sagemaker:DeleteProject",
"sagemaker:DeleteRecord",
"sagemaker:DeleteTags",
"sagemaker:DeleteTrial",
"sagemaker:DeleteTrialComponent",
"sagemaker:DeleteUserProfile",
"sagemaker:DeleteWorkforce",
"sagemaker:DeleteWorkteam",
"sagemaker:DeregisterDevices",
"sagemaker:DescribeAction",
"sagemaker:DescribeAlgorithm"
"sagemaker:DescribeApp",
"sagemaker:DescribeAppImageConfig",
"sagemaker:DescribeArtifact",
"sagemaker:DescribeAutoMLJob",
"sagemaker:DescribeCodeRepository",
"sagemaker:DescribeCompilationJob",
"sagemaker:DescribeContext",
"sagemaker:DescribeDataQualityJobDefinition",
"sagemaker:DescribeDevice",
"sagemaker:DescribeDeviceFleet",
"sagemaker:DescribeDomain",
"sagemaker:DescribeEdgePackagingJob",
"sagemaker:DescribeEndpoint",
"sagemaker:DescribeEndpointConfig",
"sagemaker:DescribeExperiment",
"sagemaker:DescribeFeatureGroup",
"sagemaker:DescribeFlowDefinition",
"sagemaker:DescribeHumanLoop",
"sagemaker:DescribeHumanTaskUi",
"sagemaker:DescribeHyperParameterTuningJob",
"sagemaker:DescribeImage",
"sagemaker:DescribeImageVersion",
"sagemaker:DescribeInferenceRecommendationsJob",
"sagemaker:DescribeLabelingJob",
"sagemaker:DescribeLineageGroup",
"sagemaker:DescribeModel",
"sagemaker:DescribeModelBiasJobDefinition",
"sagemaker:DescribeModelExplainabilityJobDefinition",
"sagemaker:DescribeModelPackage",
"sagemaker:DescribeModelPackageGroup",
"sagemaker:DescribeModelQualityJobDefinition",
"sagemaker:DescribeMonitoringSchedule",
"sagemaker:DescribeNotebookInstance",
"sagemaker:DescribeNotebookInstanceLifecycleConfig",
"sagemaker:DescribePipeline",
"sagemaker:DescribePipelineDefinitionForExecution",
"sagemaker:DescribePipelineExecution",
"sagemaker:DescribeProcessingJob",
"sagemaker:DescribeProject",
"sagemaker:DescribeSubscribedWorkteam",
"sagemaker:DescribeTrainingJob",
"sagemaker:DescribeTransformJob",
"sagemaker:DescribeTrial",
"sagemaker:DescribeTrialComponent"
"sagemaker:DescribeUserProfile",
"sagemaker:DescribeWorkforce",
"sagemaker:DescribeWorkteam",
"sagemaker:DisableSagemakerServicecatalogPortfolio",
"sagemaker:DisassociateTrialComponent",
"sagemaker:EnableSagemakerServicecatalogPortfolio",
"sagemaker:GetDeviceFleetReport",
"sagemaker:GetDeviceRegistration",
"sagemaker:GetLineageGroupPolicy",
"sagemaker:GetModelPackageGroupPolicy",
"sagemaker:GetRecord",
"sagemaker:GetSagemakerServicecatalogPortfolioStatus",
"sagemaker:GetSearchSuggestions",
"sagemaker:InvokeEndpoint",
"sagemaker:InvokeEndpointAsync",
"sagemaker:ListActions",
"sagemaker:ListAlgorithms",
"sagemaker:ListAppImageConfigs",
"sagemaker:ListApps",
"sagemaker:ListArtifacts",
"sagemaker:ListAssociations",
"sagemaker:ListAutoMLJobs",
"sagemaker:ListCandidatesForAutoMLJob",
"sagemaker:ListCodeRepositories",
"sagemaker:ListCompilationJobs",
"sagemaker:ListContexts",
"sagemaker:ListDataQualityJobDefinitions",
"sagemaker:ListDeviceFleets",
"sagemaker:ListDevices",
"sagemaker:ListDomains",
"sagemaker:ListEdgePackagingJobs",
"sagemaker:ListEndpointConfigs",
"sagemaker:ListEndpoints",
"sagemaker:ListExperiments",
"sagemaker:ListFeatureGroups",
"sagemaker:ListFlowDefinitions",
"sagemaker:ListHumanLoops",
"sagemaker:ListHumanTaskUis",
"sagemaker:ListHyperParameterTuningJobs",
"sagemaker:ListImageVersions",
"sagemaker:ListImages",
"sagemaker:ListInferenceRecommendationsJobs",
"sagemaker:ListLabelingJobs",
"sagemaker:ListLabelingJobsForWorkteam",
"sagemaker:ListLineageGroupPolicy",
"sagemaker:ListModelPackageGroupPolicy",
"sagemaker:GetRecord",
"sagemaker:GetSagemakerServicecatalogPortfolioStatus",
"sagemaker:GetSearchSuggestions",
"sagemaker:InvokeEndpoint",
"sagemaker:InvokeEndpointAsync",
"sagemaker:ListActions",
"sagemaker:ListAlgorithms",
"sagemaker:ListAppImageConfigs",
"sagemaker:ListApps",
"sagemaker:ListArtifacts",
"sagemaker:ListAssociations",
"sagemaker:ListAutoMLJobs",
"sagemaker:ListCandidatesForAutoMLJob",
"sagemaker:ListCodeRepositories",
"sagemaker:ListCompilationJobs",
"sagemaker:ListContexts",
"sagemaker:ListDataQualityJobDefinitions",
"sagemaker:ListDeviceFleets",
"sagemaker:ListDevices",
"sagemaker:ListDomains",
"sagemaker:ListEdgePackagingJobs",
"sagemaker:ListEndpointConfigs",
"sagemaker:ListEndpoints",
"sagemaker:ListExperiments",
"sagemaker:ListFeatureGroups",
"sagemaker:ListFlowDefinitions",
"sagemaker:ListHumanLoops",
"sagemaker:ListHumanTaskUis",
"sagemaker:ListHyperParameterTuningJobs",
"sagemaker:ListImageVersions",
"sagemaker:ListImages",
"sagemaker:ListInferenceRecommendationsJobs",
"sagemaker:ListLabelingJobs",
"sagemaker:ListLabelingJobsForWorkteam",
"sagemaker:ListLineageGroupPolicy",
"sagemaker:ListModelPackageGroupPolicy"
"sagemaker:ListLineageGroups",
"sagemaker:ListModelBiasJobDefinitions",
"sagemaker:ListModelExplainabilityJobDefinitions",
"sagemaker:ListModelMetadata",
"sagemaker:ListModelPackageGroups",
"sagemaker:ListModelPackages",
"sagemaker:ListModelQualityJobDefinitions",
"sagemaker:ListModels",
"sagemaker:ListMonitoringExecutions",
"sagemaker:ListMonitoringSchedules",
"sagemaker:ListNotebookInstanceLifecycleConfigs",
"sagemaker:ListNotebookInstances",
"sagemaker:ListPipelineExecutionSteps",
"sagemaker:ListPipelineExecutions",
"sagemaker:ListPipelineParametersForExecution",
"sagemaker:ListPipelines",
"sagemaker:ListProcessingJobs",
"sagemaker:ListProjects",
"sagemaker:ListSubscribedWorkteams",
"sagemaker:ListTags",
"sagemaker:ListTrainingJobs",
"sagemaker:ListTrainingJobsForHyperParameterTuningJob",
"sagemaker:ListTransformJobs",
"sagemaker:ListTrialComponents",
"sagemaker:ListTrials",
"sagemaker:ListUserProfiles",
"sagemaker:ListWorkforces",
"sagemaker:ListWorkteams",
"sagemaker:PutLineageGroupPolicy",
"sagemaker:PutModelPackageGroupPolicy",
"sagemaker:PutRecord",
"sagemaker:QueryLineage",
"sagemaker:RegisterDevices",
"sagemaker:RenderUiTemplate",
"sagemaker:Search",
"sagemaker:SendHeartbeat",
"sagemaker:SendPipelineExecutionStepFailure",
"sagemaker:SendPipelineExecutionStepSuccess",
"sagemaker:StartHumanLoop",
"sagemaker:StartMonitoringSchedule",
"sagemaker:StartNotebookInstance",
"sagemaker:StartPipelineExecution",
"sagemaker:StopAutoMLJob",
"sagemaker:StopCompilationJob",
"sagemaker:StopEdgePackagingJob",
"sagemaker:StopHumanLoop",
"sagemaker:StopHyperParameterTuningJob",
"sagemaker:StopInferenceRecommendationsJob",
"sagemaker:StopLabelingJob",
"sagemaker:StopMonitoringSchedule",
"sagemaker:StopNotebookInstance",
"sagemaker:StopPipelineExecution",
"sagemaker:StopProcessingJob",
"sagemaker:StopTrainingJob",
"sagemaker:StopTransformJob",
"sagemaker:UpdateAction",
"sagemaker:UpdateAppImageConfig",
"sagemaker:UpdateArtifact",
"sagemaker:UpdateCodeRepository",
"sagemaker:UpdateContext",
"sagemaker:UpdateDeviceFleet",
"sagemaker:UpdateDevices",
"sagemaker:UpdateDomain",
"sagemaker:UpdateEndpoint",
"sagemaker:UpdateEndpointWeightsAndCapacities",
"sagemaker:UpdateExperiment",
"sagemaker:UpdateImage",
"sagemaker:UpdateModelPackage",
"sagemaker:UpdateMonitoringSchedule",
"sagemaker:UpdateNotebookInstance",
"sagemaker:UpdateNotebookInstanceLifecycleConfig",
"sagemaker:UpdatePipeline",
"sagemaker:UpdatePipelineExecution",
"sagemaker:UpdateProject",
"sagemaker:UpdateTrainingJob",
"sagemaker:UpdateTrial",
"sagemaker:UpdateTrialComponent",
"sagemaker:UpdateUserProfile",
"sagemaker:UpdateWorkforce",
"sagemaker:UpdateWorkteam"
],
"NotResource": [
    "arn:aws:sagemaker:*:*:domain/*",
    "arn:aws:sagemaker:*:*:user-profile/*",
    "arn:aws:sagemaker:*:*:app/*",
    "arn:aws:sagemaker:*:*:flow-definition/*"
]}
}

AWS managed policy: AmazonSageMakerServiceCatalogProductsCodeBuildServiceRolePolicy

This policy is used by AWS CodeBuild within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by CodeBuild that require a role.

Permissions details

This policy includes the following permissions.

- `sagemaker` – Allow access to various SageMaker resources.
- `codecommit` – Upload CodeCommit archives to CodeBuild pipelines, get upload status, and cancel uploads; get branch and commit information. These permissions are limited to resources whose name starts with "sagemaker-".
- `ecr` – Create Amazon ECR repositories and container images; upload image layers. These permissions are limited to repositories whose name starts with "sagemaker-".
- `ecr` – Read all resources.
- `iam` – Pass the following roles:
  - AmazonSageMakerServiceCatalogProductsCloudformationRole to AWS CloudFormation.
  - AmazonSageMakerServiceCatalogProductsCodeBuildRole to AWS CodeBuild.
  - AmazonSageMakerServiceCatalogProductsCodePipelineRole to AWS CodePipeline.
• AmazonSageMakerServiceCatalogProductsEventsRole to Amazon EventBridge.
• AmazonSageMakerServiceCatalogProductsExecutionRole to Amazon SageMaker.
• logs – Create and read CloudWatch Logs groups, streams, and events; update events; describe various resources.

These permissions are limited to resources whose name prefix starts with "aws/codebuild/".
• s3 – Create, read, and list Amazon S3 buckets. These permissions are limited to buckets whose name starts with "sagemaker-".

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": [
        "codecommit:CancelUploadArchive",
        "codecommit:GetBranch",
        "codecommit:GetCommit",
        "codecommit:GetUploadArchiveStatus",
        "codecommit:UploadArchive"
      ],
      "Resource": "arn:aws:codecommit::*::*:sagemaker-*"
    },
    {
      "Effect": "Allow",
      "Action": [
        "ecr:BatchCheckLayerAvailability",
        "ecr:BatchGetImage",
        "ecr:DescribeImageScanFindings",
        "ecr:DescribeRegistry",
        "ecr:DescribeImageReplicationStatus",
        "ecr:DescribeRepositories",
        "ecr:GetAuthorizationToken",
        "ecr:GetDownloadUrlForLayer"
      ],
      "Resource": [
        "*"
      ]
    }
  ]
}
```
["Effect": "Allow",
"Action": [
"ecr:CompleteLayerUpload",
"ecr:CreateRepository",
"ecr:InitiateLayerUpload",
"ecr:PutImage",
"ecr:UploadLayerPart"
],
"Resource": [
"arn:aws:ecr::*:*:repository/sagemaker-*"
]
],
[
"Effect": "Allow",
"Action": [
"iam:PassRole"
],
"Resource": [
"arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsEventsRole",
"arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsCodePipelineRole",
"arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsCloudformationRole",
"arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsCodeBuildRole",
"arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsExecutionRole"
]
],
"Condition": {"StringEquals": {
"iam:PassedToService": [
"events.amazonaws.com",
"codepipeline.amazonaws.com",
"cloudformation.amazonaws.com",
"codebuild.amazonaws.com",
"sagemaker.amazonaws.com"
]}}]
"Action": [
    "logs:CreateLogDelivery",
    "logs:CreateLogGroup",
    "logs:CreateLogStream",
    "logs:DeleteLogDelivery",
    "logs:DescribeLogGroups",
    "logs:DescribeLogStreams",
    "logs:DescribeResourcePolicies",
    "logs:DescribeDestinations",
    "logs:DescribeExportTasks",
    "logs:DescribeMetricFilters",
    "logs:DescribeQueries",
    "logs:DescribeQueryDefinitions",
    "logs:DescribeSubscriptionFilters",
    "logs:GetLogDelivery",
    "logs:GetLogEvents",
    "logs:ListLogDeliveries",
    "logs:PutLogEvents",
    "logs:PutResourcePolicy",
    "logs:UpdateLogDelivery"
],
"Resource": "arn:aws:logs::*:*:log-group:/aws/codebuild/*"
},
{
    "Effect": "Allow",
    "Action": [
        "s3:CreateBucket",
        "s3:DeleteBucket",
        "s3:GetBucketAcl",
        "s3:GetBucketCors",
        "s3:GetBucketLocation",
        "s3:ListAllMyBuckets",
        "s3:ListBucket",
        "s3:ListBucketMultipartUploads",
        "s3:PutBucketCors",
        "s3:AbortMultipartUpload",
        "s3:DeleteObject",
        "s3:GetObject",
        "s3:GetObjectVersion",
        "s3:PutObject"
    ],
    "Resource": [
        "arn:aws:s3:::aws-glue-*",
        "arn:aws:s3:::sagemaker-*"
    ]}
"Effect": "Allow",
"Action": [
"sagemaker:AddAssociation",
"sagemaker:AddTags",
"sagemaker:AssociateTrialComponent",
"sagemaker:BatchDescribeModelPackage",
"sagemaker:BatchGetMetrics",
"sagemaker:BatchGetRecord",
"sagemaker:BatchPutMetrics",
"sagemaker:CreateAction",
"sagemaker:CreateAlgorithm",
"sagemaker:CreateApp",
"sagemaker:CreateAppImageConfig",
"sagemaker:CreateArtifact",
"sagemaker:CreateAutoMLJob",
"sagemaker:CreateCodeRepository",
"sagemaker:CreateCompilationJob",
"sagemaker:CreateContext",
"sagemaker:CreateDataQualityJobDefinition",
"sagemaker:CreateDeviceFleet",
"sagemaker:CreateDomain",
"sagemaker:CreateEdgePackagingJob",
"sagemaker:CreateEndpoint",
"sagemaker:CreateEndpointConfig",
"sagemaker:CreateExperiment",
"sagemaker:CreateFeatureGroup",
"sagemaker:CreateFlowDefinition",
"sagemaker:CreateHumanTaskUi",
"sagemaker:CreateHyperParameterTuningJob",
"sagemaker:CreateImage",
"sagemaker:CreateImageVersion",
"sagemaker:CreateInferenceRecommendationsJob",
"sagemaker:CreateLabelingJob",
"sagemaker:CreateLineageGroupPolicy",
"sagemaker:CreateModel",
"sagemaker:CreateModelBiasJobDefinition",
"sagemaker:CreateModelExplainabilityJobDefinition",
"sagemaker:CreateModelPackage",
"sagemaker:CreateModelPackageGroup",
"sagemaker:CreateModelQualityJobDefinition",
"sagemaker:CreateMonitoringSchedule"],
"sagemaker:CreateNotebookInstance",
"sagemaker:CreateNotebookInstanceLifecycleConfig",
"sagemaker:CreatePipeline",
"sagemaker:CreatePresignedDomainUrl",
"sagemaker:CreatePresignedNotebookInstanceUrl",
"sagemaker:CreateProcessingJob",
"sagemaker:CreateProject",
"sagemaker:CreateTrainingJob",
"sagemaker:CreateTransformJob",
"sagemaker:CreateTrial",
"sagemaker:CreateTrialComponent",
"sagemaker:CreateUserProfile",
"sagemaker:CreateWorkforce",
"sagemaker:CreateWorkteam",
"sagemaker:DeleteAction",
"sagemaker:DeleteAlgorithm",
"sagemaker:DeleteApp",
"sagemaker:DeleteAppImageConfig",
"sagemaker:DeleteArtifact",
"sagemaker:DeleteAssociation",
"sagemaker:DeleteCodeRepository",
"sagemaker:DeleteContext",
"sagemaker:DeleteDataQualityJobDefinition",
"sagemaker:DeleteDeviceFleet",
"sagemaker:DeleteDomain",
"sagemaker:DeleteEndpoint",
"sagemaker:DeleteEndpointConfig",
"sagemaker:DeleteExperiment",
"sagemaker:DeleteFeatureGroup",
"sagemaker:DeleteFlowDefinition",
"sagemaker:DeleteHumanLoop",
"sagemaker:DeleteHumanTaskUi",
"sagemaker:DeleteImage",
"sagemaker:DeleteImageVersion",
"sagemaker:DeleteLineageGroupPolicy",
"sagemaker:DeleteModel",
"sagemaker:DeleteModelBiasJobDefinition",
"sagemaker:DeleteModelExplainabilityJobDefinition",
"sagemaker:DeleteModelPackage",
"sagemaker:DeleteModelPackageGroup",
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"sagemaker:StopTransformJob",
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"sagemaker:UpdateImage",
"sagemaker:UpdateModelPackage",
"sagemaker:UpdateMonitoringSchedule",
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"sagemaker:UpdateNotebookInstanceLifecycleConfig",
"sagemaker:UpdatePipeline",
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"sagemaker:UpdateTrainingJob",
AWS managed policy:

AmazonSageMakerServiceCatalogProductsCodePipelineServiceRolePolicy

This policy is used by AWS CodePipeline within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by CodePipeline that require a role.

Permissions details

This policy includes the following permissions.

- **cloudformation** – Create, read, delete, and update CloudFormation stacks; create, read, delete, and execute change sets; set stack policy. These permissions are limited to resources whose name starts with "sagemaker-".
- **s3** – Create, read, list, and delete Amazon S3 buckets; add, read, and delete objects from the buckets; read and set the CORS configuration; read the access control list (ACL); and read the AWS Region the bucket resides in.

  These permissions are limited to buckets whose name starts with "sagemaker-" or "aws-glue-.
- **iam** – Pass the AmazonSageMakerServiceCatalogProductsCloudformationRole role.
- **codebuild** – Get CodeBuild build information and start builds. These permissions are limited to project and build resources whose name starts with "sagemaker-".
• codecommit – Upload CodeCommit archives to CodeBuild pipelines, get upload status, and cancel uploads; get branch and commit information.

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "cloudformation:CreateChangeSet",
                "cloudformation:CreateStack",
                "cloudformation:DescribeChangeSet",
                "cloudformation:DeleteChangeSet",
                "cloudformation:DeleteStack",
                "cloudformation:DescribeStacks",
                "cloudformation:ExecuteChangeSet",
                "cloudformation:SetStackPolicy",
                "cloudformation:UpdateStack"
            ],
            "Resource": "arn:aws:cloudformation::*::*:stack/sagemaker-*"
        },
        {
            "Effect": "Allow",
            "Action": [
                "s3:AbortMultipartUpload",
                "s3:DeleteObject",
                "s3:GetObject",
                "s3:GetObjectVersion",
                "s3:PutObject"
            ],
            "Resource": ['arn:aws:s3:::sagemaker-*']
        },
        {
            "Effect": "Allow",
            "Action": [
                "iam:PassRole"
            ],
            "Resource": ['arn:aws:iam::*:role/service-role/AmazonSageMakerServiceCatalogProductsCloudformationRole']
        }
    ]
}
```
AWS managed policy: AmazonSageMakerServiceCatalogProductsEventsServiceRolePolicy

This policy is used by Amazon EventBridge within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by EventBridge that require a role.

Permissions details

This policy includes the following permissions.

• codepipeline – Start a CodeBuild execution. These permissions are limited to pipelines whose name starts with "sagemaker-".
AWS managed policy: AmazonSageMakerServiceCatalogProductsFirehoseServiceRolePolicy

This policy is used by Amazon Data Firehose within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by Firehose that require a role.

Permissions details

This policy includes the following permissions.

- firehose – Send Firehose records. These permissions are limited to resources whose delivery stream name starts with "sagemaker-".

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "VisualEditor0",
         "Effect": "Allow",
         "Action": ["firehose:PutRecord", "firehose:PutRecordBatch"],
         "Resource": "arn:aws:firehose:*:*:deliverystream/sagemaker-*"
      }
   ]
}
```
AWS managed policy: AmazonSageMakerServiceCatalogProductsGlueServiceRolePolicy

This policy is used by AWS Glue within the AWS Service Catalog provisioned products from the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by Glue that require a role.

Permissions details

This policy includes the following permissions.

- **glue** – Create, read, and delete AWS Glue partitions, tables, and table versions. These permissions are limited to those resources whose name starts with "sagemaker-". Create and read AWS Glue databases. These permissions are limited to databases whose name is "default", "global_temp", or starts with "sagemaker-". Get user defined functions.
- **s3** – Create, read, list, and delete Amazon S3 buckets; add, read, and delete objects from the buckets; read and set the CORS configuration; read the access control list (ACL), and read the AWS Region the bucket resides in.

  These permissions are limited to buckets whose name starts with "sagemaker-" or "aws-glue-".
- **logs** – Create, read, and delete CloudWatch Logs log group, streams, and deliveries; and create a resource policy.

  These permissions are limited to resources whose name prefix starts with "aws/glue/".

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "glue:BatchCreatePartition",
                "glue:BatchDeletePartition",
                "glue:BatchDeleteTable",
                "glue:BatchDeleteTableVersion",
                "glue:BatchGetPartition",
                "glue:CreateDatabase",
                "glue:CreatePartition",
                "glue:CreateTable",
                "glue:DeletePartition",
                "s3:GetObject",
                "s3:ListBucket",
                "s3:PutObject",
                "logs:GetLogEvents",
                "logs:PutLogEvents",
                "logs:CreateLogGroup"
            ],
```

AWS Managed Policies for SageMaker

5970
"glue:DeleteTable",
"glue:DeleteTableVersion",
"glue:GetDatabase",
"glue:GetPartition",
"glue:GetPartitions",
"glue:GetTable",
"glue:GetTables",
"glue:GetTableVersion",
"glue:GetTableVersions",
"glue:SearchTables",
"glue:UpdatePartition",
"glue:UpdateTable",
"glue:GetUserDefinedFunctions"
],
"Resource": [
  "arn:aws:glue:*::*:catalog",
  "arn:aws:glue:*::*:database/default",
  "arn:aws:glue:*::*:database/global_temp",
  "arn:aws:glue:*::*:database/sagemaker-*",
  "arn:aws:glue:*::*:table/sagemaker-*",
  "arn:aws:glue:*::*:tableVersion/sagemaker-*"
]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:CreateBucket",
    "s3:DeleteBucket",
    "s3:GetBucketAcl",
    "s3:GetBucketCors",
    "s3:GetBucketLocation",
    "s3:ListAllMyBuckets",
    "s3:ListBucket",
    "s3:ListBucketMultipartUploads",
    "s3:PutBucketCors"
  ],
  "Resource": [
    "arn:aws:s3:::aws-glue-*",
    "arn:aws:s3:::sagemaker-*"
  ]
},
{
  "Effect": "Allow",
  "Action": [}
AWS managed policy: AmazonSageMakerServiceCatalogProductsLambdaServiceRolePolicy

This policy is used by AWS Lambda within the AWS Service Catalog provisioned products from
the Amazon SageMaker portfolio. The policy is intended to be attached to an IAM role that the
AmazonSageMakerServiceCatalogProductsLaunchRole passes to the AWS resources created by
Lambda that require a role.

Permissions details

This policy includes the following permissions.

- sagemaker – Allow access to various SageMaker resources.
- **ecr** – Create and delete Amazon ECR repositories; create, read, and delete container images; upload image layers. These permissions are limited to repositories whose name starts with "sagemaker-".

- **events** – Create, read, and delete Amazon EventBridge rules; and create and remove targets. These permissions are limited to rules whose name starts with "sagemaker-".

- **s3** – Create, read, list, and delete Amazon S3 buckets; add, read, and delete objects from the buckets; read and set the CORS configuration; read the access control list (ACL), and read the AWS Region the bucket resides in.

  These permissions are limited to buckets whose name starts with "sagemaker-" or "aws-glue-".

- **iam** – Pass the AmazonSageMakerServiceCatalogProductsExecutionRole role.

- **logs** – Create, read, and delete CloudWatch Logs log group, streams, and deliveries; and create a resource policy.

  These permissions are limited to resources whose name prefix starts with "aws/lambda/".

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "ecr:DescribeImages",
                "ecr:BatchDeleteImage",
                "ecr:CompleteLayerUpload",
                "ecr:CreateRepository",
                "ecr:DeleteRepository",
                "ecr:InitiateLayerUpload",
                "ecr:PutImage",
                "ecr:UploadLayerPart"
            ],
            "Resource": [
                "arn:aws:ecr:*:*:repository/sagemaker-*"
            ]
        },
        {
            "Effect": "Allow",
            "Action": [
                "events:DeleteRule",
            ]
        }
    ]
}
```
"events:DescribeRule",
"events:PutRule",
"events:PutTargets",
"events:RemoveTargets"
],
"Resource": [
  "arn:aws:events::*:rule/sagemaker-*"
]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:CreateBucket",
    "s3:DeleteBucket",
    "s3:GetBucketAcl",
    "s3:GetBucketCors",
    "s3:GetBucketLocation",
    "s3:ListAllMyBuckets",
    "s3:ListBucket",
    "s3:ListBucketMultipartUploads",
    "s3:PutBucketCors"
  ],
  "Resource": [
    "arn:aws:s3:::aws-glue-*",
    "arn:aws:s3:::sagemaker-*"
  ]
},
{
  "Effect": "Allow",
  "Action": [
    "s3:AbortMultipartUpload",
    "s3:DeleteObject",
    "s3:GetObject",
    "s3:GetObjectVersion",
    "s3:PutObject"
  ],
  "Resource": [
    "arn:aws:s3:::aws-glue-*",
    "arn:aws:s3:::sagemaker-*"
  ]
},
{
  "Effect": "Allow",
  "Action": [
"sagemaker:AddAssociation",
"sagemaker:AddTags",
"sagemaker:AssociateTrialComponent",
"sagemaker:BatchDescribeModelPackage",
"sagemaker:BatchGetMetrics",
"sagemaker:BatchGetRecord",
"sagemaker:BatchPutMetrics",
"sagemaker:CreateAction",
"sagemaker:CreateAlgorithm",
"sagemaker:CreateApp",
"sagemaker:CreateAppImageConfig",
"sagemaker:CreateArtifact",
"sagemaker:CreateAutoMLJob",
"sagemaker:CreateCodeRepository",
"sagemaker:CreateCompilationJob",
"sagemaker:CreateContext",
"sagemaker:CreateDataQualityJobDefinition",
"sagemaker:CreateDeviceFleet",
"sagemaker:CreateDomain",
"sagemaker:CreateEdgePackagingJob",
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"sagemaker:CreateEndpointConfig",
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"sagemaker:CreateFeatureGroup",
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"sagemaker:CreateTrainingJob",
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"sagemaker:SendPipelineExecutionStepSuccess",
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"sagemaker:StartNotebookInstance",
"sagemaker:StartPipelineExecution",
"sagemaker:StopAutoMLJob",
"sagemaker:StopCompilationJob",
"sagemaker:StopEdgePackagingJob",
"sagemaker:StopHumanLoop",
"sagemaker:StopHyperParameterTuningJob",
"sagemaker:StopInferenceRecommendationsJob",
"sagemaker:StopLabelingJob",
"sagemaker:StopMonitoringSchedule",
"sagemaker:StopNotebookInstance",
"sagemaker:StopPipelineExecution",
"sagemaker:StopProcessingJob",
"sagemaker:StopTrainingJob",
"sagemaker:StopTransformJob",
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"sagemaker:UpdateAppImageConfig",
"sagemaker:UpdateArtifact",
"sagemaker:UpdateCodeRepository",
"sagemaker:UpdateContext",
"sagemaker:UpdateDeviceFleet",
"sagemaker:UpdateDevices",
"sagemaker:UpdateDomain",
"sagemaker:UpdateEndpoint",
"sagemaker:UpdateEndpointWeightsAndCapacities",
"sagemaker:UpdateExperiment",
"sagemaker:UpdateImage",
"sagemaker:UpdateModelPackage",
"sagemaker:UpdateMonitoringSchedule",
"sagemaker:UpdateNotebookInstance",
"sagemaker:UpdateNotebookInstanceLifecycleConfig",
"sagemaker:UpdatePipeline",
"sagemaker:UpdatePipelineExecution",
"sagemaker:UpdateProject",
"sagemaker:UpdateTrainingJob",
"sagemaker:UpdateTrial",
"sagemaker:UpdateTrialComponent",
"sagemaker:UpdateUserProfile",
"sagemaker:UpdateWorkforce",
"sagemaker:UpdateWorkteam"
],
"Resource": [
  "arn:aws:sagemaker:*:::action/**",
  "arn:aws:sagemaker:*:::algorithm/**",
  "arn:aws:sagemaker:*:::app-image-config/**",
  "arn:aws:sagemaker:*:::artifact/**",
  "arn:aws:sagemaker:*:::automl-job/**",
  "arn:aws:sagemaker:*:::code-repository/**",
  "arn:aws:sagemaker:*:::compilation-job/**",
  "arn:aws:sagemaker:*:::context/**",
  "arn:aws:sagemaker:*:::data-quality-job-definition/**",
  "arn:aws:sagemaker:*:::device-fleet/**/device/**",
  "arn:aws:sagemaker:*:::device-fleet/**",
  "arn:aws:sagemaker:*:::edge-packaging-job/**",
  "arn:aws:sagemaker:*:::endpoint/**",
  "arn:aws:sagemaker:*:::endpoint-config/**",
  "arn:aws:sagemaker:*:::experiment/**",
  "arn:aws:sagemaker:*:::experiment-trial/**",
  "arn:aws:sagemaker:*:::experiment-trial-component/**",
  "arn:aws:sagemaker:*:::feature-group/**",
  "arn:aws:sagemaker:*:::human-loop/**",
  "arn:aws:sagemaker:*:::human-task-ui/**",
  "arn:aws:sagemaker:*:::hyper-parameter-tuning-job/**",
  "arn:aws:sagemaker:*:::image/**",
  "arn:aws:sagemaker:*:::image-version/**/*",
  "arn:aws:sagemaker:*:::inference-recommendations-job/**",
  "arn:aws:sagemaker:*:::labeling-job/**",
  "arn:aws:sagemaker:*:::model/**",
  "arn:aws:sagemaker:*:::model-bias-job-definition/**",
  "arn:aws:sagemaker:*:::model-explainability-job-definition/**",
  "arn:aws:sagemaker:*:::model-package/**",
  "arn:aws:sagemaker:*:::model-package-group/**",
  "arn:aws:sagemaker:*:::model-quality-job-definition/**",
  "arn:aws:sagemaker:*:::monitoring-schedule/**",
  "arn:aws:sagemaker:*:::notebook-instance/**",
  "arn:aws:sagemaker:*:::notebook-instance-lifecycle-config/**",
  "arn:aws:sagemaker:*:::pipeline/**",
  "arn:aws:sagemaker:*:::pipeline/**/execution/**",
  "arn:aws:sagemaker:*:::processing-job/**",
  "arn:aws:sagemaker:*:::project/**",
  "arn:aws:sagemaker:*:::training-job/**",
  "arn:aws:sagemaker:*:::transform-job/**",
  "arn:aws:sagemaker:*:::workforce/**",
  "arn:aws:sagemaker:*:::workteam/**"


```json
{
    "Effect": "Allow",
    "Action": [
        "iam:PassRole"
    ],
    "Resource": [
        "arn:aws:iam::*:service-role/AmazonSageMakerServiceCatalogProductsExecutionRole"
    ]
}

{
    "Effect": "Allow",
    "Action": [
        "logs:CreateLogDelivery",
        "logs:CreateLogGroup",
        "logs:CreateLogStream",
        "logs:DeleteLogDelivery",
        "logs:DescribeLogGroups",
        "logs:DescribeLogStreams",
        "logs:DescribeResourcePolicies",
        "logs:DescribeDestinations",
        "logs:DescribeExportTasks",
        "logs:DescribeMetricFilters",
        "logs:DescribeQueries",
        "logs:DescribeQueryDefinitions",
        "logs:DescribeSubscriptionFilters",
        "logs:GetLogDelivery",
        "logs:GetLogEvents",
        "logs:ListLogDeliveries",
        "logs:PutLogEvents",
        "logs:PutResourcePolicy",
        "logs:UpdateLogDelivery"
    ],
    "Resource": "arn:aws:logs::*:*:log-group:/aws/lambda/*"
}
```
Amazon SageMaker updates to AWS Service Catalog AWS managed policies

View details about updates to AWS managed policies for Amazon SageMaker since this service began tracking these changes.

<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmazonSageMakerPartnerServiceCatalog</td>
<td>1</td>
<td>Initial policy</td>
<td>August 1, 2023</td>
</tr>
<tr>
<td>ProductsApiGatewayServiceRolePolicy</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>AmazonSageMakerPartnerServiceCatalog</td>
<td>1</td>
<td>Initial policy</td>
<td>August 1, 2023</td>
</tr>
<tr>
<td>ProductsCloudFormationServiceRolePolicy</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>AmazonSageMakerPartnerServiceCatalog</td>
<td>1</td>
<td>Initial policy</td>
<td>August 1, 2023</td>
</tr>
<tr>
<td>ProductsLambdaServiceRolePolicy</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>AmazonSageMakerServiceCatalogProductsGlueServiceRolePolicy</td>
<td>2</td>
<td>Add permission for glue:GetUserDefinedFunctions</td>
<td>August 26, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy</td>
<td>7</td>
<td>Add permission for sagemaker:AddTags</td>
<td>August 2, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy</td>
<td>6</td>
<td>Add permission for lambda:TagResource</td>
<td>July 14, 2022</td>
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<tr>
<td>AmazonSageMakerServiceCatalogProductsServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>April 4, 2022</td>
</tr>
<tr>
<td>Policy</td>
<td>Version</td>
<td>Change</td>
<td>Date</td>
</tr>
<tr>
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<tr>
<td>AmazonSageMakerServiceCatalogProductsApiGatewayServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>March 24, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerServiceCatalogProductsCloudformationServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>March 24, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerServiceCatalogProductsCodeBuildServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>March 24, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy</td>
<td>5</td>
<td>Add new permission for ecr-idp:TagResource</td>
<td>March 21, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerServiceCatalogProductsCodePipelineServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>February 22, 2022</td>
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<tr>
<td>AmazonSageMakerServiceCatalogProductsEventsServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>February 22, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerServiceCatalogProductsFirehoseServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>February 22, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerServiceCatalogProductsGlueServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>February 22, 2022</td>
</tr>
<tr>
<td>Policy</td>
<td>Version</td>
<td>Change</td>
<td>Date</td>
</tr>
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<td>-------------------------</td>
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<tr>
<td>AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy</td>
<td>4</td>
<td>Add permissions for cognito-idp:TagResource and s3:PutBucketCORS.</td>
<td>February 16, 2022</td>
</tr>
<tr>
<td>AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy</td>
<td>3</td>
<td>Add new permissions for sagemaker. Create, read, update, and delete SageMaker Images.</td>
<td>September 15, 2021</td>
</tr>
<tr>
<td>AmazonSageMakerAdmin-ServiceCatalogProductsServiceRolePolicy</td>
<td>1</td>
<td>Initial policy</td>
<td>November 27, 2020</td>
</tr>
</tbody>
</table>

**SageMaker Updates to AWS Managed Policies**

View details about updates to AWS managed policies for SageMaker since this service began tracking these changes.
<table>
<thead>
<tr>
<th>Policy</th>
<th>Version</th>
<th>Change</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AmazonSageMakerFullAccess</strong> - Update to an existing policy</td>
<td>25</td>
<td>Add sagemaker:CreateApp, sagemaker:DescribeApp, sagemaker:DeleteApp, sagemaker:CreateSpace, sagemaker:UpdateSpace, sagemaker:DeleteSpace, s3express:CreateSession, s3express:CreateBucket, and s3express:ListAllMyDirectoryBuckets permissions.</td>
<td>November 30, 2023</td>
</tr>
<tr>
<td><strong>AmazonSageMakerFullAccess</strong> - Update to an existing policy</td>
<td>24</td>
<td>Add sagemaker-geospatial:*, sagemaker:AddTags, sagemaker-ListTags, sagemaker-DescribeSpace, and sagemaker:ListSpaces permissions.</td>
<td>November 30, 2022</td>
</tr>
<tr>
<td><strong>AmazonSageMakerFullAccess</strong> - Update to an existing policy</td>
<td>23</td>
<td>Add glue:UpdateTable.</td>
<td>June 29, 2022</td>
</tr>
<tr>
<td><strong>AmazonSageMakerFullAccess</strong> - Update to an existing policy</td>
<td>22</td>
<td>Add cloudformation:ListStackResources.</td>
<td>May 1, 2022</td>
</tr>
<tr>
<td>Policy</td>
<td>Version</td>
<td>Change</td>
<td>Date</td>
</tr>
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<td>--------</td>
<td>---------</td>
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<td>--------------------</td>
</tr>
<tr>
<td>AmazonSageMakerReadOnly - Update to an existing policy</td>
<td>11</td>
<td>Add <code>sagemaker:QueryLineage</code>, <code>sagemaker:GetLineageGroupPolicy</code>, <code>sagemaker:BatchDescribeMode</code>, <code>sagemaker:BatchDescribePackaging</code>, <code>sagemaker:GetModelPackageGroupPolicy</code> permissions.</td>
<td>December 1, 2021</td>
</tr>
<tr>
<td>AmazonSageMakerFullAccess - Update to an existing policy</td>
<td>21</td>
<td>Add <code>sns:Publish</code> permissions for endpoints with Async Inference enabled.</td>
<td>September 8, 2021</td>
</tr>
<tr>
<td>AmazonSageMakerFullAccess - Update to an existing policy</td>
<td>20</td>
<td>Update <code>iam:PassRole</code> resources and permissions.</td>
<td>July 15, 2021</td>
</tr>
<tr>
<td>AmazonSageMakerReadOnly - Update to an existing policy</td>
<td>10</td>
<td>New API <code>BatchGetRecord</code> added for SageMaker Feature Store.</td>
<td>June 10, 2021</td>
</tr>
</tbody>
</table>

**Troubleshooting Amazon SageMaker Identity and Access**

Use the following information to help you diagnose and fix common issues that you might encounter when working with SageMaker and IAM.
Topics

- I Am Not Authorized to Perform an Action in SageMaker
- I Am Not Authorized to Perform iam:PassRole
- I Want to Allow People Outside of My AWS Account to Access My SageMaker Resources

I Am Not Authorized to Perform an Action in SageMaker

If the AWS Management Console tells you that you're not authorized to perform an action, then you must contact your administrator for assistance. Your administrator is the person that provided you with your sign-in credentials.

The following example error occurs when the mateojackson IAM user tries to use the console to view details about a training job but does not have sagemaker:sagemaker:DescribeTrainingJob permissions.

User: arn:aws:iam::123456789012:user/mateojackson is not authorized to perform: sagemaker:DescribeTrainingJob on resource: my-example-widget

In this case, Mateo asks his administrator to update his policies to allow him to access the TrainingJob resource using the sagemaker:DescribeTrainingJob action.

I Am Not Authorized to Perform iam:PassRole

If you receive an error that you're not authorized to perform the iam:PassRole action, your policies must be updated to allow you to pass a role to SageMaker.

Some AWS services allow you to pass an existing role to that service instead of creating a new service role or service-linked role. To do this, you must have permissions to pass the role to the service.

The following example error occurs when an IAM user named marymajor tries to use the console to perform an action in SageMaker. However, the action requires the service to have permissions that are granted by a service role. Mary does not have permissions to pass the role to the service.

User: arn:aws:iam::123456789012:user/marymajor is not authorized to perform: iam:PassRole
In this case, Mary's policies must be updated to allow her to perform the `iam:PassRole` action.

If you need help, contact your AWS administrator. Your administrator is the person who provided you with your sign-in credentials.

**I Want to Allow People Outside of My AWS Account to Access My SageMaker Resources**

You can create a role that users in other accounts or people outside of your organization can use to access your resources. You can specify who is trusted to assume the role. For services that support resource-based policies or access control lists (ACLs), you can use those policies to grant people access to your resources.

To learn more, consult the following:

- To learn whether SageMaker supports these features, see [How Amazon SageMaker Works with IAM](#).
- To learn how to provide access to your resources across AWS accounts that you own, see [Providing access to an IAM user in another AWS account that you own](#) in the [IAM User Guide](#).
- To learn how to provide access to your resources to third-party AWS accounts, see [Providing access to AWS accounts owned by third parties](#) in the [IAM User Guide](#).
- To learn how to provide access through identity federation, see [Providing access to externally authenticated users (identity federation)](#) in the [IAM User Guide](#).
- To learn the difference between using roles and resource-based policies for cross-account access, see [How IAM roles differ from resource-based policies](#) in the [IAM User Guide](#).

**Logging and Monitoring**

You can monitor Amazon SageMaker using Amazon CloudWatch, which collects raw data and processes it into readable, near real-time metrics. These statistics are kept for 15 months, so that you can access historical information and gain a better perspective on how your web application or service is performing. You can also set alarms that watch for certain thresholds and send notifications or take actions when those thresholds are met. For more information, see [Monitor Amazon SageMaker with Amazon CloudWatch](#).

Amazon CloudWatch Logs enables you to monitor, store, and access your log files from Amazon EC2 instances, AWS CloudTrail, and other sources. You can collect and track metrics, create
customized dashboards, and set alarms that notify you or take actions when a specified metric reaches a threshold that you specify. CloudWatch Logs can monitor information in the log files and notify you when certain thresholds are met. You can also archive your log data in highly durable storage. For more information, see Log Amazon SageMaker Events with Amazon CloudWatch.

AWS CloudTrail provides a record of actions taken by a user, role, or an AWS service in SageMaker. Using the information collected by CloudTrail, you can determine the request that was made to SageMaker, the IP address from which the request was made, who made the request, when it was made, and additional details. For more information, Log Amazon SageMaker API Calls with AWS CloudTrail.

**Note**
CloudTrail does not monitor calls to `runtime_InvokeEndpoint`.

You can create rules in Amazon CloudWatch Events to react to status changes in status in a SageMaker training, hyperparameter tuning, or batch transform job. For more information, see Automating Amazon SageMaker with Amazon EventBridge.

**Compliance validation for Amazon SageMaker**

To learn whether an AWS service is within the scope of specific compliance programs, see AWS services in Scope by Compliance Program and choose the compliance program that you are interested in. For general information, see AWS Compliance Programs.

You can download third-party audit reports using AWS Artifact. For more information, see Downloading Reports in AWS Artifact.

Your compliance responsibility when using AWS services is determined by the sensitivity of your data, your company's compliance objectives, and applicable laws and regulations. AWS provides the following resources to help with compliance:

- **Security and Compliance Quick Start Guides** – These deployment guides discuss architectural considerations and provide steps for deploying baseline environments on AWS that are security and compliance focused.
- **Architecting for HIPAA Security and Compliance on Amazon Web Services** – This whitepaper describes how companies can use AWS to create HIPAA-eligible applications.
• **AWS Compliance Resources** – This collection of workbooks and guides might apply to your industry and location.

• **AWS Customer Compliance Guides** – Understand the shared responsibility model through the lens of compliance. The guides summarize the best practices for securing AWS services and map the guidance to security controls across multiple frameworks (including National Institute of Standards and Technology (NIST), Payment Card Industry Security Standards Council (PCI), and International Organization for Standardization (ISO)).

• **Evaluating Resources with Rules** in the *AWS Config Developer Guide* – The AWS Config service assesses how well your resource configurations comply with internal practices, industry guidelines, and regulations.

• **AWS Security Hub** – This AWS service provides a comprehensive view of your security state within AWS. Security Hub uses security controls to evaluate your AWS resources and to check your compliance against security industry standards and best practices. For a list of supported services and controls, see *Security Hub controls reference*.

• **AWS Audit Manager** – This AWS service helps you continuously audit your AWS usage to simplify how you manage risk and compliance with regulations and industry standards.

### Resilience in Amazon SageMaker

The AWS global infrastructure is built around AWS Regions and Availability Zones. AWS Regions provide multiple physically separated and isolated Availability Zones, which are connected with low-latency, high-throughput, and highly redundant networking. With Availability Zones, you can design and operate applications and databases that automatically fail over between Availability Zones without interruption. Availability Zones are more highly available, fault tolerant, and scalable than traditional single or multiple data center infrastructures.

For more information about AWS Regions and Availability Zones, see *AWS Global Infrastructure*.

In addition to the AWS global infrastructure, Amazon SageMaker offers several features to help support your data resiliency and backup needs.
Infrastructure Security in Amazon SageMaker

As a managed service, Amazon SageMaker is protected by AWS global network security. For information about AWS security services and how AWS protects infrastructure, see AWS Cloud Security. To design your AWS environment using the best practices for infrastructure security, see Infrastructure Protection in Security Pillar AWS Well-Architected Framework.

You use AWS published API calls to access Amazon SageMaker through the network. Clients must support the following:

- Transport Layer Security (TLS). We require TLS 1.2 and recommend TLS 1.3.
- Cipher suites with perfect forward secrecy (PFS) such as DHE (Ephemeral Diffie-Hellman) or ECDHE (Elliptic Curve Ephemeral Diffie-Hellman). Most modern systems such as Java 7 and later support these modes.

Additionally, requests must be signed by using an access key ID and a secret access key that is associated with an IAM principal. Or you can use the AWS Security Token Service (AWS STS) to generate temporary security credentials to sign requests.

Topics

- SageMaker Scans AWS Marketplace Training and Inference Containers for Security Vulnerabilities
- Connect to Resources From Within a VPC
- Run Training and Inference Containers in Internet-Free Mode
- Connect to SageMaker Within your VPC
- Give SageMaker Access to Resources in your Amazon VPC

SageMaker Scans AWS Marketplace Training and Inference Containers for Security Vulnerabilities

To meet our security requirements, all the pre-built SageMaker images, including AWS Deep Learning Containers, the SageMaker machine learning framework containers, and the SageMaker built-in algorithm containers, and algorithms and model packages listed in AWS Marketplace are scanned for Common Vulnerabilities and Exposures (CVE). CVE is a list of publicly known information about security vulnerability and exposure. The National Vulnerability Database (NVD) provides CVE details such as severity, impact rating, and fix information. Both CVE and NVD
are available for public consumption and free for security tools and services to use. For more information, see [CVE Frequently Asked Questions (FAQs)](#).

## Connect to Resources From Within a VPC

⚠️ **Important**

The following information applies to both Amazon SageMaker Studio and Amazon SageMaker Studio Classic. The same concepts of connecting to resources within a VPC apply to both Studio and Studio Classic.

Amazon SageMaker Studio and SageMaker notebook instances allow direct internet access by default. This allows you to download popular packages and notebooks, customize your development environment, and work efficiently. However, this could provide an additional avenue for unauthorized access to your data. For example, if you install malicious code on your computer in the form of a publicly available notebook or a publicly available source code library, it could access your data. You can choose to restrict which traffic can access the internet by launching your Amazon SageMaker Studio and SageMaker notebook instances in a [Amazon Virtual Private Cloud (Amazon VPC)](#) of your choosing.

An Amazon Virtual Private Cloud is a virtual network dedicated to your AWS account. With an Amazon VPC, you can control the network access and internet connectivity of your Amazon SageMaker Studio and notebook instances. You can disable direct internet access to add an additional layer of security.

The following topics describe how to connect your SageMaker Studio instances and notebook instances to resources in a VPC.

### Topics

- [Connect Amazon SageMaker Studio in a VPC to External Resources](#)
- [Connect SageMaker Studio Notebooks in a VPC to External Resources](#)
- [Connect a Notebook Instance in a VPC to External Resources](#)
Connect Amazon SageMaker Studio in a VPC to External Resources

⚠️ Important

As of November 30, 2023, the previous Amazon SageMaker Studio experience is now named Amazon SageMaker Studio Classic. The following section is specific to using the updated Studio experience. For information about using the Studio Classic application, see Amazon SageMaker Studio Classic.

The following topic gives information on how to connect Amazon SageMaker Studio in a VPC to external resources.

Topics

- Default communication with the internet
- VPC only communication with the internet

Default communication with the internet

By default, Amazon SageMaker Studio provides a network interface that allows communication with the internet through a VPC managed by SageMaker. Traffic to AWS services like Amazon S3 and CloudWatch goes through an internet gateway, as does traffic that accesses the SageMaker API and SageMaker runtime. Traffic between the domain and your Amazon EFS volume goes through the VPC that you specified when you onboarded to the domain or called the CreateDomain API.

VPC only communication with the internet

To prevent SageMaker from providing internet access to Studio, you can disable internet access by specifying the VPC only network access type when you onboard to Studio or call the CreateDomain API. As a result, you won’t be able to run Studio unless your VPC has an interface endpoint to the SageMaker API and runtime, or a NAT gateway with internet access, and your security groups allow outbound connections.

Note

The network access type can be changed after domain creation using the --app-network-access-type parameter of the update-domain command.
Requirements to use VPC only mode

When you choose VpcOnly, follow these steps:

1. You must use private subnets only. You cannot use public subnets in VpcOnly mode.
2. Ensure your subnets have the required number of IP addresses needed. The expected number of IP addresses needed per user can vary based on use case. We recommend between 2 and 4 IP addresses per user. The total IP address capacity for a domain is the sum of available IP addresses for each subnet provided when the domain is created. Ensure that your estimated IP address usage does not exceed the capacity supported by the number of subnets you provide. Additionally, using subnets distributed across many availability zones can aid in IP address availability. For more information, see VPC and subnet sizing for IPv4.

⚠️ Note

You can configure only subnets with a default tenancy VPC in which your instance runs on shared hardware. For more information on the tenancy attribute for VPCs, see Dedicated Instances.

⚠️ Warning

When using VpcOnly mode, you partly own the networking configuration for the domain. We recommend the security best practice of applying least-privilege permissions to the inbound and outbound access that security group rules provide. Overly permissive inbound rule configurations could allow users with access to the VPC to interact with the applications of other user profiles without authentication.

Set up one or more security groups with inbound and outbound rules that allow the following traffic:

- **NFS traffic over TCP on port 2049** between the domain and the Amazon EFS volume.
- **TCP traffic within the security group**. This is required for connectivity between the Jupyter Server application and the Kernel Gateway applications. You must allow access to at least ports in the range 8192–65535.
Create a distinct security group for each user profile and add inbound access from that same security group. We do not recommend reusing a domain-level security group for user profiles. If the domain-level security group allows inbound access to itself, then all applications in the domain would have access to all other applications in the domain.

4. If you want to allow internet access, you must use a [NAT gateway](#) with access to the internet, for example through an [internet gateway](#).

5. If you don’t want to allow internet access, [create interface VPC endpoints](#) (AWS PrivateLink) to allow Studio to access the following services with the corresponding service names. You must also associate the security groups for your VPC with these endpoints.

   • SageMaker API: com.amazonaws.\{region\}.sagemaker.api.
   • SageMaker runtime: com.amazonaws.\{region\}.sagemaker.runtime. This is required to run Studio notebooks and to train and host models.
   • Amazon S3: com.amazonaws.\{region\}.s3.
   • SageMaker Projects: com.amazonaws.\{region\}.servicecatalog.
   • SageMaker Studio: aws.sagemaker.\{region\}.studio.
   • Any other AWS services you require.

If you use the [SageMaker Python SDK](#) to run remote training jobs, you must also create the following Amazon VPC endpoints.

   • AWS Security Token Service: com.amazonaws.\{region\}.sts
   • Amazon CloudWatch: com.amazonaws.\{region\}.logs. This is required to allow SageMaker Python SDK to get the remote training job status from Amazon CloudWatch.

6. If using the domain in VpcOnly mode from an on-premises network, establish private connectivity from the network of the host running Studio in the browser and the target Amazon VPC. This is required because the Studio UI invokes AWS endpoints using API calls with temporary AWS credentials. These temporary credentials are associated with the execution role of the logged user profile. If the domain is configured in VpcOnly mode in an on-premises network, the execution role might define IAM policy conditions that enforce the execution of AWS service API calls only through the configured Amazon VPC endpoints. This causes API calls executed from the Studio UI to fail. We recommend resolving this using an [AWS Site-to-Site VPN](#) or [AWS Direct Connect](#) connection.
Note

For a customer working within VPC mode, company firewalls can cause connection issues with Studio or applications. Make the following checks if you encounter one of these issues when using Studio from behind a firewall.

- Verify that the Studio URL and URLs for all of your applications are in your network's allowlist. For example:

  *.*.studio.region.sagemaker.aws
  *.*.console.aws.a2z.com

- Verify that the websocket connections are not blocked. Jupyter uses websockets.

For more information

- [Security groups for your VPC](#)
- [Connect to SageMaker Within your VPC](#)
- [VPC with public and private subnets (NAT)](#)

Connect SageMaker Studio Notebooks in a VPC to External Resources

The following topic gives information on how to connect Studio Notebooks in a VPC to external resources.

Default communication with the internet

By default, SageMaker Studio provides a network interface that allows communication with the internet through a VPC managed by SageMaker. Traffic to AWS services like Amazon S3 and CloudWatch goes through an internet gateway, as does traffic that accesses the SageMaker API and SageMaker runtime. Traffic between the domain and your Amazon EFS volume goes through the VPC that you specified when you onboarded to Studio or called the [CreateDomain](#) API. The following diagram shows the default configuration.
To prevent SageMaker from providing internet access to your Studio notebooks, you can disable internet access by specifying the `VPC only` network access type when you onboard to Studio or call the `CreateDomain` API. As a result, you won’t be able to run a Studio notebook unless your VPC has an interface endpoint to the SageMaker API and runtime, or a NAT gateway with internet access, and your security groups allow outbound connections. The following diagram shows a configuration for using VPC-only mode.
Requirements to use VPC only mode

When you choose VpcOnly, follow these steps:

1. You must use private subnets only. You cannot use public subnets in VpcOnly mode.

2. Ensure your subnets have the required number of IP addresses needed. The expected number of IP addresses needed per user can vary based on use case. We recommend between 2 and 4 IP addresses per user. The total IP address capacity for a Studio domain is the sum of available IP addresses for each subnet provided when the domain is created. Ensure that your estimated IP address usage does not exceed the capacity supported by the number of subnets you provide. Additionally, using subnets distributed across many availability zones can aid in IP address availability. For more information, see VPC and subnet sizing for IPv4.

3. **Warning**

   When using VpcOnly mode, you partly own the networking configuration for the domain. We recommend the security best practice of applying least-privilege permissions to the inbound and outbound access that security group rules provide. Overly permissive inbound rule configurations could allow users with access to the VPC to interact with the applications of other user profiles without authentication.

   Set up one or more security groups with inbound and outbound rules that allow the following traffic:

   - **NFS traffic over TCP on port 2049** between the domain and the Amazon EFS volume.
   - **TCP traffic within the security group.** This is required for connectivity between the Jupyter Server application and the Kernel Gateway applications. You must allow access to at least ports in the range 8192-65535.
Create a distinct security group for each user profile and add inbound access from that same security group. We do not recommend reusing a domain-level security group for user profiles. If the domain-level security group allows inbound access to itself, then all applications in the domain would have access to all other applications in the domain.

4. If you want to allow internet access, you must use a **NAT gateway** with access to the internet, for example through an **internet gateway**.

5. If you don't want to allow internet access, create interface VPC endpoints (AWS PrivateLink) to allow Studio to access the following services with the corresponding service names. You must also associate the security groups for your VPC with these endpoints.

   - **SageMaker API**: `com.amazonaws.region.sagemaker.api`
   - **SageMaker runtime**: `com.amazonaws.region.sagemaker.runtime`. This is required to run Studio notebooks and to train and host models.
   - **Amazon S3**: `com.amazonaws.region.s3`
   - **To use SageMaker Projects**: `com.amazonaws.region.servicecatalog`
   - **Any other AWS services you require**.

If you use the [SageMaker Python SDK](https://sagemaker.readthedocs.io/en/stable/) to run remote training jobs, you must also create the following Amazon VPC endpoints.

   - **AWS Security Token Service**: `com.amazonaws.region.sts`
   - **Amazon CloudWatch**: `com.amazonaws.region.logs`. This is required to allow SageMaker Python SDK to get the remote training job status from Amazon CloudWatch.

---

**Note**

For a customer working within VPC mode, company firewalls can cause connection issues with SageMaker Studio or between JupyterServer and the KernelGateway. Make the following checks if you encounter one of these issues when using SageMaker Studio from behind a firewall.

- Check that the Studio URL is in your networks allowlist.
- Check that the websocket connections are not blocked. Jupyter uses websocket under the hood. If the KernelGateway application is InService, JupyterServer may not be able
to connect to the KernelGateway. You should see this problem when opening System Terminal as well.

For more information

- Securing Amazon SageMaker Studio connectivity using a private VPC.
- Security groups for your VPC
- Connect to SageMaker Within your VPC
- VPC with public and private subnets (NAT)

Connect a Notebook Instance in a VPC to External Resources

The following topic gives information on how to connect your notebook instance in a VPC to external resources.

Default communication with the internet

When your notebook allows direct internet access, SageMaker provides a network interface that allows the notebook to communicate with the internet through a VPC managed by SageMaker. Traffic within your VPC's CIDR goes through elastic network interface created in your VPC. All the other traffic goes through the network interface created by SageMaker, which is essentially through the public internet. Traffic to gateway VPC endpoints like Amazon S3 and DynamoDB goes through the public internet, while traffic to interface VPC interface endpoints still goes through your VPC. If you want to use gateway VPC endpoints, you might want to disable direct internet access.

VPC communication with the internet

To disable direct internet access, you can specify a VPC for your notebook instance. By doing so, you prevent SageMaker from providing internet access to your notebook instance. As a result, the notebook instance can't train or host models unless your VPC has an interface endpoint (AWS PrivateLink) or a NAT gateway and your security groups allow outbound connections.

For information about creating a VPC interface endpoint to use AWS PrivateLink for your notebook instance, see Connect to a Notebook Instance Through a VPC Interface Endpoint. For information about setting up a NAT gateway for your VPC, see VPC with Public and Private Subnets (NAT) in the Amazon Virtual Private Cloud User Guide. For information about security groups, see Security Groups for Your VPC. For more information about networking configurations in each networking
mode and configuring network on premise, see [Understanding Amazon SageMaker notebook instance networking configurations and advanced routing options](#).

**Security and Shared Notebook Instances**

A SageMaker notebook instance is designed to work best for an individual user. It is designed to give data scientists and other users the most power for managing their development environment.

A notebook instance user has root access for installing packages and other pertinent software. We recommend that you exercise judgement when granting individuals access to notebook instances that are attached to a VPC that contains sensitive information. For example, you might grant a user access to a notebook instance with an IAM policy, as shown in the following example:

```json
{
  "Version": "2012-10-17",
  "Statement": [
    {
      "Effect": "Allow",
      "Action": "sagemaker:CreatePresignedNotebookInstanceUrl",
    }
  ]
}
```

**Run Training and Inference Containers in Internet-Free Mode**

SageMaker training and deployed inference containers are internet-enabled by default. This allows containers to access external services and resources on the public internet as part of your training and inference workloads. However, this could provide an avenue for unauthorized access to your data. For example, a malicious user or code that you accidentally install on the container (in the form of a publicly available source code library) could access your data and transfer it to a remote host.

If you use an Amazon VPC by specifying a value for the VpcConfig parameter when you call `CreateTrainingJob`, `CreateHyperParameterTuningJob`, or `CreateModel`, you can protect your data and resources by managing security groups and restricting internet access from your VPC. However, this comes at the cost of additional network configuration, and has the risk of configuring your network incorrectly. If you do not want SageMaker to provide external network access to your training or inference containers, you can enable network isolation.
Network Isolation

You can enable network isolation when you create your training job or model by setting the value of the EnableNetworkIsolation parameter to True when you call CreateTrainingJob, CreateHyperParameterTuningJob, or CreateModel.

⚠️ Note

Network isolation is required to run training jobs and models using resources from AWS Marketplace. For additional security, AWS Marketplace images run within an Amazon VPC. They only have access to data within their local file systems.

If you enable network isolation, the containers can't make any outbound network calls, even to other AWS services such as Amazon S3. Additionally, no AWS credentials are made available to the container runtime environment. In the case of a training job with multiple instances, network inbound and outbound traffic is limited to the peers of each training container. SageMaker still performs download and upload operations against Amazon S3 using your SageMaker execution role in isolation from the training or inference container.

The following managed SageMaker containers do not support network isolation because they require access to Amazon S3:

- Chainer
- SageMaker Reinforcement Learning

Network isolation with a VPC

Network isolation can be used in conjunction with a VPC. In this scenario, the download and upload of customer data and model artifacts are routed through your VPC subnet. However, the training and inference containers themselves continue to be isolated from the network, and do not have access to any resource within your VPC or on the internet.

Connect to SageMaker Within your VPC

You can connect directly to the SageMaker API or to Amazon SageMaker Runtime through an interface endpoint in your virtual private cloud (VPC) instead of connecting over the internet. When you use a VPC interface endpoint, communication between your VPC and the SageMaker API or Runtime is conducted entirely and securely within an AWS network.
Connect to SageMaker through a VPC interface endpoint

The SageMaker API and SageMaker Runtime support Amazon Virtual Private Cloud (Amazon VPC) interface endpoints that are powered by AWS PrivateLink. Each VPC endpoint is represented by one or more Elastic Network Interfaces with private IP addresses in your VPC subnets. For example, an application inside your VPC uses AWS PrivateLink to communicate with SageMaker Runtime. SageMaker Runtime in turn communicates with the SageMaker endpoint. Using AWS PrivateLink allows you to invoke your SageMaker endpoint from within your VPC, as shown in the following diagram.

The VPC interface endpoint connects your VPC directly to the SageMaker API or SageMaker Runtime using AWS PrivateLink without using an internet gateway, NAT device, VPN connection, or AWS Direct Connect connection. The instances in your VPC do not need to connect to the public internet in order to communicate with the SageMaker API or SageMaker Runtime.

You can create an AWS PrivateLink interface endpoint to connect to SageMaker or to SageMaker Runtime using either the AWS Management Console or AWS Command Line Interface (AWS CLI). For instructions, see Access an AWS service using an interface VPC endpoint.

If you haven't enabled a private Domain Name System (DNS) hostname for your VPC endpoint, after you have created a VPC endpoint, specify the internet endpoint URL to the SageMaker API or SageMaker Runtime. Example code using AWS CLI commands to specify the endpoint-url parameter follows.
If you enable private DNS hostnames for your VPC endpoint, you don't need to specify the endpoint URL because the default hostname (https://api.sagemaker.Region.amazonaws.com) resolves to your VPC endpoint. Similarly, the default SageMaker Runtime DNS hostname (https://runtime.sagemaker.Region.amazonaws.com) also resolves to your VPC endpoint.

The SageMaker API and SageMaker Runtime support VPC endpoints in all AWS Regions where both Amazon VPC and SageMaker are available. SageMaker supports making calls to all of its Operations inside your VPC. If you use the AuthorizedUrl from the CreatePresignedNotebookInstanceUrl command, your traffic will go over the public internet. You can't only use a VPC endpoint to access the presigned URL, the request must go through the internet gateway.

By default, your users can share the presigned URL to people outside of your corporate network. For additional security, you must add IAM permissions to restrict the URL only be usable within your network. For information about IAM permissions, see How AWS PrivateLink works with IAM.

Note

When setting up a VPC interface endpoint for the SageMaker Runtime service (https://runtime.sagemaker.Region.amazonaws.com), you must ensure that the VPC interface endpoint is activated in the Availability Zone of your client in order for private DNS resolution to work. Otherwise, you may see DNS failures when attempting to resolve the URL.

To learn more about AWS PrivateLink, see the AWS PrivateLink documentation. Refer to AWS PrivateLink Pricing for the price of VPC endpoints. To learn more about VPC and endpoints,
see Amazon VPC. For information about how to use identity-based AWS Identity and Access Management policies to restrict access to the SageMaker API and SageMaker Runtime, see Control Access to the SageMaker API by Using Identity-based Policies.

Using SageMaker training and hosting with resources inside your VPC

SageMaker uses your execution role to download and upload information from an Amazon S3 bucket and Amazon Elastic Container Registry (Amazon ECR), in isolation from your training or inference container. If you have resources that are located inside your VPC, you can still grant SageMaker access to those resources. The following sections explain how to make your resources available to SageMaker with or without network isolation.

Without network isolation enabled

If you haven't set network isolation on your training job or model, SageMaker can access resources using either of the following methods.

- SageMaker training and deployed inference containers can access the internet by default. SageMaker containers are able to access external services and resources on the public internet as part of your training and inference workloads. SageMaker containers are not able to access resources inside your VPC without a VPC configuration, as shown in the following illustration.
- Use a VPC configuration to communicate with resources inside your VPC through an elastic network interface (ENI). The communication between the container and the resources in your VPC takes place securely within your VPC network, as shown in the following illustration. In this case, you manage networking access to your VPC resources and internet.
With network isolation

If you employ network isolation, the SageMaker container can't communicate with resources inside your VPC or make any network calls, as shown in the following illustration. If you provide a VPC configuration, the download and upload operations will be run through your VPC. For more information about hosting and training with network isolation while using a VPC, see Network Isolation.
Create a VPC Endpoint Policy for SageMaker

You can create a policy for Amazon VPC endpoints for SageMaker to specify the following:

- The principal that can perform actions.
- The actions that can be performed.
- The resources on which actions can be performed.

For more information, see [Controlling Access to Services with VPC Endpoints](https://docs.aws.amazon.com/vpc/latest/userguide/vpc-access-control-vpce.html) in the *Amazon VPC User Guide*. 
Note

VPC endpoint policies aren't supported for Federal Information Processing Standard (FIPS) SageMaker runtime endpoints for `runtime_InvokeEndpoint`.

The following example VPC endpoint policy specifies that all users who have access to the VPC interface endpoint are allowed to invoke the SageMaker hosted endpoint named `myEndpoint`.

```
{
    "Statement": [
        {
            "Action": "sagemaker:InvokeEndpoint",
            "Effect": "Allow",
            "Principal": "*
        }
    ]
}
```

In this example, the following are denied:

- Other SageMaker API actions, such as `sagemaker:CreateEndpoint` and `sagemaker:CreateTrainingJob`.
- Invoking SageMaker hosted endpoints other than `myEndpoint`.

Note

In this example, users can still take other SageMaker API actions from outside the VPC. For information about how to restrict API calls to those from within the VPC, see Control Access to the SageMaker API by Using Identity-based Policies.

Create a VPC Endpoint Policy for Amazon SageMaker Feature Store

To create a VPC Endpoint for Amazon SageMaker Feature Store, use the following endpoint template, substituting your `VPC_Endpoint_ID.api` and `Region`:

```
Connect to SageMaker Studio Classic Through an Interface VPC Endpoint

You can connect to Amazon SageMaker Studio Classic from your Amazon Virtual Private Cloud (Amazon VPC) through an interface endpoint in your VPC instead of connecting over the internet. When you use an interface VPC endpoint (interface endpoint), communication between your VPC and Studio Classic is conducted entirely and securely within the AWS network.

SageMaker Studio Classic supports interface endpoints that are powered by AWS PrivateLink. Each interface endpoint is represented by one or more Elastic network interfaces with private IP addresses in your VPC subnets.

Studio Classic supports interface endpoints in all AWS Regions where both Amazon SageMaker and Amazon VPC are available.

Topics

- Create a VPC Endpoint
- Create a VPC Endpoint Policy for SageMaker Studio Classic
- Allow Access Only from Within Your VPC

Create a VPC Endpoint

You can create an interface endpoint to connect to Studio Classic with either the AWS console or the AWS Command Line Interface (AWS CLI). For instructions, see Creating an interface endpoint. Make sure that you create interface endpoints for all of the subnets in your VPC from which you want to connect to Studio Classic.

When you create an interface endpoint, ensure that the security groups on your endpoint allow inbound access for HTTPS traffic from the security groups associated with SageMaker Studio Classic. For more information, see Control access to services with VPC endpoints.

Note

In addition to creating an interface endpoint to connect to SageMaker Studio Classic, create an interface endpoint to connect to the Amazon SageMaker API. When users call
CreatePresignedDomainUrl to get the URL to connect to Studio Classic, that call goes through the interface endpoint used to connect to the SageMaker API.

When you create the interface endpoint, specify `aws.sagemaker.Region.studio` as the service name. After you create the interface endpoint, enable private DNS for your endpoint. When you connect to SageMaker Studio Classic from within the VPC using the SageMaker API, the AWS CLI, or the console, you connect through the interface endpoint instead of the public internet. You also need to set up a custom DNS with private hosted zones for the Amazon VPC endpoint so SageMaker Studio Classic can access the SageMaker API using the `api.sagemaker.$region.amazonaws.com` endpoint rather than using the VPC endpoint URL. For instructions on setting up a private hosted zone, see [Working with private hosted zones](#).

**Create a VPC Endpoint Policy for SageMaker Studio Classic**

You can attach an Amazon VPC endpoint policy to the interface VPC endpoints that you use to connect to SageMaker Studio Classic. The endpoint policy controls access to Studio Classic. You can specify the following:

- The principal that can perform actions.
- The actions that can be performed.
- The resources on which actions can be performed.

To use a VPC endpoint with SageMaker Studio Classic, your endpoint policy must allow the CreateApp operation on the KernelGateway app type. This allows traffic that is routed to through the VPC endpoint to call the CreateApp API. The following example VPC endpoint policy shows how to allow the CreateApp operation.

```json
{
  "Statement": [
    {
      "Action": "sagemaker:CreateApp",
      "Effect": "Allow",
      "Principal": "*
    }
  ]
}
```
For more information, see [Controlling access to services with VPC endpoints](https://docs.aws.amazon.com/sagemaker/latest/dg/vpc-endpoint-auth.html).

The following example of a VPC endpoint policy specifies that all users that have access to the endpoint are allowed to access the user profiles in the SageMaker domain with the specified domain ID. Access to other domains is denied.

```json
{
    "Statement": [
        {
            "Action": "sagemaker:CreatePresignedDomainUrl",
            "Effect": "Allow",
            "Principal": "*
        }
    ]
}
```

### Allow Access Only from Within Your VPC

Users outside your VPC can connect to SageMaker Studio Classic over the internet even if you set up an interface endpoint in your VPC.

To allow access to only connections made from within your VPC, create an AWS Identity and Access Management (IAM) policy to that effect. Add that policy to every user, group, or role used to access Studio Classic. This feature is only supported in IAM mode, and is not supported in IAM Identity Center mode. The following examples demonstrate how to create such policies.

⚠️ **Important**

If you apply an IAM policy similar to one of the following examples, users can't access SageMaker Studio Classic or the specified SageMaker APIs through the SageMaker console. To access Studio Classic, users must use a presigned URL or call the SageMaker APIs directly.

### Example 1: Allow connections only within the subnet of an interface endpoint

The following policy allows connections only to callers within the subnet where you created the interface endpoint.

```json
{
    "Statement": [
        {
            "Action": "sagemaker:CreatePresignedDomainUrl",
            "Effect": "Allow",
            "Principal": "*
        }
    ]
}
```
Example 2: Allow connections only through interface endpoints using `aws:sourceVpce`

The following policy allows connections only to those made through the interface endpoints specified by the `aws:sourceVpce` condition key. For example, the first interface endpoint could allow access through the SageMaker console. The second interface endpoint could allow access through the SageMaker API.

```json
{
  "Id": "sagemaker-studio-example-2",
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "Enable SageMaker Studio Access",
      "Effect": "Allow",
      "Action": [
        "sagemaker:CreatePresignedDomainUrl",
        "sagemaker:DescribeUserProfile"
      ],
      "Resource": "*",
      "Condition": {
        "ForAnyValue:StringEquals": {
          "aws:sourceVpce": ["vpce-111bbaaa",

        "vpce-111bbccc",

      }
    }
  ]
}
```
This policy includes the `DescribeUserProfile` action. Typically you call `DescribeUserProfile` to make sure that the status of the user profile is `InService` before you try to connect to the domain. For example:

```
aws sagemaker describe-user-profile
   --domain-id domain-id
   --user-profile-name profile-name
```

Response:

```
{
    "DomainId": "domain-id",
    "UserProfileName": "profile-name",
    "HomeEfsFileSystemUid": "200001",
    "Status": "InService",
    "LastModifiedTime": 1605418785.555,
    "CreationTime": 1605418477.297
}
```

```
aws sagemaker create-presigned-domain-url
   --domain-id domain-id
   --user-profile-name profile-name
```

Response:

```
{
    "AuthorizedUrl": "https://domain-id.studio.us-west-2.sagemaker.aws/auth?
    token=AuthToken"
}
```
For both of these calls, if you are using a version of the AWS SDK that was released before August 13, 2018, you must specify the endpoint URL in the call. For example, the following example shows a call to `create-presigned-domain-url`:

```
aws sagemaker create-presigned-domain-url
   --domain-id domain-id \
   --user-profile-name profile-name \
   --endpoint-url vpc-endpoint-id.api.sagemaker.Region.vpce.amazonaws.com
```

**Example 3: Allow connections from IP addresses using `aws:SourceIp`**

The following policy allows connections only from the specified range of IP addresses using the `aws:SourceIp` condition key.

```
{
   "Id": "sagemaker-studio-example-3",
   "Version": "2012-10-17",
   "Statement": [
      {
         "Sid": "Enable SageMaker Studio Access",
         "Effect": "Allow",
         "Action": [
            "sagemaker:CreatePresignedDomainUrl",
            "sagemaker:DescribeUserProfile"
         ],
         "Resource": "*",
         "Condition": {
            "IpAddress": {
               "aws:SourceIp": [
                  "192.0.2.0/24",
                  "203.0.113.0/24"
               ]
            }
         }
      }
   ]
}
```

**Example 4: Allow connections from IP addresses through an interface endpoint using `aws:VpcSourceIp`**

Connect to SageMaker Within your VPC
If you are accessing SageMaker Studio Classic through an interface endpoint, you can use the `aws:VpcSourceIp` condition key to allow connections only from the specified range of IP addresses within the subnet where you created the interface endpoint as shown in the following policy:

```json
{
  "Id": "sagemaker-studio-example-4",
  "Version": "2012-10-17",
  "Statement": [
    {
      "Sid": "Enable SageMaker Studio Access",
      "Effect": "Allow",
      "Action": [
        "sagemaker:CreatePresignedDomainUrl",
        "sagemaker:DescribeUserProfile"
      ],
      "Resource": "*",
      "Condition": {
        "IpAddress": {
          "aws:VpcSourceIp": [
            "192.0.2.0/24",
            "203.0.113.0/24"
          ],
          "StringEquals": {
            "aws:SourceVpc": "vpc-111bbaaa"
          }
        }
      }
    }
  ]
}
```

**Connect to a Notebook Instance Through a VPC Interface Endpoint**

You can connect to your notebook instance from your VPC through an [interface endpoint](https://aws.amazon.com/sagemaker/) in your Virtual Private Cloud (VPC) instead of connecting over the public internet. When you use a VPC interface endpoint, communication between your VPC and the notebook instance is conducted entirely and securely within the AWS network.

SageMaker notebook instances support [Amazon Virtual Private Cloud](https://aws.amazon.com/vpc/) (Amazon VPC) interface endpoints that are powered by [AWS PrivateLink](https://aws.amazon.com/privatelink/). Each VPC endpoint is represented by one or more Elastic Network Interfaces with private IP addresses in your VPC subnets.
Note
Before you create an interface VPC endpoint to connect to a notebook instance, create an interface VPC endpoint to connect to the SageMaker API. That way, when users call `CreatePresignedNotebookInstanceUrl` to get the URL to connect to the notebook instance, that call also goes through the interface VPC endpoint. For information, see Connect to SageMaker Within your VPC.

You can create an interface endpoint to connect to your notebook instance with either the AWS Management Console or AWS Command Line Interface (AWS CLI) commands. For instructions, see Creating an Interface Endpoint. Make sure that you create an interface endpoint for all of the subnets in your VPC from which you want to connect to the notebook instance.

When you create the interface endpoint, specify `aws.sagemaker.Region.notebook` as the service name. After you create a VPC endpoint, enable private DNS for your VPC endpoint. Anyone using the SageMaker API, the AWS CLI, or the console to connect to the notebook instance from within the VPC connects to the notebook instance through the VPC endpoint instead of the public internet.

SageMaker notebook instances support VPC endpoints in all AWS Regions where both Amazon VPC and SageMaker are available.

Topics
- Connect Your Private Network to Your VPC
- Create a VPC Endpoint Policy for SageMaker Notebook Instances
- Restrict Access to Connections from Within Your VPC

Connect Your Private Network to Your VPC

To connect to your notebook instance through your VPC, you either have to connect from an instance that is inside the VPC, or connect your private network to your VPC by using an AWS Virtual Private Network (AWS VPN) or AWS Direct Connect. For information about AWS VPN, see VPN Connections in the Amazon Virtual Private Cloud User Guide. For information about AWS Direct Connect, see Creating a Connection in the AWS Direct Connect User Guide.
Create a VPC Endpoint Policy for SageMaker Notebook Instances

You can create a policy for Amazon VPC endpoints for SageMaker notebook instances to specify the following:

- The principal that can perform actions.
- The actions that can be performed.
- The resources on which actions can be performed.

For more information, see Controlling Access to Services with VPC Endpoints in the Amazon VPC User Guide.

The following example of a VPC endpoint policy specifies that all users that have access to the endpoint are allowed to access the notebook instance named myNotebookInstance.

```
{
   "Statement": [
      {
         "Action": "sagemaker:CreatePresignedNotebookInstanceUrl",
         "Effect": "Allow",
         "Principal": "*"
      }
   ]
}
```

Access to other notebook instances is denied.

Restrict Access to Connections from Within Your VPC

Even if you set up an interface endpoint in your VPC, individuals outside the VPC can connect to the notebook instance over the internet.

⚠️ Important

If you apply an IAM policy similar to one of the following, users can't access the specified SageMaker APIs or the notebook instance through the console.
To restrict access to only connections made from within your VPC, create an AWS Identity and Access Management policy that restricts access to only calls that come from within your VPC. Then add that policy to every AWS Identity and Access Management user, group, or role used to access the notebook instance.

**Note**

This policy allows connections only to callers within a subnet where you created an interface endpoint.

```json
{
    "Id": "notebook-example-1",
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "Enable Notebook Access",
            "Effect": "Allow",
            "Action": ["sagemaker:CreatePresignedNotebookInstanceUrl", "sagemaker:DescribeNotebookInstance"],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "aws:SourceVpc": "vpc-111bbaaa"
                }
            }
        }
    ]
}
```

If you want to restrict access to the notebook instance to only connections made using the interface endpoint, use the `aws:SourceVpce` condition key instead of `aws:SourceVpc`:

```json
{
    "Id": "notebook-example-1",
    "Version": "2012-10-17",
    "Statement": [
        {
            "Sid": "Enable Notebook Access",
            "Effect": "Allow",
            "Action": ["sagemaker:CreatePresignedNotebookInstanceUrl", "sagemaker:DescribeNotebookInstance"],
            "Resource": "*",
            "Condition": {
                "StringEquals": {
                    "aws:SourceVpce": "vpc-111bbaaa"
                }
            }
        }
    ]
}
```
Both of these policy examples assume that you have also created an interface endpoint for the SageMaker API. For more information, see [Connect to SageMaker Within your VPC](#). In the second example, one of the values for `aws:SourceVpce` is the ID of the interface endpoint for the notebook instance. The other is the ID of the interface endpoint for the SageMaker API.

The policy examples here include [DescribeNotebookInstance](#), because typically you would call `DescribeNotebookInstance` to make sure that the `NotebookInstanceStatus` is `InService` before you try to connect to it. For example:

```
aws sagemaker describe-notebook-instance
    --notebook-instance-name myNotebookInstance
```

```json
{
    "NotebookInstanceArn":
    "NotebookInstanceName": "myNotebookInstance",
    "NotebookInstanceStatus": "InService",
    "Url": "mynotebookinstance.notebook.us-west-2.sagemaker.aws",
    "InstanceType": "ml.m4.xlarge",
    "RoleArn":
    "arn:aws:iam::1234567890ab:role/service-role/AmazonSageMaker-ExecutionRole-12345678T123456",
```
Note

The `presigned-notebook-instance-url`, `AuthorizedUrl`, generated can be used from anywhere on the internet.

For both of these calls, if you did not enable private DNS hostnames for your VPC endpoint, or if you are using a version of the AWS SDK that was released before August 13, 2018, you must specify the endpoint URL in the call. For example, the call to `create-presigned-notebook-instance-url` is:

```
aws sagemaker create-presigned-notebook-instance-url --notebook-instance-name myNotebookInstance --endpoint-url VPC_Endpoint_ID.api.sagemaker.Region.vpce.amazonaws.com
```

### Connect Your Private Network to Your VPC

To call the SageMaker API and SageMaker Runtime through your VPC, you have to connect from an instance that is inside the VPC or connect your private network to your VPC by using an AWS Virtual Private Network (AWS VPN) or AWS Direct Connect. For information about AWS VPN, see [VPN Connections](#) in the [Amazon Virtual Private Cloud User Guide](#). For information about AWS Direct Connect, see [Creating a Connection](#) in the [AWS Direct Connect User Guide](#).

### Give SageMaker Access to Resources in your Amazon VPC

SageMaker runs the following job types in an Amazon Virtual Private Cloud by default.
However, containers for these jobs access AWS resources—such as the Amazon Simple Storage Service (Amazon S3) buckets where you store training data and model artifacts—over the internet.

To control access to your data and job containers, we recommend that you create a private VPC and configure it so that they aren't accessible over the internet. For information about creating and configuring a VPC, see Getting Started With Amazon VPC in the Amazon VPC User Guide. Using a VPC helps to protect your job containers and data because you can configure your VPC so that it is not connected to the internet. Using a VPC also allows you to monitor all network traffic in and out of your job containers by using VPC flow logs. For more information, see VPC Flow Logs in the Amazon VPC User Guide.

You specify your private VPC configuration when you create jobs by specifying subnets and security groups. When you specify the subnets and security groups, SageMaker creates elastic network interfaces that are associated with your security groups in one of the subnets. Network interfaces allow your job containers to connect to resources in your VPC. For information about network interfaces, see Elastic Network Interfaces in the Amazon VPC User Guide.

You specify a VPC configuration within the VpcConfig object of the CreateProcessingJob operation or CreateTrainingJob operation. Specifying a VPC configuration when you create a training job gives your model access to resources within your VPC.

Specifying a VPC configuration alone doesn't change the invocation path. To connect to Amazon SageMaker within a VPC, create a VPC endpoint and invoke it. For more information, see Connect to SageMaker Within your VPC.

Topics

- Give SageMaker Processing Jobs Access to Resources in Your Amazon VPC
- Give SageMaker Training Jobs Access to Resources in Your Amazon VPC
- Give SageMaker Hosted Endpoints Access to Resources in Your Amazon VPC
To control access to your data and processing jobs, create a Amazon VPC with private subnets. For information about creating and configuring a VPC, see Get Started With Amazon VPC in the Amazon VPC User Guide.

You can monitor all network traffic in and out of your processing containers by using VPC flow logs. For more information, see VPC Flow Logs in the Amazon VPC User Guide.

This document explains how to add Amazon VPC configurations for processing jobs.

Configure a Processing Job for Amazon VPC Access

You configure the processing job by specifying the subnets and security group IDs within the VPC. You don’t need to specify the subnet for the processing container. Amazon SageMaker automatically pulls the processing container from Amazon ECR. For more information about processing containers, see Process data.

When creating a processing job, you can specify subnets and security groups in your VPC using either the SageMaker console or the API.

To use the API, you specify the subnets and security group IDs in the NetworkConfig.VpcConfig parameter of the CreateProcessingJob operation. SageMaker uses the subnet and security group details to create the network interfaces and attaches them to the processing containers. The network interfaces provide the processing containers with a network connection within your VPC. This allows the processing job to connect to resources that exist in your VPC.

The following is an example of the VpcConfig parameter that you include in your call to the CreateProcessingJob operation:

```json
VpcConfig: {
    "Subnets": [ 
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2"
    ]
}
```
Configure Your Private VPC for SageMaker Processing

When configuring the private VPC for your SageMaker processing jobs, use the following
guidelines. For information about setting up a VPC, see Working with VPCs and Subnets in the
Amazon VPC User Guide.

Topics

- Ensure That Subnets Have Enough IP Addresses
- Create an Amazon S3 VPC Endpoint
- Use a Custom Endpoint Policy to Restrict Access to S3
- Configure Route Tables
- Configure the VPC Security Group
- Connect to Resources Outside Your VPC
- Monitor Amazon SageMaker Processing Jobs with CloudWatch Logs and Metrics

Ensure That Subnets Have Enough IP Addresses

Your VPC subnets should have at least two private IP addresses for each instance in a processing
job. For more information, see VPC and Subnet Sizing for IPv4 in the Amazon VPC User Guide.

Create an Amazon S3 VPC Endpoint

If you configure your VPC so that processing containers don't have access to the internet, they
can't connect to the Amazon S3 buckets that contain your data unless you create a VPC endpoint
that allows access. By creating a VPC endpoint, you allow your processing containers to access
the buckets where you store your data. We recommend that you also create a custom policy that
allows only requests from your private VPC to access to your S3 buckets. For more information, see
Endpoints for Amazon S3.

To create an S3 VPC endpoint:

1. Open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
2. In the navigation pane, choose **Endpoints**, then choose **Create Endpoint**

3. For **Service Name**, choose `com.amazonaws.region.s3`, where `region` is the name of the region where your VPC resides.

4. For **VPC**, choose the VPC you want to use for this endpoint.

5. For **Configure route tables**, select the route tables to be used by the endpoint. The VPC service automatically adds a route to each route table you select that points any S3 traffic to the new endpoint.

6. For **Policy**, choose **Full Access** to allow full access to the S3 service by any user or service within the VPC. Choose **Custom** to restrict access further. For information, see **Use a Custom Endpoint Policy to Restrict Access to S3**.

### Use a Custom Endpoint Policy to Restrict Access to S3

The default endpoint policy allows full access to S3 for any user or service in your VPC. To further restrict access to S3, create a custom endpoint policy. For more information, see **Using Endpoint Policies for Amazon S3**. You can also use a bucket policy to restrict access to your S3 buckets to only traffic that comes from your Amazon VPC. For information, see **Using Amazon S3 Bucket Policies**.

### Restrict Package Installation on the Processing Container

The default endpoint policy allows users to install packages from the Amazon Linux and Amazon Linux 2 repositories on the processing container. If you don't want users to install packages from that repository, create a custom endpoint policy that explicitly denies access to the Amazon Linux and Amazon Linux 2 repositories. The following is an example of a policy that denies access to these repositories:

```json
{
    "Statement": [
        {
            "Sid": "AmazonLinuxAMIRepositoryAccess",
            "Principal": "*",
            "Action": [
                "s3:GetObject"
            ],
            "Effect": "Deny",
            "Resource": [
                "arn:aws:s3:::packages.*.amazonaws.com/*",
                "arn:aws:s3:::repo.*.amazonaws.com/*"
            ]
        }
    ]
}
```
Configure Route Tables

Use default DNS settings for your endpoint route table, so that standard Amazon S3 URLs (for example, http://s3-aws-region.amazonaws.com/MyBucket) resolve. If you don't use default DNS settings, ensure that the URLs that you use to specify the locations of the data in your processing jobs resolve by configuring the endpoint route tables. For information about VPC endpoint route tables, see Routing for Gateway Endpoints in the Amazon VPC User Guide.

Configure the VPC Security Group

In distributed processing, you must allow communication between the different containers in the same processing job. To do that, configure a rule for your security group that allows inbound connections between members of the same security group. For more information, see Security Group Rules.

Connect to Resources Outside Your VPC

If you're connecting your models to resources outside the VPC that they're running in, do one of the following:

- **Connect to other AWS services** – If your model needs access to an AWS service that supports interface Amazon VPC endpoints, create an endpoint to connect to that service. For a list of
services that support interface endpoints, see AWS services that integrate with AWS PrivateLink in the AWS PrivateLink User Guide. For information about creating an interface VPC endpoint, see Access an AWS service using an interface VPC endpoint in the AWS PrivateLink User Guide.

- **Connect to resources over the internet** – If your models are running on instances in an Amazon VPC that does not have a subnet with access to the internet, the models won't have access to resources on the internet. If your model needs access to an AWS service that doesn't support interface VPC endpoints, or to a resource outside of AWS, ensure that you are running your models in a private subnet that has access to the internet using a public NAT gateway in a public subnet. After you have your models running in the private subnet, configure your security groups and network access control lists (NACLs) to allow outbound connections from the private subnet to the public NAT gateway in the public subnet. For information, see NAT gateways in the Amazon VPC User Guide.

**Monitor Amazon SageMaker Processing Jobs with CloudWatch Logs and Metrics**

Amazon SageMaker provides Amazon CloudWatch logs and metrics to monitor training jobs. CloudWatch provides CPU, GPU, memory, GPU memory, and disk metrics, and event logging. For more information about monitoring Amazon SageMaker processing jobs, see Monitor Amazon SageMaker with Amazon CloudWatch and SageMaker Jobs and Endpoint Metrics.

**Give SageMaker Training Jobs Access to Resources in Your Amazon VPC**

- **Note**
  
  For training jobs, you can configure only subnets with a default tenancy VPC in which your instance runs on shared hardware. For more information on the tenancy attribute for VPCs, see Dedicated Instances.

**Configure a Training Job for Amazon VPC Access**

To control access to your training jobs, run them in an Amazon VPC with private subnets that don’t have internet access.

You configure the training job to run in the VPC by specifying its subnets and security group IDs. You don’t need to specify the subnet for the container of the training job. Amazon SageMaker automatically pulls the training container image from Amazon ECR.
When you create a training job, you can specify the subnets and security groups in your VPC using the Amazon SageMaker console or the API.

To use the API, you specify the subnets and security group IDs in the VpcConfig parameter of the CreateTrainingJob operation. SageMaker uses the subnet and security group details to create the network interfaces and attaches them to the training containers. The network interfaces provide the training containers with a network connection within your VPC. This allows the training job to connect to resources that exist in your VPC.

The following is an example of the VpcConfig parameter that you include in your call to the CreateTrainingJob operation:

```json
VpcConfig: {
    "Subnets": [  
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2"
    ],
    "SecurityGroupIds": [  
        "sg-0123456789abcdef0"
    ]
}
```

**Configure Your Private VPC for SageMaker Training**

When configuring the private VPC for your SageMaker training jobs, use the following guidelines. For information about setting up a VPC, see [Working with VPCs and Subnets](#) in the Amazon VPC User Guide.

**Topics**

- [Ensure That Subnets Have Enough IP Addresses](#)
- [Create an Amazon S3 VPC Endpoint](#)
- [Use a Custom Endpoint Policy to Restrict Access to S3](#)
- [Configure Route Tables](#)
- [Configure the VPC Security Group](#)
- [Connect to Resources Outside Your VPC](#)
- [Monitor Amazon SageMaker Training Jobs with CloudWatch Logs and Metrics](#)
Ensure That Subnets Have Enough IP Addresses

Training instances that don't use an Elastic Fabric Adapter (EFA) should have at least 2 private IP addresses. Training instances that use an EFA should have at least 5 private IP addresses. For more information, see Multiple IP addresses in the Amazon EC2 User Guide.

Your VPC subnets should have at least two private IP addresses for each instance in a training job. For more information, see VPC and Subnet Sizing for IPv4 in the Amazon VPC User Guide.

Create an Amazon S3 VPC Endpoint

If you configure your VPC so that training containers don't have access to the internet, they can't connect to the Amazon S3 buckets that contain your training data unless you create a VPC endpoint that allows access. By creating a VPC endpoint, you allow your training containers to access the buckets where you store your data and model artifacts. We recommend that you also create a custom policy that allows only requests from your private VPC to access to your S3 buckets. For more information, see Endpoints for Amazon S3.

To create an S3 VPC endpoint:

1. Open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
2. In the navigation pane, choose Endpoints, then choose Create Endpoint
3. For Service Name, search for com.amazonaws.region.s3, where region is the name of the region where your VPC resides.
4. Choose the Gateway type.
5. For VPC, choose the VPC you want to use for this endpoint.
6. For Configure route tables, select the route tables to be used by the endpoint. The VPC service automatically adds a route to each route table you select that points any S3 traffic to the new endpoint.
7. For Policy, choose Full Access to allow full access to the S3 service by any user or service within the VPC. Choose Custom to restrict access further. For information, see Use a Custom Endpoint Policy to Restrict Access to S3.

Use a Custom Endpoint Policy to Restrict Access to S3

The default endpoint policy allows full access to S3 for any user or service in your VPC. To further restrict access to S3, create a custom endpoint policy. For more information, see Using Endpoint Policies for Amazon S3. You can also use a bucket policy to restrict access to your S3 buckets to
only traffic that comes from your Amazon VPC. For information, see Using Amazon S3 Bucket Policies.

**Restrict Package Installation on the Training Container**

The default endpoint policy allows users to install packages from the Amazon Linux and Amazon Linux 2 repositories on the training container. If you don't want users to install packages from that repository, create a custom endpoint policy that explicitly denies access to the Amazon Linux and Amazon Linux 2 repositories. The following is an example of a policy that denies access to these repositories:

```json
{
   "Statement": [
       {
           "Sid": "AmazonLinuxAMIRepositoryAccess",
           "Principal": "*",
           "Action": ["s3:GetObject"],
           "Effect": "Deny",
           "Resource": [
               "arn:aws:s3:::packages.*.amazonaws.com/*",
               "arn:aws:s3:::repo.*.amazonaws.com/*"
           ]
       }
   ]
}

{
   "Statement": [
       {
           "Sid": "AmazonLinux2AMIRepositoryAccess",
           "Principal": "*",
           "Action": ["s3:GetObject"],
           "Effect": "Deny",
           "Resource": ["arn:aws:s3:::amazonlinux.*.amazonaws.com/*"]
       }
   ]
}
```
Configure Route Tables

Use default DNS settings for your endpoint route table, so that standard Amazon S3 URLs (for example, http://s3-aws-region.amazonaws.com/MyBucket) resolve. If you don't use default DNS settings, ensure that the URLs that you use to specify the locations of the data in your training jobs resolve by configuring the endpoint route tables. For information about VPC endpoint route tables, see Routing for Gateway Endpoints in the Amazon VPC User Guide.

Configure the VPC Security Group

In distributed training, you must allow communication between the different containers in the same training job. To do that, configure a rule for your security group that allows inbound connections between members of the same security group. For EFA-enabled instances, ensure that both inbound and outbound connections allow all traffic from the same security group. For information, see Security Group Rules in the Amazon Virtual Private Cloud User Guide.

Connect to Resources Outside Your VPC

If you configure your VPC so that it doesn't have internet access, training jobs that use that VPC do not have access to resources outside your VPC. If your training job needs access to resources outside your VPC, provide access with one of the following options:

- If your training job needs access to an AWS service that supports interface VPC endpoints, create an endpoint to connect to that service. For a list of services that support interface endpoints, see VPC Endpoints in the Amazon Virtual Private Cloud User Guide. For information about creating an interface VPC endpoint, see Interface VPC Endpoints (AWS PrivateLink) in the Amazon Virtual Private Cloud User Guide.

  - If your training job needs access to an AWS service that doesn't support interface VPC endpoints or to a resource outside of AWS, create a NAT gateway and configure your security groups to allow outbound connections. For information about setting up a NAT gateway for your VPC, see Scenario 2: VPC with Public and Private Subnets (NAT) in the Amazon Virtual Private Cloud User Guide.

Monitor Amazon SageMaker Training Jobs with CloudWatch Logs and Metrics

Amazon SageMaker provides Amazon CloudWatch logs and metrics to monitor training jobs. CloudWatch provides CPU, GPU, memory, GPU memory, and disk metrics, and event logging. For more information about monitoring Amazon SageMaker training jobs, see Monitor Amazon SageMaker with Amazon CloudWatch and SageMaker Jobs and Endpoint Metrics.
Give SageMaker Hosted Endpoints Access to Resources in Your Amazon VPC

Configure a Model for Amazon VPC Access

To specify subnets and security groups in your private VPC, use the VpcConfig request parameter of the CreateModel API, or provide this information when you create a model in the SageMaker console. SageMaker uses this information to create network interfaces and attach them to your model containers. The network interfaces provide your model containers with a network connection within your VPC that is not connected to the internet. They also enable your model to connect to resources in your private VPC.

⚠️ Note

You must create at least two subnets in different availability zones in your private VPC, even if you have only one hosting instance.

The following is an example of the VpcConfig parameter that you include in your call to CreateModel:

```json
VpcConfig: {
  "Subnets": [
    "subnet-0123456789abcdef0",
    "subnet-0123456789abcdef1",
    "subnet-0123456789abcdef2"
  ],
  "SecurityGroupIds": [
    "sg-0123456789abcdef0"
  ]
}
```

Configure Your Private VPC for SageMaker Hosting

When configuring the private VPC for your SageMaker models, use the following guidelines. For information about setting up a VPC, see Working with VPCs and Subnets in the Amazon VPC User Guide.

Topics

- Ensure That Subnets Have Enough IP Addresses
• Create an Amazon S3 VPC Endpoint
• Use a Custom Endpoint Policy to Restrict Access to Amazon S3
• Add Permissions for Endpoint Access for Containers Running in a VPC to Custom IAM Policies
• Configure Route Tables
• Connect to Resources Outside Your VPC

Ensure That Subnets Have Enough IP Addresses

Training instances that don't use an Elastic Fabric Adapter (EFA) should have at least 2 private IP addresses. Training instances that use an EFA should have at least 5 private IP addresses. For more information, see Multiple IP addresses in the Amazon EC2 User Guide.

Create an Amazon S3 VPC Endpoint

If you configure your VPC so that model containers don't have access to the internet, they can't connect to the Amazon S3 buckets that contain your data unless you create a VPC endpoint that allows access. By creating a VPC endpoint, you allow your model containers to access the buckets where you store your data and model artifacts. We recommend that you also create a custom policy that allows only requests from your private VPC to access to your S3 buckets. For more information, see Endpoints for Amazon S3.

To create an Amazon S3 VPC endpoint:

1. Open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
2. In the navigation pane, choose Endpoints, then choose Create Endpoint
3. For Service Name, choose com.amazonaws.region.s3, where region is the name of the AWS Region where your VPC resides.
4. For VPC, choose the VPC that you want to use for this endpoint.
5. For Configure route tables, choose the route tables for the endpoint to use. The VPC service automatically adds a route to each route table that you choose that points Amazon S3 traffic to the new endpoint.
6. For Policy, choose Full Access to allow full access to the Amazon S3 service by any user or service within the VPC. To restrict access further, choose Custom. For more information, see Use a Custom Endpoint Policy to Restrict Access to Amazon S3.
Use a Custom Endpoint Policy to Restrict Access to Amazon S3

The default endpoint policy allows full access to Amazon Simple Storage Service (Amazon S3) for any user or service in your VPC. To further restrict access to Amazon S3, create a custom endpoint policy. For more information, see Using Endpoint Policies for Amazon S3.

You can also use a bucket policy to restrict access to your S3 buckets to only traffic that comes from your Amazon VPC. For information, see Using Amazon S3 Bucket Policies.

Restrict Package Installation on the Model Container with a Custom Endpoint Policy

The default endpoint policy allows users to install packages from the Amazon Linux and Amazon Linux 2 repositories on the model container. If you don't want users to install packages from those repositories, create a custom endpoint policy that explicitly denies access to the Amazon Linux and Amazon Linux 2 repositories. The following is an example of a policy that denies access to these repositories:

```
{
    "Statement": [

        {
            "Sid": "AmazonLinuxAMIRepositoryAccess",
            "Principal": "*",
            "Action": [
                "s3:GetObject"
            ],
            "Effect": "Deny",
            "Resource": [
                "arn:aws:s3:::packages.*.amazonaws.com/*",
                "arn:aws:s3:::repo.*.amazonaws.com/*"
            ]
        }
    ]
}
{
    "Statement": [
        {
            "Sid": "AmazonLinux2AMIRepositoryAccess",
            "Principal": "*",
            "Action": [
                "s3:GetObject"
            ],
            "Effect": "Deny",
            "Resource": [
                "arn:aws:s3:::packages.*.amazonaws.com/*",
                "arn:aws:s3:::repo.*.amazonaws.com/*"
            ]
        }
    ]
}
```
Add Permissions for Endpoint Access for Containers Running in a VPC to Custom IAM Policies

The SageMakerFullAccess managed policy includes the permissions that you need to use models configured for Amazon VPC access with an endpoint. These permissions allow SageMaker to create an elastic network interface and attach it to model containers running in a VPC. If you use your own IAM policy, you must add the following permissions to that policy to use models configured for VPC access.

```
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Allow",
            "Action": [
                "ec2:DescribeVpcEndpoints",
                "ec2:DescribeDhcpOptions",
                "ec2:DescribeVpcs",
                "ec2:DescribeSubnets",
                "ec2:DescribeSecurityGroups",
                "ec2:DescribeNetworkInterfaces",
                "ec2:DeleteNetworkInterfacePermission",
                "ec2:DeleteNetworkInterface",
                "ec2:CreateNetworkInterfacePermission",
                "ec2:CreateNetworkInterface"
            ],
            "Resource": "*
        }
    ]
}
```

For more information about the SageMakerFullAccess managed policy, see AWS managed policy: AmazonSageMakerFullAccess.
Configure Route Tables

Use default DNS settings for your endpoint route table, so that standard Amazon S3 URLs (for example, http://s3-aws-region.amazonaws.com/MyBucket) resolve. If you don't use default DNS settings, ensure that the URLs that you use to specify the locations of the data in your models resolve by configuring the endpoint route tables. For information about VPC endpoint route tables, see Routing for Gateway Endpoints in the Amazon VPC User Guide.

Connect to Resources Outside Your VPC

If you configure your VPC so that it doesn't have internet access, models that use that VPC do not have access to resources outside your VPC. If your model needs access to resources outside your VPC, provide access with one of the following options:

- If your model needs access to an AWS service that supports interface VPC endpoints, create an endpoint to connect to that service. For a list of services that support interface endpoints, see VPC Endpoints in the Amazon VPC User Guide. For information about creating an interface VPC endpoint, see Interface VPC Endpoints (AWS PrivateLink) in the Amazon VPC User Guide.

- If your model needs access to an AWS service that doesn't support interface VPC endpoints or to a resource outside of AWS, create a NAT gateway and configure your security groups to allow outbound connections. For information about setting up a NAT gateway for your VPC, see Scenario 2: VPC with Public and Private Subnets (NAT) in the Amazon Virtual Private Cloud User Guide.

Give Batch Transform Jobs Access to Resources in Your Amazon VPC

To control access to your data and batch transform jobs, we recommend that you create a private Amazon VPC and configure it so that your jobs aren't accessible over the public internet. You specify your private VPC configuration when you create a model by specifying subnets and security groups. You then specify the same model when you create a batch transform job. When you specify the subnets and security groups, SageMaker creates elastic network interfaces that are associated with your security groups in one of the subnets. Network interfaces allow your model containers to connect to resources in your VPC. For information about network interfaces, see Elastic Network Interfaces in the Amazon VPC User Guide.

This document explains how to add Amazon VPC configurations for batch transform jobs.
Configure a Batch Transform Job for Amazon VPC Access

To specify subnets and security groups in your private VPC, use the VpcConfig request parameter of the CreateModel API, or provide this information when you create a model in the SageMaker console. Then specify the same model in the ModelName request parameter of the CreateTransformJob API, or in the Model name field when you create a transform job in the SageMaker console. SageMaker uses this information to create network interfaces and attach them to your model containers. The network interfaces provide your model containers with a network connection within your VPC that is not connected to the internet. They also enable your transform job to connect to resources in your private VPC.

The following is an example of the VpcConfig parameter that you include in your call to CreateModel:

```json
VpcConfig: {
    "Subnets": [
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2"
    ],
    "SecurityGroupIds": [
        "sg-0123456789abcdef0"
    ]
}
```

If you are creating a model using the CreateModel API operation, the IAM execution role that you use to create your model must include the permissions described in CreateModel API: Execution Role Permissions, including the following permissions required for a private VPC.

When creating a model in the console, if you select Create a new role in the Model Settings section, the AmazonSageMakerFullAccess policy used to create the role already contains these permissions. If you select Enter a custom IAM role ARN or Use existing role, the role ARN that you specify must have an execution policy attached with the following permissions.

```json
{
    "Effect": "Allow",
    "Action": [
        "ec2:CreateNetworkInterface",
        "ec2:CreateNetworkInterfacePermission",
        "ec2:DeleteNetworkInterface"
    ]
}
```
Configure Your Private VPC for SageMaker Batch Transform

When configuring the private VPC for your SageMaker batch transform jobs, use the following guidelines. For information about setting up a VPC, see Working with VPCs and Subnets in the Amazon VPC User Guide.

Topics

- Ensure That Subnets Have Enough IP Addresses
- Create an Amazon S3 VPC Endpoint
- Use a Custom Endpoint Policy to Restrict Access to S3
- Configure Route Tables
- Configure the VPC Security Group
- Connect to Resources Outside Your VPC

Ensure That Subnets Have Enough IP Addresses

Your VPC subnets should have at least two private IP addresses for each instance in a transform job. For more information, see VPC and Subnet Sizing for IPv4 in the Amazon VPC User Guide.

Create an Amazon S3 VPC Endpoint

If you configure your VPC so that model containers don't have access to the internet, they can't connect to the Amazon S3 buckets that contain your data unless you create a VPC endpoint that allows access. By creating a VPC endpoint, you allow your model containers to access the buckets where you store your data and model artifacts. We recommend that you also create a custom policy that allows only requests from your private VPC to access your S3 buckets. For more information, see Endpoints for Amazon S3.

To create an S3 VPC endpoint:

1. Open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
2. In the navigation pane, choose **Endpoints**, then choose **Create Endpoint**

3. For **Service Name**, choose `com.amazonaws.region.s3`, where `region` is the name of the region where your VPC resides.

4. For **VPC**, choose the VPC you want to use for this endpoint.

5. For **Configure route tables**, select the route tables to be used by the endpoint. The VPC service automatically adds a route to each route table you select that points any S3 traffic to the new endpoint.

6. For **Policy**, choose **Full Access** to allow full access to the S3 service by any user or service within the VPC. Choose **Custom** to restrict access further. For information, see [Use a Custom Endpoint Policy to Restrict Access to S3](#).

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### Use a Custom Endpoint Policy to Restrict Access to S3

The default endpoint policy allows full access to S3 for any user or service in your VPC. To further restrict access to S3, create a custom endpoint policy. For more information, see [Using Endpoint Policies for Amazon S3](#). You can also use a bucket policy to restrict access to your S3 buckets to only traffic that comes from your Amazon VPC. For information, see [Using Amazon S3 Bucket Policies](#).

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### Restrict Package Installation on the Model Container

The default endpoint policy allows users to install packages from the Amazon Linux and Amazon Linux 2 repositories on the training container. If you don't want users to install packages from that repository, create a custom endpoint policy that explicitly denies access to the Amazon Linux and Amazon Linux 2 repositories. The following is an example of a policy that denies access to these repositories:

```json
{
    "Statement": [
        {
            "Sid": "AmazonLinuxAMIRepositoryAccess",
            "Principal": "*",
            "Action": ["s3:GetObject"],
            "Effect": "Deny",
            "Resource": [
                "arn:aws:s3:::packages.*.amazonaws.com/*",
                "arn:aws:s3:::packages2.*.amazonaws.com/*",
                "arn:aws:s3:::packages3.*.amazonaws.com/*"
            ]
        }
    ]
}
```
Configure Route Tables

Use default DNS settings for your endpoint route table, so that standard Amazon S3 URLs (for example, http://s3-aws-region.amazonaws.com/MyBucket) resolve. If you don't use default DNS settings, ensure that the URLs that you use to specify the locations of the data in your batch transform jobs resolve by configuring the endpoint route tables. For information about VPC endpoint route tables, see Routing for Gateway Endpoints in the Amazon VPC User Guide.

Configure the VPC Security Group

In distributed batch transform, you must allow communication between the different containers in the same batch transform job. To do that, configure a rule for your security group that allows inbound and outbound connections between members of the same security group. Members of the same security group should be able to communicate with each other across all ports. For more information, see Security Group Rules.

Connect to Resources Outside Your VPC

If you configure your VPC so that it doesn't have internet access, batch transform jobs that use that VPC do not have access to resources outside your VPC. If your batch transform job needs access to resources outside your VPC, provide access with one of the following options:
• If your batch transform job needs access to an AWS service that supports interface VPC endpoints, create an endpoint to connect to that service. For a list of services that support interface endpoints, see VPC Endpoints in the Amazon VPC User Guide. For information about creating an interface VPC endpoint, see Interface VPC Endpoints (AWS PrivateLink) in the Amazon VPC User Guide.

• If your batch transform job needs access to an AWS service that doesn't support interface VPC endpoints or to a resource outside of AWS, create a NAT gateway and configure your security groups to allow outbound connections. For information about setting up a NAT gateway for your VPC, see Scenario 2: VPC with Public and Private Subnets (NAT) in the Amazon Virtual Private Cloud User Guide.

Give Amazon SageMaker Clarify Jobs Access to Resources in Your Amazon VPC

To control access to your data and SageMaker Clarify jobs, we recommend that you create a private Amazon VPC and configure it so that your jobs aren't accessible over the public internet. For information about creating and configuring an Amazon VPC for processing jobs, see Give SageMaker Processing Jobs Access to Resources in Your Amazon VPC.

This document explains how to add additional Amazon VPC configurations that meet the requirements for SageMaker Clarify jobs.

Topics

• Configure a SageMaker Clarify Job for Amazon VPC Access
• Configure Your Private Amazon VPC for SageMaker Clarify jobs

Configure a SageMaker Clarify Job for Amazon VPC Access

You need to specify subnets and security groups when configuring your private Amazon VPC for SageMaker Clarify jobs and to enable the job to get inferences from the SageMaker model when computing post-training bias metrics and feature contributions that help explain model predictions.

Topics

• SageMaker Clarify Job Amazon VPC Subnets and Security Groups
• Configure a Model Amazon VPC for Inference
SageMaker Clarify Job Amazon VPC Subnets and Security Groups

Subnets and security groups in your private Amazon VPC can be assigned to a SageMaker Clarify job in various ways, depending on how you create the job.

- **SageMaker console**: Provide this information when you create the job in the SageMaker Dashboard. From the Processing menu, choose Processing jobs, then choose Create processing job. Select the VPC option in the Network panel and provide the subnets and security groups using the drop-down lists. Make sure network isolation option provided in this panel is turned off.

- **SageMaker API**: Use the NetworkConfig.VpcConfig request parameter of the CreateProcessingJob API, as shown in the following example:

```
"NetworkConfig": {  
  "VpcConfig": {  
    "Subnets": [  
      "subnet-0123456789abcdef0",  
      "subnet-0123456789abcdef1",  
      "subnet-0123456789abcdef2"  
    ],  
    "SecurityGroupIds": [  
      "sg-0123456789abcdef0"  
    ]  
  }  
}
```

- **SageMaker Python SDK**: Use the NetworkConfig parameter of the SageMakerClarifyProcessor API or Processor API, as shown in the following example:

```python
from sagemaker.network import NetworkConfig
network_config = NetworkConfig(
    subnets=[
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2",
    ],
    security_group_ids=[
        "sg-0123456789abcdef0",
    ],
)
```
SageMaker uses the information to create network interfaces and attach them to the SageMaker Clarify job. The network interfaces provide a SageMaker Clarify job with a network connection within your Amazon VPC that is not connected to the public internet. They also enable the SageMaker Clarify job to connect to resources in your private Amazon VPC.

**Note**

The network isolation option of the SageMaker Clarify job must be turned off (by default the option is turned off) so that the SageMaker Clarify job can communicate with the shadow endpoint.

**Configure a Model Amazon VPC for Inference**

In order to compute post-training bias metrics and explainability, the SageMaker Clarify job needs to get inferences from the SageMaker model that is specified by the `model_name` parameter of the analysis configuration for the SageMaker Clarify processing job. Alternatively, if you use the SageMakerClarifyProcessor API in the SageMaker Python SDK, the job needs to get the `model_name` specified by the `ModelConfig` class. To accomplish this, the SageMaker Clarify job creates an ephemeral endpoint with the model, known as a *shadow endpoint*, and then applies the Amazon VPC configuration of the model to the shadow endpoint.

To specify subnets and security groups in your private Amazon VPC to the SageMaker model, use the `VpcConfig` request parameter of the `CreateModel` API or provide this information when you create the model using the SageMaker dashboard in the console. The following is an example of the `VpcConfig` parameter that you include in your call to `CreateModel`:

```
"VpcConfig": {
    "Subnets": [
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2"
    ],
    "SecurityGroupIds": [
        "sg-0123456789abcdef0"
    ]
}
```

You can specify the number of instances of the shadow endpoint to launch with the `initial_instance_count` parameter of the analysis configuration for the SageMaker Clarify
processing job. Alternatively, if you use the `SageMakerClarifyProcessor` API in the SageMaker Python SDK, the job needs to get the `instance_count` specified by the `ModelConfig` class.

**Note**

Even if you only request one instance when creating the shadow endpoint, you need at least two subnets in the model's `ModelConfig` in distinct availability zones. Otherwise the shadow endpoint creation fails with the following error:

```
ClientError: Error hosting endpoint sagemaker-clarify-endpoint-XXX: Failed. Reason: Unable to locate at least 2 availability zone(s) with the requested instance type YYY that overlap with SageMaker subnets.
```

If your model requires model files in Amazon S3, then the model Amazon VPC needs to have an Amazon S3 VPC endpoint. For more information about creating and configuring an Amazon VPC for SageMaker models, see [Give SageMaker Hosted Endpoints Access to Resources in Your Amazon VPC](#).

### Configure Your Private Amazon VPC for SageMaker Clarify jobs

In general, you can follow the steps in [Configure Your Private VPC for SageMaker Processing](#) to configure your private Amazon VPC for SageMaker Clarify jobs. Here are some highlights and special requirements for SageMaker Clarify jobs.

**Topics**

- [Connect to Resources Outside Your Amazon VPC](#)
- [Configure the Amazon VPC Security Group](#)

### Connect to Resources Outside Your Amazon VPC

If you configure your Amazon VPC so that it does not have public internet access, then some additional setup is required to grant SageMaker Clarify jobs access to resources and services outside of your Amazon VPC. For example, an Amazon S3 VPC endpoint is required because a SageMaker Clarify job needs to load a dataset from an S3 bucket as well as save the analysis results to an S3 bucket. For more information, see [Create an Amazon S3 VPC Endpoint](#) for the creation guide. In addition, if a SageMaker Clarify job needs to get inferences from the shadow endpoint, then it needs to call several more AWS services.
• **Create an Amazon SageMaker API service VPC endpoint:** The SageMaker Clarify job needs to call the Amazon SageMaker API service to manipulate the shadow endpoint, or to describe a SageMaker model for Amazon VPC validation. You can follow the guidance provided in the [Securing all Amazon SageMaker API calls with AWS PrivateLink](https://aws.amazon.com/blogs/opensource/securing-all-amazon-sagemaker-api-calls-with-aws-privatelink) blog to create an Amazon SageMaker API VPC endpoint that allows the SageMaker Clarify job to make the service calls. Note that the service name of Amazon SageMaker API service is `com.amazonaws.region.sagemaker.api`, where `region` is the name of the Region where your Amazon VPC resides.

• **Create an Amazon SageMaker Runtime VPC Endpoint:** The SageMaker Clarify job needs to call the Amazon SageMaker runtime service, which routes the invocations to the shadow endpoint. The setup steps are similar to those for the Amazon SageMaker API service. Note that the service name of Amazon SageMaker Runtime service is `com.amazonaws.region.sagemaker.runtime`, where `region` is the name of the Region where your Amazon VPC resides.

### Configure the Amazon VPC Security Group

SageMaker Clarify jobs support distributed processing when two or more processing instances are specified in one of the following ways:

• **SageMaker console:** The **Instance count** is specified in the **Resource configuration** part of the **Job settings** panel on the **Create processing job** page.

• **SageMaker API:** The **InstanceCount** is specified when you create the job with the [CreateProcessingJob](https://docs.aws.amazon.com/sagemaker/latest/dg/sageMaker-console-create-processing-job.html) API.

• **SageMaker Python SDK:** The **instance_count** is specified when using the [SageMakerClarifyProcessor](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-python-sdk.html) API or the [Processor](https://docs.aws.amazon.com/sagemaker/latest/dg/sagemaker-python-sdk.html) API.

In distributed processing, you must allow communication between the different instances in the same processing job. To do that, configure a rule for your security group that allows inbound connections between members of the same security group. For information, see [Security group rules](https://docs.aws.amazon.com/elasticloadbalancing/latest/LoadBalancer/creating-security-group.html).
Give SageMaker Compilation Jobs Access to Resources in Your Amazon VPC

Note
For compilation jobs, you can configure only subnets with a default tenancy VPC in which your job runs on shared hardware. For more information on the tenancy attribute for VPCs, see Dedicated Instances.

Configure a Compilation Job for Amazon VPC Access

To specify subnets and security groups in your private VPC, use the VpcConfig request parameter of the CreateCompilationJob API, or provide this information when you create a compilation job in the SageMaker console. SageMaker Neo uses this information to create network interfaces and attach them to your compilation jobs. The network interfaces provide compilation jobs with a network connection within your VPC that is not connected to the internet. They also enable your compilation job to connect to resources in your private VPC. The following is an example of the VpcConfig parameter that you include in your call to CreateCompilationJob:

```
VpcConfig: {
    "Subnets": [
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2"
    ],
    "SecurityGroupIds": [
        "sg-0123456789abcdef0"
    ]
}
```

Configure Your Private VPC for SageMaker Compilation

When configuring the private VPC for your SageMaker compilation jobs, use the following guidelines. For information about setting up a VPC, see Working with VPCs and Subnets in the Amazon VPC User Guide.

Topics
- Ensure That Subnets Have Enough IP Addresses
- Create an Amazon S3 VPC Endpoint
- Use a Custom Endpoint Policy to Restrict Access to S3
• Configure Route Tables
• Configure the VPC Security Group

Ensure That Subnets Have Enough IP Addresses

Your VPC subnets should have at least two private IP addresses for each instance in a compilation job. For more information, see VPC and Subnet Sizing for IPv4 in the Amazon VPC User Guide.

Create an Amazon S3 VPC Endpoint

If you configure your VPC to block access to the internet, SageMaker Neo can't connect to the Amazon S3 buckets that contain your models unless you create a VPC endpoint that allows access. By creating a VPC endpoint, you allow your SageMaker Neo compilation jobs to access the buckets where you store your data and model artifacts. We recommend that you also create a custom policy that allows only requests from your private VPC to access to your S3 buckets. For more information, see Endpoints for Amazon S3.

To create an S3 VPC endpoint:

1. Open the Amazon VPC console at https://console.aws.amazon.com/vpc/.
2. In the navigation pane, choose Endpoints, then choose Create Endpoint.
3. For Service Name, search for com.amazonaws.region.s3, where region is the name of the region where your VPC resides.
4. Choose the Gateway type.
5. For VPC, choose the VPC you want to use for this endpoint.
6. For Configure route tables, select the route tables to be used by the endpoint. The VPC service automatically adds a route to each route table you select that points any S3 traffic to the new endpoint.
7. For Policy, choose Full Access to allow full access to the S3 service by any user or service within the VPC. Choose Custom to restrict access further. For information, see Use a Custom Endpoint Policy to Restrict Access to S3.

Use a Custom Endpoint Policy to Restrict Access to S3

The default endpoint policy allows full access to S3 for any user or service in your VPC. To further restrict access to S3, create a custom endpoint policy. For more information, see Using Endpoint.
Policies for Amazon S3. You can also use a bucket policy to restrict access to your S3 buckets to only traffic that comes from your Amazon VPC. For information, see Using Amazon S3 Bucket Policies. The following is a sample customized policy:

```json
{
    "Version": "2012-10-17",
    "Statement": [
        {
            "Effect": "Deny",
            "Principal": {
                "AWS": "*"
            },
            "Action": "s3:GetObject",
            "Resource": [
                "arn:aws:s3:::your-sample-bucket",
                "arn:aws:s3:::your-sample-bucket/*"
            ],
            "Condition": {
                "StringNotEquals": {
                    "aws:SourceVpce": [
                        "vpce-01234567890123456"
                    ]
                }
            }
        }
    ]
}
```

Add Permissions for Compilation Job Running in a Amazon VPC to Custom IAM Policies

The SageMakerFullAccess managed policy includes the permissions that you need to use models configured for Amazon VPC access with an endpoint. These permissions allow SageMaker Neo to create an elastic network interface and attach it to compilation job running in a Amazon VPC. If you use your own IAM policy, you must add the following permissions to that policy to use models configured for Amazon VPC access.

```json
{"Version": "2012-10-17",
 "Statement": [
     {
         "Effect": "Allow",
         "Action": [
             "ec2:DescribeVpcEndpoints",
             "ec2:DescribeDhcpOptions",
             "ec2:RunInstance",
             "ec2:AuthorizeSecurityGroupIngress",
             "ec2:CreateInternetGateway",
             "ec2:CreateNetworkAcl",
             "ec2:CreateNetworkAclEntry",
             "ec2:CreateSubnet",
             "ec2:CreateVpc",
             "ec2:DeleteNetworkAcl",
             "ec2:DeleteNetworkAclEntry",
             "ec2:DeleteRouteTable",
             "ec2:DeleteSubnet",
             "ec2:DeleteVpc",
             "ec2:DescribeAccountAttributes",
             "ec2:DescribeAvailabilityZones",
             "ec2:DescribeInternetGateways",
             "ec2:DescribeNetworkAclEntries",
             "ec2:DescribeNetworkAcls",
             "ec2:DescribeRouteTables",
             "ec2:DescribeSubnets",
             "ec2:DescribeVpcs",
             "ec2:DescribeVpcPeeringConnections",
             "ec2:DisassociateAddress",
             "ec2:ModifyInstanceAttribute",
             "ec2:ModifyNetworkAclEntry",
             "ec2:ModifyNetworkAcl",
             "ec2:ModifySecurityGroupIngress",
             "ec2:ModifySubnetAttribute",
             "ec2:ModifyVpcAttribute",
             "ec2:ModifyVpcPeeringConnectionAttribute",
             "ec2:StartInstances",
             "ec2:StopInstances",
             "ec2:TerminateInstances"
         ]
     }
 ]
}
```
Configure Route Tables

Use default DNS settings for your endpoint route table, so that standard Amazon S3 URLs (for example, http://s3-aws-region.amazonaws.com/MyBucket) resolve. If you don't use default DNS settings, ensure that the URLs that you use to specify the locations of the data in your compilation jobs resolve by configuring the endpoint route tables. For information about VPC endpoint route tables, see Routing for Gateway Endpoints in the Amazon VPC User Guide.

Configure the VPC Security Group

In your security group for the compilation job, you must allow outbound communication to your Amazon S3 Amazon VPC endpoints and the subnet CIDR ranges used for the compilation job. For information, see Security Group Rules and Control access to services with Amazon VPC endpoints.

Give Inference Recommender Jobs Access to Resources in Your Amazon VPC

Note

Inference Recommender requires you to register your model with Model Registry. Note that Model Registry doesn't allow your model artifacts or Amazon ECR image to be VPC restricted.

Inference Recommender also has a requirement that your sample payload Amazon S3 object is not VPC restricted. For inference recommendation jobs, you can't create a custom
To specify subnets and security groups in your private VPC, use the RecommendationJobVpcConfig request parameter of the CreateInferenceRecommendationsJob API, or specify your subnets and security groups when you create a recommendation job in the SageMaker console.

Inference Recommender uses this information to create endpoints. When provisioning endpoints, SageMaker creates network interfaces and attaches them to your endpoints. The network interfaces provide your endpoints with a network connection to your VPC. The following is an example of the VpcConfig parameter that you include in a call to CreateInferenceRecommendationsJob:

```json
VpcConfig: {
    "Subnets": [
        "subnet-0123456789abcdef0",
        "subnet-0123456789abcdef1",
        "subnet-0123456789abcdef2"
    ],
    "SecurityGroupIds": [
        "sg-0123456789abcdef0"
    ]
}
```

Refer to the following topics for more information on configuring your Amazon VPC for use with Inference Recommender jobs.

**Topics**

- [Ensure that subnets have enough IP addresses](#)
- [Create an Amazon S3 VPC endpoint](#)
- [Add permissions for Inference Recommender jobs running in an Amazon VPC to custom IAM policies](#)
- [Configure route tables](#)
- [Configure the VPC security group](#)
Ensure that subnets have enough IP addresses

Your VPC subnets should have at least two private IP addresses for each instance in an inference recommendation job. For more information about subnets and private IP addresses, see How Amazon VPC works in the Amazon VPC User Guide.

Create an Amazon S3 VPC endpoint

If you configure your VPC to block access to the internet, Inference Recommender can't connect to the Amazon S3 buckets that contain your models unless you create a VPC endpoint that allows access. By creating a VPC endpoint, you allow your SageMaker inference recommendation jobs to access the buckets where you store your data and model artifacts.

To create an Amazon S3 VPC endpoint, use the following procedure:

1. Open the Amazon VPC console.
2. In the navigation pane, choose Endpoints, and then choose Create Endpoint.
3. For Service Name, search for com.amazonaws.region.s3, where region is the name of the Region where your VPC resides.
4. Choose the Gateway type.
5. For VPC, choose the VPC you want to use for this endpoint.
6. For Configure route tables, select the route tables to be used by the endpoint. The VPC service automatically adds a route to each route table you select that points any Amazon S3 traffic to the new endpoint.
7. For Policy, choose Full Access to allow full access to the Amazon S3 service by any user or service within the VPC.

Add permissions for Inference Recommender jobs running in an Amazon VPC to custom IAM policies

The AmazonSageMakerFullAccess managed policy includes the permissions that you need to use models configured for Amazon VPC access with an endpoint. These permissions allow Inference Recommender to create an elastic network interface and attach it to the inference recommendation job running in an Amazon VPC. If you use your own IAM policy, you must add the following permissions to that policy to use models configured for Amazon VPC access.

```json
{
    "Version": "2012-10-17",
```
Configure route tables

Use the default DNS settings for your endpoint route table, so that standard Amazon S3 URLs (for example: http://s3-aws-region.amazonaws.com/MyBucket) resolve. If you don't use the default DNS settings, ensure that the URLs that you use to specify the locations of the data in your inference recommendation jobs resolve by configuring the endpoint route tables. For information about VPC endpoint route tables, see Routing gateway endpoints in the Amazon VPC User Guide.

Configure the VPC security group

In your security group for the inference recommendation job, you must allow outbound communication to your Amazon S3 VPC endpoints and the subnet CIDR ranges used for the inference recommendation job. For information, see Security Group Rules and Control access to services with Amazon VPC endpoints in the Amazon VPC User Guide.
Sell algorithms and packages in the AWS Marketplace

Amazon SageMaker integrates with AWS Marketplace, enabling developers to charge other SageMaker users for the use of their algorithms and model packages. AWS Marketplace is a curated digital catalog that makes it easy for customers to find, buy, deploy, and manage third-party software and services that customers need to build solutions and run their businesses. AWS Marketplace includes thousands of software listings in popular categories, such as security, networking, storage, machine learning, business intelligence, database, and DevOps. It simplifies software licensing and procurement with flexible pricing options and multiple deployment methods.

For information, see AWS Marketplace Documentation.

Topics

- SageMaker Algorithms
- SageMaker Model Packages
- Sell Amazon SageMaker Algorithms and Model Packages
- Find and Subscribe to Algorithms and Model Packages on AWS Marketplace
- Use Algorithm and Model Package Resources

SageMaker Algorithms

An algorithm enables you to perform end-to-end machine learning. It has two logical components: training and inference. Buyers can use the training component to create training jobs in SageMaker and build a machine learning model. SageMaker saves the model artifacts generated by the algorithm during training to an Amazon S3 bucket. For more information, see Train a Model with Amazon SageMaker.

Buyers use the inference component with the model artifacts generated during a training job to create a deployable model in their SageMaker account. They can use the deployable model for real-time inference by using SageMaker hosting services. Or, they can get inferences for an entire dataset by running batch transform jobs. For more information, see Deploy a Model in Amazon SageMaker.
SageMaker Model Packages

Buyers use a model package to build a deployable model in SageMaker. They can use the deployable model for real-time inference by using SageMaker hosting services. Or, they can get inferences for an entire dataset by running batch transform jobs. For more information, see Deploy a Model in Amazon SageMaker. As a seller, you can build your model artifacts by training in SageMaker, or you can use your own model artifacts from a model that you trained outside of SageMaker. You can charge buyers for inference.

Use your own algorithms and models with the AWS Marketplace

The following sections show how to create algorithm and model package resources that you can use locally and publish to the AWS Marketplace.

Topics

- Create Algorithm and Model Package Resources
- Use Algorithm and Model Package Resources

Create Algorithm and Model Package Resources

After your training and/or inference code is packaged in Docker containers, create algorithm and model package resources that you can use in your Amazon SageMaker account and, optionally, publish on AWS Marketplace.

Topics

- Create an Algorithm Resource
- Create a Model Package Resource

Create an Algorithm Resource

To create an algorithm resource that you can use to run training jobs in Amazon SageMaker and publish on AWS Marketplace specify the following information:

- The Docker containers that contains the training and, optionally, inference code.
- The configuration of the input data that your algorithm expects for training.
- The hyperparameters that your algorithm supports.
- Metrics that your algorithm sends to Amazon CloudWatch during training jobs.
- The instance types that your algorithm supports for training and inference, and whether it supports distributed training across multiple instances.
- Validation profiles, which are training jobs that SageMaker uses to test your algorithm's training code and batch transform jobs that SageMaker runs to test your algorithm's inference code.

To ensure that buyers and sellers can be confident that products work in SageMaker, we require that you validate your algorithms before listing them on AWS Marketplace. You can list products in the AWS Marketplace only if validation succeeds. To validate your algorithms, SageMaker uses your validation profile and sample data to run the following validations tasks:

1. Create a training job in your account to verify that your training image works with SageMaker.
2. If you included inference code in your algorithm, create a model in your account using the algorithm's inference image and the model artifacts produced by the training job.
3. If you included inference code in your algorithm, create a transform job in your account using the model to verify that your inference image works with SageMaker.

When you list your product on AWS Marketplace, the inputs and outputs of this validation process persist as part of your product and are made available to your buyers. This helps buyers understand and evaluate the product before they buy it. For example, buyers can inspect the input data that you used, the outputs generated, and the logs and metrics emitted by your code. The more comprehensive your validation specification, the easier it is for customers to evaluate your product.

**Note**

In your validation profile, provide only data that you want to expose publicly.

Validation can take up to a few hours. To see the status of the jobs in your account, in the SageMaker console, see the **Training jobs** and **Transform jobs** pages. If validation fails, you can access the scan and validation reports from the SageMaker console. If any issues are found, you will have to create the algorithm again.
You can create an algorithm by using either the SageMaker console or the SageMaker API.

Topics

- **Create an Algorithm Resource (Console)**
- **Create an Algorithm Resource (API)**

Create an Algorithm Resource (Console)

To create an algorithm resource (console)

2. From the left menu, choose **Training**.
3. From the dropdown menu, choose **Algorithms**, then choose **Create algorithm**.
4. On the **Training specifications** page, provide the following information:
   a. For **Algorithm name**, type a name for your algorithm. The algorithm name must be unique in your account and in the AWS region. The name must have 1 to 64 characters. Valid characters are a-z, A-Z, 0-9, and - (hyphen).
   b. Type a description for your algorithm. This description appears in the SageMaker console and in the AWS Marketplace.
   c. For **Training image**, type the path in Amazon ECR where your training container is stored.
   d. For **Support distributed training**, Choose Yes if your algorithm supports training on multiple instances. Otherwise, choose No.
   e. For **Support instance types for training**, choose the instance types that your algorithm supports.
   f. For **Channel specification**, specify up to 8 channels of input data for your algorithm. For example, you might specify 3 input channels named train, validation, and test. For each channel, specify the following information:
1. For **Channel name**, type a name for the channel. The name must have 1 to 64 characters. Valid characters are a-z, A-Z, 0-9, and - (hyphen).

2. To require the channel for your algorithm, choose **Channel required**.

3. Type a description for the channel.

4. For **Supported input modes**, choose **Pipe mode** if your algorithm supports streaming the input data, and **File mode** if your algorithm supports downloading the input data as a file. You can choose both.

5. For **Supported content types**, type the MIME type that your algorithm expects for input data.

6. For **Supported compression type**, choose **Gzip** if your algorithm supports Gzip compression. Otherwise, choose **None**.

7. Choose **Add channel** to add another data input channel, or choose **Next** if you are done adding channels.

5. On the **Tuning specifications** page, provide the following information:

   a. For **Hyperparameter specification**, specify the hyperparameters that your algorithm supports by editing the JSON object. For each hyperparameter that your algorithm supports, construct a JSON block similar to the following:

   ```json
   {
   "DefaultValue": "5",
   "Description": "The first hyperparameter",
   "IsRequired": true,
   "IsTunable": false,
   "Name": "intRange",
   "Range": {
   "IntegerParameterRangeSpecification": {
   "MaxValue": "10",
   "MinValue": "1"
   },
   "Type": "Integer"
   }
   }
   
   In the JSON, supply the following:

   i. For **DefaultValue**, specify a default value for the hyperparameter, if there is one.

   ii. For **Description**, specify a description for the hyperparameter.
iii. For **IsRequired**, specify whether the hyperparameter is required.

iv. For **IsTunable**, specify `true` if this hyperparameter can be tuned when a user runs a hyperparameter tuning job that uses this algorithm. For information, see [Perform Automatic Model Tuning with SageMaker](#).

v. For **Name**, specify a name for the hyperparameter.

vi. For **Range**, specify one of the following:

   - **IntegerParameterRangeSpecification** - the values of the hyperparameter are integers. Specify minimum and maximum values for the hyperparameter.

   - **ContinuousParameterRangeSpecification** - the values of the hyperparameter are floating-point values. Specify minimum and maximum values for the hyperparameter.

   - **CategoricalParameterRangeSpecification** - the values of the hyperparameter are categorical values. Specify a list of all of the possible values.

vii. For **Type**, specify **Integer**, **Continuous**, or **Categorical**. The value must correspond to the type of **Range** that you specified.

b. For **Metric definitions**, specify any training metrics that you want your algorithm to emit. SageMaker uses the regular expression that you specify to find the metrics by parsing the logs from your training container during training. Users can view these metrics when they run training jobs with your algorithm, and they can monitor and plot the metrics in Amazon CloudWatch. For information, see [Monitor and Analyze Training Jobs Using Amazon CloudWatch Metrics](#). For each metric, provide the following information:

i. For **Metric name**, type a name for the metric.

ii. For **Regex**, type the regular expression that SageMaker uses to parse training logs so that it can find the metric value.

iii. For **Objective metric support** choose **Yes** if this metric can be used as the objective metric for a hyperparameter tuning job. For information, see [Perform Automatic Model Tuning with SageMaker](#).

iv. Choose **Add metric** to add another metric, or choose **Next** if you are done adding metrics.

6. On the **Inference specifications** page, provide the following information if your algorithm supports inference:
a. For **Location of inference image**, type the path in Amazon ECR where your inference container is stored.

b. For **Container DNS host name**, type the name of a DNS host for your image.

c. For **Supported instance types for real-time inference**, choose the instance types that your algorithm supports for models deployed as hosted endpoints in SageMaker. For information, see [Deploy models for inference](#).

d. For **Supported instance types for batch transform jobs**, choose the instance types that your algorithm supports for batch transform jobs. For information, see [Use Batch Transform](#).

e. For **Supported content types**, type the type of input data that your algorithm expects for inference requests.

f. For **Supported response MIME types**, type the MIME types that your algorithm supports for inference responses.

g. Choose **Next**.

7. On the **Validation specifications** page, provide the following information:

   a. For **Publish this algorithm on AWS Marketplace**, choose **Yes** to publish the algorithm on AWS Marketplace.

   b. For **Validate this resource**, choose **Yes** if you want SageMaker to run training jobs and/or batch transform jobs that you specify to test the training and/or inference code of your algorithm.

   c. For **IAM role**, choose an IAM role that has the required permissions to run training jobs and batch transform jobs in SageMaker, or choose **Create a new role** to allow SageMaker to create a role that has the AmazonSageMakerFullAccess managed policy attached. For information, see [SageMaker Roles](#).

   d. For **Validation profile**, specify the following:

      - A name for the validation profile.
• A **Training job definition**. This is a JSON block that describes a training job. This is in the same format as the `TrainingJobDefinition` input parameter of the `CreateAlgorithm` API.

• A **Transform job definition**. This is a JSON block that describes a batch transform job. This is in the same format as the `TransformJobDefinition` input parameter of the `CreateAlgorithm` API.

e. Choose **Create algorithm**.

### Create an Algorithm Resource (API)

To create an algorithm resource by using the SageMaker API, call the `CreateAlgorithm` API.

### Create a Model Package Resource

To create a model package resource that you can use to create deployable models in Amazon SageMaker and publish on AWS Marketplace specify the following information:

- The Docker container that contains the inference code, or the algorithm resource that was used to train the model.
- The location of the model artifacts. Model artifacts can either be packaged in the same Docker container as the inference code or stored in Amazon S3.
- The instance types that your model package supports for both real-time inference and batch transform jobs.
- Validation profiles, which are batch transform jobs that SageMaker runs to test your model package's inference code.

Before listing model packages on AWS Marketplace, you must validate them. This ensures that buyers and sellers can be confident that products work in Amazon SageMaker. You can list products on AWS Marketplace only if validation succeeds.

The validation procedure uses your validation profile and sample data to run the following validations tasks:

1. Create a model in your account using the model package's inference image and the optional model artifacts that are stored in Amazon S3.
A model package is specific to the region in which you create it. The S3 bucket where the model artifacts are stored must be in the same region where your created the model package.

2. Create a transform job in your account using the model to verify that your inference image works with SageMaker.

3. Create a validation profile.

Note
In your validation profile, provide only data that you want to expose publicly.

Validation can take up to a few hours. To see the status of the jobs in your account, in the SageMaker console, see the Transform jobs pages. If validation fails, you can access the scan and validation reports from the SageMaker console. After fixing issues, recreate the algorithm. When the status of the algorithm is COMPLETED, find it in the SageMaker console and start the listing process.

Note
To publish your model package on AWS Marketplace, at least one validation profile is required.

You can create a model package either by using the SageMaker console or by using the SageMaker API.

Topics
- Create a Model Package Resource (Console)
- Create a Model Package Resource (API)
Create a Model Package Resource (Console)

To create a model package in the SageMaker console:

2. From the left menu, choose Inference.
3. Choose Marketplace model packages, then choose Create marketplace model package.
4. On the Inference specifications page, provide the following information:
   a. For Model package name, type a name for your model package. The model package name must be unique in your account and in the AWS region. The name must have 1 to 64 characters. Valid characters are a-z, A-Z, 0-9, and - (hyphen).
   b. Type a description for your model package. This description appears in the SageMaker console and in the AWS Marketplace.
   c. For Inference specification options, choose Provide the location of the inference image and model artifacts to create a model package by using an inference container and model artifacts. Choose Provide the algorithm used for training and its model artifacts to create a model package from an algorithm resource that you created or subscribe to from AWS Marketplace.
   d. If you chose Provide the location of the inference image and model artifacts for Inference specification options, provide the following information for Container definition and Supported resources:
      i. For Location of inference image, type the path to the image that contains your inference code. The image must be stored as a Docker container in Amazon ECR.
      ii. For Location of model data artifacts, type the location in S3 where your model artifacts are stored.
      iii. For Container DNS host name, type the name of the DNS host to use for your container.
      iv. For Supported instance types for real-time inference, choose the instance types that your model package supports for real-time inference from SageMaker hosted endpoints.
      v. For Supported instance types for batch transform jobs, choose the instance types that your model package supports for batch transform jobs.
      vi. Supported content types, type the content types that your model package expects for inference requests.
vii. For **Supported response MIME types**, type the MIME types that your model package uses to provide inferences.

e. If you chose **Provide the algorithm used for training and its model artifacts** for **Inference specification options**, provide the following information:

i. For **Algorithm ARN**, type the Amazon Resource Name (ARN) of the algorithm resource to use to create the model package.

ii. For **Location of model data artifacts**, type the location in S3 where your model artifacts are stored.

f. Choose **Next**.

5. On the **Validation and scanning** page, provide the following information:

a. For **Publish this model package on AWS Marketplace**, choose **Yes** to publish the model package on AWS Marketplace.

b. For **Validate this resource**, choose **Yes** if you want SageMaker to run batch transform jobs that you specify to test the inference code of your model package.

> **Note**

To publish your model package on AWS Marketplace, your model package must be validated.

c. For **IAM role**, choose an IAM role that has the required permissions to run batch transform jobs in SageMaker, or choose **Create a new role** to allow SageMaker to create a role that has the AmazonSageMakerFullAccess managed policy attached. For information, see [SageMaker Roles](#).

d. For **Validation profile**, specify the following:

- A name for the validation profile.

- A **Transform job definition**. This is a JSON block that describes a batch transform job. This is in the same format as the **TransformJobDefinition** input parameter of the **CreateAlgorithm** API.

6. Choose **Create marketplace model package**.
Create a Model Package Resource (API)

To create a model package by using the SageMaker API, call the `CreateModelPackage` API.

Use Algorithm and Model Package Resources

You can create algorithms and model packages as resources in your Amazon SageMaker account, and you can find and subscribe to algorithms and model packages on AWS Marketplace.

Use algorithms to:

- Run training jobs. For information, see [Use an Algorithm to Run a Training Job](#).
- Run hyperparameter tuning jobs. For information, see [Use an Algorithm to Run a Hyperparameter Tuning Job](#).
- Create model packages. After you use an algorithm resource to run a training job or a hyperparameter tuning job, you can use the model artifacts that these jobs output along with the algorithm to create a model package. For information, see [Create a Model Package Resource](#).

⚠️ Note

If you subscribe to an algorithm on AWS Marketplace, you must create a model package before you can use it to get inferences by creating hosted endpoint or running a batch transform job.
Use model packages to:

- Create models that you can use to get real-time inference or run batch transform jobs. For information, see [Use a Model Package to Create a Model](#).
- Create hosted endpoints to get real-time inference. For information, see [Deploy the Model to SageMaker Hosting Services](#).
- Create batch transform jobs. For information, see [Optional] Make Prediction with Batch Transform.

**Topics**

- [Use an Algorithm to Run a Training Job](#)
- [Use an Algorithm to Run a Hyperparameter Tuning Job](#)
- [Use a Model Package to Create a Model](#)
Use an Algorithm to Run a Training Job

You can create use an algorithm resource to create a training job by using the Amazon SageMaker console, the low-level Amazon SageMaker API, or the Amazon SageMaker Python SDK.

Topics

- Use an Algorithm to Run a Training Job (Console)
- Use an Algorithm to Run a Training Job (API)
- Use an Algorithm to Run a Training Job (Amazon SageMaker Python SDK)

Use an Algorithm to Run a Training Job (Console)

To use an algorithm to run a training job (console)

2. Choose Algorithms.
3. Choose an algorithm that you created from the list on the My algorithms tab or choose an algorithm that you subscribed to on the AWS Marketplace subscriptions tab.
4. Choose Create training job.

The algorithm you chose will automatically be selected.
5. On the Create training job page, provide the following information:

   a. For Job name, type a name for the training job.
   b. For IAM role, choose an IAM role that has the required permissions to run training jobs in SageMaker, or choose Create a new role to allow SageMaker to create a role that has the AmazonSageMakerFullAccess managed policy attached. For information, see SageMaker Roles.
   c. For Resource configuration, provide the following information:

      i. For Instance type, choose the instance type to use for training.
      ii. For Instance count, type the number of ML instances to use for the training job.
      iii. For Additional volume per instance (GB), type the size of the ML storage volume that you want to provision. ML storage volumes store model artifacts and incremental states.
iv. For **Encryption key**, if you want Amazon SageMaker to use an AWS Key Management Service key to encrypt data in the ML storage volume attached to the training instance, specify the key.

v. For **Stopping condition**, specify the maximum amount of time in seconds, minutes, hours, or days, that you want the training job to run.

d. For **VPC**, choose a Amazon VPC that you want to allow your training container to access. For more information, see Give SageMaker Training Jobs Access to Resources in Your Amazon VPC.

e. For **Hyperparameters**, specify the values of the hyperparameters to use for the training job.

f. For **Input data configuration**, specify the following values for each channel of input data to use for the training job. You can see what channels the algorithm you’re using for training support, and the content type, supported compression type, and supported input modes for each channel, under **Channel specification** section of the **Algorithm summary** page for the algorithm.

i. For **Channel name**, type the name of the input channel.

ii. For **Content type**, type the content type of the data that the algorithm expects for the channel.

iii. For **Compression type**, choose the data compression type to use, if any.

iv. For **Record wrapper**, choose RecordIO if the algorithm expects data in the RecordIO format.

v. For **S3 data type**, **S3 data distribution type**, and **S3 location**, specify the appropriate values. For information about what these values mean, see S3DataSource.

vi. For **Input mode**, choose **File** to download the data from to the provisioned ML storage volume, and mount the directory to a Docker volume. Choose **Pipe** to stream data directly from Amazon S3 to the container.

vii. To add another input channel, choose **Add channel**. If you are finished adding input channels, choose **Done**.

g. For **Output location**, specify the following values:

i. For **S3 output path**, choose the S3 location where the training job stores output, such as model artifacts.
Note

You use the model artifacts stored at this location to create a model or model package from this training job.

ii. For **Encryption key**, if you want SageMaker to use a AWS KMS key to encrypt output data at rest in the S3 location.

h. For **Tags**, specify one or more tags to manage the training job. Each tag consists of a key and an optional value. Tag keys must be unique per resource.

i. Choose **Create training job** to run the training job.

Use an Algorithm to Run a Training Job (API)

To use an algorithm to run a training job by using the SageMaker API, specify either the name or the Amazon Resource Name (ARN) as the `AlgorithmName` field of the `AlgorithmSpecification` object that you pass to `CreateTrainingJob`. For information about training models in SageMaker, see [Train a Model with Amazon SageMaker](#).

Use an Algorithm to Run a Training Job ([Amazon SageMaker Python SDK](#))

Use an algorithm that you created or subscribed to on AWS Marketplace to create a training job, create an `AlgorithmEstimator` object and specify either the Amazon Resource Name (ARN) or the name of the algorithm as the value of the `algorithm_arn` argument. Then call the `fit` method of the estimator. For example:

```python
from sagemaker import AlgorithmEstimator

data_path = os.path.join(DATA_DIR, 'marketplace', 'training')

algo = AlgorithmEstimator(
    role='SageMakerRole',
    instance_count=1,
    instance_type='ml.c4.xlarge',
    sagemaker_session=sagemaker_session,
    base_job_name='test-marketplace')

train_input = algo.sagemaker_session.upload_data(
    path=data_path, key_prefix='integ-test-data/marketplace/train')
```
Use an Algorithm to Run a Hyperparameter Tuning Job

A hyperparameter tuning job finds the best version of a model by running many training jobs on your dataset using the algorithm and ranges of hyperparameters that you specify. It then chooses the hyperparameter values that result in a model that performs the best, as measured by a metric that you choose. For more information, see Perform Automatic Model Tuning with SageMaker.

You can create use an algorithm resource to create a hyperparameter tuning job by using the Amazon SageMaker console, the low-level Amazon SageMaker API, or the Amazon SageMaker Python SDK.

Topics

• Use an Algorithm to Run a Hyperparameter Tuning Job (Console)
• Use an Algorithm to Run a Hyperparameter Tuning Job (API)
• Use an Algorithm to Run a Hyperparameter Tuning Job (Amazon SageMaker Python SDK)

Use an Algorithm to Run a Hyperparameter Tuning Job (Console)

To use an algorithm to run a hyperparameter tuning job (console)

2. Choose Algorithms.
3. Choose an algorithm that you created from the list on the My algorithms tab or choose an algorithm that you subscribed to on the AWS Marketplace subscriptions tab.
4. Choose Create hyperparameter tuning job.

The algorithm you chose will automatically be selected.

5. On the Create hyperparameter tuning job page, provide the following information:

   a. For Warm start, choose Enable warm start to use the information from previous hyperparameter tuning jobs as a starting point for this hyperparameter tuning job. For more information, see Run a Warm Start Hyperparameter Tuning Job.

   i. Choose Identical data and algorithm if your input data is the same as the input data for the parent jobs of this hyperparameter tuning job, or choose Transfer learning to use additional or different input data for this hyperparameter tuning job.
ii. For **Parent hyperparameter tuning job(s)**, choose up to 5 hyperparameter tuning jobs to use as parents to this hyperparameter tuning job.

b. For **Hyperparameter tuning job name**, type a name for the tuning job.

c. For **IAM role**, choose an IAM role that has the required permissions to run hyperparameter tuning jobs in SageMaker, or choose **Create a new role** to allow SageMaker to create a role that has the AmazonSageMakerFullAccess managed policy attached. For information, see **SageMaker Roles**.

d. For **VPC**, choose a Amazon VPC that you want to allow the training jobs that the tuning job launches to access. For more information, see **Give SageMaker Training Jobs Access to Resources in Your Amazon VPC**.

e. Choose **Next**.

f. For **Objective metric**, choose the metric that the hyperparameter tuning job uses to determine the best combination of hyperparameters, and choose whether to minimize or maximize this metric. For more information, see **View the Best Training Job**.

g. For **Hyperparameter configuration**, choose ranges for the tunable hyperparameters that you want the tuning job to search, and set static values for hyperparameters that you want to remain constant in all training jobs that the hyperparameter tuning job launches. For more information, see **Define Hyperparameter Ranges**.

h. Choose **Next**.

i. For **Input data configuration**, specify the following values for each channel of input data to use for the hyperparameter tuning job. You can see what channels the algorithm you're using for hyperparameter tuning supports, and the content type, supported compression type, and supported input modes for each channel, under **Channel specification** section of the **Algorithm summary** page for the algorithm.

   i. For **Channel name**, type the name of the input channel.

   ii. For **Content type**, type the content type of the data that the algorithm expects for the channel.

   iii. For **Compression type**, choose the data compression type to use, if any.

   iv. For **Record wrapper**, choose RecordIO if the algorithm expects data in the RecordIO format.

   v. For **S3 data type**, **S3 data distribution type**, and **S3 location**, specify the appropriate values. For information about what these values mean, see **S3DataSource**.
vi. For **Input mode**, choose **File** to download the data from to the provisioned ML storage volume, and mount the directory to a Docker volume. Choose **Pipe** to stream data directly from Amazon S3 to the container.

vii. To add another input channel, choose **Add channel**. If you are finished adding input channels, choose **Done**.

j. For **Output** location, specify the following values:

i. For **S3 output path**, choose the S3 location where the training jobs that this hyperparameter tuning job launches store output, such as model artifacts.

   ![Note]
   You use the model artifacts stored at this location to create a model or model package from this hyperparameter tuning job.

ii. For **Encryption key**, if you want SageMaker to use a AWS KMS key to encrypt output data at rest in the S3 location.

k. For **Resource configuration**, provide the following information:

i. For **Instance type**, choose the instance type to use for each training job that the hyperparameter tuning job launches.

ii. For **Instance count**, type the number of ML instances to use for each training job that the hyperparameter tuning job launches.

iii. For **Additional volume per instance (GB)**, type the size of the ML storage volume that you want to provision each training job that the hyperparameter tuning job launches. ML storage volumes store model artifacts and incremental states.

iv. For **Encryption key**, if you want Amazon SageMaker to use an AWS Key Management Service key to encrypt data in the ML storage volume attached to the training instances, specify the key.

l. For **Resource limits**, provide the following information:

i. For **Maximum training jobs**, specify the maximum number of training jobs that you want the hyperparameter tuning job to launch. A hyperparameter tuning job can launch a maximum of 500 training jobs.
ii. For **Maximum parallel training jobs**, specify the maximum number of concurrent training jobs that the hyperparameter tuning job can launch. A hyperparameter tuning job can launch a maximum of 10 concurrent training jobs.

iii. For **Stopping condition**, specify the maximum amount of time in seconds, minutes, hours, or days, that you want each training job that the hyperparameter tuning job launches to run.

m. For **Tags**, specify one or more tags to manage the hyperparameter tuning job. Each tag consists of a key and an optional value. Tag keys must be unique per resource.

n. Choose **Create jobs** to run the hyperparameter tuning job.

### Use an Algorithm to Run a Hyperparameter Tuning Job (API)

To use an algorithm to run a hyperparameter tuning job by using the SageMaker API, specify either the name or the Amazon Resource Name (ARN) of the algorithm as the AlgorithmName field of the **AlgorithmSpecification** object that you pass to **CreateHyperParameterTuningJob**.

For information about hyperparameter tuning in SageMaker, see [Perform Automatic Model Tuning with SageMaker](#).

### Use an Algorithm to Run a Hyperparameter Tuning Job (**Amazon SageMaker Python SDK**)  

Use an algorithm that you created or subscribed to on AWS Marketplace to create a hyperparameter tuning job, create an **AlgorithmEstimator** object and specify either the Amazon Resource Name (ARN) or the name of the algorithm as the value of the **algorithm_arn** argument. Then initialize a **HyperparameterTuner** object with the **AlgorithmEstimator** you created as the value of the **estimator** argument. Finally, call the `fit` method of the **AlgorithmEstimator**. For example:

```python
from sagemaker import AlgorithmEstimator
from sagemaker.tuner import HyperparameterTuner

data_path = os.path.join(DATA_DIR, 'marketplace', 'training')

algo = AlgorithmEstimator(
    role='SageMakerRole',
    instance_count=1,
    instance_type='ml.c4.xlarge',
    sagemaker_session=sagemaker_session,
)``
Use Algorithm and Model Package Resources

Use a Model Package to Create a Model

Use a model package to create a deployable model that you can use to get real-time inferences by creating a hosted endpoint or to run batch transform jobs. You can create a deployable model from a model package by using the Amazon SageMaker console, the low-level SageMaker API, or the Amazon SageMaker Python SDK.

Topics

• Use a Model Package to Create a Model (Console)
• Use a Model Package to Create a Model (API)
• Use a Model Package to Create a Model (Amazon SageMaker Python SDK)

Use a Model Package to Create a Model (Console)

To create a deployable model from a model package (console)

2. Choose Model packages.
3. Choose a model package that you created from the list on the My model packages tab or choose a model package that you subscribed to on the AWS Marketplace subscriptions tab.
4. Choose Create model.
5. For Model name, type a name for the model.
6. For **IAM role**, choose an IAM role that has the required permissions to call other services on your behalf, or choose **Create a new role** to allow SageMaker to create a role that has the AmazonSageMakerFullAccess managed policy attached. For information, see [SageMaker Roles](#).

7. For **VPC**, choose a Amazon VPC that you want to allow the model to access. For more information, see [Give SageMaker Hosted Endpoints Access to Resources in Your Amazon VPC](#).

8. Leave the default values for **Container input options** and **Choose model package**.

9. For environment variables, provide the names and values of environment variables you want to pass to the model container.

10. For **Tags**, specify one or more tags to manage the model. Each tag consists of a key and an optional value. Tag keys must be unique per resource.

11. Choose **Create model**.

After you create a deployable model, you can use it to set up an endpoint for real-time inference or create a batch transform job to get inferences on entire datasets. For information about hosting endpoints in SageMaker, see [Deploy Models for Inference](#).

### Use a Model Package to Create a Model (API)

To use a model package to create a deployable model by using the SageMaker API, specify the name or the Amazon Resource Name (ARN) of the model package as the `ModelPackageName` field of the `ContainerDefinition` object that you pass to the `CreateModel` API.

After you create a deployable model, you can use it to set up an endpoint for real-time inference or create a batch transform job to get inferences on entire datasets. For information about hosted endpoints in SageMaker, see [Deploy Models for Inference](#).

### Use a Model Package to Create a Model ([Amazon SageMaker Python SDK](#))

To use a model package to create a deployable model by using the SageMaker Python SDK, initialize a `ModelPackage` object, and pass the Amazon Resource Name (ARN) of the model package as the `model_package_arn` argument. For example:

```python
from sagemaker import ModelPackage
model = ModelPackage(role='SageMakerRole',
                     model_package_arn='training-job-scikit-decision-trees-1542660466-6f92',
                     sagemaker_session=sagemaker_session)
```
After you create a deployable model, you can use it to set up an endpoint for real-time inference or create a batch transform job to get inferences on entire datasets. For information about hosting endpoints in SageMaker, see [Deploy Models for Inference](#).

**Sell Amazon SageMaker Algorithms and Model Packages**

Selling Amazon SageMaker algorithms and model packages is a three-step process:

1. Develop your algorithm or model, and package it in a Docker container. For information, see [Develop Algorithms and Models in Amazon SageMaker](#).
2. Create an algorithm or model package resource in SageMaker. For information, see [Create Algorithm and Model Package Resources](#).
3. Register as a seller on AWS Marketplace and list your algorithm or model package on AWS Marketplace. For information about registering as a seller, see [Getting Started as a Seller](#) in the [User Guide for AWS Marketplace Providers](#). For information about listing and monetizing your algorithms and model packages, see [Listing Algorithms and Model Packages in AWS Marketplace for Machine Learning](#) in the [User Guide for AWS Marketplace Providers](#).

<table>
<thead>
<tr>
<th>Seller’s environment/SageMaker</th>
<th>SageMaker</th>
<th>AWS Marketplace</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="#" alt="Package your code" /> Build your algorithm or model and package the code in a Docker container.</td>
<td><img src="#" alt="Create algorithm or model package in Amazon SageMaker" /> Ingest your algorithm or model into Amazon SageMaker. Set training and inference specifications and validation profiles, then upload sample data.</td>
<td><img src="#" alt="List and monetize your product" /> Add a description, promotions, and pricing information, and distribute your product globally.</td>
</tr>
</tbody>
</table>

**Topics**

- [Develop Algorithms and Models in Amazon SageMaker](#)
- [Create Algorithm and Model Package Resources](#)
- [List Your Algorithm or Model Package on AWS Marketplace](#)
Develop Algorithms and Models in Amazon SageMaker

Before you can create algorithm and model package resources to use in Amazon SageMaker or list on AWS Marketplace, you have to develop them and package them in Docker containers.

**Note**

When algorithms and model packages are created for listing on AWS Marketplace, SageMaker scans the containers for security vulnerabilities on supported operating systems.

Only the following operating system versions are supported:

- Debian: 6.0, 7, 8, 9, 10
- CentOS: 5, 6, 7
- Oracle Linux: 5, 6, 7
- Alpine: 3.3, 3.4, 3.5
- Amazon Linux

**Topics**

- Develop Algorithms in SageMaker
- Develop Models in SageMaker

**Develop Algorithms in SageMaker**

An algorithm should be packaged as a docker container and stored in Amazon ECR to use it in SageMaker. The Docker container contains the training code used to run training jobs and, optionally, the inference code used to get inferences from models trained by using the algorithm.

For information about developing algorithms in SageMaker and packaging them as containers, see [Use Docker containers to build models](https://sagemaker-examples.readthedocs.io/en/latest/advanced_functionality/scikitbring_your_own/scikitbring_your_own.html). For a complete example of how to create an algorithm container, see the sample notebook at [https://sagemaker-examples.readthedocs.io/en/latest/advanced_functionality/scikitbring_your_own/scikitbring_your_own.html](https://sagemaker-examples.readthedocs.io/en/latest/advanced_functionality/scikitbring_your_own/scikitbring_your_own.html). You can also find the sample notebook in a SageMaker notebook instance. The notebook is in the **Advanced**
**Functionality** section, and is named `scikitBringYourOwn.ipynb`. For information about using the sample notebooks in a notebook instance, see [Example Notebooks](#).

Always thoroughly test your algorithms before you create algorithm resources to publish on AWS Marketplace.

**Note**

When a buyer subscribes to your containerized product, the Docker containers run in an isolated (internet-free) environment. When you create your containers, do not rely on making outgoing calls over the internet. Calls to AWS services are also not allowed.

---

**Develop Models in SageMaker**

A deployable model in SageMaker consists of inference code, model artifacts, an IAM role that is used to access resources, and other information required to deploy the model in SageMaker. Model artifacts are the results of training a model by using a machine learning algorithm. The inference code must be packaged in a Docker container and stored in Amazon ECR. You can either package the model artifacts in the same container as the inference code, or store them in Amazon S3.

You create a model by running a training job in SageMaker, or by training a machine learning algorithm outside of SageMaker. If you run a training job in SageMaker, the resulting model artifacts are available in the `ModelArtifacts` field in the response to a call to the `DescribeTrainingJob` operation. For information about how to develop a SageMaker model container, see [Use your own inference code](#). For a complete example of how to create a model container from a model trained outside of SageMaker, see the sample notebook at [https://sagemaker-examples.readthedocs.io/en/latest/advanced_functionality/xgboostbringyourown_model/xgboostbringyourown_model.html](https://sagemaker-examples.readthedocs.io/en/latest/advanced_functionality/xgboostbringyourown_model/xgboostbringyourown_model.html). You can also find the sample notebook in a SageMaker notebook instance. The notebook is in the **Advanced Functionality** section, and is named `xgboostBringYourOwnModel.ipynb`. For information about using the sample notebooks in a notebook instance, see [Example Notebooks](#).

Always thoroughly test your models before you create model packages to publish on AWS Marketplace.
Note
When a buyer subscribes to your containerized product, the Docker containers run in an isolated (internet-free) environment. When you create your containers, do not rely on making outgoing calls over the internet. Calls to AWS services are also not allowed.

List Your Algorithm or Model Package on AWS Marketplace

After creating and validating your algorithm or model in Amazon SageMaker, list your product on AWS Marketplace. The listing process makes your products available in the AWS Marketplace and the SageMaker console.

To list products on AWS Marketplace, you must be a registered seller. To register, use the self-registration process from the AWS Marketplace Management Portal (AMMP). For information, see Getting Started as a Seller in the User Guide for AWS Marketplace Providers. When you start the product listing process from the Amazon SageMaker console, we check your seller registration status. If you have not registered, we direct you to do so.

To start the listing process, do one of the following:

- From the SageMaker console, choose the product, choose Actions, and choose Publish new ML Marketplace listing. This carries over your product reference, the Amazon Resource Name (ARN), and directs you to the AMMP to create the listing.

- Go to ML listing process, manually enter the Amazon Resource Name (ARN), and start your product listing. This process carries over the product metadata that you entered when creating the product in SageMaker. For an algorithm listing, the information includes the supported instance types and hyperparameters. In addition, you can enter a product description, promotional information, and support information as you would with other AWS Marketplace products.

Find and Subscribe to Algorithms and Model Packages on AWS Marketplace

With AWS Marketplace, you can browse and search for hundreds of machine learning algorithms and models in a broad range of categories, such as computer vision, natural language processing,
speech recognition, text, data, voice, image, video analysis, fraud detection, predictive analysis, and more.

To find algorithms on AWS Marketplace

1. Open the Amazon SageMaker console at https://console.aws.amazon.com/sagemaker/.
2. Choose Algorithms, then choose Find algorithms.

   This takes you to the AWS Marketplace algorithms page. For information about finding and subscribing to algorithms on AWS Marketplace, see Machine Learning Products in the AWS Marketplace User Guide for AWS Consumers.

To find model packages on AWS Marketplace

2. Choose Model packages, then choose Find model packages.

   This takes you to the AWS Marketplace model packages page. For information about finding and subscribing to model packages on AWS Marketplace, see Machine Learning Products in the AWS Marketplace User Guide for AWS Consumers.

Use Algorithms and Model Packages

For information about using algorithms and model packages that you subscribe to in SageMaker, see Use Algorithm and Model Package Resources.
Note

When you create a training job, inference endpoint, and batch transform job from an algorithm or model package that you subscribe to on AWS Marketplace, the training and inference containers do not have access to the internet. Because the containers do not have access to the internet, the seller of the algorithm or model package does not have access to your data.
Monitor AWS resources provisioned while using Amazon SageMaker

Monitoring is an important part of maintaining the reliability, availability, and performance of SageMaker and your other AWS solutions. AWS provides the following monitoring tools to watch SageMaker, report when something is wrong, and take automatic actions when appropriate:

- **Amazon CloudWatch** monitors your AWS resources and the applications that you run on AWS in real time. You can collect and track metrics, create customized dashboards, and set alarms that notify you or take actions when a specified metric reaches a threshold that you specify. For example, you can have CloudWatch track CPU usage or other metrics of your Amazon EC2 instances and automatically launch new instances when needed. For more information, see the [Amazon CloudWatch User Guide](https://docs.aws.amazon.com/AmazonCloudWatch/latest/userguide/)

- **Amazon CloudWatch Logs** enables you to monitor, store, and access your log files from EC2 instances, AWS CloudTrail, and other sources. CloudWatch Logs can monitor information in the log files and notify you when certain thresholds are met. You can also archive your log data in highly durable storage. For more information, see the [Amazon CloudWatch Logs User Guide](https://docs.aws.amazon.com/AmazonCloudWatch/latest/monitoring/cloudwatchlogs.html).

- **AWS CloudTrail** captures API calls and related events made by or on behalf of your AWS account and delivers the log files to an Amazon S3 bucket that you specify. You can identify which users and accounts called AWS, the source IP address from which the calls were made, and when the calls occurred. For more information, see the [AWS CloudTrail User Guide](https://docs.aws.amazon.com/AmazonCloudWatch/latest/userguide/).

- **CloudWatch Events** delivers a near real-time stream of system events that describe changes in AWS resources. Create CloudWatch Events rules react to a status change in a SageMaker training, hyperparameter tuning, or batch transform job

Topics

- [Monitor Amazon SageMaker with Amazon CloudWatch](https://docs.aws.amazon.com/sagemaker/latest/dg/monitoring-sagemaker-cloudwatch.html)
- [Log Amazon SageMaker Events with Amazon CloudWatch](https://docs.aws.amazon.com/sagemaker/latest/dg/monitoring-sagemaker-cloudwatch-events.html)
- [Log Amazon SageMaker API Calls with AWS CloudTrail](https://docs.aws.amazon.com/sagemaker/latest/dg/monitoring-sagemaker-cloudtrail.html)
- [Monitoring user resource access from Amazon SageMaker Studio Classic](https://docs.aws.amazon.com/sagemaker/latest/dg/monitoring-sagemaker-access.html)
- [Automating Amazon SageMaker with Amazon EventBridge](https://docs.aws.amazon.com/sagemaker/latest/dg/automate-sagemaker-events.html)
Monitor Amazon SageMaker with Amazon CloudWatch

You can monitor Amazon SageMaker using Amazon CloudWatch, which collects raw data and processes it into readable, near real-time metrics. These statistics are kept for 15 months, so that you can access historical information and gain a better perspective on how your web application or service is performing. However, the Amazon CloudWatch console limits the search to metrics that were updated in the last 2 weeks. This limitation ensures that the most current jobs are shown in your namespace. To graph metrics without using a search, specify its exact name in the source view. You can also set alarms that watch for certain thresholds, and send notifications or take actions when those thresholds are met. For more information, see the Amazon CloudWatch User Guide.

SageMaker Metrics and Dimensions

- SageMaker Endpoint Invocation Metrics
- SageMaker Inference Component Metrics
- SageMaker Multi-Model Endpoint Metrics
- SageMaker Jobs and Endpoint Metrics
- SageMaker Inference Recommender Jobs Metrics
- SageMaker Ground Truth Metrics
- Amazon SageMaker Feature Store Metrics
- SageMaker Pipelines Metrics

SageMaker Endpoint Invocation Metrics

The AWS/SageMaker namespace includes the following request metrics from calls to InvokeEndpoint.

Metrics are available at a 1-minute frequency.

The following illustration shows how a SageMaker endpoint interacts with the Amazon SageMaker Runtime API. The overall time between sending a request to an endpoint and receiving a response depends on the following three components.

- Network latency – the time that it takes between making a request to the SageMaker Runtime API and receiving a response back from the SageMaker Runtime Runtime API.
- Overhead latency – the time that it takes to transport a request to the model container from the SageMaker Runtime Runtime API and transport the response back to the SageMaker Runtime Runtime API.
- Model latency – the time that it takes the model container to process the request and return a response.

Total time (end-to-end) from request to response = network latency + overhead latency + model latency

For more information about total latency, see Best practices for load testing Amazon SageMaker real-time inference endpoints. For information about how long CloudWatch metrics are retained for, see GetMetricStatistics in the Amazon CloudWatch API Reference.

### Endpoint Invocation Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invocation4XXErrors</td>
<td>The number of InvokeEndpoint requests where the model returned a 4xx HTTP response code. For each 4xx response, 1 is sent; otherwise, 0 is sent.</td>
</tr>
<tr>
<td>Units: None</td>
<td></td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Invocations5XXErrors</td>
<td>The number of <code>InvokeEndpoint</code> requests where the model returned a 5xx HTTP response code. For each 5xx response, 1 is sent; otherwise, 0 is sent.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>InvocationsModelErrors</td>
<td>The number of model invocation requests which did not result in 2XX HTTP response. This includes 4XX/5XX status codes, low-level socket errors, malformed HTTP responses, and request timeouts. For each error response, 1 is sent; otherwise, 0 is sent.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>Invocations</td>
<td>The number of <code>InvokeEndpoint</code> requests sent to a model endpoint. To get the total number of requests sent to a model endpoint, use the Sum statistic.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>InvocationsPerCopy</td>
<td>The number of invocations normalized by each copy of an inference component.</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>InvocationsPerInstance</td>
<td>The number of invocations sent to a model, normalized by InstanceCount in each ProductionVariant. (1/\text{numberOfInstances}) is sent as the value on each request, where (\text{numberOfInstances}) is the number of active instances for the ProductionVariant behind the endpoint at the time of the request.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>ModelLatency</td>
<td>The interval of time taken by a model to respond to a SageMaker Runtime API request. This interval includes the local communication times taken to send the request and to fetch the response from the model container and the time taken to complete the inference in the container.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelSetupTime</td>
<td>The time it takes to launch new compute resources for a serverless endpoint. The time can vary depending on the model size, how long it takes to download the model, and the start-up time of the container.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Min, Max, Sample Count, Percentiles</td>
</tr>
</tbody>
</table>
## Amazon SageMaker

### Metric Description

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>OverheadLatency</td>
<td>The interval of time added to the time taken to respond to a client request by SageMaker overheads. This interval is measured from the time SageMaker receives the request until it returns a response to the client, minus the ModelLatency. Overhead latency can vary depending on multiple factors, including request and response payload sizes, request frequency, and authentication/authorization of the request. Units: Microseconds Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>

### Dimensions for Endpoint Invocation Metrics

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EndpointName, VariantName</td>
<td>Filters endpoint invocation metrics for a ProductionVariant of the specified endpoint and variant.</td>
</tr>
<tr>
<td>Inference ComponentName</td>
<td>Filters inference component invocation metrics.</td>
</tr>
</tbody>
</table>

### SageMaker Inference Component Metrics

The `/aws/sagemaker/InferenceComponents` namespace includes the following metrics from calls to `InvokeEndpoint` for endpoints that host inference components.

Metrics are available at a 1-minute frequency.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPUUtilizationNormalized</td>
<td>The value of the CPUUtilizationNormalized metric reported by each copy of the inference component. The value ranges between 0%–100%. If you set the NumberOfCpuCoresRequired parameter in the settings for the inference component copy, the metric presents</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>GPUMemoryUtilizationNormalized</td>
<td>The value of the GPUMemoryUtilizationNormalized metric reported by each copy of the inference component.</td>
</tr>
<tr>
<td>GPUUtilizationNormalized</td>
<td>The value of the GPUUtilizationNormalized metric reported by each copy of the inference component. If you set the NumberOfAcceleratorDevicesRequired parameter in the settings for the inference component copy, the metric presents the utilization over the reservation. Otherwise, the metric presents the utilization over the limit.</td>
</tr>
<tr>
<td>MemoryUtilizationNormalized</td>
<td>The value of MemoryUtilizationNormalized reported by each copy of the inference component. If you set the MinMemoryRequiredInMb parameter in the settings for the inference component copy, the metrics present the utilization over the reservation. Otherwise, the metrics present the utilization over the limit.</td>
</tr>
</tbody>
</table>

**Dimensions for Inference Component Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>InferenceComponentName</td>
<td>Filters inference component metrics.</td>
</tr>
</tbody>
</table>

**SageMaker Multi-Model Endpoint Metrics**

The AWS/SageMaker namespace includes the following model loading metrics from calls to **InvokeEndpoint**.

Metrics are available at a 1-minute frequency.
For information about how long CloudWatch metrics are retained for, see [GetMetricStatistics](https://docs.aws.amazon.com/AmazonCloudWatch/latest/APIReference/API_GetMetricStatistics.html) in the *Amazon CloudWatch API Reference*.

### Multi-Model Endpoint Model Loading Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelLoadingWaitTime</td>
<td>The interval of time that an invocation request has waited for the target model to be downloaded, or loaded, or both in order to perform inference.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelUnloadingTime</td>
<td>The interval of time that it took to unload the model through the container's <code>UnloadModel</code> API call.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelDownloadTime</td>
<td>The interval of time that it took to download the model from Amazon Simple Storage Service (Amazon S3).</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelLoadingTime</td>
<td>The interval of time that it took to load the model through the container's <code>LoadModel</code> API call.</td>
</tr>
<tr>
<td></td>
<td>Units: Microseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
<tr>
<td>ModelCacheHit</td>
<td>The number of <code>InvokeEndpoint</code> requests sent to the multi-model endpoint for which the model was already loaded.</td>
</tr>
<tr>
<td></td>
<td>The Average statistic shows the ratio of requests for which the model was already loaded.</td>
</tr>
</tbody>
</table>
## Metric

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Sample Count</td>
</tr>
</tbody>
</table>

## Dimensions for Multi-Model Endpoint Model Loading Metrics

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EndpointName, VariantName</td>
<td>Filters endpoint invocation metrics for a ProductionVariant of the specified endpoint and variant.</td>
</tr>
</tbody>
</table>

The /aws/sagemaker/Endpoints namespaces include the following instance metrics from calls to **InvokeEndpoint**.

Metrics are available at a 1-minute frequency.

For information about how long CloudWatch metrics are retained for, see [GetMetricStatistics](https://docs.aws.amazon.com/AmazonCloudWatch/latest/APIReference/API_GetMetricStatistics.html) in the Amazon CloudWatch API Reference.

## Multi-Model Endpoint Model Instance Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>LoadedModelCount</td>
<td>The number of models loaded in the containers of the multi-model endpoint. This metric is emitted per instance.</td>
</tr>
</tbody>
</table>

  * The Average statistic with a period of 1 minute tells you the average number of models loaded per instance.  
  * The Sum statistic tells you the total number of models loaded across all instances in the endpoint.  
  * The models that this metric tracks are not necessarily unique because a model might be loaded in multiple containers at the endpoint.  

Units: None
<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>

### Dimensions for Multi-Model Endpoint Model Loading Metrics

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EndpointName, VariantName</td>
<td>Filters endpoint invocation metrics for a ProductionVariant of the specified endpoint and variant.</td>
</tr>
</tbody>
</table>

### SageMaker Jobs and Endpoint Metrics

The `/aws/sagemaker/ProcessingJobs, /aws/sagemaker/TrainingJobs, /aws/sagemaker/TransformJobs, and /aws/sagemaker/Endpoints` namespaces include the following metrics for the training jobs and endpoint instances.

Metrics are available at a 1-minute frequency.

#### Note

Amazon CloudWatch supports [high-resolution custom metrics](https://docs.aws.amazon.com/AmazonCloudWatch/latest/monitoring/HighResOverview.html) and its finest resolution is 1 second. However, the finer the resolution, the shorter the lifespan of the CloudWatch metrics. For the 1-second frequency resolution, the CloudWatch metrics are available for 3 hours. For more information about the resolution and the lifespan of the CloudWatch metrics, see `GetMetricStatistics` in the [Amazon CloudWatch API Reference](https://docs.aws.amazon.com/AmazonCloudWatch/latest/APIReference/API_GetMetricStatistics.html).

#### Tip

If you want to profile your training job with a finer resolution down to 100-millisecond (0.1 second) granularity and store the training metrics indefinitely in Amazon S3 for custom analysis at any time, consider using [Amazon SageMaker Debugger](https://aws.amazon.com/sagemaker-debugger/). SageMaker Debugger provides built-in rules to automatically detect common training issues; it detects hardware resource utilization issues (such as CPU, GPU, and I/O bottlenecks) and non-converging model issues (such as overfit, vanishing gradients, and exploding tensors). SageMaker
Debugger also provides visualizations through Studio Classic and its profiling report. To explore the Debugger visualizations, see [SageMaker Debugger Insights Dashboard Walkthrough](#), [Debugger Profiling Report Walkthrough](#), and [Analyze Data Using the SMDebug Client Library](#).

## Processing Job, Training Job, Batch Transform Job, and Endpoint Instance Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPUReservation</td>
<td>The sum of CPUs reserved by containers on an instance. The value ranges between 0%–100%. In the settings for an inference component, you set the CPU reservation with the <code>NumberOfCpuCoresRequired</code> parameter. For example, if there 4 CPUs, and 2 are reserved, the CPUReservation metric is 50%.</td>
</tr>
<tr>
<td>CPUUtilization</td>
<td>The sum of each individual CPU core's utilization. The CPU utilization of each core range is 0–100. For example, if there are four CPUs, the CPUUtilization range is 0%–400%. For processing jobs, the value is the CPU utilization of the processing container on the instance. For training jobs, the value is the CPU utilization of the algorithm container on the instance. For batch transform jobs, the value is the CPU utilization of the transform container on the instance. For endpoint variants, the value is the sum of the CPU utilization of the primary and supplementary containers on the instance.</td>
</tr>
</tbody>
</table>

**Note**

For multi-instance jobs, each instance reports CPU utilization metrics. However, the default view in CloudWatch shows the average CPU utilization across all instances.

Units: Percent
<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPUUtilizationNormalized</td>
<td>The normalized sum of the utilization of each individual CPU core. The value ranges between 0%–100%. For example, if there are four CPUs, and the CPUUtilization metric is 200%, then the CPUUtilizationNormalized metric is 50%.</td>
</tr>
<tr>
<td>DiskUtilization</td>
<td>The percentage of disk space used by the containers on an instance uses. This value range is 0%–100%. This metric is not supported for batch transform jobs. For processing jobs, the value is the disk space utilization of the processing container on the instance. For training jobs, the value is the disk space utilization of the algorithm container on the instance. For endpoint variants, the value is the sum of the disk space utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
</tbody>
</table>

**Note**

For multi-instance jobs, each instance reports disk utilization metrics. However, the default view in CloudWatch shows the average disk utilization across all instances.
<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPUMemory Utilization</td>
<td>The percentage of GPU memory used by the containers on an instance. The value range is 0–100 and is multiplied by the number of GPUs. For example, if there are four GPUs, the GPUMemoryUtilization range is 0%–400%. For processing jobs, the value is the GPU memory utilization of the processing container on the instance. For training jobs, the value is the GPU memory utilization of the algorithm container on the instance. For batch transform jobs, the value is the GPU memory utilization of the transform container on the instance. For endpoint variants, the value is the sum of the GPU memory utilization of the primary and supplementary containers on the instance.</td>
</tr>
<tr>
<td>Note</td>
<td>For multi-instance jobs, each instance reports GPU memory utilization metrics. However, the default view in CloudWatch shows the average GPU memory utilization across all instances.</td>
</tr>
<tr>
<td>Units: Percent</td>
<td></td>
</tr>
<tr>
<td>GPUMemoryUtilizationNormalized</td>
<td>The normalized percentage of GPU memory used by the containers on an instance. The value ranges between 0%–100%. For example, if there are four GPUs, and the GPUMemoryUtilization metric is 200%, then the GPUMemoryUtilizationNormalized metric is 50%.</td>
</tr>
<tr>
<td>GPUReservation</td>
<td>The sum of GPUs reserved by containers on an instance. The value ranges between 0%–100%. In the settings for an inference component, you set the GPU reservation by NumberOfAcceleratorDevicesRequired. For example, if there are 4 GPUs and 2 are reserved, the GPUReservation metric is 50%.</td>
</tr>
</tbody>
</table>
### Metric Description

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPUUtilization</td>
<td>The percentage of GPU units that are used by the containers on an instance. The value can range between 0–100 and is multiplied by the number of GPUs. For example, if there are four GPUs, the GPUUtilization range is 0%–400%. For processing jobs, the value is the GPU utilization of the processing container on the instance. For training jobs, the value is the GPU utilization of the algorithm container on the instance. For batch transform jobs, the value is the GPU utilization of the transform container on the instance. For endpoint variants, the value is the sum of the GPU utilization of the primary and supplementary containers on the instance.</td>
</tr>
<tr>
<td>GPUUtilizationNormalized</td>
<td>The normalized percentage of GPU units that are used by the containers on an instance. The value ranges between 0%–100%. For example, if there are four GPUs, and the GPUUtilization metric is 200%, then the GPUUtilizationNormalized metric is 50%.</td>
</tr>
<tr>
<td>MemoryReservation</td>
<td>The sum of memory reserved by containers on an instance. The value ranges between 0%–100%. In the settings for an inference component, you set the memory reservation with the MinMemoryRequiredInMb parameter. For example, if a 32 GiB instance reserved 1024 MB, the MemoryReservation metric is 29.8%.</td>
</tr>
</tbody>
</table>

**Note**

For multi-instance jobs, each instance reports GPU utilization metrics. However, the default view in CloudWatch shows the average GPU utilization across all instances.
<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MemoryUtilization</td>
<td>The percentage of memory that is used by the containers on an instance. This value range is 0%–100%. For processing jobs, the value is the memory utilization of the processing container on the instance. For training jobs, the value is the memory utilization of the algorithm container on the instance. For batch transform jobs, the value is the memory utilization of the transform container on the instance. For endpoint variants, the value is the sum of the memory utilization of the primary and supplementary containers on the instance. Units: Percent</td>
</tr>
</tbody>
</table>

**Note**

For multi-instance jobs, each instance reports memory utilization metrics. However, the default view in CloudWatch shows the average memory utilization across all instances.

---

**Dimensions for Processing Job, Training Job and Batch Transform Job Instance Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Host</td>
<td>For processing jobs, the value for this dimension has the format [processing-job-name]/algo-[instance-number-in-cluster]. Use this dimension to filter instance metrics for the specified processing job and instance. This dimension format is present only in the /aws/sagemaker/ProcessingJobs namespace. For training jobs, the value for this dimension has the format [training-job-name]/algo-[instance-number-in-cluster]. Use this dimension to filter instance metrics for the training job.</td>
</tr>
</tbody>
</table>

---

Jobs and Endpoint Metrics
<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>specified training job and instance. This dimension format is present only in the /aws/sagemaker/TrainingJobs namespace.</td>
</tr>
<tr>
<td></td>
<td>For batch transform jobs, the value for this dimension has the format [transform-job-name]/[instance-id]. Use this dimension to filter instance metrics for the specified batch transform job and instance. This dimension format is present only in the /aws/sagemaker/TransformJobs namespace.</td>
</tr>
</tbody>
</table>

**SageMaker Inference Recommender Jobs Metrics**

The /aws/sagemaker/InferenceRecommendationsJobs namespace includes the following metrics for inference recommendation jobs.

**Inference Recommender Metrics**

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ClientInvocations</td>
<td>The number of InvokeEndpoint requests sent to a model endpoint, as observed by Inference Recommender.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>ClientInvocationErrors</td>
<td>The number of InvokeEndpoint requests that failed, as observed by Inference Recommender.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum</td>
</tr>
<tr>
<td>ClientLatency</td>
<td>The interval of time taken between sending an InvokeEndpoint call and receiving a response as observed by Inference Recommender. Note that the time is in milliseconds, whereas the ModelLatency endpoint invocation metric is in microseconds.</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Metrics</td>
<td>Description</td>
</tr>
<tr>
<td>Units: Milliseconds</td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count, Percentiles</td>
</tr>
<tr>
<td>NumberOfUsers</td>
<td>The number of concurrent users sending <code>InvokeEndpoint</code> requests to the model endpoint.</td>
</tr>
<tr>
<td>Units: None</td>
<td>Valid statistics: Max, Min, Average</td>
</tr>
</tbody>
</table>

**Dimensions for Inference Recommender Job Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>JobName</td>
<td>Filters Inference Recommender job metrics for the specified Inference Recommender job.</td>
</tr>
<tr>
<td>EndpointName</td>
<td>Filters Inference Recommender job metrics for the specified endpoint.</td>
</tr>
</tbody>
</table>

**SageMaker Ground Truth Metrics**

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ActiveWorkers</td>
<td>A single active worker on a private work team submitted, released, or declined a task. To get the total number of active workers, use the Sum statistic. Ground Truth attempts to deliver each individual <code>ActiveWorkers</code> event once. If this delivery is unsuccessful, this metric may not report the total number of active workers</td>
</tr>
<tr>
<td>Units: None</td>
<td>Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>DatasetObjectsAutoAnnotated</td>
<td>The number of dataset objects auto-annotated in a labeling job. This metric is only emitted when automated labeling is enabled. To view the labeling job progress, use the Max metric. Units: None Valid statistics: Max</td>
</tr>
<tr>
<td>DatasetObjectsHumanAnnotated</td>
<td>The number of dataset objects annotated by a human in a labeling job. To view the labeling job progress, use the Max metric.                                                                                     Units: None Valid statistics: Max</td>
</tr>
<tr>
<td>DatasetObjectsLabelingFailed</td>
<td>The number of dataset objects that failed labeling in a labeling job. To view the labeling job progress, use the Max metric.                                                                                      Units: None Valid statistics: Max</td>
</tr>
<tr>
<td>JobsFailed</td>
<td>A single labeling job failed. To get the total number of labeling jobs that failed, use the Sum statistic.                                                                                                       Units: None Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>JobsSucceeded</td>
<td>A single labeling job succeeded. To get the total number of labeling jobs that succeeded, use the Sum statistic.                                                                                                  Units: None Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>JobsStopped</td>
<td>A single labeling jobs was stopped. To get the total number of labeling jobs that were stopped, use the Sum statistic.</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>TasksAccepted</td>
<td>A single task was accepted by a worker. To get the total number of tasks accepted by workers, use the Sum statistic. Ground Truth attempts to deliver each individual TaskAccepted event once. If this delivery is unsuccessful, this metric may not report the total number of tasks accepted.</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>TasksDeclined</td>
<td>A single task was declined by a worker. To get the total number of tasks declined by workers, use the Sum statistic. Ground Truth attempts to deliver each individual TasksDeclined event once. If this delivery is unsuccessful, this metric may not report the total number of tasks declined.</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Valid Statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>TasksReturned</td>
<td>A single task was returned. To get the total number of tasks returned, use the Sum statistic. Ground Truth attempts to deliver each individual TasksReturned event once. If this delivery is unsuccessful, this metric may not report the total number of tasks returned.</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>Metric</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>TasksSubmitted</td>
<td>A single task was submitted/completed by a private worker. To get the total number of tasks submitted by workers, use the Sum statistic. Ground Truth attempts to deliver each individual TasksSubmitted event once. If this delivery is unsuccessful, this metric may not report the total number of tasks submitted.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>TimeSpent</td>
<td>Time spent on a task completed by a private worker. This metric does not include time when a worker paused or took a break. Ground Truth attempts to deliver each TimeSpent event once. If this delivery is unsuccessful, this metric may not report the total amount of time spent.</td>
</tr>
<tr>
<td></td>
<td>Units: Seconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Sum, Sample Count</td>
</tr>
<tr>
<td>TotalDataSetObjectsLabeled</td>
<td>The number of dataset objects labeled successfully in a labeling job. To view the labeling job progress, use the Max metric.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Max</td>
</tr>
</tbody>
</table>

**Dimensions for Dataset Object Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>LabelingJobName</td>
<td>Filters dataset object count metrics for a labeling job.</td>
</tr>
</tbody>
</table>

**Amazon SageMaker Feature Store Metrics**

**Feature Store Consumption Metrics**
### Metric Description

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ConsumedReadRequestsUnits</td>
<td>The number of consumed read units over the specified time period. You can retrieve the consumed read units for a feature store runtime operation and its corresponding feature group.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: All</td>
</tr>
<tr>
<td>ConsumedWriteRequestsUnits</td>
<td>The number of consumed write units over the specified time period. You can retrieve the consumed write units for a feature store runtime operation and its corresponding feature group.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: All</td>
</tr>
<tr>
<td>ConsumedReadCapacityUnits</td>
<td>The number of provisioned read capacity units consumed over the specified time period. You can retrieve the consumed read capacity units for a feature store runtime operation and its corresponding feature group.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: All</td>
</tr>
<tr>
<td>ConsumedWriteCapacityUnits</td>
<td>The number of provisioned write capacity units consumed over the specified time period. You can retrieve the consumed write capacity units for a feature store runtime operation and its corresponding feature group.</td>
</tr>
<tr>
<td></td>
<td>Units: None</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: All</td>
</tr>
</tbody>
</table>

**Dimensions for Feature Store Consumption Metrics**
<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeatureGroupName, OperationName</td>
<td>Filters feature store runtime consumption metrics of the feature group and the operation that you've specified.</td>
</tr>
</tbody>
</table>

**Feature Store Operational Metrics**

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
</table>
| Invocations | The number of requests made to the feature store runtime operations over the specified time period.  
Units: None  
Valid statistics: Sum |
| Operation 4XXErrors | The number of requests made to the Feature Store runtime operations where the operation returned a 4xx HTTP response code. For each 4xx response, 1 is sent; otherwise, 0 is sent.  
Units: None  
Valid statistics: Average, Sum |
| Operation 5XXErrors | The number of requests made to the feature store runtime operations where the operation returned a 5xx HTTP response code. For each 5xx response, 1 is sent; otherwise, 0 is sent.  
Units: None  
Valid statistics: Average, Sum |
| Throttled Requests | The number of requests made to the feature store runtime operations where the request got throttled. For each throttled request, 1 is sent; otherwise, 0 is sent.  
Units: None |
<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valid statistics: Average, Sum</td>
<td></td>
</tr>
</tbody>
</table>

**Latency**
The time interval to process requests made to the Feature Store runtime operations. This interval is measured from the time SageMaker receives the request until it returns a response to the client.

Units: Microseconds

Valid statistics: Average, Sum, Min, Max, Sample Count, Percentiles

**Dimensions for Feature Store Operational Metrics**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeatureGroupName, OperationName</td>
<td>Filters feature store runtime operational metrics of the feature group and the operation that you've specified. You can use these dimensions for non batch operations, such as GetRecord, PutRecord, and DeleteRecord.</td>
</tr>
<tr>
<td>OperationName</td>
<td>Filters feature store runtime operational metrics for the operation that you've specified. You can use this dimension for batch operations such as BatchGetRecord.</td>
</tr>
</tbody>
</table>

**SageMaker Pipelines Metrics**
The AWS/Sagemaker/ModelBuildingPipeline namespace includes the following metrics for pipeline executions.

Two categories of Pipelines execution metrics are available:

- **Execution Metrics across All Pipelines** – Account level pipeline execution metrics (for all pipelines in the current account)
- **Execution Metrics by Pipeline** – Pipeline execution metrics per pipeline

Metrics are available at a 1-minute frequency.
## Pipelines Execution Metrics

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Execution Started</td>
<td>The number of pipeline executions that started.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>ExecutionFailed</td>
<td>The number of pipeline executions that failed.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>Execution Succeeded</td>
<td>The number of pipeline executions that succeeded.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>Execution Stopped</td>
<td>The number of pipeline executions that stopped.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>Execution Duration</td>
<td>The duration in milliseconds that the pipeline execution ran.</td>
</tr>
<tr>
<td></td>
<td>Units: Milliseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>

## Dimensions for Execution Metrics by Pipeline

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PipelineName</td>
<td>Filters pipeline execution metrics for a specified pipeline.</td>
</tr>
</tbody>
</table>
**Pipelines Step Metrics**

The AWS/Sagemaker/ModelBuildingPipeline namespace includes the following metrics for pipeline steps.

Metrics are available at a 1-minute frequency.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>StepStarted</td>
<td>The number of steps that started.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>StepFailed</td>
<td>The number of steps that failed.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>StepSucceeded</td>
<td>The number of steps that succeeded.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>StepStopped</td>
<td>The number of steps that stopped.</td>
</tr>
<tr>
<td></td>
<td>Units: Count</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum</td>
</tr>
<tr>
<td>StepDuration</td>
<td>The duration in milliseconds that the step ran.</td>
</tr>
<tr>
<td></td>
<td>Units: Milliseconds</td>
</tr>
<tr>
<td></td>
<td>Valid statistics: Average, Sum, Min, Max, Sample Count</td>
</tr>
</tbody>
</table>

**Dimensions for Pipelines Step Metrics**
<table>
<thead>
<tr>
<th>Dimension</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PipelineName</td>
<td>Filters step metrics for a specified pipeline and step.</td>
</tr>
<tr>
<td>StepName</td>
<td></td>
</tr>
</tbody>
</table>

Log Amazon SageMaker Events with Amazon CloudWatch

To help you debug your compilation jobs, processing jobs, training jobs, endpoints, transform jobs, notebook instances, and notebook instance lifecycle configurations, anything an algorithm container, a model container, or a notebook instance lifecycle configuration sends to stdout or stderr is also sent to Amazon CloudWatch Logs. In addition to debugging, you can use these for progress analysis.

Logs

The following table lists all of the logs provided by Amazon SageMaker.

Logs

<table>
<thead>
<tr>
<th>Log Group Name</th>
<th>Log Stream Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>/aws/sagemaker/CompilationJobs</td>
<td>[compilation-job-name]</td>
</tr>
<tr>
<td>/aws/sagemaker/Endpoints/([EndpointName]</td>
<td>([production-variant-name]/[instance-id])</td>
</tr>
<tr>
<td></td>
<td>(For Asynchronous Inference endpoints) [production-variant-name]/[instance-id]/data-log</td>
</tr>
<tr>
<td></td>
<td>(For Inference Pipelines) [production-variant-name]/[instance-id]/[container-name provided in SageMaker model]</td>
</tr>
<tr>
<td>/aws/sagemaker/groundtruth/WorkerActivity</td>
<td>aws/sagemaker/groundtruth/worker-activity/[requester-AWS-Id]-[region]/[timestamp]</td>
</tr>
<tr>
<td>Log Group Name</td>
<td>Log Stream Name</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>/aws/sagemaker/InferenceRecommendationsJobs</td>
<td>[inference-recommendations-job-name]/execution</td>
</tr>
<tr>
<td></td>
<td>[inference-recommendations-job-name]/CompilationJob/[compilation-job-name]</td>
</tr>
<tr>
<td></td>
<td>[inference-recommendations-job-name]/Endpoint/[endpoint-name]</td>
</tr>
<tr>
<td>/aws/sagemaker/LabelingJobs</td>
<td>[labeling-job-name]</td>
</tr>
<tr>
<td>/aws/sagemaker/NotebookInstances</td>
<td>[notebook-instance-name]/[LifecycleConfigHook]</td>
</tr>
<tr>
<td></td>
<td>[notebook-instance-name]/jupyter.log</td>
</tr>
<tr>
<td>/aws/sagemaker/ProcessingJobs</td>
<td>[processing-job-name]/[hostname]-[epoch_timestamp]</td>
</tr>
<tr>
<td>/aws/sagemaker/studio</td>
<td>[domain-id]/[user-profile-name]/[app-type]/[app-name]</td>
</tr>
<tr>
<td></td>
<td>[domain-id]/domain-shared/rstudioserverpro/default</td>
</tr>
<tr>
<td>/aws/sagemaker/TrainingJobs</td>
<td>[training-job-name]/algo-[instance-number-in-cluster]-[epoch_timestamp]</td>
</tr>
<tr>
<td>/aws/sagemaker/TransformJobs</td>
<td>[transform-job-name]/[instance-id]-[epoch_timestamp]</td>
</tr>
<tr>
<td></td>
<td>[transform-job-name]/[instance-id]-[epoch_timestamp]/data-log</td>
</tr>
<tr>
<td></td>
<td>[transform-job-name]/[instance-id]-[epoch_timestamp]/[container-name provided in SageMaker model] (For Inference Pipelines)</td>
</tr>
</tbody>
</table>
Note

1. The /aws/sagemaker/NotebookInstances/[LifecycleConfigHook] log stream is created when you create a notebook instance with a lifecycle configuration. For more information, see Customize a Notebook Instance Using a Lifecycle Configuration Script.
2. For Inference Pipelines, if you don't provide container names, the platform uses **container-1, container-2**, and so on, corresponding to the order provided in the SageMaker model.

For more information about logging events with CloudWatch logging, see What is Amazon CloudWatch Logs? in the Amazon CloudWatch User Guide.

Log Amazon SageMaker API Calls with AWS CloudTrail

Amazon SageMaker is integrated with AWS CloudTrail, a service that provides a record of actions taken by a user, role, or an AWS service in SageMaker. CloudTrail captures all API calls for SageMaker, with the exception of InvokeEndpoint and InvokeEndpointAsync, as events. The calls captured include calls from the SageMaker console and code calls to the SageMaker API operations. If you create a trail, you can enable continuous delivery of CloudTrail events to an Amazon S3 bucket, including events for SageMaker. If you don't configure a trail, you can still view the most recent events in the CloudTrail console in Event history. Using the information collected by CloudTrail, you can determine the request that was made to SageMaker, the IP address from which the request was made, who made the request, when it was made, and additional details.

To learn more about CloudTrail, see the AWS CloudTrail User Guide.

By default, log data is stored in CloudWatch Logs indefinitely. However, you can configure how long to store log data in a log group. For information, see Change Log Data Retention in CloudWatch Logs in the Amazon CloudWatch Logs User Guide.

SageMaker Information in CloudTrail

CloudTrail is enabled on your AWS account when you create the account. When activity occurs in Amazon SageMaker, that activity is recorded in a CloudTrail event along with other AWS service events in Event history. You can view, search, and download recent events in your AWS account. For more information, see Viewing Events with CloudTrail Event History.
For an ongoing record of events in your AWS account, including events for Amazon SageMaker, create a trail. A trail enables CloudTrail to deliver log files to an Amazon S3 bucket. By default, when you create a trail in the console, the trail applies to all AWS Regions. The trail logs events from all Regions in the AWS partition and delivers the log files to the Amazon S3 bucket that you specify. Additionally, you can configure other AWS services to further analyze and act upon the event data collected in CloudTrail logs. For more information, see the following:

- **Overview for Creating a Trail**
- **CloudTrail Supported Services and Integrations**
- **Configuring Amazon SNS Notifications for CloudTrail**
- **Receiving CloudTrail Log Files from Multiple Regions** and **Receiving CloudTrail Log Files from Multiple Accounts**

All SageMaker actions, with the exception of `InvokeEndpoint` and `InvokeEndpointAsync`, are logged by CloudTrail and are documented in the **Operations**. For example, calls to the `CreateTrainingJob`, `CreateEndpoint` and `CreateNotebookInstance` actions generate entries in the CloudTrail log files.

Every event or log entry contains information about who generated the request. The identity information helps you determine the following:

- Whether the request was made with root or IAM user credentials.
- Whether the request was made with temporary security credentials for a role or federated user.
- Whether the request was made by another AWS service.

For more information, see the **CloudTrail userIdentity Element**.

**Operations Performed by Automatic Model Tuning**

SageMaker supports logging non-API service events to your CloudTrail log files for automatic model tuning jobs. These events are related to your tuning jobs but, are not the direct result of a customer request to the public AWS API. For example, when you create a hyperparameter tuning job by calling `CreateHyperParameterTuningJob`, SageMaker creates training jobs to evaluate various combinations of hyperparameters to find the best result. Similarly, when you call `StopHyperParameterTuningJob` to stop a hyperparameter tuning job, SageMaker might stop any of the associated running training jobs. Non-API events for your tuning jobs are logged to
CloudTrail to help you improve governance, compliance, and operational and risk auditing of your AWS account.

Log entries that result from non-API service events have an eventType of AwsServiceEvent instead of AwsApiCall.

**Understanding SageMaker Log File Entries**

A trail is a configuration that enables delivery of events as log files to an S3 bucket that you specify. CloudTrail log files contain one or more log entries. An event represents a single request from any source and includes information about the requested action, the date and time of the action, request parameters, and so on. CloudTrail log files are not an ordered stack trace of the public API calls, so they do not appear in any specific order.

The following examples a log entry for the CreateEndpoint action, which creates an endpoint to deploy a trained model.

```json
{
    "eventVersion":"1.05",
    "userIdentity": { 
        "type":"IAMUser",
        "principalId":"AIXDAYQEXAMPLEUMLYNL",
        "arn":"arn:aws:iam::123456789012:user/intern",
        "accountId":"123456789012",
        "accessKeyId":"ASXIAGXEXAMPLEQULKNXV",
        "userName":"intern"
    },
    "eventTime":"2018-01-02T13:39:06Z",
    "eventSource":"sagemaker.amazonaws.com",
    "eventName":"CreateEndpoint",
    "awsRegion":"us-west-2",
    "sourceIPAddress":"127.0.0.1",
    "userAgent":"USER_AGENT",
    "requestParameters": { 
        "endpointName":"ExampleEndpoint",
        "endpointConfigName":"ExampleEndpointConfig"
    },
    "responseElements": { 
        "endpointArn":"arn:aws:sagemaker:us-west-2:123456789012:endpoint/exampleendpoint"
    },
    "requestID":"6b1b42b9-EXAMPLE"
}
```
The following example is a log entry for the CreateModel action, which creates one or more containers to host a previously trained model.

```json
{
    "eventVersion":"1.05",
    "userIdentity": {
        "type":"IAMUser",
        "principalId":"AIXDAYQEXAMPLEUMLYNGL",
        "arn":"arn:aws:iam::123456789012:user/intern",
        "accountId":"123456789012",
        "accessKeyId":"ASXIAGXEXAMPLEQULKNXV",
        "userName":"intern"
    },
    "eventTime":"2018-01-02T15:23:46Z",
    "eventSource":"sagemaker.amazonaws.com",
    "eventName":"CreateModel",
    "awsRegion":"us-west-2",
    "sourceIPAddress":"127.0.0.1",
    "userAgent":"USER_AGENT",
    "requestParameters": {
        "modelName":"ExampleModel",
        "primaryContainer": {
            "image":"174872318107.dkr.ecr.us-west-2.amazonaws.com/kmeans:latest"
        },
        "executionRoleArn":"arn:aws:iam::123456789012:role/EXAMPLEARN"
    },
    "responseElements": {
    },
    "requestID":"417b8dab-EXAMPLE",
    "eventID":"0f2b3e81-EXAMPLE",
    "eventType":"AwsApiCall",
    "recipientAccountId":"444455556666"
}
```
Monitoring user resource access from Amazon SageMaker Studio Classic

With Amazon SageMaker Studio Classic, you can monitor user resource access. To view resource access activity, you can configure AWS CloudTrail to monitor and record user activities by following the steps in Log Amazon SageMaker API Calls with AWS CloudTrail.

However, the AWS CloudTrail logs for resource access only list the Studio Classic execution IAM role as the identifier. This level of logging is enough to audit user activity when each user profile has a distinct execution role. However, when a single execution IAM role is shared between several user profiles, you can't get information about the specific user that accessed the AWS resources.

You can get information about which specific user performed an action in an AWS CloudTrail log when using a shared execution role, using the sourceIdentity configuration to propagate the Studio Classic user profile name. For more information about source identity, see Monitor and control actions taken with assumed roles.

Prerequisites

- Install and configure the AWS Command Line Interface following the steps in Installing or updating the latest version of the AWS CLI.
- Ensure that Studio Classic users in your domain don't have a policy that allows them to update or modify the domain.
- To turn on or turn off sourceIdentity propagation, all apps in the domain must be in the Stopped or Deleted state. For more information about how to stop and shut down apps, see Shut down and Update Studio Classic Apps.
- All execution roles must have the following trust policy permissions:
  - Any role that the domain's execution role assumes must have the sts:SetSourceIdentity permission in the trust policy as follows.

```json
{
   "Version": "2012-10-17",
   "Statement": [
      {
         "Effect": "Allow",
         "Principal": {
            "Service": "sagemaker.amazonaws.com"
         }
      }
   ]
}
```
• When you assume a role with another role, called role chaining, do the following:
  • Permissions for sts:SetSourceIdentity are required in both the permissions policy of the principal that is assuming the role, and in the role trust policy of the target role. Otherwise, the assume role operation will fail.
  • This role chaining can happen in Studio Classic or any other downstream service, such as Amazon EMR. For more information about role chaining, see Roles terms and concepts.

Considerations when using sourceIdentity

When you make AWS API calls from Studio Classic notebooks, SageMaker Canvas, or Amazon SageMaker Data Wrangler, the sourceIdentity is only recorded in CloudTrail if those calls are made using the Studio Classic execution role session or any chained role from that session.

When these API calls invoke other services to perform additional operations, sourceIdentity logging depends on the specific implementation of the invoked services.

• Amazon SageMaker Training, Processing, and Pipeline: When you create a job using these features, the job creation APIs are not able to ingest the sourceIdentity that exists in the session. As a result, any AWS API calls made from these jobs do not record sourceIdentity in CloudTrail logs.

• Amazon EMR: When connecting to Amazon EMR from Studio Classic using runtime roles, administrators must explicitly set the PropagateSourceIdentity field. This ensures that Amazon EMR applies the sourceIdentity from the calling credentials to a job or query session. The sourceIdentity is then recorded in CloudTrail logs.

Note

The following exceptions apply when using sourceIdentity.
- SageMaker Studio Classic shared spaces do not support `sourceIdentity` passthrough. AWS API calls made from SageMaker shared spaces do not record `sourceIdentity` in CloudTrail logs.

- If AWS API calls are made from sessions that are created by users or other services and the sessions are not based on the Studio Classic execution role session, then the `sourceIdentity` is not recorded in CloudTrail logs.

### Turn on `sourceIdentity`

The ability to propagate the user profile name as the `sourceIdentity` in Studio Classic is turned off by default.

To enable the ability to propagate the user profile name as the `sourceIdentity`, use the AWS CLI during domain creation and domain update. This feature is enabled at the domain level and not at the user profile level.

After you enable this configuration, administrators can view the user profile in the AWS CloudTrail log for the service accessed. The user profile is given as the `sourceIdentity` value in the `userIdentity` section. For more information about using AWS CloudTrail logs with SageMaker, see [Log Amazon SageMaker API Calls with AWS CloudTrail](#).

You can use the following code to enable the propagation of the user profile name as the `sourceIdentity` during domain creation using the `create-domain` API.

```bash
create-domain
--domain-name <value>
--auth-mode <value>
--default-user-settings <value>
--subnet-ids <value>
--vpc-id <value>
[--tags <value>]
[--app-network-access-type <value>]
[--home-efs-file-system-kms-key-id <value>]
[--kms-key-id <value>]
[--app-security-group-management <value>]
[--domain-settings "ExecutionRoleIdentityConfig=USER_PROFILE_NAME"]
[--cli-input-json <value>]
```
You can enable the propagation of the user profile name as the `sourceIdentity` during domain update using the `update-domain` API.

To update this configuration, all apps in the domain must be in the Stopped or Deleted state. For more information about how to stop and shut down apps, see [Shut down and Update Studio Classic Apps](#).

Use the following code to enable the propagation of the user profile name as the `sourceIdentity`.

```
update-domain
  --domain-id <value>
  [--default-user-settings <value>]
  [--domain-settings-for-update "ExecutionRoleIdentityConfig=USER_PROFILE_NAME"]
  [--cli-input-json <value>]
  [--generate-cli-skeleton <value>]
```

**Turn off `sourceIdentity`**

You can also turn off the propagation of the user profile name as the `sourceIdentity` using the AWS CLI. This occurs during domain update by passing the `ExecutionRoleIdentityConfig=DISABLED` value for the `--domain-settings-for-update` parameter as part of the `update-domain` API call.

In the AWS CLI, use the following code to disable the propagation of the user profile name as the `sourceIdentity`.

```
update-domain
  --domain-id <value>
  [--default-user-settings <value>]
  [--domain-settings-for-update "ExecutionRoleIdentityConfig=DISABLED"]
  [--cli-input-json <value>]
  [--generate-cli-skeleton <value>]
```
Automating Amazon SageMaker with Amazon EventBridge

Amazon EventBridge monitors status change events in Amazon SageMaker. EventBridge enables you to automate SageMaker and respond automatically to events such as a training job status change or endpoint status change. Events from SageMaker are delivered to EventBridge in near real time. You can write simple rules to indicate which events are of interest to you, and what automated actions to take when an event matches a rule. For an example of how to create a rule, see Schedule a Pipeline with Amazon EventBridge.

**Note**

SageMaker may send multiple events to EventBridge for each state change. This behavior is expected and does not necessarily indicate an error.

Some examples of the actions that can be automatically triggered include the following:

- Invoking an AWS Lambda function
- Invoking Amazon EC2 Run Command
- Relaying the event to Amazon Kinesis Data Streams
- Activating an AWS Step Functions state machine
- Notifying an Amazon SNS topic or an AWS SMS queue

**SageMaker events monitored by EventBridge**

- SageMaker model state change
- Training job state change
- Hyperparameter tuning job state change
- Transform job state change
- Endpoint state change
- Feature group state change
- Model package state change
- Pipeline execution state change
- Pipeline step state change
- Processing job state change
• **SageMaker image state change**
• **SageMaker image version state change**
• **Endpoint deployment state change**
• **Model card state change**

## SageMaker model state change

Indicates a change in the state of a SageMaker model. The state changes when a SageMaker model is either created or deleted.

```
{
  "source": ["aws.sagemaker"],
  "detail-type": ["SageMaker Model State Change"],
  "Resources": ["arn:aws:sagemaker:us-east-1:123456789012:model/model-name"]
}
```

If a model is specified under `Resources`, an event will be generated and sent to EventBridge when the state of this model changes. If you do not specify a value for `Resources`, an event will generate when the status of any of the SageMaker models associated with your account changes.

## Training job state change

Indicates a change in the status of a SageMaker training job.

If the value of `TrainingJobStatus` is Failed, the event contains the `FailureReason` field, which provides a description of why the training job failed.

```
{
  "version": "0",
  "id": "844e2571-85d4-695f-b930-0153b71dcb42",
  "detail-type": "SageMaker Training Job State Change",
  "source": "aws.sagemaker",
  "account": "123456789012",
  "time": "2018-10-06T12:26:13Z",
  "region": "us-east-1",
  "resources": [
    "arn:aws:sagemaker:us-east-1:123456789012:training-job/kmeans-1"
  ],
  "detail": {
    "TrainingJobName": "89c96cc8-dded-4739-afcc-6f1dc936701d",
  }
}
```
  "TrainingJobStatus": "Completed",
  "SecondaryStatus": "Completed",
  "HyperParameters": {
    "Hyper": "Parameters"
  },
  "AlgorithmSpecification": {
    "TrainingImage": "TrainingImage",
    "TrainingInputMode": "TrainingInputMode"
  },
  "RoleArn": "arn:aws:iam::123456789012:role/SMRole",
  "InputDataConfig": [
    {
      "ChannelName": "Train",
      "DataSource": {
        "S3DataSource": {
          "S3DataType": "S3DataType",
          "S3Uri": "S3Uri",
          "S3DataDistributionType": "S3DataDistributionType"
        }
      },
      "ContentType": "ContentType",
      "CompressionType": "CompressionType",
      "RecordWrapperType": "RecordWrapperType"
    }
  ],
  "OutputDataConfig": {
    "KmsKeyId": "KmsKeyId",
    "S3OutputPath": "S3OutputPath"
  },
  "ResourceConfig": {
    "InstanceType": "InstanceType",
    "InstanceCount": 3,
    "VolumeSizeInGB": 20,
    "VolumeKmsKeyId": "VolumeKmsKeyId"
  },
  "VpcConfig": {

  },
  "StoppingCondition": {
    "MaxRuntimeInSeconds": 60
  },
  "CreationTime": "1583831889050"
Hyperparameter tuning job state change

Indicates a change in the status of a SageMaker hyperparameter tuning job.
"RoleArn": "roleArn",
"InputDataConfig": [
  {
    "ChannelName": "channelName",
    "DataSource": {
      "S3DataSource": {
        "S3DataType": "s3DataType",
        "S3Uri": "s3Uri",
        "S3DataDistributionType": "s3DistributionType"
      }
    },
    "ContentType": "contentType",
    "CompressionType": "gz",
    "RecordWrapperType": "RecordWrapper"
  }
],
"VpcConfig": {
  "SecurityGroupIds": ["securityGroupIds"],
  "Subnets": ["subnets"
],
"OutputDataConfig": {
  "KmsKeyId": "kmsKeyId",
  "S3OutputPath": "s3OutputPath"
},
"ResourceConfig": {
  "InstanceType": "instanceType",
  "InstanceCount": 10,
  "VolumeSizeInGB": 500,
  "VolumeKmsKeyId": "volumeKeyId"
},
"StoppingCondition": {
  "MaxRuntimeInSeconds": 3600
},
"HyperParameterTuningJobStatus": "status",
"CreationTime": "1583831889050",
"LastModifiedTime": "1583831889050",
"TrainingJobStatusCounters": {
  "Completed": 1,
  "InProgress": 0,
Transform job state change

Indicates a change in the status of a SageMaker batch transform job.

If the value of TransformJobStatus is Failed, the event contains the FailureReason field, which provides a description of why the training job failed.
Endpoint state change

Indicates a change in the status of a SageMaker hosted real-time inference endpoint.

The following shows an event with an endpoint in the IN_SERVICE state.

```json
{
  "version": "0",
  "id": "d2921b5a-b0ad-cace-a8e3-0f159d018e06",
  "detail-type": "SageMaker Endpoint State Change",
  "source": "aws.sagemaker",
  "account": "123456789012",
  "time": "1583831889050",
  "region": "us-west-2",
  "resources": [
  ]
}
```
Feature group state change

Indicates a change either in the FeatureGroupStatus or the OfflineStoreStatus of a SageMaker feature group.
Model package state change

Indicates a change in the status of a SageMaker model package.
"source": [
  "aws.sagemaker"
],
"detail": {
  "ModelPackageGroupName": "versionedmp-p-idy6c3e1fiqj",
  "ModelPackageVersion": 2,
  "CreationTime": "2021-02-24T17:00:14Z",
  "InferenceSpecification": {
    "Containers": [
      {
        "Image": "257758044811.dkr.ecr.us-east-2.amazonaws.com/sagemaker-xgboost:1.0-1-cpu-py3",
        "ImageDigest": "sha256:4dc8a7e4a010a19bb9e0a6b063f355393f6e623603361bd8b105f554d4f0c004",
        "ModelDataUrl": "s3://sagemaker-project-p-idy6c3e1fiqj/versionedmp-p-idy6c3e1fiqj/AbaloneTrain/pipelines-4r83jejmhov-TrainAbaloneModel-xw869y8C4a/output/model.tar.gz"
      }
    ],
    "SupportedContentTypes": ["text/csv"],
    "SupportedResponseMIMETypes": ["text/csv"],
  },
  "ModelPackageStatus": "Completed",
  "ModelPackageStatusDetails": {
    "ValidationStatuses": [],
    "ImageScanStatuses": []
  },
  "CertifyForMarketplace": false,
  "ModelApprovalStatus": "Rejected",
  "MetadataProperties": {
    "GeneratedBy": "arn:aws:sagemaker:us-east-2:123456789012:pipeline/versionedmp-p-idy6c3e1fiqj/execution/4r83jejmhov"
  },
  "ModelMetrics": {
    "ModelQuality": {
      "Model package state change": 6126
    }
  }
}
Pipeline execution state change

Indicates a change in the status of a SageMaker pipeline execution.

currentPipelineExecutionStatus and previousPipelineExecutionStatus can be one of the following values:

- Executing
- Succeeded
- Failed
- Stopping
- Stopped

```json
{
  "version": "0",
  "id": "315c1398-40ff-a850-213b-158f73kd93ir",
  "detail-type": "SageMaker Model Building Pipeline Execution Status Change",
  "source": "aws.sagemaker",
  "account": "123456789012",
  "time": "2021-03-15T16:10:11Z",
  "region": "us-east-1",
  "resources": ["arn:aws:sagemaker:us-east-1:123456789012:pipeline/myPipeline-123",
  "arn:aws:sagemaker:us-east-1:123456789012:pipeline/myPipeline-123/execution/p4jn9xou8a8s"],
  "detail": {
    "pipelineExecutionDisplayName": "SomeDisplayName",
    "currentPipelineExecutionStatus": "Succeeded",
    "previousPipelineExecutionStatus": "Executing",
    "executionStartTime": "2021-03-15T16:03:13Z",
```

Pipeline step state change

Indicates a change in the status of a SageMaker pipeline step.

If there is a cache hit, the event contains the cacheHitResult field. currentStepStatus and previousStepStatus can be one of the following values:

- Starting
- Executing
- Succeeded
- Failed
- Stopping
- Stopped

If the value of currentStepStatus is Failed, the event contains the failureReason field, which provides a description of why the step failed.
Processing job state change

Indicates a change in the status of a SageMaker Processing job.

The following example event is for a failed Processing job, where the ProcessingJobStatus value is Failed.

{
  "version": "0",
  "id": "0a15f67d-aa23-0123-0123-01a23w89r01t",
  "detail-type": "SageMaker Processing Job State Change",
  "source": "aws.sagemaker",
  "account": "123456789012",
  "time": "2019-05-31T21:49:54Z",
  "region": "us-east-1",
  "resources": ["arn:aws:sagemaker:us-west-2:037210630506:processing-job/integ-test-analytics-algo-54ee3282-5899-4aa3-afc2-7ce1d02"],
  "detail": {
    "ProcessingInputs": [{
      "InputName": "InputName",
      "S3Input": {
        "S3Uri": "s3://input/s3/uri",
        "LocalPath": "/opt/ml/processing/input/local/path",
        "S3DataType": "MANIFEST_FILE",
        "S3InputMode": "PIPE",
        "S3DataDistributionType": "FULLYREPLICATED"
      }
    }]
  }
}
Amazon SageMaker

Developer Guide

"ProcessingOutputConfig": {
  "Outputs": [
    {
      "OutputName": "OutputName",
      "S3Output": {
        "S3Uri": "s3://output/s3/uri",
        "LocalPath": "/opt/ml/processing/output/local/path",
        "S3UploadMode": "CONTINUOUS"
      }
    }
  ],
  "KmsKeyId": "KmsKeyId"
},
"ProcessingJobName": "integ-test-analytics-algo-54ee3282-5899-4aa3-afc2-7ce1d02",
"ProcessingResources": {
  "ClusterConfig": {
    "InstanceCount": 3,
    "InstanceType": "ml.c5.xlarge",
    "VolumeSizeInGB": 5,
    "VolumeKmsKeyId": "VolumeKmsKeyId"
  }
},
"StoppingCondition": {
  "MaxRuntimeInSeconds": 2000
},
"AppSpecification": {
  "ImageUri": "012345678901.dkr.ecr.us-west-2.amazonaws.com/processing-uri:latest"
},
"NetworkConfig": {
  "EnableInterContainerTrafficEncryption": true,
  "EnableNetworkIsolation": false,
  "VpcConfig": {
    "SecurityGroupIds": ["SecurityGroupId1", "SecurityGroupId2", "SecurityGroupId3"],
    "Subnets": ["Subnet1", "Subnet2"]
  }
},
"RoleArn": "arn:aws:iam::037210630506:role/SageMakerPowerUser",
"ExperimentConfig": {},
"ProcessingJobArn": "arn:aws:sagemaker:us-west-2:037210630506:processing-job/integ-test-analytics-algo-54ee3282-5899-4aa3-afc2-7ce1d02",
"ProcessingJobStatus": "Failed",
"FailureReason": "InternalServerError: We encountered an internal error. Please try again."
"ProcessingEndTime": 1704320746000,
SageMaker image state change

Indicates a change in the status of a SageMaker image.

```
{
    "version": "0",
    "id": "cee033a3-17d8-49f8-865f-b9ebf485d9ee",
    "detail-type": "SageMaker Image State Change",
    "source": "aws.sagemaker",
    "account": "123456789012",
    "time": "2021-04-29T01:29:59Z",
    "region": "us-east-1",
    "resources": ["arn:aws:sagemaker:us-west-2:123456789012:image/cee033a3-17d8-49f8-865f-b9ebf485d9ee"],
    "detail": {
        "ImageName": "cee033a3-17d8-49f8-865f-b9ebf485d9ee",
        "ImageArn": "arn:aws:sagemaker:us-west-2:123456789012:image/cee033a3-17d8-49f8-865f-b9ebf485d9ee",
        "ImageStatus": "Creating",
        "Version": 1.0,
        "Tags": {}
    }
}
```

SageMaker image version state change

Indicates a change in the status of a SageMaker image version.

```
{
    "version": "0",
    "id": "07fc4615-ebd7-15fc-1746-243411f09f04",
    "detail-type": "SageMaker Image Version State Change",
    "source": "aws.sagemaker",
    "account": "123456789012",
    "time": "2021-04-29T01:29:59Z",
    "region": "us-east-1",
}
```
For more information about the status values and their meanings for SageMaker jobs, endpoints, and pipelines, see the following links:

- AlgorithmStatus
- EndpointStatus
- FeatureGroupStatus
- HyperParameterTuningJobStatus
- LabelingJobStatus
- ModelPackageStatus
- NotebookInstanceStatus
- PipelineExecutionStatus
- StepStatus
- ProcessingJobStatus
- TrainingJobStatus
- TransformJobStatus

For more information, see the Amazon EventBridge User Guide.
Endpoint deployment state change

⚠️ Important

The following examples may not work for all endpoints. For a list of features that may exclude your endpoint, see the [Exclusions](#) page.

Indicates a state change for an endpoint deployment. The following example shows an endpoint updating with a blue/green canary deployment.

```json
{
    "version": "0",
    "id": "0bd4a141-0a02-9d8a-f977-3924c3fb259c",
    "detail-type": "SageMaker Endpoint Deployment State Change",
    "source": "aws.sagemaker",
    "account": "123456789012",
    "time": "2021-10-25T01:52:12Z",
    "region": "us-west-2",
    "resources": [null],
    "detail": {
        "EndpointName": "sample-endpoint",
        "EndpointConfigName": "sample-endpoint-config-1",
        "ProductionVariants": [
            {
                "VariantName": "AllTraffic",
                "CurrentWeight": 1,
                "DesiredWeight": 1,
                "CurrentInstanceCount": 3,
                "DesiredInstanceCount": 3
            }
        ],
        "EndpointStatus": "UPDATING",
        "CreationTime": 1635195148181,
        "LastModifiedTime": 1635195148181,
        "Tags": [],
        "PendingDeploymentSummary": {
            "EndpointConfigName": "sample-endpoint-config-2",
```
The following example indicates a state change for an endpoint deployment, which is being updated with new capacity on an existing endpoint configuration.

```json
{
  "version": "0",
  "id": "0bd4a141-0a02-9d8a-f977-3924c3fb259c",
  "detail-type": "SageMaker Endpoint Deployment State Change",
  "source": "aws.sagemaker",
  "account": "123456789012",
  "time": "2021-10-25T01:52:12Z",
  "region": "us-west-2",
  "resources": [
  ],
  "detail": {
    "EndpointName": "sample-endpoint",
    "EndpointConfigName": "sample-endpoint-config-1",
    "ProductionVariants": [
      {
```
The following secondary deployment statuses are also available for endpoints (found in the VariantStatus object.

- **Creating**: creating instances for the production variant.

  Example message: "Launching X instance(s)."

- **Deleting**: terminating instances for the production variant.

  Example message: "Terminating X instance(s)."

- **Updating**: updating capacity for the production variant.

  Example messages: "Launching X instance(s).", "Scaling out desired instance count to X."

- **ActivatingTraffic**: turning on traffic for the production variant.

  Example message: "Activating traffic on canary capacity of X instance(s)."

- **Baking**: waiting period to monitor the CloudWatch alarms in the auto-rollback configuration.

  Example message: "Baking for X seconds (TerminationWaitInSeconds) with traffic enabled on full capacity of Y instance(s)."
Model card state change

Indicates a change in the status of an Amazon SageMaker Model Card. For more information about model cards, see Amazon SageMaker Model Cards.

```json
{
    "version": "0",
    "id": "aa7a9c4f-2caa-4d04-a6de-e67227ba4302",
    "detail-type": "SageMaker Model Card State Change",
    "source": "aws.sagemaker",
    "account": "123456789012",
    "time": "2022-11-30T00:00:00Z",
    "region": "us-east-1",
    "resources": [
        "arn:aws:sagemaker:us-east-1:123456789012:model-card/example-card"
    ],
    "detail": {
        "ModelCardVersion": 2,
        "LastModifiedTime": "2022-12-03T00:09:44.893854735Z",
        "LastModifiedBy": {
            "DomainId": "us-east-1",
            "UserProfileArn": "arn:aws:sagemaker:us-east-1:123456789012:user-profile/user",
            "UserProfileName": "user"
        },
        "CreationTime": "2022-12-03T00:09:33.084Z",
        "CreatedBy": {
            "DomainId": "us-east-1",
            "UserProfileArn": "arn:aws:sagemaker:us-east-1:123456789012:user-profile/user",
            "UserProfileName": "user"
        },
        "ModelCardName": "example-card",
        "ModelId": "example-model",
        "ModelCardStatus": "Draft",
        "AccountId": "123456789012",
        "SecurityConfig": {}  
    }
}
```
Amazon SageMaker Reference

Topics

- Machine Learning Frameworks and Languages
- API Reference
- SageMaker Distribution Images
- Document History for Amazon SageMaker
- Docker Registry Paths and Example Code

Machine Learning Frameworks and Languages

You can use Python and R natively in Amazon SageMaker notebook kernels. There are also kernels that support specific frameworks. A very popular way to get started with SageMaker is to use the Amazon SageMaker Python SDK. It provides open source Python APIs and containers that make it easy to train and deploy models in SageMaker, as well as examples for use with several different machine learning and deep learning frameworks.

For information about using specific frameworks or how to use R in SageMaker, see the following topics.

Languages SDKs and user guides:

- Amazon SageMaker Python SDK
- R
- API Reference

Machine learning and deep learning frameworks guides:

- Apache MXNet
- Apache Spark
- Chainer
- Hugging Face
Use Apache MXNet with Amazon SageMaker

You can use SageMaker to train and deploy a model using custom MXNet code. The Amazon SageMaker Python SDK MXNet estimators and models and the SageMaker open-source MXNet container make writing a MXNet script and running it in SageMaker easier.

What do you want to do?

I want to train a custom MXNet model in SageMaker.

For documentation, see Train a Model with MXNet.

I have an MXNet model that I trained in SageMaker, and I want to deploy it to a hosted endpoint.

For more information, see Deploy MXNet models.

I have an MXNet model that I trained outside of SageMaker, and I want to deploy it to a SageMaker endpoint.

For more information, see Deploy Endpoints from Model Data.

I want to see the API documentation for Amazon SageMaker Python SDK MXNet classes.

For more information, see MXNet Classes.

I want to find the SageMaker MXNet container repository.

For more information, see SageMaker MXNet Container GitHub repository.

I want to find information about MXNet versions supported by AWS Deep Learning Containers.

For more information, see Available Deep Learning Container Images.

For general information about writing MXNet script mode training scripts and using MXNet script mode estimators and models with SageMaker, see Using MXNet with the SageMaker Python SDK.
Use Apache Spark with Amazon SageMaker

Amazon SageMaker Spark is an open source Spark library that helps you construct Spark machine learning (ML) pipelines with SageMaker. This simplifies the integration of Spark ML stages with SageMaker stages, like model training and hosting. For information about SageMaker Spark, see the [SageMaker Spark](https://github.com/aws-sagemaker/sagemaker-spark) GitHub repository.

The SageMaker Spark library is available in Python and Scala. You can use SageMaker Spark to train models in SageMaker using `org.apache.spark.sql.DataFrame` data frames in your Spark clusters. After model training, you can also host the model using SageMaker hosting services.

The SageMaker Spark library, `com.amazonaws.services.sagemaker.sparksdk`, provides the following classes, among others:

- **SageMakerEstimator**—Extends the `org.apache.spark.ml.Estimator` interface. You can use this estimator for model training in SageMaker.


- **SageMakerModel**—Extends the `org.apache.spark.ml.Model` class. You can use this `SageMakerModel` for model hosting and obtaining inferences in SageMaker.

You can download the source code for both Python Spark (PySpark) and Scala libraries from the [SageMaker Spark](https://github.com/aws-sagemaker/sagemaker-spark) GitHub repository.

For installation and examples of the SageMaker Spark library, see [SageMaker Spark for Scala examples](https://github.com/aws-sagemaker/sagemaker-spark) or [SageMaker Spark for Python (PySpark) examples](https://github.com/aws-sagemaker/sagemaker-spark).

If you use Amazon EMR on AWS to manage Spark clusters, see [Apache Spark](https://aws.amazon.com/emr). For more information on using Amazon EMR in SageMaker, see [Prepare data using Amazon EMR](https://docs.aws.amazon.com/sagemaker/latest/dg/prepare-data-using-amazon-emr.html).

**Topics**

- [Integrate Your Apache Spark Application with SageMaker](https://docs.aws.amazon.com/sagemaker/latest/dg/use-apache-spark.html)

- [SageMaker Spark for Scala examples](https://github.com/aws-sagemaker/sagemaker-spark)

- [SageMaker Spark for Python (PySpark) examples](https://github.com/aws-sagemaker/sagemaker-spark)
Integrate Your Apache Spark Application with SageMaker

The following is a high-level summary of the steps for integrating your Apache Spark application with SageMaker.

1. Continue data preprocessing using the Apache Spark library that you are familiar with. Your dataset remains a DataFrame in your Spark cluster. Load your data into a DataFrame and preprocess it so that you have a features column with `org.apache.spark.ml.linalg.Vector` of Doubles, and an optional label column with values of Double type.

2. Use the estimator in the SageMaker Spark library to train your model. For example, if you choose the k-means algorithm provided by SageMaker for model training, you call the `KMeansSageMakerEstimator.fit` method.

Provide your DataFrame as input. The estimator returns a `SageMakerModel` object.

>Note

`SageMakerModel` extends the `org.apache.spark.ml.Model`.

The fit method does the following:

a. Converts the input DataFrame to the protobuf format by selecting the features and label columns from the input DataFrame and uploading the protobuf data to an Amazon S3 bucket. The protobuf format is efficient for model training in SageMaker.

b. Starts model training in SageMaker by sending a SageMaker `CreateTrainingJob` request. After model training has completed, SageMaker saves the model artifacts to an S3 bucket. SageMaker assumes the IAM role that you specified for model training to perform tasks on your behalf. For example, it uses the role to read training data from an S3 bucket and to write model artifacts to a bucket.

c. Creates and returns a `SageMakerModel` object. The constructor does the following tasks, which are related to deploying your model to SageMaker.

   i. Sends a `CreateModel` request to SageMaker.

   ii. Sends a `CreateEndpointConfig` request to SageMaker.
iii. Sends a CreateEndpoint request to SageMaker, which then launches the specified resources, and hosts the model on them.

3. You can get inferences from your model hosted in SageMaker with the SageMakerModel.transform.

Provide an input DataFrame with features as input. The transform method transforms it to a DataFrame containing inferences. Internally, the transform method sends a request to the InvokeEndpoint SageMaker API to get inferences. The transform method appends the inferences to the input DataFrame.

SageMaker Spark for Scala examples

Amazon SageMaker provides an Apache Spark library (SageMaker Spark) that you can use to integrate your Apache Spark applications with SageMaker. For example, you might use Apache Spark for data preprocessing and SageMaker for model training and hosting. For information about the SageMaker Apache Spark library, see Use Apache Spark with Amazon SageMaker.

Download Spark for Scala

You can download the source code and examples for both Python Spark (PySpark) and Scala libraries from the SageMaker Spark GitHub repository.

For detailed instructions on installing the SageMaker Spark library, see SageMaker Spark.

SageMaker Spark SDK for Scala is available in the Maven central repository. Add the Spark library to your project by adding the following dependency to your pom.xml file:

- If your project is built with Maven, add the following to your pom.xml file:

  ```xml
  <dependency>
    <groupId>com.amazonaws</groupId>
    <artifactId>sagemaker-spark_2.11</artifactId>
    <version>spark_2.2.0-1.0</version>
  </dependency>
  ```

- If your project depends on Spark 2.1, add the following to your pom.xml file:

  ```xml
  <dependency>
    <groupId>com.amazonaws</groupId>
    <artifactId>sagemaker-spark_2.11</artifactId>
  </dependency>
  ```
Spark for Scala example

This section provides example code that uses the Apache Spark Scala library provided by SageMaker to train a model in SageMaker using DataFrames in your Spark cluster. This is then followed by examples on how to Use Custom Algorithms for Model Training and Hosting on Amazon SageMaker with Apache Spark and Use the SageMakerEstimator in a Spark Pipeline.

The following example hosts the resulting model artifacts using SageMaker hosting services. For more details on this example, see Getting Started: K-Means Clustering on SageMaker with SageMaker Spark SDK. Specifically, this example does the following:

- Uses the KMeansSageMakerEstimator to fit (or train) a model on data

Because the example uses the k-means algorithm provided by SageMaker to train a model, you use the KMeansSageMakerEstimator. You train the model using images of handwritten single-digit numbers (from the MNIST dataset). You provide the images as an input DataFrame. For your convenience, SageMaker provides this dataset in an Amazon S3 bucket.

In response, the estimator returns a SageMakerModel object.

- Obtains inferences using the trained SageMakerModel

To get inferences from a model hosted in SageMaker, you call the SageMakerModel.transform method. You pass a DataFrame as input. The method transforms the input DataFrame to another DataFrame containing inferences obtained from the model.

For a given input image of a handwritten single-digit number, the inference identifies a cluster that the image belongs to. For more information, see K-Means Algorithm.

```scala
import org.apache.spark.sql.SparkSession
import com.amazonaws.services.sagemaker.sparksdk.IAMRole
import com.amazonaws.services.sagemaker.sparksdk.algorithms
import com.amazonaws.services.sagemaker.sparksdk.algorithms.KMeansSageMakerEstimator

val spark = SparkSession.builder.getOrCreate
```
// load mnist data as a dataframe from libsvm
val region = "us-east-1"
val trainingData = spark.read.format("libsvm")
  .option("numFeatures", "784")
  .load(s"s3://sagemaker-sample-data-$region/spark/mnist/train/")
val testData = spark.read.format("libsvm")
  .option("numFeatures", "784")
  .load(s"s3://sagemaker-sample-data-$region/spark/mnist/test/")

val roleArn = "arn:aws:iam::account-id:role/rolename"

val estimator = new KMeansSageMakerEstimator(
  sagemakerRole = IAMRole(roleArn),
  trainingInstanceType = "ml.p2.xlarge",
  trainingInstanceCount = 1,
  endpointInstanceType = "ml.c4.xlarge",
  endpointInitialInstanceCount = 1)
.setK(10).setFeatureDim(784)

// train
val model = estimator.fit(trainingData)
val transformedData = model.transform(testData)
transformedData.show

The example code does the following:

- Loads the MNIST dataset from an S3 bucket provided by SageMaker (awsai-sparksdk-dataset) into a Spark DataFrame (mnistTrainingDataFrame):
val roleArn = "arn:aws:iam::account-id:role/rolename"

trainingData.show()

The show method displays the first 20 rows in the data frame:

| label | features                  |
+-------+---------------------------|
| 5.0   | (784, [152, 153, 154...  |
| 0.0   | (784, [127, 128, 129...  |
| 4.0   | (784, [160, 161, 162...  |
| 1.0   | (784, [158, 159, 160...  |
| 9.0   | (784, [208, 209, 210...  |
| 2.0   | (784, [155, 156, 157...  |
| 1.0   | (784, [124, 125, 126...  |
| 3.0   | (784, [151, 152, 153...  |
| 1.0   | (784, [152, 153, 154...  |
| 4.0   | (784, [134, 135, 161...  |
| 3.0   | (784, [123, 124, 125...  |
| 5.0   | (784, [216, 217, 218...  |
| 3.0   | (784, [143, 144, 145...  |
| 6.0   | (784, [72, 73, 74, 99... |
| 1.0   | (784, [151, 152, 153...  |
| 7.0   | (784, [211, 212, 213...  |
| 2.0   | (784, [151, 152, 153...  |
| 8.0   | (784, [159, 160, 161...  |
| 6.0   | (784, [100, 101, 102...  |
| 9.0   | (784, [209, 210, 211...  |

only showing top 20 rows

In each row:

- The **label** column identifies the image's label. For example, if the image of the handwritten number is the digit 5, the label value is 5.

- The **features** column stores a vector (org.apache.spark.ml.linalg.Vector) of Double values. These are the 784 features of the handwritten number. (Each handwritten number is a 28 x 28-pixel image, making 784 features.)

- Creates a SageMaker estimator (KMeansSageMakerEstimator)
The `fit` method of this estimator uses the k-means algorithm provided by SageMaker to train models using an input DataFrame. In response, it returns a `SageMakerModel` object that you can use to get inferences.

**Note**

The `KMeansSageMakerEstimator` extends the `SageMakerSageMakerEstimator`, which extends the Apache Spark Estimator.

```scala
val estimator = new KMeansSageMakerEstimator(
  sagemakerRole = IAMRole(roleArn),
  trainingInstanceType = "ml.p2.xlarge",
  trainingInstanceCount = 1,
  endpointInstanceType = "ml.c4.xlarge",
  endpointInitialInstanceCount = 1)
  .setK(10).setFeatureDim(784)
```

The constructor parameters provide information that is used for training a model and deploying it on SageMaker:

- **trainingInstanceType** and **trainingInstanceCount**—Identify the type and number of ML compute instances to use for model training.
- **endpointInstanceType**—Identifies the ML compute instance type to use when hosting the model in SageMaker. By default, one ML compute instance is assumed.
- **endpointInitialInstanceCount**—Identifies the number of ML compute instances initially backing the endpoint hosting the model in SageMaker.
- **sagemakerRole**—SageMaker assumes this IAM role to perform tasks on your behalf. For example, for model training, it reads data from S3 and writes training results (model artifacts) to S3.

**Note**

This example implicitly creates a SageMaker client. To create this client, you must provide your credentials. The API uses these credentials to authenticate requests to
After the `KMeansSageMakerEstimator` object has been created, you set the following parameters, are used in model training:

- The number of clusters that the k-means algorithm should create during model training. You specify 10 clusters, one for each digit, 0 through 9.
- Identifies that each input image has 784 features (each handwritten number is a 28 x 28-pixel image, making 784 features).

Calls the estimator `fit` method

```scala
// train
val model = estimator.fit(trainingData)
```

You pass the input DataFrame as a parameter. The model does all the work of training the model and deploying it to SageMaker. For more information see, [Integrate Your Apache Spark Application with SageMaker](#). In response, you get a `SageMakerModel` object, which you can use to get inferences from your model deployed in SageMaker.

You provide only the input DataFrame. You don't need to specify the registry path to the k-means algorithm used for model training because the `KMeansSageMakerEstimator` knows it.

Calls the `SageMakerModel`'s `transform` method to get inferences from the model deployed in SageMaker.

The `transform` method takes a DataFrame as input, transforms it, and returns another DataFrame containing inferences obtained from the model.

```scala
val transformedData = model.transform(testData)
transformedData.show
```

For simplicity, we use the same DataFrame as input to the `transform` method that we used for model training in this example. The `transform` method does the following:

- Serializes the `features` column in the input DataFrame to protobuf and sends it to the SageMaker endpoint for inference.
- Deserializes the protobuf response into the two additional columns (`distance_to_cluster` and `closest_cluster`) in the transformed DataFrame.
The show method gets inferences to the first 20 rows in the input DataFrame:

<table>
<thead>
<tr>
<th>label</th>
<th>features</th>
<th>distance_to_cluster</th>
<th>closest_cluster</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.0</td>
<td>(784, [152, 153, 154, ...])</td>
<td>1767.897705078125</td>
<td>4.0</td>
</tr>
<tr>
<td>0.0</td>
<td>(784, [127, 128, 129, ...])</td>
<td>1392.157470703125</td>
<td>5.0</td>
</tr>
<tr>
<td>4.0</td>
<td>(784, [160, 161, 162, ...])</td>
<td>1671.571669921875</td>
<td>9.0</td>
</tr>
<tr>
<td>1.0</td>
<td>(784, [158, 159, 160, ...])</td>
<td>1182.6082763671875</td>
<td>6.0</td>
</tr>
<tr>
<td>9.0</td>
<td>(784, [208, 209, 210, ...])</td>
<td>1390.4002685546875</td>
<td>0.0</td>
</tr>
<tr>
<td>2.0</td>
<td>(784, [155, 156, 157, ...])</td>
<td>1713.988037109375</td>
<td>1.0</td>
</tr>
<tr>
<td>1.0</td>
<td>(784, [124, 125, 126, ...])</td>
<td>1246.3016357421875</td>
<td>2.0</td>
</tr>
<tr>
<td>3.0</td>
<td>(784, [151, 152, 153, ...])</td>
<td>1753.229248046875</td>
<td>4.0</td>
</tr>
<tr>
<td>1.0</td>
<td>(784, [152, 153, 154, ...])</td>
<td>978.8394165039062</td>
<td>2.0</td>
</tr>
<tr>
<td>4.0</td>
<td>(784, [134, 135, 161, ...])</td>
<td>1623.176513671875</td>
<td>3.0</td>
</tr>
<tr>
<td>3.0</td>
<td>(784, [123, 124, 125, ...])</td>
<td>1533.863525390625</td>
<td>4.0</td>
</tr>
<tr>
<td>5.0</td>
<td>(784, [216, 217, 218, ...])</td>
<td>1469.3571777347375</td>
<td>6.0</td>
</tr>
<tr>
<td>3.0</td>
<td>(784, [143, 144, 145, ...])</td>
<td>1736.765869140625</td>
<td>4.0</td>
</tr>
<tr>
<td>6.0</td>
<td>(784, [72, 73, 74, 99, ...])</td>
<td>1473.69384765625</td>
<td>8.0</td>
</tr>
<tr>
<td>1.0</td>
<td>(784, [151, 152, 153, ...])</td>
<td>944.88720703125</td>
<td>2.0</td>
</tr>
<tr>
<td>7.0</td>
<td>(784, [211, 212, 213, ...])</td>
<td>1285.9071044921875</td>
<td>3.0</td>
</tr>
<tr>
<td>2.0</td>
<td>(784, [151, 152, 153, ...])</td>
<td>1635.0125732421875</td>
<td>1.0</td>
</tr>
<tr>
<td>8.0</td>
<td>(784, [159, 160, 161, ...])</td>
<td>1436.3162841796875</td>
<td>6.0</td>
</tr>
<tr>
<td>6.0</td>
<td>(784, [100, 101, 102, ...])</td>
<td>1499.7366943359375</td>
<td>7.0</td>
</tr>
<tr>
<td>9.0</td>
<td>(784, [209, 210, 211, ...])</td>
<td>1364.631958078125</td>
<td>6.0</td>
</tr>
</tbody>
</table>

You can interpret the data, as follows:

- A handwritten number with the label 5 belongs to cluster 4 (closest_cluster).
- A handwritten number with the label 0 belongs to cluster 5.
- A handwritten number with the label 4 belongs to cluster 9.
- A handwritten number with the label 1 belongs to cluster 6.

**Topics**

- [Use Custom Algorithms for Model Training and Hosting on Amazon SageMaker with Apache Spark](#)
- [Use the SageMakerEstimator in a Spark Pipeline](#)
Use Custom Algorithms for Model Training and Hosting on Amazon SageMaker with Apache Spark

In SageMaker Spark for Scala examples, you use the kMeansSageMakerEstimator because the example uses the k-means algorithm provided by Amazon SageMaker for model training. You might choose to use your own custom algorithm for model training instead. Assuming that you have already created a Docker image, you can create your own SageMakerEstimator and specify the Amazon Elastic Container Registry path for your custom image.

The following example shows how to create a KMeansSageMakerEstimator from the SageMakerEstimator. In the new estimator, you explicitly specify the Docker registry path to your training and inference code images.

```scala
import com.amazonaws.services.sagemaker.sparksdk.IAMRole
import com.amazonaws.services.sagemaker.sparksdk.SageMakerEstimator
import com.amazonaws.services.sagemaker.sparksdk.transformation.serializers.ProtobufRequestRowSerializer
import com.amazonaws.services.sagemaker.sparksdk.transformation.deserializers.KMeansProtobufResponseRowDeserializer

val estimator = new SageMakerEstimator(
  trainingImage =
  "811284229777.dkr.ecr.us-east-1.amazonaws.com/kmeans:1",
  modelImage =
  "811284229777.dkr.ecr.us-east-1.amazonaws.com/kmeans:1",
  requestRowSerializer = new ProtobufRequestRowSerializer(),
  responseRowDeserializer = new KMeansProtobufResponseRowDeserializer(),
  hyperParameters = Map("k" -> "10", "feature_dim" -> "784"),
  sagemakerRole = IAMRole(roleArn),
  trainingInstanceType = "ml.p2.xlarge",
  trainingInstanceCount = 1,
  endpointInstanceType = "ml.c4.xlarge",
  endpointInitialInstanceCount = 1,
  trainingSparkDataFormat = "sagemaker")
```

In the code, the parameters in the SageMakerEstimator constructor include:

- **trainingImage** — Identifies the Docker registry path to the training image containing your custom code.
- **modelImage** — Identifies the Docker registry path to the image containing inference code.
• **requestRowSerializer** — Implements
  com.amazonaws.services.sagemaker.sparksdk.transformation.RequestRowSerializer.

  This parameter serializes rows in the input DataFrame to send them to the model hosted in SageMaker for inference.

• **responseRowDeserializer** — Implements
  com.amazonaws.services.sagemaker.sparksdk.transformation.ResponseRowDeserializer.

  This parameter deserializes responses from the model, hosted in SageMaker, back into a DataFrame.

• **trainingSparkDataFormat** — Specifies the data format that Spark uses when uploading training data from a DataFrame to S3. For example, "sagemaker" for protobuf format, "csv" for comma-separated values, and "libsvm" for LibSVM format.

You can implement your own RequestRowSerializer and ResponseRowDeserializer to serialize and deserialize rows from a data format that your inference code supports, such as .libsvm or .csv.

**Use the SageMakerEstimator in a Spark Pipeline**

You can use org.apache.spark.ml.Estimators and org.apache.spark.ml.Model models, and SageMakerEstimator estimators and SageMakerModel models in org.apache.spark.ml.Pipeline pipelines, as shown in the following example:

```scala
import org.apache.spark.ml.Pipeline
import org.apache.spark.ml.feature.PCA
import org.apache.spark.sql.SparkSession
import com.amazonaws.services.sagemaker.sparksdk.IAMRole
import com.amazonaws.services.sagemaker.sparksdk.algorithms
import com.amazonaws.services.sagemaker.sparksdk.algorithms.KMeansSageMakerEstimator

val spark = SparkSession.builder.getOrCreate

// load mnist data as a dataframe from libsvm
val region = "us-east-1"
val trainingData = spark.read.format("libsvm")
  .option("numFeatures", "784")
  .load(s"s3://sagemaker-sample-data-$region/spark/mnist/train/"
```
val testData = spark.read.format("libsvm")
  .option("numFeatures", "784")
  .load(s"s3://sagemaker-sample-data-$region/spark/mnist/test/")

// substitute your SageMaker IAM role here
val roleArn = "arn:aws:iam::account-id:role/rolename"

val pcaEstimator = new PCA()
  .setInputCol("features")
  .setOutputCol("projectedFeatures")
  .setK(50)

val kMeansSageMakerEstimator = new KMeansSageMakerEstimator(
  sagemakerRole = IAMRole(integTestingRole),
  requestRowSerializer =
    new ProtobufRequestRowSerializer(featuresColumnName = "projectedFeatures"),
  trainingSparkDataFormatOptions = Map("featuresColumnName" -> "projectedFeatures"),
  trainingInstanceType = "ml.p2.xlarge",
  trainingInstanceCount = 1,
  endpointInstanceType = "ml.c4.xlarge",
  endpointInitialInstanceCount = 1)
  .setK(10).setFeatureDim(50)

val pipeline = new Pipeline().setStages(Array(pcaEstimator, kMeansSageMakerEstimator))

// train
val pipelineModel = pipeline.fit(trainingData)

val transformedData = pipelineModel.transform(testData)
transformedData.show()

The parameter trainingSparkDataFormatOptions configures Spark to serialize to protobuf the "projectedFeatures" column for model training. Additionally, Spark serializes to protobuf the "label" column by default.

Because we want to make inferences using the "projectedFeatures" column, we pass the column name into the ProtobufRequestRowSerializer.

The following example shows a transformed DataFrame:

+-----+--------------------+--------------------+-------------------+---------------+
| label|            features|   projectedFeatures|distance_to_cluster|closest_cluster|
+-----+--------------------+--------------------+-------------------+---------------+
SageMaker Spark for Python (PySpark) examples

Amazon SageMaker provides an Apache Spark Python library (SageMaker PySpark) that you can use to integrate your Apache Spark applications with SageMaker. For example, you might use Apache Spark for data preprocessing and SageMaker for model training and hosting. For information about the SageMaker Apache Spark library, see Use Apache Spark with Amazon SageMaker.

Download PySpark

You can download the source code for both Python Spark (PySpark) and Scala libraries from the SageMaker Spark GitHub repository.

For instructions on installing the SageMaker Spark library, use any of the following options or visit SageMaker PySpark.

- Install using pip:

  ```bash
  pip install sagemaker_pyspark
  ```
• Install from the source:

```bash
git clone git@github.com:aws/sagemaker-spark.git
cd sagemaker-pyspark-sdk
python setup.py install
```

• You can also create a new notebook in a notebook instance that uses either the Sparkmagic (PySpark) or the Sparkmagic (PySpark3) kernel and connect to a remote Amazon EMR cluster.

> **Note**

The Amazon EMR cluster must be configured with an IAM role that has the `AmazonSageMakerFullAccess` policy attached. For information about configuring roles for an EMR cluster, see [Configure IAM Roles for Amazon EMR Permissions to AWS Services](#) in the *Amazon EMR Management Guide*.

**PySpark examples**

For examples on using SageMaker PySpark, see:

- [Using Amazon SageMaker with Apache Spark](#) in Read the Docs.
- [SageMaker Spark](#) GitHub repository.

To run the notebooks on a notebook instance, see [Example Notebooks](#). To run the notebooks on Studio, see [Create or Open an Amazon SageMaker Studio Classic Notebook](#).

**Use Chainer with Amazon SageMaker**

You can use SageMaker to train and deploy a model using custom Chainer code. The SageMaker Python SDK Chainer estimators and models and the SageMaker open-source Chainer container make writing a Chainer script and running it in SageMaker easier.

**What do you want to do?**

I want to train a custom Chainer model in SageMaker.

For a sample Jupyter notebook, see the [Chainer example notebooks](#) in the Amazon SageMaker Examples GitHub repository.
For documentation, see Train a Model with Chainer.

I have a Chainer model that I trained in SageMaker, and I want to deploy it to a hosted endpoint.

For more information, see Deploy Chainer models.

I have a Chainer model that I trained outside of SageMaker, and I want to deploy it to a SageMaker endpoint

For more information, see Deploy Endpoints from Model Data.

I want to see the API documentation for Amazon SageMaker Python SDK Chainer classes.

For more information, see Chainer Classes.

I want to find information about SageMaker Chainer containers.

For more information, see the SageMaker Chainer Container GitHub repository.

For information about supported Chainer versions, and for general information about writing Chainer training scripts and using Chainer estimators and models with SageMaker, see Using Chainer with the SageMaker Python SDK.

Use Hugging Face with Amazon SageMaker

Amazon SageMaker enables customers to train, fine-tune, and run inference using Hugging Face models for Natural Language Processing (NLP) on SageMaker. You can use Hugging Face for both training and inference. This functionality is available through the development of Hugging Face AWS Deep Learning Containers. These containers include Hugging Face Transformers, Tokenizers and the Datasets library, which allows you to use these resources for your training and inference jobs. For a list of the available Deep Learning Containers images, see Available Deep Learning Containers Images. These Deep Learning Containers images are maintained and regularly updated with security patches.

To use the Hugging Face Deep Learning Containers with the SageMaker Python SDK for training, see the Hugging Face SageMaker Estimator. With the Hugging Face Estimator, you can use the Hugging Face models as you would any other SageMaker Estimator. However, using the SageMaker Python SDK is optional. You can also orchestrate your use of the Hugging Face Deep Learning Containers with the AWS CLI and AWS SDK for Python (Boto3).

For more information on Hugging Face and the models available in it, see the Hugging Face documentation.
Training

To run training, you can use any of the thousands of models available in Hugging Face and fine-tune them for your specific use case with additional training. With SageMaker, you can use standard training or take advantage of SageMaker Distributed Data and Model Parallel training. As with other SageMaker training jobs using custom code, you can capture your own metrics by passing a metrics definition to the SageMaker Python SDK as shown in Defining Training Metrics (SageMaker Python SDK). The captured metrics are then accessible via CloudWatch and as a Pandas DataFrame via the TrainingJobAnalytics method. Once your model is trained and fine-tuned, you can use it like any other model to run inference jobs.

How to run training with the Hugging Face Estimator

You can implement the Hugging Face Estimator for training jobs using the SageMaker Python SDK. The SageMaker Python SDK is an open source library for training and deploying machine learning models on SageMaker. For more information on the Hugging Face Estimator, see the SageMaker Python SDK documentation.

With the SageMaker Python SDK, you can run training jobs using the Hugging Face Estimator in the following environments:

- **SageMaker Studio**: Amazon SageMaker Studio is the first fully integrated development environment (IDE) for machine learning (ML). SageMaker Studio provides a single, web-based visual interface where you can perform all ML development steps required to prepare, build, train and tune, deploy and manage models. For information on using Jupyter Notebooks in Studio, see Use Amazon SageMaker Studio Notebooks.

- **SageMaker Notebook Instances**: An Amazon SageMaker notebook instance is a machine learning (ML) compute instance running the Jupyter Notebook App. This app lets you run Jupyter Notebooks in your notebook instance to prepare and process data, write code to train models, deploy models to SageMaker hosting, and test or validate your models without SageMaker Studio features like Debugger, Model Monitoring, and a web-based IDE.

- Locally: If you have connectivity to AWS and have appropriate SageMaker permissions, you can use the SageMaker Python SDK locally to launch remote training and inference jobs for Hugging Face in SageMaker on AWS. This works on your local machine, as well as other AWS services with a connected SageMaker Python SDK and appropriate permissions.
Inference

For inference, you can use your trained Hugging Face model or one of the pretrained Hugging Face models to deploy an inference job with SageMaker. With this collaboration, you only need one line of code to deploy both your trained models and pre-trained models with SageMaker. You can also run inference jobs without having to write any custom inference code. With custom inference code, you can customize the inference logic by providing your own Python script.

How to deploy an inference job using the Hugging Face Deep Learning Containers

You have two options for running inference with SageMaker. You can run inference using a model that you trained, or deploy a pre-trained Hugging Face model.

- **Run inference with your trained model:** You have two options for running inference with your own trained model. You can run inference with a model that you trained using an existing Hugging Face model with the SageMaker Hugging Face Deep Learning Containers, or you can bring your own existing Hugging Face model and deploy it using SageMaker. When you run inference with a model that you trained with the SageMaker Hugging Face Estimator, you can deploy the model immediately after training completes or you can upload the trained model to an Amazon S3 bucket and ingest it when running inference later. If you bring your own existing Hugging Face model, you must upload the trained model to an Amazon S3 bucket and ingest that bucket when running inference as shown in [Deploy your Hugging Face Transformers for inference example](#).

- **Run inference with a pre-trained HuggingFace model:** You can use one of the thousands of pre-trained Hugging Face models to run your inference jobs with no additional training needed. To run inference, you select the pre-trained model from the list of [Hugging Face models](#), as outlined in [Deploy pre-trained Hugging Face Transformers for inference example](#).

What do you want to do?

The following Jupyter Notebooks in the Hugging Face notebooks repository illustrate how to use the Hugging Face Deep Learning Containers with SageMaker in various use cases.

I want to train and deploy a text classification model using Hugging Face in SageMaker with PyTorch.

For a sample Jupyter Notebook, see the [PyTorch Getting Started Demo](#).
I want to train and deploy a text classification model using Hugging Face in SageMaker with TensorFlow.

   For a sample Jupyter Notebook, see the TensorFlow Getting Started example.

I want to run distributed training with data parallelism using Hugging Face and SageMaker Distributed.

   For a sample Jupyter Notebook, see the Distributed Training example.

I want to run distributed training with model parallelism using Hugging Face and SageMaker Distributed.

   For a sample Jupyter Notebook, see the Model Parallelism example.

I want to use a spot instance to train and deploy a model using Hugging Face in SageMaker.

   For a sample Jupyter Notebook, see the Spot Instances example.

I want to capture custom metrics and use SageMaker Checkpointing when training a text classification model using Hugging Face in SageMaker.

   For a sample Jupyter Notebook, see the Training with Custom Metrics example.

I want to train a distributed question-answering TensorFlow model using Hugging Face in SageMaker.

   For a sample Jupyter Notebook, see the Distributed TensorFlow Training example.

I want to train a distributed summarization model using Hugging Face in SageMaker.

   For a sample Jupyter Notebook, see the Distributed Summarization Training example.

I want to train an image classification model using Hugging Face in SageMaker.

   For a sample Jupyter Notebook, see the Vision Transformer Training example.

I want to deploy my trained Hugging Face model in SageMaker.

   For a sample Jupyter Notebook, see the Deploy your Hugging Face Transformers for inference example.

I want to deploy a pre-trained Hugging Face model in SageMaker.

   For a sample Jupyter Notebook, see the Deploy pre-trained Hugging Face Transformers for inference example.
Use PyTorch with Amazon SageMaker

You can use Amazon SageMaker to train and deploy a model using custom PyTorch code. The SageMaker Python SDK PyTorch estimators and models and the SageMaker open-source PyTorch container make writing a PyTorch script and running it in SageMaker easier.

What do you want to do?

I want to train a custom PyTorch model in SageMaker.

For a sample Jupyter notebook, see the PyTorch example notebook in the Amazon SageMaker Examples GitHub repository.

For documentation, see Train a Model with PyTorch.

I have a PyTorch model that I trained in SageMaker, and I want to deploy it to a hosted endpoint.

For more information, see Deploy PyTorch models.

I have a PyTorch model that I trained outside of SageMaker, and I want to deploy it to a SageMaker endpoint.

For more information, see Deploy your own PyTorch model.

I want to see the API documentation for Amazon SageMaker Python SDK PyTorch classes.

For more information, see PyTorch Classes.

I want to find the SageMaker PyTorch container repository.

For more information, see SageMaker PyTorch Container GitHub repository.

I want to find information about PyTorch versions supported by AWS Deep Learning Containers.

For more information, see Available Deep Learning Container Images.

For general information about writing PyTorch training scripts and using PyTorch estimators and models with SageMaker, see Using PyTorch with the SageMaker Python SDK.

R User Guide to Amazon SageMaker

This document will walk you through ways of leveraging Amazon SageMaker features using R. This guide introduces SageMaker's built-in R kernel, how to get started with R on SageMaker, and finally several example notebooks.
The examples are organized in three levels, Beginner, Intermediate, and Advanced. They start from Getting Started with R on SageMaker, continue to end-to-end machine learning with R on SageMaker, and then finish with more advanced topics such as SageMaker Processing with R script, and Bring-Your-Own (BYO) R algorithm to SageMaker.

For information on how to bring your own custom R image to Studio, see Bring your own SageMaker image. For a similar blog article, see Bringing your own R environment to Amazon SageMaker Studio.

RStudio Support in SageMaker

Amazon SageMaker supports RStudio as a fully-managed integrated development environment (IDE) integrated with Amazon SageMaker domain. With RStudio integration, you can launch an RStudio environment in the domain to run your RStudio workflows on SageMaker resources. For more information, see RStudio on Amazon SageMaker.

R Kernel in SageMaker

SageMaker notebook instances support R using a pre-installed R kernel. Also, the R kernel has the reticulate library, an R to Python interface, so you can use the features of SageMaker Python SDK from within an R script.

- reticulatelibrary: provides an R interface to the Amazon SageMaker Python SDK. The reticulate package translates between R and Python objects.

Get Started with R in SageMaker

- Create a Notebook Instance using the t2.medium instance type and default storage size. You can pick a faster instance and more storage if you plan to continue using the instance for more advanced examples, or create a bigger instance later.
- Wait until the status of the notebook is In Service, and then click Open Jupyter.
• Create a new notebook with R kernel from the list of available environments.

• When the new notebook is created, you should see an R logo in the upper right corner of the notebook environment, and also R as the kernel under that logo. This indicates that SageMaker has successfully launched the R kernel for this notebook.

• Alternatively, when you are in a Jupyter notebook, you can use Kernel menu, and then select R from Change Kernel option.
Example Notebooks

Prerequisites

Getting Started with R on SageMaker: This sample notebook describes how you can develop R scripts using Amazon SageMaker’s R kernel. In this notebook you set up your SageMaker environment and permissions, download the abalone dataset from the UCI Machine Learning Repository, do some basic processing and visualization on the data, then save the data as .csv format to S3.

Beginner Level

SageMaker Batch Transform using R Kernel: This sample notebook describes how to conduct a batch transform job using SageMaker’s Transformer API and the XGBoost algorithm. The notebook also uses the Abalone dataset.

Intermediate Level

Hyperparameter Optimization for XGBoost in R: This sample notebook extends the previous beginner notebooks that use the abalone dataset and XGBoost. It describes how to do model tuning with hyperparameter optimization. You will also learn how to use batch transform for batching predictions, as well as how to create a model endpoint to make real-time predictions.

Amazon SageMaker Processing with R: SageMaker Processing lets you preprocess, post-process and run model evaluation workloads. This example shows you how to create an R script to orchestrate a Processing job.

Advanced Level

Train and Deploy Your Own R Algorithm in SageMaker: Do you already have an R algorithm, and you want to bring it into SageMaker to tune, train, or deploy it? This example walks you through
how to customize SageMaker containers with custom R packages, all the way to using a hosted endpoint for inference on your R-origin model.

**Use Scikit-learn with Amazon SageMaker**

You can use Amazon SageMaker to train and deploy a model using custom Scikit-learn code. The SageMaker Python SDK Scikit-learn estimators and models and the SageMaker open-source Scikit-learn containers make writing a Scikit-learn script and running it in SageMaker easier.

**Requirements**

Scikit-learn 1.2 has the following dependencies.

<table>
<thead>
<tr>
<th>Dependency</th>
<th>Minimum version</th>
</tr>
</thead>
<tbody>
<tr>
<td>Python</td>
<td>3.8</td>
</tr>
<tr>
<td>NumPy</td>
<td>1.17.3</td>
</tr>
<tr>
<td>SciPy</td>
<td>1.3.2</td>
</tr>
<tr>
<td>joblib</td>
<td>1.1.1</td>
</tr>
<tr>
<td>threadpoolctl</td>
<td>2.0.0</td>
</tr>
</tbody>
</table>

The SageMaker Scikit-learn container supports the following Scikit-learn versions.

<table>
<thead>
<tr>
<th>Supported Scikit-learn version</th>
<th>Minimum Python version</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.2-1</td>
<td>3.8</td>
</tr>
<tr>
<td>1.0-1</td>
<td>3.7</td>
</tr>
<tr>
<td>0.23-1</td>
<td>3.6</td>
</tr>
<tr>
<td>0.20.0</td>
<td>2.7 or 3.4</td>
</tr>
</tbody>
</table>

For general information about writing Scikit-learn training scripts and using Scikit-learn estimators and models with SageMaker, see [Using Scikit-learn with the SageMaker Python SDK](#).
What do you want to do?

Note
Matplotlib v2.2.3 or newer is required to run the SageMaker Scikit-learn example notebooks.

I want to use Scikit-learn for data processing, feature engineering, or model evaluation in SageMaker.

For a sample Jupyter notebook, see https://github.com/awslabs/amazon-sagemaker-examples/tree/master/sagemaker_processing/scikit_learn_data_processing_and_model_evaluation.

For documentation, see ReadTheDocs.

I want to train a custom Scikit-learn model in SageMaker.


For documentation, see Train a Model with Scikit-learn.

I have a Scikit-learn model that I trained in SageMaker, and I want to deploy it to a hosted endpoint.

For more information, see Deploy Scikit-learn models.

I have a Scikit-learn model that I trained outside of SageMaker, and I want to deploy it to a SageMaker endpoint

For more information, see Deploy Endpoints from Model Data.

I want to see the API documentation for Amazon SageMaker Python SDK Scikit-learn classes.

For more information, see Scikit-learn Classes.

I want to see information about SageMaker Scikit-learn containers.

For more information, see SageMaker Scikit-learn Container GitHub repository.
Use SparkML Serving with Amazon SageMaker

The Amazon SageMaker Python SDK SparkML Serving model and predictor and the Amazon SageMaker open-source SparkML Serving container support deploying Apache Spark ML pipelines serialized with MLeap in SageMaker to get inferences.

For information about using the SparkML Serving container to deploy models to SageMaker, see SageMaker Spark ML Container GitHub repository. For information about the Amazon SageMaker Python SDK SparkML Serving model and predictors, see the SparkML Serving Model and Predictor API documentation.

Use TensorFlow with Amazon SageMaker


Use TensorFlow Version 1.11 and Later

For TensorFlow versions 1.11 and later, the Amazon SageMaker Python SDK supports script mode training scripts.

What do you want to do?

I want to train a custom TensorFlow model in SageMaker.

    For a sample Jupyter notebook, see TensorFlow script mode training and serving.

    For documentation, see Train a Model with TensorFlow.

I have a TensorFlow model that I trained in SageMaker, and I want to deploy it to a hosted endpoint.

    For more information, see Deploy TensorFlow Serving models.

I have a TensorFlow model that I trained outside of SageMaker, and I want to deploy it to a SageMaker endpoint.

    For more information, see Deploying directly from model artifacts.

I want to see the API documentation for Amazon SageMaker Python SDK TensorFlow classes.

    For more information, see TensorFlow Estimator.
I want to find the SageMaker TensorFlow container repository.

For more information, see SageMaker TensorFlow Container GitHub repository.

I want to find information about TensorFlow versions supported by AWS Deep Learning Containers.

For more information, see Available Deep Learning Container Images.

For general information about writing TensorFlow script mode training scripts and using TensorFlow script mode estimators and models with SageMaker, see Using TensorFlow with the SageMaker Python SDK.

**Use TensorFlow Legacy Mode for Versions 1.11 and Earlier**

The Amazon SageMaker Python SDK provides a legacy mode that supports TensorFlow versions 1.11 and earlier. Use legacy mode TensorFlow training scripts to run TensorFlow jobs in SageMaker if:

- You have existing legacy mode scripts that you do not want to convert to script mode.
- You want to use a TensorFlow version earlier than 1.11.

For information about writing legacy mode TensorFlow scripts to use with the SageMaker Python SDK, see TensorFlow SageMaker Estimators and Models.

**Use Triton Inference Server with Amazon SageMaker**

SageMaker enables customers to deploy a model using custom code with NVIDIA Triton Inference Server. This functionality is available through the development of Triton Inference Server Containers. These containers include NVIDIA Triton Inference Server, support for common ML frameworks, and useful environment variables that let you optimize performance on SageMaker. For a list of all available Deep Learning Containers images, see Available Deep Learning Containers Images. Deep Learning Containers images are maintained and regularly updated with security patches.

You can use the Triton Inference Server Container with SageMaker Python SDK as you would any other container in your SageMaker models. However, using the SageMaker Python SDK is optional. You can use Triton Inference Server Containers with the AWS CLI and AWS SDK for Python (Boto3).

For more information on NVIDIA Triton Inference Server see the Triton documentation.
Inference

Note

The Triton Python backend uses shared memory (SHMEM) to connect your code to Triton. SageMaker Inference provides up to half of the instance memory as SHMEM so you can use an instance with more memory for larger SHMEM size.

For inference, you can use your trained ML models with Triton Inference Server to deploy an inference job with SageMaker.

Some of the key features of Triton Inference Server Container are:

- **Support for multiple frameworks**: Triton can be used to deploy models from all major ML frameworks. Triton supports TensorFlow GraphDef and SavedModel, ONNX, PyTorch TorchScript, TensorRT, and custom Python/C++ model formats.

- **Model pipelines**: Triton model ensemble represents a pipeline of one model with pre/post processing logic and the connection of input and output tensors between them. A single inference request to an ensemble triggers the execution of the entire pipeline.

- **Concurrent model execution**: Multiple instances of the same model can run simultaneously on the same GPU or on multiple GPUs.

- **Dynamic batching**: For models that support batching, Triton has multiple built-in scheduling and batching algorithms that combine individual inference requests together to improve inference throughput. These scheduling and batching decisions are transparent to the client requesting inference.

- **Diverse CPU and GPU support**: The models can be executed on CPUs or GPUs for maximum flexibility and to support heterogeneous computing requirements.

What do you want to do?

I want to deploy my trained PyTorch model in SageMaker.

For a sample Jupyter Notebook, see the [Deploy your PyTorch Resnet50 model with Triton Inference Server example](#).
I want to deploy my trained Hugging Face model in SageMaker.

For a sample Jupyter Notebook, see the [Deploy your PyTorch BERT model with Triton Inference Server example](https://example.com).

### API Reference

Making API calls directly from code is cumbersome, and requires you to write code to authenticate your requests. Amazon SageMaker provides the following alternatives:

#### Topics

- Programming Model for Amazon SageMaker
- APIs, CLI, and SDKs

### Programming Model for Amazon SageMaker

Making API calls directly from code is cumbersome, and requires you to write code to authenticate your requests. Amazon SageMaker provides the following alternatives:

- **Use the SageMaker console**–With the console, you don't write any code. You use the console UI to start model training or deploy a model. The console works well for simple jobs, where you use a built-in training algorithm and you don't need to preprocess training data.

- **Modify the example Jupyter notebooks**–SageMaker provides several Jupyter notebooks that train and deploy models using specific algorithms and datasets. Start with a notebook that has a suitable algorithm and modify it to accommodate your data source and specific needs.

- **Write model training and inference code from scratch**–SageMaker provides multiple AWS SDK languages (listed in the overview) and the [Amazon SageMaker Python SDK](https://aws.amazon.com/sdk-for-python/), a high-level Python library that you can use in your code to start model training jobs and deploy the resulting models.
• **The SageMaker Python SDK**—This Python library simplifies model training and deployment. In addition to authenticating your requests, the library abstracts platform specifics by providing simple methods and default parameters. For example:

  • To deploy your model, you call only the `deploy()` method. The method creates a SageMaker model artifact, an endpoint configuration, then deploys the model on an endpoint.

  • If you use a custom framework script for model training, you call the `fit()` method. The method creates a `.gzip` file of your script, uploads it to an Amazon S3 location, and then runs it for model training, and other tasks. For more information, see Machine Learning Frameworks and Languages.

  • To set defaults for SageMaker API calls made by the SageMaker Python SDK, you use a default configuration dictionary. For more information, see Configuring and using defaults with the SageMaker Python SDK.

• **The AWS SDKs**—The SDKs provide methods that correspond to the SageMaker API (see Operations). Use the SDKs to programmatically start a model training job and host the model in SageMaker. SDK clients handle authentication for you, so you don’t need to write authentication code. They are available in multiple languages and platforms. For more information, see the preceding list in the overview.

In Get started, you train and deploy a model using an algorithm provided by SageMaker. That exercise shows how to use both of these libraries. For more information, see Get started.

• **Integrate SageMaker into your Apache Spark workflow**—SageMaker provides a library for calling its APIs from Apache Spark. With it, you can use SageMaker-based estimators in an Apache Spark pipeline. For more information, see Use Apache Spark with Amazon SageMaker.
APIs, CLI, and SDKs

Amazon SageMaker provides APIs, SDKs, and a command line interface that you can use to create and manage notebook instances and train and deploy models.

- Amazon SageMaker Python SDK (Recommended)
- Amazon SageMaker API Reference
- Amazon Augmented AI API Reference
- AWS Command Line Interface
- AWS SDK for .NET
- AWS SDK for C++
- AWS SDK for Go
- AWS SDK for Java
- AWS SDK for JavaScript
- AWS SDK for PHP
- AWS SDK for Python (Boto)
- AWS SDK for Ruby
- Amazon SageMaker Spark

You can also get code examples from the Amazon SageMaker example notebooks GitHub repository.

- Example notebooks

SageMaker Distribution Images

SageMaker Distribution is a collection of Docker images, which includes popular libraries and packages for machine learning, data science, and data analytics visualization. The Docker images include deep learning frameworks such as the following:

- PyTorch
- TensorFlow
- Keras
It also includes popular Python packages such as the following:

- numpy
- scikit-learn
- pandas

Within the container, you can use the following IDEs:

- JupyterLab
- Code Editor, based on Code-OSS (Visual Studio Code Open Source)

Each SageMaker Distribution image has a GPU variant and a CPU variant.

SageMaker Distribution is available in:

- Studio
- Studio Lab

The packages included in the container are guaranteed to be compatible with each other and the runtime is built to work anywhere. You can use the container to run Amazon SageMaker Studio notebooks or SageMaker training jobs. You can also run the container on a local laptop. Use SageMaker Distribution to quickly get started with ML development in your local environment. Seamlessly transition to tasks such as the batch execution of training jobs without needing to reconfigure your runtime environment.

For the list of all supported libraries within SageMaker distribution and their corresponding versions, see the [SageMaker Distribution GitHub](https://github.com/aws/sagemaker). You can also use the pre-built and ready-to-use SageMaker Distribution images from the [Amazon Elastic Container Registry Gallery](https://aws.amazon.com/elastic-containers-repository/)

### Supported packages and versions

For the list of the packages that are installed in a version of SageMaker Distribution, see the RELEASE.md file in the `build_artifacts` directory of the [SageMaker Distribution GitHub](https://github.com/aws/sagemaker) repository.

### SageMaker Distribution Image Support Policy

<table>
<thead>
<tr>
<th>Major</th>
<th>A major version release of Amazon</th>
<th>Half-yearly</th>
</tr>
</thead>
</table>

**Supported packages and versions**

For the list of the packages that are installed in a version of SageMaker Distribution, see the RELEASE.md file in the `build_artifacts` directory of the [SageMaker Distribution GitHub](https://github.com/aws/sagemaker) repository.

### SageMaker Distribution Image Support Policy

<table>
<thead>
<tr>
<th>Major</th>
<th>A major version release of Amazon</th>
<th>Half-yearly</th>
</tr>
</thead>
</table>
SageMaker Distribution upgrades all of its core dependencies to the latest compatible version. SageMaker Distribution can add or remove packages in a major version release. Major versions are denoted by the first number in the version string. For example, 1.0, 2.0, 3.0.

| Minor | A minor version release of Amazon SageMaker Distribution ensures that all of its core dependencies are updated to the latest compatible minor version within the same major version. SageMaker Distribution can add new packages during a minor version release. Minor versions are denoted by the second number in the version string. For example, 1.1, 1.2, or 2.1 | Monthly (additional minor versions released on an as needed basis as well) |
A patch version release of Amazon SageMaker Distribution ensures that all its core dependencies are updated to the latest compatible patch version within the same minor version. SageMaker Distribution does not add or remove packages during a patch version release.

7 days (overnight fixes also deployed based on the severity)

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⚠️ **Important**

- SageMaker Distribution v0.x.y is only used in Studio Classic. SageMaker Distribution v1.x.y is only used in JupyterLab.
- We try to update the Studio images with new versions regularly. If the packages in the Distribution image are out of date, we recommend waiting for the next update.
- Some dependencies, such as Python, are treated differently. Amazon SageMaker Distribution allows for a minor upgrade of Python with a release. For example, you can upgrade Python 3.10 to Python 3.11 when you upgrade from version 4.8 to 5.0.

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**Document History for Amazon SageMaker**

<table>
<thead>
<tr>
<th>Change</th>
<th>Description</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>AWS managed policy updates - New policy</td>
<td>SageMaker added the following new AWS managed policy.</td>
<td>February 2, 2024</td>
</tr>
</tbody>
</table>
- **AmazonSageMakerCanVASBedrockAccess**

AWS managed policy updates
- Updates to existing policies

SageMaker updated the following AWS managed policy.

- **AmazonSageMakerCanVASFullAccess**

AWS managed policy updates
- Updates to existing policies

SageMaker updated the following AWS managed policy.

- **AmazonSageMakerCanVASFullAccess**

AWS managed policy updates
- Updates to existing policies

SageMaker updated the following AWS managed policy.

- **AmazonSageMakerCanVASDataPrepFullAccess**
New features re:Invent 2023

The following new features were introduced at re:Invent 2023.

- SageMaker Canvas Chat for data prep
- Code Editor
- Deep learning containers for large model inference
- Deploy models for real-time inference
- SageMaker Distribution Images
- Domain onboarding simplification
- Amazon S3 Express One Zone
- Foundation model evaluations (FMEval)
- SageMaker HyperPod
- JupyterAI
- JupyterLab in Studio
- SageMaker Notebook Jobs
- SageMaker Pipelines
- SageMakersmart sifting
- SageMaker Studio

AWS managed policy updates - Updates to existing policies

SageMaker updated the following AWS managed policy at re:Invent 2023.

- AmazonSageMakerFullAccess

November 30, 2023
AWS managed policy updates
- Updates to existing policies

SageMaker updated the following AWS managed policies at re:Invent 2023.

- `AmazonSageMakerCanVASAI_servicesAccess`
- `AmazonSageMakerCanVASDataPrepFullAccess`

November 29, 2023

AWS managed policy updates
- New policies

SageMaker added the following new AWS managed policy at re:Invent 2023.

- `AmazonSageMakerClusterInstanceRolePolicy`

November 29, 2023

AWS managed policy updates
- New policy

SageMaker added the following new AWS managed policy.

- `AmazonSageMakerCanVASDataPrepFullAccess`

October 26, 2023

AWS managed policy updates
- New policy

SageMaker added the following new AWS managed policy.

- `AmazonSageMakerCanVASDirectDeployAccess`

October 6, 2023

AWS managed policy updates
- Updates to existing policies

SageMaker updated the following AWS managed policies.

- `AmazonSageMakerCanVASFullAccess`
- `AmazonSageMakerCanVASAI_servicesAccess`

September 29, 2023
### AWS managed policy updates

**- Updates to existing policies**

SageMaker updated the following AWS managed policy.

- [AmazonSageMakerCanvasFullAccess](#)

**August 29, 2023**

**- New policies**

SageMaker added the following new AWS managed policies.

- [AmazonSageMakerPartnerServiceCatalogProductsApiGatewayServiceRolePolicy](#)
- [AmazonSageMakerPartnerServiceCatalogProductsCloudFormationServiceRolePolicy](#)
- [AmazonSageMakerPartnerServiceCatalogProductsLambdaServiceRolePolicy](#)

**August 1, 2023**

**- Updates to existing policies**

SageMaker updated the following AWS managed policy.

- [AmazonSageMakerCanvasFullAccess](#)

**July 24, 2023**

**- Updates to existing policies**

SageMaker updated the following AWS managed policy.

- [AmazonSageMakerModelGovernanceUseAccess](#)

**July 17, 2023**
<table>
<thead>
<tr>
<th>Section</th>
<th>Description</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Refactored Table of Contents</td>
<td>Table of Contents refactored to better reflect the new content.</td>
<td>June 1, 2023</td>
</tr>
<tr>
<td>SageMaker ECR Paths</td>
<td>Docker Registry Paths and Example Code published.</td>
<td>May 25, 2023</td>
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<tr>
<td>AWS managed policy updates</td>
<td>SageMaker updated the following AWS managed policy.</td>
<td>May 10, 2023</td>
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<tr>
<td>- Updates to existing policies</td>
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<td></td>
<td>• AmazonSageMakerGeoSpatialExecutionRole.</td>
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<td>AWS managed policy updates</td>
<td>SageMaker updated the following AWS managed policy.</td>
<td>May 4, 2023</td>
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<td>- Updates to existing policies</td>
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<td>• AmazonSageMakerCanvasFullAccess.</td>
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<tr>
<td>AWS managed policy updates</td>
<td>SageMaker added the following new AWS managed policy.</td>
<td>April 12, 2023</td>
</tr>
<tr>
<td>- New policy</td>
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<td></td>
<td>• AmazonSageMakerModelRegistryFullAccess</td>
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<tr>
<td>AWS managed policy updates</td>
<td>SageMaker updated the following AWS managed policy.</td>
<td>March 24, 2023</td>
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<td>- Updates to existing policies</td>
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<td>• AmazonSageMakerCanvasFullAccess.</td>
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<tr>
<td>AWS managed policy updates</td>
<td>SageMaker added the following new AWS managed policy.</td>
<td>March 23, 2023</td>
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<tr>
<td>---------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------</td>
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<td>- New policy</td>
<td>- AmazonSageMakerCanVasAIServicesAccess</td>
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<table>
<thead>
<tr>
<th>AWS managed policy updates</th>
<th>SageMaker updated the following AWS managed policy.</th>
<th>March 9, 2023</th>
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<tbody>
<tr>
<td>- Updates to existing policies</td>
<td>- AmazonSageMakerNotEbooksServiceRolePolicy</td>
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<table>
<thead>
<tr>
<th>AWS managed policy updates</th>
<th>SageMaker updated the following AWS managed policy.</th>
<th>January 12, 2023</th>
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<tbody>
<tr>
<td>- Updates to existing policies</td>
<td>- AmazonSageMakerNotEbooksServiceRolePolicy</td>
<td></td>
</tr>
</tbody>
</table>
New features re:Invent 2022

The following new features were introduced at re:Invent 2022.

- **SageMaker geospatial capabilities**
- **SageMaker Model Cards**
- **SageMaker Model Dashboard**
- **SageMaker Role Manager**
- **Collaboration with shared spaces**
- **Inference shadow tests**
- **Notebook-based Workflows**
- **Data Wrangler data preparation widget**
- **AutoML step** in Amazon SageMaker Model Building Pipelines
- **Studio Classic Git extension**

AWS managed policy updates - Updates to existing policies

SageMaker updated the following AWS managed policies at re:Invent 2022.

- **AmazonSageMakerFullAccess**
- **AmazonSageMakerFeatureStoreAccess**
- **AmazonSageMakerCanvasFullAccess**

November 30, 2022
AWS managed policy updates
- New policies

SageMaker added the following new AWS managed policies at re:Invent 2022.

- AmazonSageMakerGeoSpatialFullAccess
- AmazonSageMakerGeoSpatialExecutionRole
- AmazonSageMakerModelGovernanceUseAccess

New features re:Invent 2021

The following new features were introduced at re:Invent 2021.

- SageMaker Canvas
- SageMaker Ground Truth Plus
- SageMaker Inference Recommender
- SageMaker Serverless Endpoints
- SageMaker Studio Lab
- SageMaker Studio Notebooks and Amazon EMR
- SageMaker Training Compiler

Autopilot time series data

Amazon SageMaker Autopilot accepts time series as model inputs. For more information, see Amazon SageMaker Autopilot data and problem types.
<table>
<thead>
<tr>
<th>AWS managed policies</th>
<th>Started tracking changes for SageMaker managed policies.</th>
<th>June 10, 2021</th>
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<tbody>
<tr>
<td>New features re:Invent 2020</td>
<td>The following new features were introduced at re:Invent 2020.</td>
<td>December 1, 2020</td>
</tr>
<tr>
<td>• Amazon SageMaker Model Building Pipelines</td>
<td></td>
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<tr>
<td>• Automate MLOps with SageMaker Projects</td>
<td></td>
<td></td>
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<tr>
<td>• SageMaker Edge Manager</td>
<td></td>
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<tr>
<td>• SageMaker Clarify</td>
<td></td>
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<tr>
<td>• SageMaker Data Wrangler</td>
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<tr>
<td>• SageMaker Feature Store</td>
<td></td>
<td></td>
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<tr>
<td>• SageMaker Studio JumpStart</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Register and Deploy Models with Model Registry</td>
<td></td>
<td></td>
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<tr>
<td>• SageMaker Distributed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Deep Profiling with SageMaker Debugger</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Studio Notebooks</td>
<td>SageMaker Studio Notebooks</td>
<td>April 28, 2020</td>
</tr>
</tbody>
</table>
New features re:Invent 2019

The following new features were introduced at re:Invent 2019.

- SageMaker Studio
- SageMaker Studio Notebooks (preview)
- SageMaker Experiments
- SageMaker Autopilot
- SageMaker Debugger
- SageMaker Model Monitor

December 3, 2019

New features re:Invent 2018

The following new features were introduced at re:Invent 2018.

- Amazon SageMaker Ground Truth
- Amazon Elastic Inference
- SageMaker Resources in AWS Marketplace
- SageMaker Inference Pipelines
- SageMaker Neo
- Search Amazon SageMaker Experiments
- Reinforcement Learning
- Associate Git Repositories with SageMaker Notebook Instances
- Semantic Segmentation Algorithm
- Augmented Manifest Files in Training Jobs

November 28, 2018
<table>
<thead>
<tr>
<th>Feature</th>
<th>Description</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Configuring notebook instances</td>
<td>Use shell scripts to configure notebook instances when you create or start them. For more information, see Customize a Notebook Instance.</td>
<td>May 1, 2018</td>
</tr>
<tr>
<td>Application Auto Scaling support</td>
<td>Amazon SageMaker now supports Application Auto Scaling for production variants. For information, see Automatically Scaling SageMaker Models.</td>
<td>February 28, 2018</td>
</tr>
<tr>
<td>TensorFlow 1.5 and MXNet 1.0 support</td>
<td>Amazon SageMaker Deep Learning containers now support TensorFlow 1.5 and Apache MXNet 1.0.</td>
<td>February 27, 2018</td>
</tr>
<tr>
<td>BlazingText algorithm</td>
<td>Amazon SageMaker now supports the BlazingText algorithm.</td>
<td>January 18, 2018</td>
</tr>
<tr>
<td>KMS encryption</td>
<td>Amazon SageMaker now supports KMS encryption for hosting instances and training model artifacts at rest.</td>
<td>January 17, 2018</td>
</tr>
<tr>
<td>CloudTrail support</td>
<td>Amazon SageMaker now supports logging with AWS CloudTrail.</td>
<td>January 11, 2018</td>
</tr>
<tr>
<td>DeepAR Forecasting algorithm</td>
<td>Amazon SageMaker now supports the DeepAR algorithm for time series forecasting.</td>
<td>January 8, 2018</td>
</tr>
</tbody>
</table>
AWS Glossary

For the latest AWS terminology, see the AWS glossary in the AWS Glossary Reference.